Post-transcriptional regulation of microRNAs in cancer: From prediction to validation

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Abstract

MicroRNA (miRNA) is a small non-coding RNA with an established function to regulate genes at the post-transcriptional level leading to suppression or degradation of its messenger RNA expression (mRNA). Its dysregulation plays a vital role in a variety of biological and pathological processes including cancer. A lot of algorithms have been established to predict the target sites of miRNA, but experimentally identifying and validating its target region is still lacking. Guidance in experimental procedures is really needed to find genuine miRNA targets. Therefore, in this review, we provide an outline on the workflow in predicting and validating the targeted sites of miRNA using several methods as a guideline for the scientists. The final outcome of this type of experiment is essential to explore the major impact of miRNA-mRNA interaction involved in the biological processes and to assist miRNA-based drug development in the future.

Introduction

MicroRNA (miRNA) is a short single-stranded non-coding RNA, approximately 18 to 25 nucleotides in length. It is generated from a primary miRNA transcript (pri-miRNA) which is transcribed from non-protein-coding genes or from the introns of protein-coding genes in the nucleus. Then, pri-miRNA is cleaved by RNAse III enzyme Drosha to form hairpin precursor miRNA (pre-miRNA) with ~70 nucleotides in length. This is followed by the pre-miRNA is transported into cytoplasm by a Ran-GTP-dependent nuclear export factor, Exportin-5. The pre-miRNA in the cytoplasm is further processed by RNAse III-like nuclease Dicer to generate a mature miRNA strand which is roughly 22 nucleotides in duplex form.1

More than a decade after miRNA was first discovered in 1993, it has started to draw the attention of the science community as it was emerged as an important regulator of gene expression by sharing its extensive sequence complimentarily.2 It was known to control in every aspect in biology such as development, proliferation, differentiation and metabolism. Dysregulation of miRNAs has been reported to lead to many human diseases, including cardiovascular disorders, complex genetic diseases and cancer.3-5

MicroRNA in cancer

It was found that cancer is the most relevant pathology in the world of miRNA-mRNA interactions, where many of the miRNA target sites are clustered in cancer-associated genomic region.6 It can take part either as an oncogene (oncomir), which is upregulated in tumor tissues to promote oncogenesis. On the other hand, it could act as a tumor-suppressor gene (anti-oncomir), which is downregulated in tumor tissues that promotes tumor growth. Dysregulation of these miRNAs could then promote oncogenesis.

MiRNAs have been attested to take part in many types of cancers such as gastric, ovary, lungs, thyroid, colon, prostate and breasts.7-13 Our previous studies on endometrioid endometrial cancer have identified several important transcripts and their related molecular pathways.14,15 The up- or down-regulation of these genes are most likely regulated by miRNAs. The relationship between miRNAs and mRNAs is complex as each miRNA could have multiple target sites on the targeted mRNA and vice versa with each mRNA could be targeted by several miRNAs.16 For example, in breast cancer, miR-29a/-b/-c has been reported to act as an oncogene, but in lung tumor, it turned out to be a tumor-suppressor gene.17,18 While in other studies, loss of miR-23b leads to bladder cancer cell migration and invasion, however in renal cell carcinoma, it reduced invasion by inducing apoptosis.19,20

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miRNA is considered as the master regulator in cellular networks and signaling cascade as it regulates about 60% of the targeted mRNAs.21 To date, about 28,465 miRNAs have been discovered (miRBase Release 21: http://www.mirbase.org, accessed in March 2nd, 2018). To recognize the targeted mRNA, they do not need to completely match as it could identify through a perfect real complement sequence, known as seed region. Usually the seed region located at position 2 to 8 from 5’miRNA, where at this binding site, a stable Watson-Crick base pairs were energetically produced with the targeted mRNAs.21 Then, it could regulate the gene expression either by repressing the mRNA translation or causing mRNA degradation.22,23

The genetic diversity in different types of tumor is associated with different miRNAs in diverse actions and makes the relationship between miRNA and mRNA become more complex. They can participate in various mechanisms and pathways, by targeting key molecules and their networks involved in apoptosis, cell cycle, cell adhesion, cell migration, chromosome stability and DNA repair.24 As the expression of genes regulated by miRNAs is very distinct, thus, it is very essential to validate which target regions of each individual miRNA will give high impact on the gene expression. This is to help to elucidate their biological functions. Table 1 shows several databases available on the reported and experimentally validated miRNA-target interactions at the moment.25-28 However, to study miRNA-target interactions is very challenging as there are many prediction models while only a few have been biologically validated.29 So in this review, we summarize the workflow of molecular network analysis in terms of the prediction methods of miRNAs targets and the experimental approaches that have been described for the identification of their targets. In Figure 1, we proposed a workflow as a guideline in predicting and validating the targeted sites of miRNAs.

**MicroRNA profiling in cancers**

To determine the target region and the biological effects of miRNA, a list of potential miRNAs is required to prepare prior to further characterization.30 Up till today, there are three methods in miRNA profiling which is quantitative polymerase chain reaction (qPCR), hybridisation-based methods such as miRNA profiling using microarray, and high-throughput sequencing such as small RNA sequencing (sRNA-Seq).

Quantitative PCR has been employed to profile miRNAs based on highly reported miRNAs related to cancers. Many companies have developed qPCR-based microarray for the miRNA expression detection. For example, miRStarTM Human Cancer Focus miRNA PCR Array (Arraystar) can test up to 184 miRNAs mostly that are related to cancer to be profiled in parallel. This array uses a panel that includes eight miRNA references to provide a better data quality and quantity, and also to normalize the PCR data. Therefore, qPCR is a powerful tool to analyze the expression level of cancer-related miRNAs quickly and conveniently where it requires less than six hours.31 The obstacle to perform this highly parallel qPCR is the optimal reaction conditions that may be different between miRNAs in primer annealing. Yet, the use of locked nucleic acid (LNAs) into primers is an effective strategy to optimize miRNA primer hybridization conditions. However, this pre-designed miRNA cancer panel could not be used to identify novel miRNAs. This method is not suitable as a discovery tool for those who are looking for a new list of miRNAs related to a cancer but it is suitable to be used as a validation method.

![Figure 1. A proposed workflow as a guideline in predicting and validating the microRNA target site.](image)

**Table 1. Databases of validated miRNA-target interactions.**

| Name          | URL                                                      | Reference |
|---------------|----------------------------------------------------------|-----------|
| MirWalk 2.0   | http://zmf.umm.uni-heidelberg.de/apps/zmf/mirwalk2       | (25)      |
| MirTarBase    | http://mirtarbase.mbc.nctu.edu.tw/php                    | (26)      |
| miRecords     | http://c1.accurascience.com/miRecords                    | (27)      |
| miRTarCLIP    | http://mirtarclip.mbc.nctu.edu.tw                        | (28)      |
MiRNA microarray is defined as a high-throughput technology used primarily in the analysis of gene expression. It was an established method used to analyze large numbers of miRNAs in the past.\textsuperscript{22,23} In this method, 3′ end of miRNAs are tagged with fluorescent-labeled nucleotides using T4 RNA ligase. There is a step whereby the labeled miRNA will hybridize to separately arrayed probes on slides or beads. After removing the non-specific binding, individual miRNAs can be detected. Till today, this method is still being chosen by many scientists due to its low price as compared to other methods, able to produce higher number of parallel measurement, and it could also compare relative abundance of specific miRNAs between two different states.\textsuperscript{27,28} However, the profiling results by this method produced lower specificity compared to small RNA-sequencing.\textsuperscript{34-37}

Small RNA-sequencing (sRNA-seq) is a new technique used to isolate and sequence many types of known and novel small RNA (sRNA) including miRNA. Usually the method starts with sRNA complementary DNA (cDNA) library preparation from the RNA sample of interest. Then, it is followed by parallel sequencing of millions of individual cDNA molecules from cDNA library on a single run. The sequence reads will be analyzed by bioinformatics to identify both known and novel microRNAs in the datasets, and specifies relative quantification by a digital approach. The major advantages of this method are it could produce precise miRNA sequence identification even for novel miRNAs, and also able to identify and distinguish between miRNA variants with high accuracy and sensitivity. Even though it is very helpful, however, it requires expertise in bioinformatics analysis and additional time to complete sequencing-based analysis.\textsuperscript{31}

Without performing miRNA profiling, the list of whole human miRNAs could also be retrieved from miRBase 21 (www.mirbase.org), as described previously.\textsuperscript{39} The miRNA expression profiling dataset can also be downloaded from Gene Expression Omnibus (GEO) repository (www.ncbi.nlm.nih.gov/geo) for focused miRNA selection, as all the data derived from experimental study performed on microarray, qPCR, and high-throughput sequencing.\textsuperscript{39}

### Validation of miRNA profiling and small RNA sequencing

Quantitative PCR is considered as a gold standard and established method in gene expression detection and quantitation to validate miRNA expression profiling and small RNA sequencing. There are two types of qPCR which is TaqMan qPCR and SYBR-green-based qPCR. In TaqMan qPCR, a stem-loop primer is used in the reverse transcription reactions that are specific to the 3′ end of the miRNA. The forward and reverse primers will subsequently be used for amplification. While the DNA polymerase working on the template, the TaqMan probe will be hydrolyzed by Taq polymerase and freed the fluorescent dye from the quencher, resulting in light emission. In SYBR-green-based qPCR, to generate miRNA cDNA, enzymatic addition of a poly(A) tail is used. For PCR amplification, both specific forward and reverse primers are used to anneal to the 3′ miRNA. Then, SYBR green dye will bind to each double-stranded DNA product to give fluorescent intensity.

### Prediction of miRNA target region

According to Bartel,\textsuperscript{21} computational approaches can accurately predict miRNAs with their targeted miRNAs based on the thermodynamic rule and the evolution of miRNA recognition element (MRE) sequence \textit{in silico}. Besides 3′UTR, target sites of miRNA can also happen in 5′UTR and open reading frame (ORF).\textsuperscript{40-42} However, in this review, we are focusing more on 3′UTR since ORF and 5′UTR appear to be less effective and less frequent, and most target mRNAs studied to date are regulated through 3′UTR interaction.\textsuperscript{21}

There are several computational algorithms that predict target MRE sequences with different measures that include base-matched, target availability and evolutionary conservation of targeted sites. Basic characteristics of prediction methods as seen in Table 2 where it can compute all the predicted miRNA targeted sites of the gene by simply enter the Gene Symbol.\textsuperscript{29,43-46} All algorithms used can predict up to hundred miRNA’s targeted sites and the predicted binding sites usually distinct from one to another. Hence, the best possible way recommended in determining predicted miRNA binding site is by combining at least two or three algorithms that predict the same miRNA binding sites and identify the overlapping results among them.\textsuperscript{47}

### Validation of the target regions

According to Martinez-Sanchez & Murphy (2013), once the predicted binding site of miRNA is determined, its functionality in biological model can be validated.\textsuperscript{48} As stated before, the computational algorithm could predict a high number of miRNA binding sites. To identify and verify these target interactions between miRNA and mRNA, luciferase reporter assay has become a favourable method among researchers. It is a recommended assay as it could produce rapid and reproducible results to determine the functional and non-functional interaction sites.

Generally, before performing this assay, 3′UTR of the transcript of interest is cloned downstream of the luciferase ORF contained in the reporter plasmid. Then, the recombinant plasmid

### Table 2. Computational methods for miRNA target prediction. 3′UTR, 3′untranslated region.

| Name         | Targets on mRNA | Species                     | URL                                                                 | Reference |
|--------------|-----------------|------------------------------|----------------------------------------------------------------------|-----------|
| mIRTar       | Full length     | Human, mouse, rat and chicken | http://mirtar.mbc.nctu.edu.tw/human/                                 | (28)      |
| TargetScan   | 3′UTR           | Human, mouse, rat, dog and chicken | http://www.targetscan.org/vert_71                                    | (44)      |
| microrna.org | 3′UTR           | Human, mouse, rat, roundworm and fruitfly | http://34.236.212.39/microrna/home.do                             | (45)      |
| miRDB        | 3′UTR           | Human, mouse, rat, dog and chicken | http://www.mirdb.org                                                   | (46)      |

(UTR, 3′untranslated region)
Validation in biological function

Once the target site of a given miRNA has been validated experimentally, it can be revealed that there is biologically gene regulation. Theoretically, when the expression of miRNA is downregulated, the mRNA is predicted to be upregulated, and vice versa. Hence, the functionally inverse relationship between miRNAs and mRNAs target site can be validated by several method.

To look for the effects of validated targets towards miRNA and mRNA co-expression, it can be simply demonstrated by Northern blot analysis or qPCR by using total RNA from specific cell type, together with their specific primers of miRNA or targeted mRNA. Since the interaction of miRNA-mRNA is negatively regulated, the overexpression of miRNA would reduce the expression of the targeted mRNA, and vice versa. Another experimental technique to demonstrate miRNA and mRNA co-expression, is in situ hybridization whereby it can be utilized in tissue or cell of relevant samples.

If the miRNA given was the real target of specific miRNA, the amount of protein expression could also change. Usually loss-of-function or gain-of-function (over-expression) experiments are the typical approach by transiently transfect miRNA mimics or inhibitors into a cell type which expressing the putative target protein. Subsequently, western blot analysis using a specific antibody against protein can be utilized to look for the protein expression. Another alternative experiment can be used to quantify difference in protein expression is ELISA or immunocytochemistry. So, with the presence of miRNA mimics or inhibitors, the target protein levels or activities should be easily detected. With the presence of miRNA mimics in cells, the endogenous expression of targeted mRNA would be reduced. However, with the presence of miRNA inhibitors, the endogenous expression of targeted miRNA would be reduced, hence, increased the expression of mRNA.

Once regulation of a target gene by a given miRNA has been confirmed experimentally, the regulation in biological function becomes inevitably to be demonstrated. Usually the assays includes signalling pathway, cell proliferation, cell differentiation, cell death, cell migration and invasion, metastasis, and others. For example, a study showed that the expression of miR-130a were up regulated in high grade serous ovarian cancer (HGSOC) tissues by using microarray and qPCR. However, with the presence of anti-miR-130a, it could reverse cell proliferation, migration and invasion.

Conclusions

The interactions of miRNA-mRNA contribute to the complexity and constitute the biological network of functionally-associated molecules in pathological events involved in cancer. A computational prediction algorithm has been developed recently. However only a small proportion of the interaction have been functionally validated. Even though there are limitations in the method discussed, following the workflow may lead to relevant target identification biologically, where understanding the functions may assist undoubted therapeutic potentials.

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