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Gene module based regulator inference identifying miR-139 as a tumor suppressor in colorectal cancer†

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Colorectal cancer is one of the most commonly diagnosed cancer types worldwide. Identification of the key regulators of the altered biological networks is crucial for understanding the complex molecular mechanisms of colorectal cancer. We proposed a gene module based approach to infer key miRNAs regulating the major gene network alterations in cancer tissues. By integrating gene differential expression and co-expression information with a protein–protein interaction network, the differential gene expression modules, which captured the major gene network changes, were identified for colorectal cancer. Then, several key miRNAs, which extensively regulate the gene modules, were inferred by analyzing their target gene enrichment in the modules. Among the inferred candidates, three miRNAs, miR-101, miR-124 and miR-139, are frequently down-regulated in colorectal cancers. The following computational and experimental analyses demonstrate that miR-139 can inhibit cell proliferation and cell cycle G1/S transition. A known oncogene ETS1, a key transcription factor in the gene module, was experimentally verified as a novel target of miR-139. miR-139 was found to be significantly down-regulated in early pathological cancer stages and its expression remained at very low levels in advanced stages. These results indicate that miR-139, inferred by the gene module based approach, should be a key tumor suppressor in early cancer development.

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Introduction

Colorectal cancer is a kind of severe bowel disease and it is the third most commonly diagnosed cancer around the world. The molecular networks are usually significantly altered in cancer cells. To understand the complex regulatory mechanism of colorectal cancer initiation and progression, it is crucial to identify the key regulators of the altered molecular networks. MicroRNAs (miRNAs), a class of ~22 nt endogenous small regulatory RNAs, are significantly differentially expressed between colorectal cancerous and adjacent normal tissues.¹ But only a few of them are linked with the major alterations of the molecular networks in colorectal cancer. To infer the key miRNAs regulating the altered molecular networks in cancer,

one simple way is to analyze the enrichment of their targets in the differentially expressed genes between cancerous and normal tissues.² But due to the complexity of the cancer transcriptome, this kind of method frequently failed to find candidates due to low statistical significances. Multiple genes usually work cooperatively as functional gene modules.³ Differential network analysis can better identify the altered molecular networks by integrating gene expression data with network data (such as a protein–protein interaction network and a literature co-citation network).^{4,5}

The differential gene expression modules (DGEMs, or called active gene sub-networks), a class of “differential networks” consisting of a set of densely connected differentially expressed genes and their neighbors, can be treated as the major network alterations in cancer.^{4,6} Then, the key miRNAs can be inferred by analyzing their target enrichment in the DGEMs. Based on this hypothesis, we developed a gene module based master regulator inference (ModMRI) approach to identify the key miRNAs in colorectal cancer: firstly, the DGEMs were identified by the integrative analysis of multiple gene expression datasets and protein–protein interaction networks using ClustEx package.⁷ Then, a network-based permutation test was used to infer the key miRNAs whose target genes are significantly overlapped with the DGEMs. Among the inferred candidates, miR-101,

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miR-124 and miR-139 are frequently down-regulated in colorectal cancer.^{1,8} The following computational and experimental analyses suggest that miR-139 should act as an important tumor suppressor in early cancer pathologic stages.

Methods

Colorectal cancer gene expression data and gene network data

A combined gene expression dataset of colorectal cancer was constructed based on four microarray data series from the NCBI GEO database with at least 15 clinical cancer/normal samples and using Affymetrix Human Genome U133 Plus 2.0 Array (GSE20916, GSE21510, GSE22598, GSE23878; 90 normal and 107 cancer samples in total). The .CEL files of the four data series were processed and normalized using RMA package independently.⁹ Probe signals were mapped to gene expression values according to the latest probe annotations. Then, the raw gene expression values were transformed to ranks. To avoid the noises of the lowly expressed genes, only the genes ranked as top 10 000 (~50% of the genes on the microarray) in at least 30% samples were left for the following analysis.

We extracted all binary interactions from HPRD (release 9)¹⁰ and BioGRID (version 3.1.86).¹¹ To reduce the high false positives in PPI data, we only used the interactions annotated in both databases. By filtering the above “expressed” genes in microarray datasets, we got the background gene network with 4181 genes and 10 261 edges. The edges were weighted by Spearman’s co-expression levels according to the combined gene expression dataset.

A gene was regarded as a target of a miRNA family, if the gene has at least one conserved target site predicted by TargetScan (v6.0).¹² The 3′-UTR lengths of genes were calculated from the 3′-UTR multiple alignment file from TargetScan resources. If a gene has multiple transcripts, the length of the longest 3′-UTR was used.

Gene module based key miRNA inference

The miRNAs, whose target genes were significantly enriched in the differential gene expression modules (DGEMs), were identified as key regulators. Firstly, differentially expressed genes were identified from the combined gene expression dataset. Secondly, the DGEMs were identified by finding the sub-networks consisting of a set of closely connected differentially expressed genes and their neighbors. Then, the key miRNAs were inferred by analyzing their target gene enrichments in the DGEMs. Finally, bootstrapping was implemented to get robust inferences. The flowchart of this gene module based master regulator inference (ModMRI) method can be found in Fig. 1.

Identification of the differentially expressed genes. The differentially expressed genes were identified by combining the top 600 genes ranked by the *t*-test between cancerous and adjacent normal samples in at least one of the four microarray datasets. 1330 non-redundant differentially expressed genes were identified by this criterion.

Identification of the differential gene expression module. ClustEx⁷ was used to identify the DGEMs of colorectal cancer by clustering and extending the differentially expressed genes in the network.

miRNA target gene enrichment analysis. For each miRNA, we counted the number of its target genes in DGEMs. Then, a degree-preserving permutation was implemented to generate 10 000 randomized DGEMs. The numbers of the miRNA target genes in the randomized DGEMs were counted to estimate the background distribution of the number of the overlapped target genes. The *p*-value was calculated by comparing the original number with the corresponding background distribution. The *p*-values were multiple-test adjusted as *q*-values using fdrtool.¹³

Resampling experiments for robust inferences. Cancer gene expressions data are usually noisy. To reduce the unstable inferences due to the gene expression variances, bootstrapping was implemented: in each run, 80% out of all samples are randomly selected to re-do the whole analysis. Only the miRNAs,

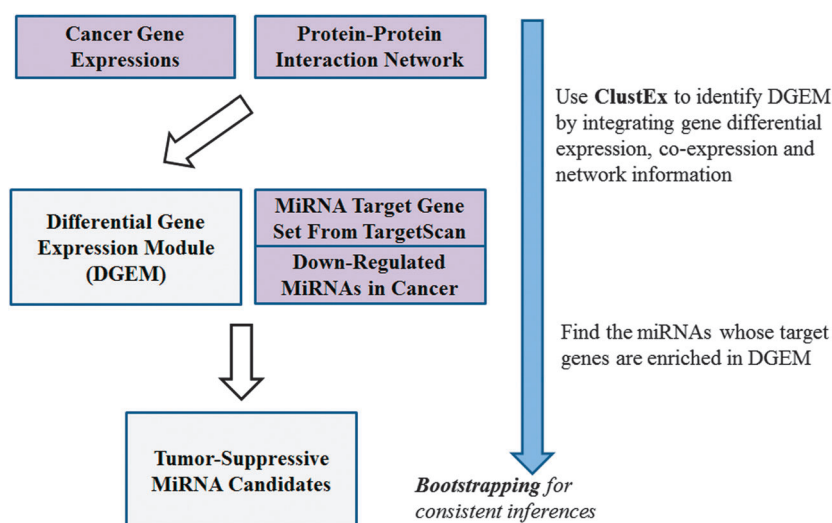


Fig. 1 The flowchart of the module-based master regulator inference (ModMRI).



which were repeatedly inferred in at least 50% re-sampling runs, were reported as the final results. To further reduce the false positives due to miRNA target predictions, we randomly added or removed 10% miRNA target genes to calculate the means and the standard variations of the empirical *p*-values.

Experimental validation of miR-139 cellular functions and target genes

A series of experiments were conducted to validate the cellular functions and target genes of miR-139 in colorectal cancer cells.

Cell culture. Human colorectal cancer cell lines, HCT-116 and SW480, were obtained from the Cell center of Peking Union Medical College (Beijing, China) and HEK293 was obtained from American Type Culture Collection (Manassas, VA). The cells were maintained at 37 °C in a humidified atmosphere of 5% CO₂ in air. Both cells were maintained in Dulbecco's Modified Eagle Medium. The media were supplemented with 10% fetal bovine serum (FBS), 5 mM L-glutamine, 100 U ml⁻¹ penicillin and 100 mg ml⁻¹ streptomycin.

Cell proliferation assay. Cells were plated onto 96-well plates and incubated overnight before the transfection. After transfection with miRNA mimics or a negative control, both 50 nM, for 48 hours, the cells were used for cell viability evaluation using a CCK8 assay kit (Biyutian, China) according to the protocol.

Cell cycle assay. Cells were harvested and fixed in 70% ethanol and stored at -20 °C overnight. Cells were washed twice with ice-cold phosphate buffer saline (PBS) and incubated with RNase and propidium iodide for 30 min and then cell cycle analysis was performed using a flow cytometer.

Luciferase reporter assay. HCT-116 or HEK293 cells were plated onto 24-well plates, 1×10^5 in each well. After 24 hours, cells were co-transfected with 1 µg of the psiCheck2 luciferase reporter vector containing the conserved binding sites in 3'-UTRs of the candidate target genes and the 50 nmol miRNA mimic. Luciferase assays were performed using the Dual Luciferase Reporter Assay System (Promega) 48 hours after transfection. The renilla luciferase activity was normalized to the firefly luciferase activity as an internal transfection control. Then, the luciferase values were normalized to the average values for the corresponding vehicle control transfections. Values represent mean + SD of at least three experimental repeats.

Western blot. Proteins from cells and tissue samples were extracted with RIPA lysis buffer (150 mM NaCl, 10 mM Tris, pH 7.5, 1% NP40, 1% deoxycholate, 0.1% SDS, protease inhibitor cocktail (Roche)). Proteins from total cell lysates were resolved by 10% SDS-PAGE gel, transferred to the nitrocellulose membrane, blocked in 5% non-fat milk in PBS/Tween-20, and blotted with the antibody against ETS1 (1:500, Abcam), and blotted with Goat anti Rabbit IgG (1:3000, Santa Cruz). Gene PCNA was used as loading control.

Results

The differential gene expression module in colorectal cancer

A large number of genes were significantly differentially expressed between colorectal cancer and noncancerous tissues. The gene

module based master regulator inference (ModMRI) is an integrative gene network analysis for identifying the key regulators mediating these significant alterations of molecular networks. At first, ClustEx program was used to identify the differential gene expression modules (DGEMs) associated with colorectal cancer. The identified biggest or the principal DGEM has 822 genes including 305 differentially expressed seed genes. The module genes are significantly enriched in KEGG annotated "pathways in cancer" (*q*-value 1.4×10^{-17}), "cell cycle" (1.4×10^{-14}), "MAKP signaling pathway" (6.7×10^{-14}) and "colorectal cancer" (2.8×10^{-10}), which suggests that the identified DGEM is highly associated with colorectal cancer. The details of the module and the enriched cellular processes can be found in Supplementary File 1 (ESI[†]).

The key miRNAs extensively regulating the differential gene expression modules

Then, several key miRNAs, including several known oncogenic and tumor-suppressive miRNA families in colorectal cancer, miR-17,¹⁴ miR-93,¹⁵ miR-101¹⁶ and miR-135,¹⁷ were identified by analyzing their target gene enrichments in the biggest DGEM (Table 1). Three of them, miR-101, miR-124 and miR-139, were found frequently down-regulated in colorectal cancer,^{1,8} which suggest that these miRNAs may act as tumor suppressors by extensively regulating the biggest DGEM. We also analyzed the miRNA target enrichments in the second biggest DGEM (609 genes), but no significant candidate was found.

miR-139 inhibited cancer cell proliferation and cell cycle progression

Among the three inferred tumor suppressors, miR-139 has the minimal *p*-value and highest bootstrapping re-sampling stability (Table 1). We chose to further study the cellular functions and target genes of miR-139 by a series of computational and experimental analyses. The 35 miR-139 target genes in the DGEM are significantly enriched in "regulation of cell proliferation" (11 out of the 35 target genes are annotated with the GO term, *q*-value < 0.05) (Supplementary File 2, ESI[†]). CCK-8 cell proliferation assays show that miR-139 significantly inhibited cancer cell proliferation (*q*-value < 0.01). The inhibition rates are comparable to the well-studied tumor suppressor miR-101

Table 1 The key miRNAs inferred by ModMRI in colorectal cancer

miRFam	<i>p</i> -value	#Target	BS (%)	Expression
miR-135	0.0005 (0.049)	57/198	50	Up
miR-874	0.0021 (0.104)	29/77	58	—
miR-139	0.0042 (0.131)	35/108	84	Down
miR-17	0.0142 (0.200)	91/358	50	Up
miR-93	0.0160 (0.206)	64/256	86	Up
miR-124	0.0175 (0.211)	116/500	52	Down
miR-101	0.0365 (0.241)	65/262	50	Down

"*p*-value": the *p*-value and the corresponding *q*-value. "#Target": the number of targets overlapped with the DGEM and the total number of predicted targets; "BS": the percentage of significant inferences in bootstrapping experiments; "Expression": the miRNA expression patterns in ref. 1 and PhenomiR database.



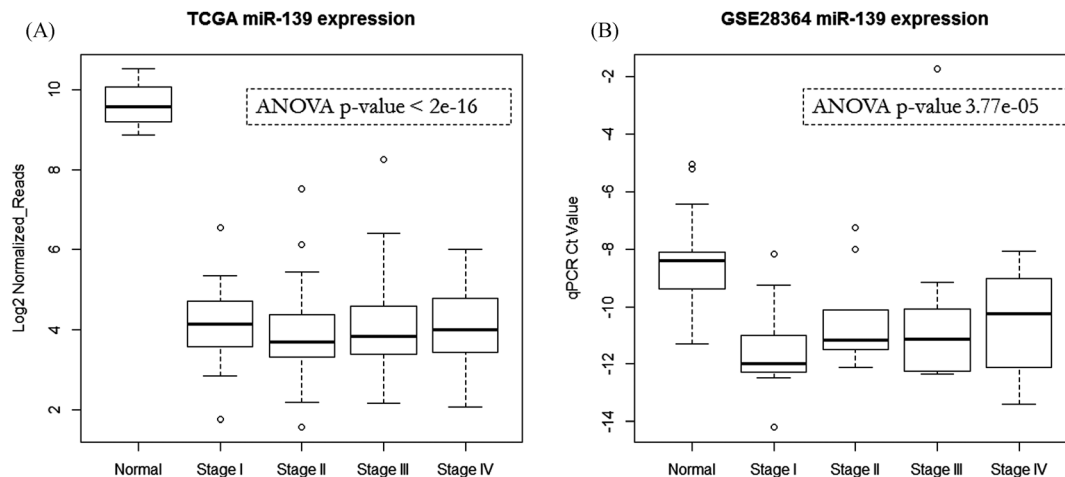


Fig. 4 miR-139 expressions in colorectal cancer tissues in different pathologic stages and adjacent normal tissues. (A) miR-139 expressions in the TCGA COAD dataset (HiSeq sequencing data). (B) miR-139 expressions in GSE28364 (qPCR data).

miR-139 expression was down-regulated in cancer tissues

We examined miR-139 expressions in two clinical studies (TCGA COAD dataset with 222 cancer and 8 non-cancerous samples;²⁰ GSE28364 with 40 cancer and 40 non-cancerous samples²¹). The data show that miR-139 is significantly down-regulated in early cancer pathologic stages compared with adjacent non-cancerous tissues, and it remains at a very low expression level in advanced pathologic stages (Fig. 4). Reid *et al.* also reported that the miR-139 region has genomic loss in cancer samples.²¹ These results suggest that the loss of miR-139 is an important indicator of colorectal cancer. Similar expression patterns can be observed in several other solid tumors (Fig. S2, ESI†).

Discussion

miR-139 is a broadly conserved miRNA in vertebrates. The module-based master regulator inference (ModMRI) predicted that miR-139 is a key regulator by targeting tens of genes in the DGEM of colorectal cancer. The following expression and functional analyses further suggested that miR-139 may act as a tumor suppressor by regulating cancer cell proliferation. Cellular functional assays validated that miR-139 can inhibit cancer cell proliferation and cell cycle G1/S transition. And ETS1, an oncogenic transcription factor promoting cell cycle progression, was verified as a direct target of miR-139. ETS1 is not differentially expressed in any of the four colorectal gene expression datasets, but it can be found by module-based analysis after incorporating the gene network information. Two large-scale expression datasets from colorectal cancer patients show that miR-139 was significantly down-regulated in early pathologic stages of colorectal cancer and remained at a very low expression level in advanced stages. A few other studies also reported that miR-139 can inhibit colorectal cancer cell proliferation.^{22,23} These results indicate that miR-139 should be a key tumor suppressor in early cancer development.

Modularity is an important property of biological networks.^{4,24} Tens of genes work cooperatively as a functional module for

different cellular processes. Inference of any single gene frequently fails to disturb the module activity due to complex network compensations and feedbacks. This study indicates that the module-based inference using miRNAs may be an alternative approach to disturb cancer cellular states.

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