A Combined Array-Based Comparative Genomic Hybridization and Functional Library Screening Approach Identifies mir-30d As an Oncomir in Cancer

Ning Li1,5, Sippy Kaur4, Joel Greshock2, Heini Lassus4, Xiaomin Zhong1, Yanling Wang1, Arto Leminen4, Zhongjun Shao1, Xiaowen Hu1, Shun Liang1, Dionyssios Katsaros6, Qihong Huang3, Ralf Butzow4, Barbara L. Weber2, George Coukos1,2, and Lin Zhang1

Abstract

Oncomirs are microRNAs (miRNA) that acts as oncogenes or tumor suppressor genes. Efficient identification of oncomirs remains a challenge. Here we report a novel, clinically guided genetic screening approach for the identification of oncomirs, identifying mir-30d through this strategy. mir-30d regulates tumor cell proliferation, apoptosis, senescence, and migration. The chromosomal locus harboring mir-30d was amplified in more than 30% of multiple types of human solid tumors (n = 1,283). Importantly, higher levels of mir-30d expression were associated significantly with poor clinical outcomes in ovarian cancer patients (n = 330, P = 0.0016). Mechanistic investigations suggested that mir-30d regulates a large number of cancer-associated genes, including the apoptotic caspase CASP3. The guided genetic screening approach validated by this study offers a powerful tool to identify oncomirs that may have utility as biomarkers or targets for drug development. Cancer Res; 72(1); 154–64. ©2011 AACR.

Introduction

Cancer is a genetic disease involving multistep changes in the genome (1). Most studies to date have focused mainly on the role of protein-coding sequences in cancer. Although noncoding sequences constitute up to 98% of the human genome and have been implicated in tumorigenesis, our knowledge of the function of these genomic sequences in cancer is still limited (2–7). MicroRNAs (miRNAs) are noncoding RNAs about 22 nucleotides in length, which regulate gene expression in a sequence-specific manner (2–4, 6–8–13).

Increasing evidence indicates that miRNAs may play a critical role in cancer (14–21). miRNAs act as key regulators of various fundamental biological processes, many of which share common pathways with the development of cancer. In addition, it has been widely reported that miRNA expression is remarkably deregulated in cancer. Several miRNAs have already been shown to play a critical role in cancer. Referred to as "oncomirs," these miRNAs can act as either tumor suppressors or as oncogenes (4). Current rapid advances in oligonucleotide/nanoparticle therapies create a realistic optimism for the establishment of miRNAs as a new and potent therapeutic target for cancer treatment (22–25). However, the efficient identification of oncomirs remains a challenge. Although both microarray based high-throughput genomic/genetic profiling (3, 18, 19, 26) and whole genome-wide genetic screening (16, 27–31) have been successfully applied to the identification of oncomirs in human cancer, both approaches have limitations. Genome-wide screening is time and labor intensive and, most importantly, can produce artifacts. The profiling strategies cannot distinguish "causal" from "bystander" genetic events. In this study, we report a novel, clinically guided genetic screening approach to identify oncomirs involved human cancer. On the basis of this approach, one promising oncomir candidate, mir-30d, was successfully identified.

Materials and Methods

Patients and specimens

All frozen ovarian cancer specimens were collected at the University of Turin. Detailed information is provided in Supporting Methods.

Note: Supplementary data for this article are available at Cancer Research Online (http://cancerres.aacrjournals.org).
Cell lines and cell culture
Cancer cell lines 2008, OVCAR5, OVCAR10, OWA42, MDA-MB-231, T47D, HCT116, and BT549 were purchased from American Type Culture Collection (ATCC) or DCTD Tumor/Cell Line Repository. Four independent immortalized human ovarian surface epithelial cell lines (IOSE) were generously provided by Dr. Nelly Auersperg. All cancer cell lines were obtained directly from ATCC or DCTD that carries out cell line characterizations and passaged in our laboratory for fewer than 6 months after receipt. The method of characterization used by ATCC or DCTD can be found in its website. Detailed information is provided in Supporting Methods.

Retroviral transductions
The retrovirus-based human miRNA expression vector (pmiRvec) was purchased from GeneService (27). Retroviral vectors containing human miRNAs were transfected into the packaging cell line PT67 (Clontech) using the FuGene6 Transfection Reagent (Roche). The growth media was changed 48 hours posttransfection, and the media containing retrovirus was collected 48 hours later. Human tumor cells were infected with retrovirus in the presence of 8 μg/mL of polybrene and selected using the antibiotic blasticidin (Invitrogen).

Transfection of miRNA mimic oligos and of miRZip miRNA inhibitors
Pre-miR miRNA precursors and control oligos were purchased from Ambion. miRZips lentiviral-based miRNA inhibition vectors and control vectors were purchased from System Biosciences. The effects of transfection or infection on endogenous mir-30d expression were examined by real-time reverse transcriptase PCR (RT-PCR; Supplementary Fig. S1). Detailed information is provided in Supporting Methods.

RNA isolation and TaqMan miRNA assays
Total RNA was isolated from 100 to 500 mg of frozen tissue or 1 × 10⁶ cultured cells using TRIzol reagent (Invitrogen). The expression of mature miRNAs was analyzed using the TaqMan miRNA Assay (Applied Biosystems). Detailed information is provided in Supporting Methods.

MTT proliferation assays, cell-cycle assays, caspase-3 activity assays, Annexin-V apoptosis assays, β-Gal senescence assays, and migration assays
Detailed information is provided in Supporting Methods.

Western blots
Anti-total caspase-3 and anti-cleaved caspase-3 primary antibodies were purchased from Cell Signaling Technology. Detailed information is provided in Supporting Methods.

Generation of the in vivo xenograft model and bioluminescence optical imaging
Six to 8-week-old female nude mice were used to generate xenograft tumor models. The IVIS Lumina II Bioluminescence and Fluorescence Imaging System (Caliper Life Sciences) was used for in vivo bioluminescent imaging of MDA-MB-231 tumors. Detailed information is provided in Supporting Methods.

Array-based comparative genomic hybridization
BAC clones included in the “1 Mb–array” platform have been previously described (32). Detailed information is provided in Supporting Methods.

Analysis of DNA copy number profiling data sets
We examined whether mir-30d exhibits a high frequency of genomic amplification in a variety of human cancers by analyzing a panel of high-resolution DNA copy number profiling data sets. These data sets consisted of 1,282 primary tumors representing 9 human cancer types. Detailed information is provided in Supporting Methods.

Single-nucleotide polymorphism arrays
A total of 192 established tumor cell lines were analyzed using the Affymetrix 500K single-nucleotide polymorphism (SNP) chip (Affymetrix; data were collected by Joel Greshock and Barbara L. Weber at GlaxoSmithKline). Detailed information is provided in Supporting Methods.

SNP data analysis
All SNP chip images (‘CEL files’), were extracted using Affymetrix Genotype software and analyzed using the dChip software package. Detailed information is provided in Supporting Methods.

Tissue microarrays
Detailed information is provided in Supporting Methods.

miRNA in situ hybridization and image analysis
In situ detection of miRNA expression was done on formalin-fixed paraffin-embedded tissue microarray sections. Detailed information is provided in Supporting Methods.

cDNA microarrays and microarray analysis
A Human Gene 1.0 ST oligonucleotide array (Affymetrix) was used in this study. Detailed information is provided in Supporting Methods.

Bioinformatic analysis
mir-30d targets were predicted using TargetScan (33). miRNA effects to the expression levels of protein-coding transcripts were identified by DIANA-miRExTra (34).

Quantitative real-time RT-PCR
Total RNA was reverse transcribed using a High-Capacity RNA-to-cDNA Kit (Applied Biosystems) under conditions provided by the supplier. cDNA was quantified by RT-PCR on an ABI Prism 7900 Sequence Detection System (Applied Biosystems).

Statistical analysis
Statistical analysis was done using the SPSS or SAS statistics software package. All results were expressed as mean ± SD, and P < 0.05 was used for significance.
Results

A clinically guided genetic screening approach to identify oncomirs

We hypothesized that the limitations of the approaches discussed above could be overcome by integrating these 2 approaches, that is, using miRNA array profiling as a ‘clinical/biological’ filter to generate a restricted/selected miRNA expression library, which could then be used for functional genetic screening. To test our hypothesis, we generated a restricted miRNA expression library as follows: (i) We used array-based comparative genomic hybridization (aCGH) profiles as the ‘clinical/biological’ filter. miRNAs that are highly amplified in human cancer were selected on the basis of our previous aCGH studies (32). In ovarian (n = 109) and breast (n = 73) cancer specimens, 55 and 166 miRNAs were identified, respectively, with DNA copy number amplification observed in more than 15% of cases (Fig. 1A; Supplementary Table S1). (ii) These potential oncomirs were further concentrated by combining the miRNAs that were coamplified in both ovarian and breast tumors (n = 41; Fig. 1B; Supplementary Table S1). (iii) We used a whole genome human miRNA expression library (27) to derive this restricted library by selecting retrovirus miRNA expression vectors for these 41 miRNAs. This restricted library was used in an ovarian cancer cell line, 2008, to screen in vitro for miRNAs that are advantageous for cell growth in long-term culture. This was done using a modification of a previously described whole genome-wide short hairpin RNA screening method (refs. 35, 36; Fig. 1C).

Several potential candidate oncomirs, including mir-30d, were identified by the above approach. mir-30d represented 42% of the viral insertions sequenced. To validate and further characterize this finding, we first transfected a chemically synthesized mir-30d mimic into 6 cancer cell lines, including 2008, and 5 independent cancer cell lines. Using an MTT assay, we found that forced mir-30d expression significantly increased cell growth in all 6 cell lines (Fig. 2A). Meanwhile, a similar experiment was done using a lentiviral mir-30d inhibitor. Consistent with our gain-of-function studies, blocking of endogenous mir-30d expression even more remarkably decreased the growth rates of cancer cell lines in vitro (Fig. 2B). In 2008, MDA-MB-231 and HCT116, mir-30d inhibitor nearly completely blocked cell growth ability in vitro (Fig. 2B). In both mimic and inhibitor experiments, total RNA was isolated from parallel transfection wells, and real-time RT-PCR was used to confirm the effect on mir-30d expression. Because blocking endogenous mir-30d significantly reduced tumor cell proliferation, we next examined the effect of mir-30d inhibitor treatment on the cell cycle, using DNA staining followed by flow cytometry. We found that mir-30d inhibitor transfection led to a significant cell-cycle arrest at the G0/G1 phase (Fig. 2C). Finally, we observed that transfection of a mir-30d inhibitor not only significantly blocked cancer cell proliferation (Fig. 2B) but also remarkably increased the number of the cells with typical senescence morphology (Fig. 2D). To confirm this observation, we examined β-galactosidase activity, a known characteristic of senescent cells that is not found in

Figure 1. A clinically guided genetic screening approach to identify oncomirs. A, aCGH frequency plots of ovarian (n = 109) and breast (n = 73) cancer specimens. Columns represent miRNAs which are ordered on the basis of location in the genome (from chromosome 1 to 22). B, Venn diagrams of miRNA genes with copy number gain shared by ovarian and breast cancers. C, a clinically guided genetic screening approach to identify oncomirs in vitro.
presenescence, quiescence, or immortal cells. Four to 6 days after transfection with a mir-30d inhibitor, the percentage of β-galactosidase–positive cancer cells was significantly increased compared with controls (Fig. 2E).

Most recent studies have suggested that the mir-30 family regulates apoptosis in cardiomyocytes (37) and cancer cells (30). It indicates that decreasing of cell death may be another mechanism to explain the advantage of mir-30d for cell growth in long-term culture (about 2 months selection in our genetic screening approach). Thus, we examined whether mir-30d regulated apoptosis in the cell lines used in this study. On the basis of a bioluminescent caspase assay, we found that transfection of a mir-30d mimic significantly reduced caspase-3 activity in camptothecin-induced apoptosis in cancer cells (Fig. 3A). This result was further confirmed via Western blot. Figure 3B shows that the transfection of a mir-30d mimic remarkably reduced both total and cleaved caspase-3 protein in both control and camptothecin-induced apoptotic cells. To prove that mir-30d blocks chemotherapy drug-induced apoptosis, the apoptotic status after camptothecin treatment was analyzed by flow cytometry using an Annexin-V assay and Western blots of PARP protein. Consistent with the above results, we found that the transfection of a mir-30d mimic significantly reduces the apoptotic cells induced by camptothecin treatment (Fig. 3C and D). Taken together, the above results show that mir-30d regulates proliferation, senescence, and apoptosis in cancer cells, suggesting that mir-30d may serve as an oncomir in human cancer.
mir-30d blockade significantly inhibits tumor growth in vivo

We next questioned whether mir-30d could regulate tumor growth in vivo. Cancer cells (MDA-MB-231) in which mir-30d had been knocked down by a lentiviral inhibitor and control cells were injected subcutaneously into female nude mice (Fig. 4). The cells in which mir-30d were knocked down lost all in vivo growth ability, whereas the control cells grew well in vivo in the same animal (Fig. 4A and B, n = 5, P < 0.05). To enhance tumorigenicity of mir-30d knocked down tumors, we repeated the above experiments using tumor cells mixed with Matrigel, a mixture of basement membrane proteins, tissue plasminogen activators, and growth factors. To increase the sensitivity of tumor visualization, a luciferase reporter expression vector was transfected into the MDA-MB-231 cells. We found that the tumors produced by cells in which mir-30d had been knocked down displayed slow growth in vivo with the Matrigel mixture, consistent with the first experiment (without the Matrigel mixture). Thus, knockdown of mir-30d significantly decreases the growth rate of tumors in vivo (Fig. 4C–E, n = 5, P < 0.05). This in vivo result provides further evidence indicating that mir-30d is an oncomir in human cancer.
mir-30d exhibits a high frequency of genomic amplification in human solid tumors

Our previous aCGH study indicated that the DNA copy number of mir-30d was amplified in about 50% of the ovarian cancer (n = 109) and 40% of the breast cancer (n = 73) specimens (32). First, we investigated whether there was a positive correlation between the DNA copy number amplification and mature mir-30d expression in primary tumors. We analyzed mature and pre-mir-30d expression in 36 primary ovarian tumor specimens by real-time RT-PCR and found that both mature and pre-mir-30d expression was detectable in all specimens and significantly higher in the tumors with a high DNA copy number of mir-30d (n = 21, mature mir-30d: P = 0.014; pre-mir-30d: P < 0.001) compared with tumors with no amplification of mir-30d DNA copy number (n = 15). We also examined the 16 ovarian cancer cell lines. The average level of mir-30d expression was higher in the cell lines with amplified mir-30d DNA copy numbers (Fig. 5A). In addition, the average level of mir-30d expression in these ovarian cancer cell lines was remarkably higher than in IOSEs (Fig. 5A), which were used as control cells for epithelial ovarian cancer (38). These results show that genomic copy number amplification of mir-30d increases mature mir-30d expression in human cancer.

Next, we asked whether mir-30d also exhibits high levels of genomic amplification in other human cancers. We analyzed a panel of high-resolution DNA copy number profiling data sets that consisted of 1,282 primary tumors representing 9 human cancer types (detailed information provided in the Methods section and Supplementary Table S2). We found that the genomic locus harboring mir-30d was significantly amplified in 8 of these 9 human cancers (Fig. 5B). For example, in breast cancer (n = 327), 8.1% of the specimens revealed that the genomic locus harboring mir-30d was amplified to greater than 5 copies, and 30.4% of the specimens indicated that mir-30d was present in 3 to 5 copies (Fig. 5B). Overall, mir-30d was amplified at a high frequency (>30%) in human solid tumors (n = 1,282). In contrast, 2.7% of tumors displayed a heterozygous deletion for mir-30d, and one specimen with a homozygous deletion of mir-30d was found (Fig. 5B). These DNA copy number profiling results were further validated using a SNP array on 6 matching human tumor types (detailed information provided in the Methods section). One hundred and ninety-two established tumor cell lines were examined in this SNP array.
The genomic locus containing mir-30d was examined in each cell line (Fig. 5C). Consistent with the aCGH findings, mir-30d displayed remarkable amplification in tumor cell lines (Fig. 5B). In breast cancer (n = 29), in 6.9% of specimens the genomic locus harboring mir-30d was present in more than 5 copies, whereas 44.8% of the specimens had 3 to 5 copies (Fig. 5B). Collectively, mir-30d was amplified in more than 30% of the cancer cell lines examined (Fig. 5B, n = 192).

Higher levels of mir-30d expression were significantly correlated with poor clinical outcomes in ovarian cancer patients

To study the expression of mature mir-30d in human cancer, in situ hybridization (ISH) was done on ovarian cancer tissue arrays (n = 330). We detected mature mir-30d in the majority of ovarian cancer specimens (81.2%, 268/330, Fig. 5D). Strong cytoplasmic hybridization signal was mainly found in tumor islet but not stroma cells (Fig. 5D). To further examine the clinical significance of mir-30d expression, the cytoplasmic ISH signals were scored as negative or weak positive (n = 62), intermediate positive (n = 161), and strong positive (n = 107, Fig. 5D). We found that higher expression levels of mature mir-30d in ovarian cancer were significantly correlated with poor clinical outcomes in these patients (Fig. 5E, P = 0.0016). The above clinical information (DNA copy number amplification and correlation with survival) provides robust evidence that mir-30d does indeed play a critical oncogenic role in the development and progression of human cancer.

mir-30d regulates a large number of transcripts/genes in cancer cells

Up to one-third of mRNAs seem to be miRNA targets (33). Each miRNA can target hundreds of mRNA transcripts (39).
affecting the production of proteins (40, 41), and more than one miRNA can converge on a single target transcript (42). Taken together, the potential regulatory circuitry afforded by specific miRNAs may be enormous, although typically this repression is relatively mild (39–41). Therefore, we believed that the oncomiric function of mir-30d would be mediated by repressing multiple signaling pathways rather than by targeting one or a few cancer-associated proteins. To test this hypothesis, we used microarray analyses to profile the transcriptional changes seen after the transfection of a mir-30d mimic into 2 cancer cell lines (Fig. 6). Consistent with other reports in the literature (39–41), the repression of genome-wide transcription caused by mir-30d was typically relatively mild. We found that the mir-30d mimic downregulated 462 and 376 transcripts in 2008 and MDA-MB-231 cells, respectively (Fig. 6A–E; Supplementary Tables S3 and S4). To confirm that the majority of the downregulated transcripts were repressed by mir-30d directly, we undertook both bioinformatic and experimental validations. First, following a method described by Krutzfeld and colleagues (43), we calculated the number of nonoverlapping occurrences of each possible hexamer in every 3′-UTR (untranslated region) sequence of those downregulated transcripts (377 and 313 annotated genes were included in 2008 and MDA-MB-231, respectively). After dividing this by the length of the 3′-UTR, we applied a 1-tailed Wilcoxon rank sum to each normalized hexamer count distribution in downregulated versus unchanged 3′-UTRs using the online program DIANA-mirExTra (ref. 34; Fig. 6B and D). The 3 hexamers that were most overrepresented in the downregulated genes were ACATTT, CATTTG, and ATTTGT (P < 10^-19) and corresponded to the seed positions of mir-30d (Fig. 6B and D). We also predicted mir-30d targets using TargetScan (33) and found that 128 of 462 and 151 of 376 downregulated transcripts in 2008 and MDA-MB-231, respectively, were mir-30d predicted targets (Fig. 6E). This bioinformatic evidence strongly suggests that a large percentage of those downregulated transcripts were indeed directly repressed by mir-30d. That is, about 30% to 60% of the downregulated transcripts were direct mir-30d targets. In addition, about 1 of 3 (n = 150) downregulated transcripts were shared by both cell lines, and 79 of these were predicted to be mir-30d targets by TargetScan (Supplementary Table S5). Finally, 11% (9 of 79) of these transcripts were selected and further validated by real-time RT-PCR in 2008 and MDA-MB-231 cells, as well as another 2 independent cell lines, HCT116 and A2780.

Figure 6. mir-30d regulates a large number of transcripts/genes in cancer cells. A and C, the genome-wide transcriptional effect of mir-30d mimic transfection compared with control mimic transfection was analyzed using an Affymetrix Human Gene 1.0 ST oligonucleotide array in ovarian cancer cell line 2008 (A) and breast cancer cell line MDA-MB-231 (C). Light gray spots indicate downregulated transcripts. B and D, histograms of negative natural logarithms of P values derived from a 1-tailed Wilcoxon rank sum test applied to the distributions of hexamers in the 3′-UTRs of all downregulated versus unchanged transcripts. E, Venn diagrams of transcript numbers shared by downregulated transcripts in 2008 and MDA-MB-231 cells and predicted targets of mir-30d by TargetScan. F, the microarray results were validated by real-time RT-PCR in 4 tumor cells. Real-time RT-PCR validation of transcripts that were downregulated in both 2008 and MDA-MB-231 cells after transfection with the mir-30d mimic and that were also predicted mir-30d targets by TargetScan (E). Validations were done in 2008 and MDA-MB-231 cells, as well as in 2 independent cell lines, HCT116 and A2780. *, P < 0.05; **, P < 0.01.
A2780 (Fig. 6F and Supplementary Fig. S2). Notably, 3 of known mir-30d targets identified by another group, GNAI2 (44) and GALNT1, GALNT7 (45) were consistently found in our 79 transcript list (Fig. 6F). Taken together, these results indicate that mir-30d represses a large number of transcripts in cancer cells.

Most importantly, miRNA target prediction programs strongly indicated that CASP3, a central protein in the execution phase of cell apoptosis, was a potential target of mir-30d (Fig. 7A). The mir-30d binding sites in the CASP3 3′-UTR predicted by TargetScan were broadly conserved among different species (Fig. 7A). In addition, we found that both protein and mRNA levels of CASP3 were significantly repressed after transfection of mir-30d into cancer cells (protein: Fig. 3B and mRNA: Fig. 6E and 7B). Thus, we used a luciferase reporter assay to experimentally examine whether mir-30d directly regulated CASP3 expression via its 3′-UTR. The mir-30d mimics and control mimic were transiently transfected into 4 cancer cell lines. At 48 hours posttransfection, total RNA were collected, and the endogenous CASP3 expression was examined by real-time RT-PCR. As indicated in Fig. 6E and 7B, mir-30d significantly reduced luciferase activity in the wild type but not the binding site mutant CASP3 3′-UTR reporters (Fig. 7C). These results show that mir-30d directly regulates expression of CASP3, a master regulator of apoptosis, in cancer cells. It is one of important molecular mechanisms to explain our observation on cellular function of mir-30d in apoptosis and long-term maintenance of tumor cell growth.

Discussion
In this study, we report a novel, clinically guided, genetic screening approach to identify oncomirs in human cancer. This approach uses array-based profiling data as a clinical/biological filter to generate a selected functional genetic screening library. Our strategy can distinguish “causal” from “bystander” genetic events to decrease noise and artificial results and also requires less time and labor than whole-genome wide genetic screening approaches. With this novel strategy, one promising oncomir candidate, mir-30d, was successfully identified. This clinically guided genetic screening approach is a powerful tool to identify oncomirs as biomarkers for diagnosis or as targets for the development of cancer treatments. There are also limitations of our strategy to identify oncomirs. For example, only oncogenic miRNAs, especially miRNAs promoting tumor cell growth or survival, are able to be identified by our current approach. This limitation can be overcome by combining with other library screening approaches such as transwell assay (to identify miRNAs promoting migration and invasion), soft agar assay (to identify miRNAs inducing malignant transformation), and negative selection approaches (to identify tumor suppressor miRNAs). Finally, because our approach is based on an in vitro positive selection strategy, the oncomirs that give a growth advantage to tumor cells in vitro, such as promoting angiogenesis and

![Image](https://cancerres.aacrjournals.org/)

Figure 7. mir-30d regulates CASP3 in cancer cells. A, the schematic diagram of the mir-30d binding site in the CASP3 3′-UTR, which was broadly conserved among different species. Hsa, Human; Prm, chimpanzee; Mmt, Rhesus; Mmu, mouse; Rno, Rat; Cpo, Pig; Srr, Shrew; Cfa, Dog; Fca, Cat; Eca, Horse; Bta, Cow; Dno, armadillo; Lep, elephant. B, the mir-30d mimics and control mimic were transiently transfected into 4 cancer cell lines. At 48 hours posttransfection, total RNA were collected, and the endogenous CASP3 expression was examined by real-time RT-PCR. C, reporter assays using wild type and mir-30d binding site mutant CASP-3 3′-UTR. The mir-30d mimic transfection significantly reduced luciferase activity in the wild type but not the binding site mutant CASP3 3′-UTR reporters. WT, wild type. *P < 0.05; **, P < 0.01.
mir-30d in Human Cancer

regulating antitumor immune response, cannot be identified by our approach. Combination with an in vivo selection approach will more closely recapitulate the pathobiology of human cancers.

mir-30d was initially identified using this screening approach, in which the cells harboring miRNAs producing a growth advantage could be expanded in vitro. Subsequently, our in vitro experiments showed that mir-30d regulates cell proliferation, senescence, and apoptosis. In agreement with our observations, Kumar and colleagues also reported that mir-30d inhibits apoptosis in human cancer via regulation of the p53 pathway (30). A similar function of mir-30d was also observed in normal cells such as cardiomyocytes (37). Furthermore, using a xenograft model, we showed that inhibition of endogenous mir-30d expression remarkably blocked tumor growth in vivo. Two independent groups have provided strong evidence indicating that mir-30d promotes tumor invasion and metastasis in hepatocellular carcinoma via regulating GNAI2 (44) and in melanoma via repressing GALNT7 (45). Consistent with these studies, we also found that mir-30d repressed GNAI2 and GALNT7 expression in ovarian and breast cancer cell lines (Fig. 6E), and using an in vitro migration assay, we observed that transfection of mir-30d significantly promoted the migration of BT549 and MDA-MB-231 cells in vitro (Supplementary Fig. S3). Our study found that the DNA copy number of mir-30d was remarkably amplified (more than 30%) in 8 of 9 human cancer types that were examined. This suggests that mir-30d amplification in human cancer is a common oncogenic event. In agreement with our conclusion, 2 independent groups reported that mir-30d was amplified in brain tumors (46) and melanoma (45). Most importantly, we found that overexpression of mir-30d was significantly associated with a poor clinical outcome in human ovarian cancer patients. In addition, a recent genome-wide serum miRNA profiling study showed that higher levels of serum mir-30d were significantly associated with shorter survival times in non–small cell lung cancer patients (47). Gaziel-Sovran and colleagues also reported that higher levels of mir-30d expression was correlated with shorter time to recurrence and lower overall survival in melanoma (45). Taken together, the above evidence strongly suggests that mir-30d serves as an oncomir in human cancer.

The results of this study indicate that mir-30d can downregulate a large number of transcripts in human cancer cells, although as with most other miRNAs, this repression is relatively mild (39–41). We identified 79 protein coding genes which were predicted by bioinformatics to be potential direct targets of mir-30d. We then showed that the expression of these genes (involved in multiple cancer-associated pathways such as apoptosis and migration) were indeed downregulated by the transfection of a mir-30d mimic into each of the 2 cell lines used in our microarray study as well as 2 independent cancer cell lines. We found that CASP3, a central protein in the execution phase of cell apoptosis, is a direct target of mir-30d. Another mir-30d targets [GNAI2 which was identified in liver cancer (44); GALNT1 and GALNT7, which were identified in melanoma (45)] were also found in our target list. Notably, we also found that a relatively large number of predicted mir-30d target genes [49 of 462 in 2008 (ovarian cancer) and 72 of 376 in MDA-MB-231 (breast cancer) cells, respectively (Fig. 6E)] were only downregulated in one cell line in our microarray study. This indicates that the function of mir-30d may be cell-context dependent or tumor-type specific. In agreement with our finding, Martines and colleagues showed that the mir-30 family regulates B-Myb expression in a cervical cancer cell line (48).

Finally, because mir-30d regulates a large number of protein coding genes, it may regulate multiple cancer-associated pathways. For example, beyond apoptosis and migration, the mir-30 family may also regulate autophagy (49), immune suppression (45), and cancer stem cell differentiation (50). Therefore, the biological function and clinical significance of mir-30d in human cancer may be complex and dependent on the context of downstream targets and molecular networks in each specific cancer type.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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