miRWalk – Database: Prediction of possible miRNA binding sites by “walking” the genes of three genomes

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A B S T R A C T
MicroRNAs are small, non-coding RNA molecules that can complementarily bind to the miRNA 3’-UTR region to regulate the gene expression by transcriptional repression or induction of miRNA degradation. Increasing evidence suggests a new mechanism by which miRNAs may regulate target gene expression by binding in promoter and amino acid coding regions. Most of the existing databases on miRNAs are restricted to miRNA 3’-UTR region. To address this issue, we present miRWalk, a comprehensive database on miRNAs, which hosts predicted as well as validated miRNA binding sites, information on all known genes of human, mouse and rat.

All miRNAs, mitochondrial genes and 10 kb upstream flanking regions of all known genes of human, mouse and rat were analyzed by using a newly developed algorithm named ‘miRWalk’ as well as with eight already established programs for putative miRNA binding sites. An automated and extensive text-mining search was performed on PubMed database to extract validated information on miRNAs. Combined information was put into a MySQL database.

miRWalk presents predicted and validated information on miRNA-target interaction. Such a resource enables researchers to validate new targets of miRNA not only on 3′-UTR, but also on the other regions of all known genes. The ‘Validated Target module’ is updated every month and the ‘Predicted Target module’ is updated every 6 months. miRWalk is freely available at http://mirwalk.uni-hd.de/.

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1. Introduction

MicroRNAs (miRNAs) are small, non-coding RNA molecules of 21–25 nucleotides in length that regulate the gene expression by base-pairing with the transcripts of their targets i.e. protein-coding genes, leading to downregulation or repression of the target genes [1]. However, target gene activation has also been described in target repression [4]. The 5′ end of miRNA could be determinant in target repression [4]. The 5′ end sequence of miRNA is called “seed” and has a length of 6–8 nucleotides which is energetically favorable for the miRNA target interaction [5]. Mutations in the seed region of a miRNA sequence leads to an inactive interaction [6]. The binding reduces the expression level of target protein by a number of mechanisms including inhibition of translational initiation [7], inhibition of elongation, and induction of deadenylation which decreases mRNA stability and increases the rate of mRNA degradation [8]. The miRNA gene family is one of the largest in higher eukaryotes: more than 700 miRNAs have been identified in the human genome [9], each of them having the potential to bind to hundreds of transcripts. miRNAs are involved in diverse regulatory pathways [10,11], as well as in disease development, progression, prognosis, diagnosis and evaluation of treatment response [12,13].

Computational prediction of miRNA targets is much more challenging in animals than in plants, because animal miRNAs often perform imperfect base-pairing with their target sites [14], unlike plant miRNAs which almost always bind their targets with near perfect complementarity [15]. In the past years, a large number of target prediction programs and databases on experimentally validated information have been developed for animal miRNAs [5,16–26].

For more than a decade, attempts to study the interaction of miRNAs with their targets were focused to the 3′-UTR region of miRNAs. But recent studies on miRNA-target interaction revealed
a new mode of action of miRNAs by which they may regulate the gene expression by targeting promoter as well as amino acid coding (CDS) regions. Tay et al. demonstrated the existence of many naturally occurring miRNA targets sites of miR-314, miR-296 and miR-470 in the CDS of the genes Nanog, Oct4 and Sox2 \[27\]. Guang et al. have shown that argonuata proteins can transport classes of small regulatory RNA to distinct cellular compartments to regulate gene expression \[28\]. In another study, Place et al. have demonstrated that miR-373 targets the promoter sequences of E-cadherin and CSDC2 genes to induce gene expression \[29\]. On the other hand, a few experiments have indicated possible target sites in the 5'-UTR for e.g. \[30\]. Thus, it is of paramount importance to design a new approach which can identify putative miRNA binding sites not only in the 3'-UTR region, but also in other regions (promoter, 5'-UTR, and CDS) of a gene.

Here we present miRWalk (http://mirwalk.uni-hd.de/), a comprehensive database that provides predicted as well as validated miRNA binding site information on miRNAs for human, mouse and rat.

2. Materials and methods

2.1. Genome data acquisition

All sequences [mRNAs, 10 kb upstream flanking regions (assumed promoter regions) and mitochondrial genes] and other necessary information (EntrezID, mRNA and CDS length, gene location and definition) of all known genes of human, mouse and rat were downloaded by submitting identifiers i.e. RefSeqID, Ensembl gene IDs and official gene symbols to Entrez at GenBank \[31\] and Ensembl \[32\] databases. miRNA sequences and other information (e.g. Sanger name, MIID, genomic location of miRNA, stem loop sequence and other accession numbers like stem loop, and mature sequence) were collected from miRBase \[33\]. For Comparative analysis, DIANA-microT (version 3.0), miRanda (August 2010), miRDB (April 2009), PicTar (March 2007), PITA (August 2008), RNA22 (May 2008) and TargetScan/TargetScanS (version 5.1) predictions files were downloaded from their databases, while in case of RNAhybrid, the prediction data was generated by running the executable (version 2.1) of RNAhybrid on bwGRID server.

2.2. Compilation and curation of keyword dictionary

The gene and miRNA dictionaries for human, mouse and rat were compiled from several databases: HUGO Gene Nomenclature Committee (HGNC), Mouse Genome Database (MGD), Rat Genome Database (RGD), gene-centered information at NCBI (Entrez Gene), Targetsan and miRBase. The names, aliases, symbols, official names, synonyms and database identifiers were merged into synonym dictionaries. In the curation step, inappropriate synonymous or expressions that would lead to ambiguous or wrong identifications were detected and removed as described \[34\]. The information on diseases, organs, and OMIM disorders was extracted from MeSH (Medical Subject Heading) and OMIM \[35\]. Whereas the keywords on proteins known to be involved in miRNA processing and cell lines were collected from PubMed database by reading the publications. Collated information was then organized in several lists. Then, these keywords were compiled, classified and stored in different dictionaries according to disease, organ, cell line,
OMIM (Online Mendelian Inheritance in Man) and miRNA processing proteins. Simultaneously, the information on pathways such as gene sets, names and pathway identifiers were retrieved from KEGG [36] and Biocarta (www.biocarta.com) databases. Thereafter, all the available abstracts that contained miRNA keywords in their title and/or abstracts were downloaded from PubMed database.

2.3. miRWalk algorithm

The miRWalk algorithm is based on a computational approach which is written in Perl programming language to identify multiple consecutive Watson–Crick complementary subsequences between miRNA and gene sequences as described in Fig. 1 (Predicted Target module). miRWalk algorithm searches for seeds based on Watson–Crick complementarity, walking on the complete sequence of a gene starting with a heptamer (seven nucleotides) seed from position 1 and 2 of miRNA sequences. As soon as it identifies a heptamer perfect base-pairing, it immediately extends the length of the miRNA seed until a mismatch arises. It then returns all possible hits with 7 or longer matches. These binding sites are then separated on the basis of their identified locations (start, and end positions and regions) in the analyzed sequences. Then it assigns the prediction results in five parts, according to promoter region, 5’-UTR, coding sequence (CDS), and 3’-UTR and mitochondrial genes.

In addition, the probability distribution of random matches of a subsequence (5’ end miRNA sequence) in the analyzed sequence is calculated by using a Poisson distribution [20]. It can be expected that the longer perfect complementation of a seed is associated with a lower probability, thus the higher are the chances of an effective miRNA-target interaction.

2.4. Incorporating other prediction databases in miRWalk

Comparative studies conducted with the earlier miRNA target prediction programs suggested that no program was consistently superior to all others [37,38]. Indeed, it has become a common practice for researchers to look at predictions produced by several miRNA-target prediction programs and focus on their intersection [39,40]. Thus, miRWalk compares the identified miRNA binding sites with the results obtained from eight established miRNA-target prediction programs. These eight established programs are chosen on the basis of their popularity. Finally miRWalk incorporates all the predicted miRNA binding sites of the miRWalk algorithm and of the eight programs as predicted information on miRNAs into its database. All the putative targets (both matched as well as unmatched with miRWalk prediction data) of other programs are stored in miRWalk database.

2.5. Automated text-mining search module

A new module was written in Perl programming language to accomplish the automated text-mining task in the downloaded titles and abstracts from PubMed database against curated dictionaries. We chose abstracts because they are not only more readily available than full text, but are also condensed descriptions focusing on what is central to the study, combining the back-
ground, results and conclusions. Shah et al. [41], demonstrated that the abstract section of articles contains the best proportion of keywords, while the other sections are a better source of biologically relevant data. In details, first the automated text-mining search module downloads all the available abstracts in XML (Extensible Markup Language) format that contained keywords such as ‘microRNA’, ‘miRNA’, ‘micro-RNA’, ‘micro_RNA’, ‘micro RNA’, ‘miR’ in their title and/or abstracts. Thereafter, the keywords from curated dictionaries are detected in the texts of downloaded titles/abstracts (Name Entity Recognition, NER) by string matching script with the help of Perl regular expressions. The identification of named entities allows identifying miRNA terms linked with genes, diseases, organs, cell lines, pathways, OMIM disorders and proteins known to be involved in miRNA processing in an abstract. Then these relationships are processed and stored along with their PubMed identifiers as validated information under Validated Target module of miRWalk database as shown in Fig. 1 (Validated Target module).

2.6. Data processing

After sequence acquisition, miRWalk algorithm was executed to produce putative miRNA binding sites on all downloaded sequences. The bwGRID Cluster (high performance cluster facility) has been used to run miRWalk algorithm in a batch mode by adopting four nodes with 32 processors for the faster identification of miRNA binding sites and computing of probability. After the identification of the miRNA binding sites with miRWalk, the predicted miRNA-target interactions data on 3′-UTR of all known genes of human, mouse and rat were obtained from eight established miRNA-target prediction programs for the comparison of the results with different algorithms. The predicted miRNA-target interaction data on more than 2000 miRNAs generated by both miRWalk and eight established miRNA-target prediction programs was documented in the Predicted Target module.

For validated information, the dictionaries for human, mouse and rat were created from several databases as described in compilation and curation section of this paper. In a keyword identification step, a newly developed automated text-mining search module was used to detect gene and miRNA names in the downloaded PubMed abstracts against curated dictionaries by using NER. Then the collated information was processed and stored under Validated Target module of miRWalk database.

3. Results

3.1. miRWalk database and web interface

miRWalk is implemented as a relational database on a MySQL database management system. The web interface of miRWalk database is divided into two modules: (i) Predicted Target module and (ii) Validated Target module.

The Predicted Target module is classified into six parts: Target Gene, miRNA, Pathway, Chromosome, OMIM and Mitochondrial Target. There are two options implemented for user input, i.e. entering the gene symbols or EntrezIDs either directly into a text-box area or by uploading a file under Target Gene and MicroRNA Target search pages. Whereas other predicted search pages can

![Fig. 3. Overview of the predictions produced by miRWalk algorithm. More information on the estimation of predictions in different databases is given in Supplementary Table 2.](image-url)
be easily queried by a simple selection of drop down lists for e.g. Pathway Target search page can be easily inquired by choosing a pathway name from the given drop down list, to determine how many genes of a selected pathway are the targets of similar and/or different miRNAs. The result tables of all searches display only 20 records and appropriate links are given, which redirect the user to NCBI, Ensembl, UCSC, miRBase, KEGG, Biocarta and OMIM databases for more annotation and further information on the results. Moreover, one can view and/or download the complete result tables by clicking on the providing links such as ‘View Complete Table’, ‘Paging View’ and ‘Download Table’.

The Validated Target module has different search pages. These search pages are organized similar to Predicted Target module and are called; Target Gene, miRNA, Pathway, Disease, Organ, Cell line, miRNA literature, OMIM disorder and miRNA Processing Proteins. The results of this module are hyperlinked to PubMed database. More information on web interface can be found in Supplementary material.

3.2. Overview of predictions produced by miRWalk and eight established databases

Since miRNA-target prediction programs are designed with different combinations of features to perform the same task, it is essential to analyse the distribution of these predictions across miRWalk and eight established miRNA-target prediction programs. We count over seven million predictions in both miRanda (36,507 genes and 2050 miRNAs) and PITA (37,714 genes and 1458 miRNAs), whereas over 4 million unique predictions (only single binding site of a miRNA per 3' UTR region) are counted on 3' UTR by miRWalk algorithm which cover over 40,491 genes (from all three species) and 2057 miRNAs (Fig. 2). An overview of miRWalk predictions on different regions is presented in Fig. 3. In a comparative study, the predictions are decreasing to a surprisingly small 79 records (73 genes and 34 miRNAs) identified in 10 datasets of nine databases considered (Fig. 4). Fig. 4 depicts overlaps of predictions among nine different databases (including miRWalk predictions on 3' UTR region) and indicates that although it shows low total overlap among all databases, in spite of that there is considerable similarity between at least five prediction databases.

3.3. Evaluation of miRWalk algorithm

We compared the performance of miRWalk with respect to eight already established programs. For evaluation we selected a set of genes (positive and negative sets) from TarBase (version 5.0) [42], miRecords (version 3) [21] and miRTarBase (release 2.1) [43] databases on which miRNA binding sites are already verified and published in the PubMed database. We analyzed 1870 positive miRNA-target and 61 negative miRNA-target pairs for the performance of different prediction programs. The positive and negative datasets were given as input to each of these programs by using ‘Target Gene’ which is implemented under Predicted Target module of miRWalk database and the output was analyzed to calculate the Accuracy (Acc), Recall and Precision. The results of the miRWalk algorithm and other are presented in Fig. 5. miRWalk algorithm successfully obtained (97.93% Accuracy, 98.88% Recall and 98.98% Precision) on the input gene sets and the results identified by 8 other programs are shown in Fig. 5.

3.4. Overview of validated data among miRWalk, TarBase, miR2Disease, miRecords, and miRTarBase databases

We compared Validated Target module of miRWalk with TarBase, miR2Disease [44], miRecords, PhenomiR [45] and miRTarBase databases in terms of relationships documented on miRNAs,
genes, diseases, organs, cell lines, OMIM disorders, pathways, miRNA processing proteins (MPP) and PubMed articles. These databases are not only chosen due to their popularity, but also for surveying pertinent literature. For comprehensive analysis, we downloaded TarBase 5.0, miR2Disease (June 2010 release), miRecords (release 3.0) and miRTarBase (release 2.1) files from their respective websites and parsed the information only on human, mouse, and rat genes along with miRNAs. The validated information of PhenomiR was taken from the database home page and its publication. The Validated Target module of miRWalk database increased the validated information more than 3-fold (1572 miRNAs linked to 5080 genes, 691 diseases, 556 organs, 121 cell lines, 2033 OMIM disorders, 375 pathways and 70 MPP) as compared to above mentioned databases.

Fig. 6 depicts an overview of the data stored in miRWalk and five other existing databases.

3.5. Evaluation of validated information of miRWalk database

For the evaluation process, a recall of EntrezGene PubMed articles was conducted. In details, first, we collected Entrez Gene Identifiers of human, mouse and rat miRNAs. Collected identifiers were compiled in a unique identifiers list. Second, a script was written in Perl programming language which automatically extracts all PubMed article Identifiers (PMID) which are marked as relevant for an input miRNA Entrez GeneID. A total of 1360 unique PMID were found to be marked as relevant. Afterwards, these PMIDs were queried against PubMed database to download their titles and abstracts. Then, the downloaded articles (only titles and abstracts) were scanned for miRNA name by using automated text-mining search module. Twelve hundred twenty-five out of 1360 abstracts were found to have at least one miRNA name present in their titles and/or abstracts. Thus, we obtained a recall of 90.07%. For the remaining 10% of articles, we randomly selected 50 PMIDs out of 135 and read their titles/abstracts for miRNA names. After reading, we observed that the randomly selected articles do not have any miRNA names present in their titles/abstract. Therefore, we can expect that the automated text-mining search module can achieve a recall of more than 90.07%.

4. Discussion

miRWalk database is different from existing miRNA resources as: (i) a newly developed algorithm is used to predict all the possible miRNA binding sites by “walking” on the genomes of three genomes (i.e. all protein coding genes, and their 10 kb upstream flanking regions and mitochondrial genes); (ii) the results of miRWalk are presented together with the results obtained from eight

Fig. 5. Evaluation of miRWalk algorithm and eight established prediction programs. In this analysis, 1870 positive and 61 negative miRNA-target pairs were chosen to calculate the performance of the miRWalk algorithm and eight established prediction programs. In order to evaluate the performance of these different prediction programs, we used the statistical parameters, viz., Accuracy (Acc), Recall and Precision. These parameters are based on number of true positives (TP), true negatives (TN), false positives (FP) and false negatives (FN) and are calculated by using the following equations: Accuracy (Acc %) = (TP + TN)/(TP + TN + FP + FN) × 100, Recall (%) = TP/(TP + FN) × 100 and Precision (%) = TP/(TP + FP) × 100.
already established miRNA-target prediction programs for a comprehensive view of predicted miRNA binding sites on mRNA 3'-UTR region; (iii) miRWalk provides a more holistic view of genetic networks of miRNA-gene-pathways and miRNA-gene-OMIM disorder interactions; and (iv) miRWalk hosts new and unique features on experimentally validated miRNAs. Besides validated information, it also offers the information on proteins known to be involved in miRNA processing and available literature on miRNAs. A comparative analysis of features of miRWalk and already established databases on miRNAs is shown in Supplementary Table 5.

The ‘Validated Target module’ is updated every month and the ‘Predicted Target module’ is updated 6 months by executing automated Perl (www.perl.org) and Bioperl (www.bioper.org) scripts on the server of bwGRID Cluster (http://www.urz.uni-heidelberg.de/server/grid/index.en.html).

Similar to miRGator, which integrates three prediction databases (miRanda, PicTar and Targetscan), miRWalk integrates eight databases for a comprehensive study of predictions obtained from different algorithms. This allows the user to take more control over the prediction data that they consider. Not only does our resource conveniently incorporate eight different databases at one place, it also allows users to choose which combinations of databases they would like to consider for their search.

Some research groups have been adopting a new approach for the identification of new targets for known miRNAs. In this approach, miRNA expression in healthy and/or diseased tissue and/or organ was profiled by using miRNA microarray and statistical significant miRNAs were selected for further validation by Northern blot and/or q-PCR experiments. Afterwards, miRNA-target prediction programs were used to identify the possible target genes of these miRNAs. Finally, cell lines and/or animals were used for the knockdown of these miRNAs to measure the expression level of predicted genes. miRWalk database is helpful for this kind of approach, as a user can retrieve information on possible miRNA binding sites on the complete sequence or specific region(s) of targets by supplying miRNA names or uploading a file under Predicted Target module.

Moreover, the basic information on miRNAs (like mature, and stem loop sequence, identifiers, chromosome, strand and band) as well as other necessary data required for a miRNA research, such as regulatory binding sites on upstream and/or downstream flanking regions of pre-miRNA, information on the host gene of miRNA, which miRNAs share a similar seed with the user input miRNAs can be easily obtained. Furthermore, links are given, which redirect the user to NCBI, UCSC (for more annotations and information on miRNAs) and PubMed database for the quick and convenient access to available literature on the user input miRNAs associated with expression, diseases, organs and cell lines.

In the past couple of years, research groups demonstrated the involvement of miRNAs in complicated biological processes/pathways, including the control of developmental timing, haematopoietic cell differentiation, apoptosis, cell proliferation, organ development [10,11,46,47], as well as cancerogenesis [48–51] and other human diseases [40,52–57]. miR2Disease and PhenomiR are well known databases which offer information on miRNA interactions involved in diseases and biological processes. miR2Disease hosts 2663 relationships linked with 347 miRNAs whereas PhenomiR offers 12,192 relationships on 675 miRNAs linked to 146 diseases. In comparison to these resources, miRWalk database hosts 98,887 relationships on 1572 miRNAs from human, mouse and rat linked to 691 diseases. Thus, miRWalk presents much more information on miRNAs linked to human diseases.

Cell lines have been established in life sciences as easy to manipulate model systems for the study of cellular processes. A number of cell lines have been used to investigate the expression of miRNAs under different conditions. Such information is important because of the therapeutic potential of miRNAs in various diseases. miRWalk documents information on 121 cell lines linked to miRNA investigations.

Several studies in mice have demonstrated an important role of Dicer in the generation of mature miRNAs [58,59]. In contrast to the linear miRNA processing pathway that was initially thought to be universal for the biogenesis of all mature miRNAs, multiple discoveries led to the recognition of miRNA-specific differences,
that open a plethora of regulatory options to express and process individual miRNAs differentially. For instance, Drosha-mediated cleavage can regulate individual miRNAs: the hnrNP A1 binds specif-
cically to pri-miR-18a and facilitates its processing [60]. Thus, information on proteins known to be involved in miRNA process-
ning is helpful for researchers investigating the role of miRNAs in or-
gan development and functional maintenance. This kind of information is lacking in the established miRNA databases hosting validated miRNA-target data, while it is implemented under the Validated Target module of miRWalk.

miRBase, TarBase, miRecords and miRDisease are well-known databases for experimentally validated miRNA targets associated with genes, diseases, organs, and pathways. The Validated Targets module of miRWalk offers all the scattered information about miRN-
A interactions in a structured and uniform format. All the infor-
mation of this module is extracted from PubMed database. Each of these entries is hyper-linked to PubMed database by using Pub-
Med identifiers and allowing the retrieval of abstracts.

5. Conclusion

In conclusion, we developed a new platform on predicted as well as validated binding sites of miRNAs on the sequence of all known protein coding and mitochondrial genes of human, mouse and rat. In the future we will incorporate the same information on other species into miRWalk database. Furthermore the web interface will be improved and new modules for additional annotations will be added. Also an online tool for a motif search will be incorporated to query against input sequences of interest.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found, in

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