Functional screening using a microRNA virus library and microarrays: a new high-throughput assay to identify tumor-suppressive microRNAs

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MicroRNA (miRNA) is a class of non-coding RNAs that represses expression of target messenger RNAs posttranscriptionally. A growing body of evidence supports their roles in various normal cellular processes, as well as in pathological conditions, such as cancer. We established a functional screening assay that enables high-throughput identification of miRNAs that have a role in cancer phenotypes of interest, via the combination of pooled lentivirus vectors expressing several hundred miRNA precursors and a custom-made microarray. Self versus self-hybridization analysis using pooled polymerase chain reaction products generated highly linear and reproducible results. To test the feasibility of the assay, we focused on miRNAs that control proliferation of pancreatic cancer cells and successfully identified five miRNAs that negatively control cell proliferation, including miRNA-34a that was previously identified as a representative tumor-suppressive miRNA. The results were further validated using lentivirus vectors expressing each of the five miRNAs or synthetic miRNAs. The function-based nature of the assay enabled identification of miRNAs that were strongly linked to cell proliferation, but the relative ease and flexibility of the assay allow for future studies of cancer stem cells, metastasis and other cancer phenotypes of interest.

Introduction

MicroRNA (miRNA) is a class of evolutionarily conserved non-coding RNAs of ~19 to 22 nucleotides that modulate expression of their target genes posttranscriptionally together with the RNA-induced silencing complex. Since the first identification of lin-4 in Caenorhabditis elegans, the number of registered miRNAs is expanding and 940 miRNA genes (1100 mature miRNAs) are annotated in humans (miRBase release 15, http://www.mirbase.org/) and more than a thousand miRNA genes are estimated to be located in the human genome (1). Also, contrary to the ‘evolutionarily conserved’ definition of miRNAs, species–specific miRNAs have been implicated and cloned, suggesting their critical role in the integrity of higher organisms (2). A growing body of evidence suggests that miRNAs have pivotal roles in normal cellular processes (differentiation, proliferation and cell death) and stress response, as well as implicating their involvement in cancer and other pathological conditions (3,4). Indeed, expression profiling studies of miRNAs have shown that miRNAs are aberrantly expressed in a variety of cancers (5). Some miRNAs are consistently upregulated or downregulated in cancers, suggesting their possible tumor-promotive or tumor-suppressive features. Moreover, miRNAs were generally downregulated in tumors compared with normal tissues, implicating a multitude of tumor-suppressive miRNAs that have not been fully recognized (5). In light of these facts, the development of functional assays for miRNAs appears to be warranted in order to better elucidate the mechanisms underlying characteristic features of cancer. Here, we have established a high-throughput functional screening assay in which hundreds of miRNAs are characterized after expression of corresponding miRNA precursors via lentivirus vectors. To test the feasibility of the assay, we screened for miRNAs that suppress proliferation of the pancreatic cancer cell line MIA PaCa-2. Five miRNAs were identified, including microRNA-34a (miR-34a) that was previously reported as one of the p53-responsive miRNAs, with strong tumor-suppressive activity in various cancers (6). Proliferation-suppressive effects of the five miRNAs identified by the new functional screening assay were individually validated by the infection of lentivirus vectors expressing each miRNA precursor. Cooperative data were obtained following transfection of synthetic miRNAs to MIA PaCa-2 cells; flow cytometry revealed that cell cycle arrest was, at least in part, an underlying mechanism for the observed phenotypic effects.

Materials and methods

Cell culture

MIA PaCa-2 cells and 293T cells (ATCC, Manassas, VA) were cultured in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum and 50 U/ml penicillin and 50 μg/ml streptomycin (Invitrogen, Carlsbad, CA). The cells were routinely incubated at 37°C in a humidified atmosphere with 5% CO2.

Custom-made oligonucleotide microarray

Sequences of miRNA precursors were obtained from Ensembl Genome Browser (Release 55, http://www.ensembl.org) and the UCSC genome browser (hg19, http://genome.ucsc.edu). Oligonucleotide (60mer) probes were designed for 445 miRNA precursors in a Lenti-miR miRNA precursor clone collection (System Biosciences, Mountain View, CA) using eArray software (Agilent Technologies, Santa Clara, CA). All probe sequences were BLAST searched against miRNA precursor sequences using BlastStation2 software (TM Software, Arcadia, CA). The specificity of the probes with the corresponding miRNA precursors was manually checked and probes with poor specificity were replaced with redesigned probes. The custom-made oligonuclotide microarray was designed so that each microarray contains 32 replicates (16 sense and 16 antisense) of specific probes for 445 miRNA precursors on an 8 × 15 k format (Agilent Technologies). Detailed information of the custom-made microarray we designed is available on request.

Infection of lentivirus library into cells and subsequent passages

MIA PaCa-2 cells were seeded at 4 × 105 cells in a six-well dish 1 day prior to viral transduction. A Lenti-miR Virus Library (System Biosciences) and polybrene (hexadimethrine bromide; Sigma–Aldrich, St Louis, MO) at a final concentration of 5 μg/ml were added to the culture medium. Two parallel infections of the Lenti-miR Virus Library (pooled virus library) were performed with a multiplicity of infection of ~3. Cells were incubated at 37°C in a humidified atmosphere for 24 h, after which medium containing the virus library was replaced with Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum and antibiotics. Half of the infected cells were transferred to a 10 cm dish and DNA was extracted from the remaining cells (P0). Afterward, cells were cultured in a 10 cm dish and passaged in the proportion of 1:8 when they reached 80–90% confluency. The cells not used for the passage were used for the extraction of DNA. Throughout the screening process, cells were monitored for green fluorescent protein (GFP)-like protein from a copedope (copGFP) by fluorescence microscopy (IX71 Inverted Microscope; Olympus, Tokyo, Japan).

Functional screening assay using the custom-made microarray

Genomic DNA (gDNA) was extracted with a DNeasy Blood and Tissue Kit (Qiagen, Germantown, MD) according to the manufacturer’s instructions. miRNA precursors were recovered from gDNA by polymerase chain
reaction (PCR) with specific primers against lentivirus vectors (forward primer: 5'-GCGTGAGGACCGCATCCAAGCTG-3'; reverse primer: 5'-GATGTGGCTCTGCGCCACTGAC-3'). The PCR amplicon is a composite of 445 miRNA precursors; each precursor is composed of the stem–loop sequence (defined in the miRBase, ~100 bp) and the flanking genomic regions (a 5 bp upstream and downstream) of the miRNA gene, making the total size of the amplicon ~500 to 700 bp (supplementary Figure S1 is available at Carcinogenesis Online). Fifty microliter of PCR reaction contained the following final concentrations: 200 ng of gDNA template, 400 µmol/l each of deoxynucleoside triphosphates, 0.3 µM of each PCR primer and 1 U of KOD FX DNA polymerase (Toyobo, Osaka, Japan). PCR was performed with the following program on a Veriti thermal cycler (Applied Biosystems, Foster City, CA): 94°C for 3 min, 25 cycles of 94°C for 35 s, 65°C for 35 s, 72°C for 1 min and a final step of 72°C for 7 min. Four independent PCR reactions using the same gDNA template were pooled and purified with the Rapid PCR Purification System (Marligen Biosciences, Rockville, MD). Hundred nanogram of purified PCR product were labeled with either Cy3- or Cy5-deoxycytidine triphosphate using a Genomic DNA Enzymatic Labeling Kit (Agilent Technologies) and purified with a Microcon YM-30 Centrifugal Filter Unit (Millipore, Billerica, MA). A pair of Cy3- and Cy5-labeled (Agilent Technologies) and purified with a Microcon YM-30 Centrifugal Filter Unit (Millipore, Billerica, MA). A pair of Cy3- and Cy5-labeled DNA was combined and hybridized to the custom-made microarray at 65°C and 20 r.p.m. for 24 h in a hybridization oven (Agilent Technologies). Washing of the microarray and data analysis were performed according to the CGH protocol version 5.0 (Agilent Technologies). The log ratio of each miRNA precursor was calculated by averaging the ratio of replicate probes for each miRNA precursor excluding the highest and the lowest values.

Production of pseudovirus particles

Cells (5 x 10⁶ 293T) (ATCC) were seeded in a 10 cm dish 1 day prior to transfection and cotransfected with packaging plasmids (pLOCK1-GAG, pPACKH1-REV and pSVS-G) and a vector plasmid containing each miRNA precursor/copGFP (System Biosciences) using Lipofectamine2000 (Invitrogen) according to the manufacturer’s instructions. Cells were cultured at 32°C for 48 h in a humidified atmosphere with 5% CO₂, after which culture medium containing pseudovirus particles was collected and stored at −80°C before infection to MIA PaCa-2 cells.

Cell proliferation assay

For the validation using lentivirus vector expressing miRNA precursors, cells were infected with pseudovirus particles using 5 µg/ml of polybrene for 24 h. Cells (0.5 x 10⁵) were subcultured into a 48-well plate after infection, and 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide assay was performed using a CellTiter 96 AQueous One Solution Cell Proliferation Assay (Promega, Madison, WI) and a 2030 Arvo X 3 multilabel plate reader (PerkinElmer, Waltham, MA) at 3 and 5 days after subculturing. Cells infected with pseudovirus that only express copGFP, but no miRNA precursor (lenti-mir-control), were used as a control of the assay. Expression of functional miRNA in cells transduced with each miRNA clone was measured by quantitative reverse transcription–polymerase chain reaction prior to 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide assay (supplementary method and supplementary Table S2 are available at Carcinogenesis Online). For the validation using synthetic mature miRNA, 1 x 10⁵ MIA PaCa-2 cells were subcultured in a 24-well plate and transfected with HiPerfect Transfection Reagent (Qiagen) and 10 nmol/l of Pre-miR miRNA Precursor Molecules (Ambion, Austin, TX). Cells transfected with Pre-miR miRNA Precursor Molecules—Negative Control (Ambion) were used as a control of the assay. Cells were counted every day after transfection up to 5 days using a Countess Automated Cell Counter (Invitrogen) according to the manufacturer’s instruction.

Flow cytometric analysis of cell cycle

2 x 10⁵ MIA PaCa-2 cells were plated per 10 cm dish 1 day prior to transfection of synthetic mature miRNAs and incubated at 37°C in a humidified atmosphere with 5% of CO₂ until transfection. Synthetic miR-29b, -34a, -222, -224, -532 and negative control miRNA (Pre-miR, Ambion) were transfected with HiPerfect Transfection Reagent (Qiagen) at a final concentration of 10 nmol/l. Cells were trypsinized and collected at 48 and 72 h after transfection, then fixed with 90% methanol and finally suspended in phosphate-buffered saline (−) with 50 µg/ml of propidium iodide (Sigma–Aldrich), 50 µg/ml of RNase A (Nippon Gene, Tokyo, Japan) and 0.1% fetal bovine serum (Invitrogen). Ten thousand stained cells were analyzed with a FACSCalibur flow cytometer and CellQuest pro software (BD Biosciences, San Jose, CA) according to the manufacturer’s instructions. The proportions of cells in sub-G₁, G₁/G₂, S and G₂/M phase were analyzed by quantifying the ratio of fluorescence according to the ratio of change in the copy number of each miRNA library. miRNAs that generate phenotypes of interest are identified according to the ratio of change in the copy number of each miRNA precursor that is detected by the ratio of differently labeled fluorescent protein cloned from a copepod, which enables fluorescent microscopic monitoring of cells infected with the vectors (Figure 1B) (7). Although a pooled library is more convenient than individually assessing phenotypic changes of cells infected with each miRNA and is thus suitable for a functional screening assay, it is virtually impossible to screen miRNAs that negatively regulate. miRNAs decrease throughout the screening process. We employed a microarray-based approach to the functional screening assay that enables quantification of changes in the copy number of each miRNA precursor to identify miRNAs that negatively as well as positively regulate phenotypes of interest. Briefly, cells infected with a pooled lentivirus vector library undergo phenotypic screening, during which miRNAs that positively or negatively regulate phenotypes of interest are enriched or excluded compared with the original library, miRNAs that generate phenotypes of interest are identified according to the ratio of change in the copy number of each miRNA precursor that is detected by the ratio of differently labeled fluorescent intensity.

In order to assess the linearity of PCR amplification of mixed miRNAs precursors, reproducibility of PCR reactions from two independent experiments was examined (Figure 2A and B). Eight independent PCR amplifications were performed, and two pooled

Statistical analysis

Pearson product-moment correlation coefficient and standard deviation were calculated using Excel software (Microsoft, Redmond, WA).

Results

Development of a miRNA functional screening assay

To establish a functional screening assay of miRNAs, we employed a pooled lentivirus vector library expressing 445 human miRNA precursors and a custom-made oligonucleotide microarray for the detection of these precursors (Figure 1A). Each lentivirus vector expresses human miRNA precursors together with copGFP, a green fluorescent protein cloned from a copepod, which enables fluorescent microscopic monitoring of cells infected with the vectors (Figure 1B) (7). Although a pooled library is more convenient than individually assessing phenotypic changes of cells infected with each miRNA and is thus suitable for a functional screening assay, it is virtually impossible to screen miRNAs that negatively regulate phenotypes of interest as the number of cells infected with such miRNAs decrease throughout the screening process. We employed a microarray-based approach to the functional screening assay that enables quantification of changes in the copy number of each miRNA precursor to identify miRNAs that negatively as well as positively regulate phenotypes of interest. Briefly, cells infected with a pooled lentivirus vector library undergo phenotypic screening, during which miRNAs that positively or negatively regulate phenotypes of interest are enriched or excluded compared with the original library, miRNAs that generate phenotypes of interest are identified according to the ratio of change in the copy number of each miRNA precursor that is detected by the ratio of differently labeled fluorescent intensity.

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PCR products, each comprising four PCR reactions (pool I and pool II), were labeled with Cy3- and Cy5-deoxycytidine triphosphate, respectively, and hybridized to the custom-made miRNA oligonucleotide microarray (self versus self-hybridization). We compared the correlation coefficient of pool I and pool II under several PCR conditions, together with the optimized annealing temperature and the number of amplification cycles. A high correlation coefficient for the assay was obtained \((R = 0.944, \text{Figure 2B})\). The correlation coefficient was comparable between PCR products purified by gel electrophoresis or silica membrane (data not shown).

**Identification of miRNAs that repress proliferation of pancreatic cancer cells**

To test the feasibility of the functional screening assay, we focused on the identification of miRNAs that negatively regulate proliferation of human pancreatic cancer cells. MIA PaCa-2 cells were infected with a pooled lentivirus library of miRNA-expressing vectors and passaged several times, during which the expression of miRNA precursors in a majority of cells was confirmed via microscopic inspection of the expression of copGFP (Figure 1B). gDNA was extracted from cells not used for the passage and integrated miRNA precursors were amplified by PCR. Subsequently, amplified miRNA precursors were labeled with Cy3- or Cy5-deoxycytidine triphosphate and hybridized to custom-made microarrays. Any change in the relative proportion of cells expressing each miRNA precursor were measured by comparing labeled miRNAs from cells passaged nine times (P9) and cells immediately after infection (P0) (Figure 2C; supplementary Table S1 is available at Carcinogenesis Online). We focused on five miRNAs, namely \(\text{miR-29b, -34a, -222, -224 and -532}\), which consistently showed a remarkably low log ratio in two independent screening assays (Table I) and individually validated the proliferation-suppressive phenotypes either using lentivirus vector expressing miRNA precursors, which are components of the pooled virus library used in the functional screening or synthetic mature miRNAs. MIA PaCa-2 cells infected with lentivirus vectors expressing each positive miRNA significantly suppressed cell proliferations (Figure 3A). Moreover, transfection of synthetic \(\text{miR-29b, -34a, -222, -224 and -532}\) also suppressed proliferation of MIA PaCa-2 cells in comparison with cells transfected with negative control miRNAs. (Figure 3B and C). The fold changes in the expression of the five proliferation-suppressive miRNAs showed variations according to the endogenous expression level of these miRNAs and the titers of the lentivirus vectors transduced (supplementary Table S2 is available at Carcinogenesis Online).

**Induction of cell cycle arrest after transfection of proliferation-suppressive miRNAs**

To gain further insight into the proliferation-suppressive effects of these five miRNAs, flow cytometric analysis was performed using MIA PaCa-2 cells 48 and 72 h after transfection with synthetic miRNAs (Figure 4; supplementary Figure S2 is available at Carcinogenesis Online). MIA PaCa-2 cells transfected with \(\text{miR-34a, -224 and -532}\) showed an increased proportion of G0/G1 phase in comparison with cells transfected with negative control miRNA, whereas the proportion of G2/M phase was increased in cells transfected with \(\text{miR-222}\). No prominent change in cell cycle was observed in MIA PaCa-2 cells transfected with \(\text{miR-29b}\). A marked increase in the proportion of the sub-G1 fraction was not observed in MIA PaCa-2 cells transfected with any of the five proliferation-suppressive miRNAs.

**Table I.** Top five ranked miRNAs that suppress proliferation of MIA PaCa-2 cells

<table>
<thead>
<tr>
<th>miRNA clones</th>
<th>Mean (\log_{10}) ratio of two independent screenings (P9 versus P0)</th>
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<tbody>
<tr>
<td>miR-532</td>
<td>-1.668</td>
</tr>
<tr>
<td>miR-224</td>
<td>-1.639</td>
</tr>
<tr>
<td>miR-29b</td>
<td>-1.211</td>
</tr>
<tr>
<td>miR-34a</td>
<td>-1.153</td>
</tr>
<tr>
<td>miR-222</td>
<td>-0.970</td>
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Fig. 3. Cell proliferation assays of MIA PaCa-2 cells. (A) 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide assay of MIA PaCa-2 cells 5 days after infection with lentivirus vectors expressing miRNA, (B) Cell proliferation curve and (C) Phase contrast micrographs 4 days after transfection with synthetic miRNAs.

Fig. 4. Distribution of cell cycle phases of MIA PaCa-2 cells 48 and 72 h after transfection with synthetic miR-29b, -34a, -222, -224, -532 and miR-negative control (miR-NC). The average ± standard deviations of three biological replicate samples are shown.
Discussion

A growing body of evidence supports the critical roles of miRNAs in a variety of normal cellular processes, including cell proliferation, cell death and development (3,4). Comprehensive profiling studies have revealed aberrant expression of miRNAs in various pathological conditions, suggesting that miRNAs have a role in the pathogenesis of some diseases, such as cancer. There is increasing interest in miRNA as therapeutic targets, as well as in the application of miRNA as therapeutic agents (5,8). The identification of miR-34a, which is trans-activated by p53 and represses cell proliferation both in vitro and in vivo, prompted us to develop an assay to systematically screen for miRNAs that are directly linked to phenotypes characteristic of cancer cells (6). We combined lentivirus vectors expressing miRNA precursors and a custom-made microarray, which we used to monitor the number of lentivalent integrations based on specific sequences within the Lenti-miR virus library.

The usefulness and advantages of such a functional screening approach has recently been shown using a retrovirus vector library expressing short hairpin RNA for several thousand genes, although these short hairpin RNAs are only targeted to knockdown protein-coding genes (9–11). Voorhoeve et al. (12) have also constructed a genetic screening assay of miRNAs using a retrovirus vector library (miR-Lib) expressing miRNA mimigenes and a barcode microarray (miR-Array). These phenotype-based approaches enabled identification of essential genes critical in cell proliferation that do not necessarily have mutated sequences or aberrant copy numbers and that do not exhibit significantly altered gene expression. Together with the aforementioned assays, the functional screening assay presented here would be a powerful complementary tool for the elucidation and identification of genes critical in establishing the phenotypic characteristics of cancers. By employing lentivirus vectors, we can perform the functional screening assay using non-dividing cells, including stem cells and neural cells, thus broadening the potential application of the assay.

The feasibility of our assay was shown by the successful identification of miRNAs that suppress proliferation of MIA PaCa-2 pancreatic cancer cell lines. We individually validated the proliferation-suppressive effect of the miRNAs that showed significant reduction in their copy numbers in our functional screening assays by transfection of synthetic miRNAs or transduction using lentivirus vectors (Figure 3). Although the fold changes in the expression of these miRNAs in comparison with the endogenous counterpart showed variations, the proliferation-suppressive effect of these miRNAs was at least evident under our experimental conditions (supplementary Table S2 is available at Carcinogenesis Online). Interestingly, miR-34a, a representative tumor-suppressive miRNA, was identified in the functional screening assay. miR-34a is located on chromosome 1p36, which is frequently lost in a variety of cancers (13–17). p53 is one of the factors transactivating miR-34a, and it also upregulates p53, suggesting a positive feedback loop formed by p53 and miR-34a (18,19). When introduced to cells, miR-34a strongly repressed proliferation of colon cancer cell lines HCT116 and RKO, and the cells underwent apoptosis or a premature-senescence phenotype depending on the experimental conditions (18,20–23). Topical application of miR-34a also inhibited growth of HCT116 cell lines transplanted to nude mice, implicating miR-34a as a potential novel therapeutic agent (18). The precise role of miR-34a in pancreatic cancers is still unclear, though its expression is downregulated in most pancreatic cancer cell lines (24). The present functional assay supports a tumour-suppressive role for miR-34a in pancreatic cancers, warranting further in vivo validation studies of this miRNA using mouse models of pancreatic cancers.

Besides miR-34a, we identified four candidate tumor-suppressive miRNAs in pancreatic cancers, namely miR-29b, -222, -224 and -532. Although a detailed function analysis of these miRNAs is beyond the scope of this study, flow cytometric analysis revealed changes in the distribution of cell cycle phases of MIA PaCa-2 cells transfected with miR-222, -224 and -532. Thus, cell cycle arrest is, at least in part, an underlying mechanism of the proliferation-suppressive effect of these miRNAs. Among the predicted targets of these miRNAs are cell division cycle 42 (GTP-binding protein, 25kDa) (CDC42) (miR-224 and -532) and p21 protein (Cdc42/Rac)-activated kinase 1 (PAK1) (miR-222) (25–28). CDC42 is a small-guanosine triphosphatase of the Rho-subfamily that contributes to G1–S phase progression through p70 S6 kinase-mediated induction of cyclin E expression, suggesting the possible role of miR-224 and -532 in G1 arrest through translational suppression of CDC42 (29,30). PAK1 is a serine/threonine p21-activated kinase and is the downstream effector of CDC42 and Rho (31). Knockdown of PAK1 in gastric cancer cells exhibited an increased proportion of cells in G2/M phase, indicating the possible role of miR-222 in G2 arrest through translational suppression of PAK1 (32). Whereas cell cycle arrest was evident in MIA PaCa-2 cells transfected with miR-222, -224 and -532, miR-29b apparently suppresses cell proliferation through other mechanisms. Park et al. (33) reported that miR-29b upregulates p53 levels through suppression of p85α and CDC42. Although TP53 is mutated in MIA PaCa-2 cells (R248W), the combination of suppression of p85α and CDC42 may synergistically confer a proliferation-suppressive phenotype independent of p53.

In conclusion, we have developed a functional screening assay of miRNAs by the combination of a pooled lentivirus vector expressing miRNA precursors and a custom-made microarray. The feasibility of the assay was shown by the successful identification of miRNAs that suppress proliferation of MIA PaCa-2 pancreatic cancer cells. Flow cytometric analysis revealed that cell cycle arrest was, at least in part, the underlying mechanism of proliferation-suppressive effects of miR-34a, -222, -224 and -532. The flexible nature of the assay should facilitate its use in the identification of miRNAs that are involved in a wide array of cancer phenotypes, including cancer stem cells or metastasis.

Supplementary material

Supplementary Tables S1 and S2 and Figures S1 and S2 can be found at http://carcin.oxfordjournals.org/.

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References


