Abstract. Long non-coding RNA MEG3 is suggested to function as a tumor suppressor. However, the activation mechanism of MEG3 is still not well understood and data are not available on its role under adenosine-induced apoptosis. In this study, HepG2 cells were treated with adenosine or 5-Aza-cdR. Methylation status of MEG3 promoter was detected by methylation specific PCR (MSP) and MEG3 expression was determined by qRT-PCR. PcDNA3.1-MEG3 recombinant plasmid was constructed and transfected to hepatoma HepG2 and Huh7 cells. Cell growth, morphological changes, cell-cycle distribution and apoptosis were analyzed by MTT assay, fluorescence microscopy and flow cytometry. The mRNA and protein expression levels were detected by qRT-PCR and western blot analysis. MEG3 binding proteins were screened by the improved MS2 biotin tagged RNA affinity purification method. The co-expression network of MEG3 was generated by GO analysis and ILF3 was identified as MEG3 binding protein by RNA pulldown and western blot analysis. Both adenosine and 5-Aza-CdR increased MEG3 mRNA expression and the CpG island of MEG3 gene in HepG2 cells was typical hypermethylation. Ectopic expression of MEG3 inhibited hepatoma cell growth in a time-dependent manner, resulted in cell cycle arrest and induced apoptosis. Ectopic expression of MEG3 increased p53, caspase-3 mRNA and protein levels, decreased MDM2 and cyclin D1 mRNA and protein levels, as well as ILF3 protein expression in HepG2 cells. These findings are the first to identify that adenosine increases MEG3 expression by inhibition of DNA methylation and its antitumor effects is involved in MEG3 activation. ILF3 may participate in the anticancer regulation of MEG3 by interacting with MEG3.

Introduction

Hepatocellular carcinoma (HCC) is the most common cancer and the third leading cause of cancer-related deaths worldwide (1). Yet, the pathogenesis mechanisms of HCC remain unclear. One of the potential molecular mechanisms involves the long non-coding RNAs (lncRNAs) (2,3).

lncRNAs play an important role in tumorigenesis and investigation of their biological functions and molecular mechanisms is important for understanding the development and progression of cancer (4). The lncRNA HOTAIR is overexpressed in HCC tissues compared to corresponding non-cancerous tissues and high HOTAIR expression tightly correlated with the presence of liver metastasis (5). lncRNA maternally-expressed gene 3 (MEG3) is an imprinted gene, located on human chromosome 14q32.3 (6). The downregulation of MEG3 has been observed in various human cancers (7-9). MEG3 might function as a tumor suppressor (10,11). However, the reasons for its downregulation and its function in HCC are not well understood.

Adenosine and their analogues can interfere with the synthesis of nucleic acids and exert genotoxic activity by incorporating with and altering the DNA or RNA macromolecules, which makes them exciting candidates for anticancer therapy (12-14). Although MEG3 plays a role in inhibiting tumor growth, no data are available on its function under genotoxic stress-induced apoptosis. In our previous study, we demonstrated that adenosine induces hepatoma cell apoptosis and the mechanism is involved in DNA demethylation and genotoxic effects (15,16). In this study, we treated HepG2 cells with adenosine or 5-Aza-2'-deoxy-cytidine (5-Aza-CdR), a selective DNA methyltransferase inhibitor, and observed the methylation status of MEG3 promoter and MEG3 mRNA expression. In addition, an improved MS2-BioTRAP (MS2 biotin tagged RNA affinity purification) technique was applied to screen MEG3-binding protein in HEK293 cell line. The antitumor effect of adenosine and MEG3 in HCC was observed and the possible molecular mechanisms were investigated.

Materials and methods

Cell culture and experimental groups. HepG2, Huh-7 and HEK293 cells, obtained from the American Type Culture
Collection, were cultured in Dulbecco's modified Eagle's medium supplemented with 10% (v/v) fetal bovine serum, penicillin (final concentration, 100 U/ml), and streptomycin (final concentration, 100 µg/ml) under a humidified atmosphere of 5% CO₂ and 95% air at 37°C. Cells were passaged at 70–80% confluence. HCC cells were divided into two groups for the experiment of overexpression MEG3: pcDNA3.1 (control group) and pcDNA3.1-MEG3 vector (experimental group).

**Treatment of HepG2 cells with adenosine or 5-Aza-CdR.** A total of 1.0x10⁵ HepG2 cells were inoculated into each well of a 6-well plate. After the cells attached, the original medium was replaced with fresh medium containing 4.0 mmol/l adenosine for 48 h or 20 µmol/l 5-Aza-cdR for 72 h. The medium with the same concentration of 5-Aza-CdR was replaced daily. Cells were collected and prepared for detection of MEG3 expression.

**Detection of DNA methylation by methylation-specific PCR (MSP).** A total of 1.0x10⁵ HCC cells were inoculated into each well of a 6-well plate and treated as described above (16). DNA from the cells was extracted using an EZNA Tissue DNA kit according to the manufacturer's instructions. The methylation status of the MEG3 gene was determined by MSP assay. For the MSP, the CpGenome Turbo Bisulfite Modification kit was used in accordance with the manufacturer's instructions. The sequences of the methylation-specific primers of MEG3 were as follows: the methylated pair (M), 5'-GTTAGTAATCGGGTTTGTCGGC(forward) and 5'-AATCATAACTCCGAACACCGCG(reverse); the unmethylated pair (U), 5'-GAGGATGGTTTGTCGGC(forward) and 5'-CCACCATGACCCGTCCCTTG-3' (reverse). MEG3 DNA was amplified in a DNA Engine Peltier thermal cycler (Bio-Rad) by 1 cycle of 95°C for 15 min followed by 40 cycles of 94°C for 30 sec, 70°C for 30 sec and 72°C for 30 sec; 5 cycles of 94°C for 30 sec, 65°C for 30 sec and 72°C for 30 sec; 30 cycles of 94°C for 30 sec, 60°C for 30 sec and 72°C for 30 sec; and 1 cycle of 72°C for 7 min. The PCR products were identified by electrophoresis through a 2.0% agarose gel and ethidium bromide staining.

**Plasmid identification and cell transfection and bacterial transformation.** PcDNA3.1-MEG3 plasmid was constructed as previously described (16). PcDNA3.1-MEG3 plasmids were extracted from E. coli according to the manufacturer's specifications. Both restriction enzymes, XhoI and BamHI, were used to cut the pcDNA3.1-MEG3 plasmid DNA for 1 h at 37°C. Subsequently, DNA fragments after digestion were identified by the agarose gel electrophoresis and quantified by capturing the ethidium bromide absorbance with a fluorescence microscope (Olympus BX51, Tokyo, Japan).

For ectopic MEG3 expression, the full-length MEG3 cDNA was subcloned into the pcDNA3.1 plasmid vector and transfected into HepG2 or HuH7 cells as previously described (16). PcDNA3.1 was transfected to cells in parallel as a control. The cells were then subjected to RNA/protein extraction and further functional assays.

**RNA extraction and qRT-PCR analysis.** HepG2 and HuH7 cells were transfected with pcDNA3.1-MEG3 vector or pcDNA3.1 for 48 h. The total RNA was isolated with TRIzol reagent, according to the manufacturer's protocol. For reverse transcriptase analysis, 1 µg of total RNA was reverse-transcribed using random primers under standard conditions using an EasyScript First-Strand cDNA Synthesis Super Mix kit. Real-time PCR amplification with one microliter of the reverse transcriptase reaction mixture was performed on the ABI 7500 real-time PCR system (Applied Biosystems, Foster City, CA, USA) using SYBR Premix Ex Taq following the manufacturer's instructions. PCR conditions were 95°C for 30 sec, followed by 40 cycles of 95°C for 3 sec and 60°C for 30 sec. Gene-specific primers are listed in Table I. 18s rRNA was used as a housekeeper gene for the qRT-PCR reactions. Fold change of MEG3, p53, MDM2, caspase-3 and cyclin D1 was calculated by the 2^{-∆∆Ct} method.

**Protein extraction and western blot analysis.** Cells were washed with ice-cold PBS, harvested and lysed in RIPA buffer supplemented with protease inhibitor cocktail. Cell lysates were centrifuged at 12,000 rpm at 4°C for 10 min. The supernatant was boiled for 5 min and subjected to 12% SDS-PAGE and transferred to a nitrocellulose membrane. The membranes were blocked with 5% fat-free milk for 1 h at room temperature for 30 min. Bands were visualized with horseradish peroxidase-conjugated secondary antibody for 1 h at room temperature, followed by incubation with primary antibodies against β-actin (1:3,000), Mdm2 (1:500), cyclin D1 (1:500), total caspase-3 (1:800), cleaved caspase-3 (1:800) and ILF3 (1:1,000) at 4°C overnight. The next day, the membranes were incubated with biotin-conjugated secondary antibody for 1 h at room temperature, followed by incubation with an HRP complex at room temperature for 30 min. Bands were visualized with an ECL detection system (Thermo Scientific, Waltham, MA, USA). Protein expression was analyzed by the Quantity One software (Bio-Rad, Hercules, CA, USA) and normalized to that of β-actin.

**Identification of proteins associated with IncRNA MEG3.** An improved MS2-BioTRAP (MS2 biotin tagged RNA affinity purification) technique was used which relies on the interaction of the bacteriophage MS2 coat-protein with a specific viral

---

**Table I. Primer sequences for quantitative real-time PCR.**

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer sequences</th>
</tr>
</thead>
<tbody>
<tr>
<td>MEG3</td>
<td>F: 5'-CTCAGGCGAGATCTGGCATATA-3'</td>
</tr>
<tr>
<td></td>
<td>R: 5'-CCTGGAGTGCTGTGGAAGA-3'</td>
</tr>
<tr>
<td>p53</td>
<td>F: 5'-TGAAGCTCCAGAGATGCG-3'</td>
</tr>
<tr>
<td></td>
<td>R: 5'-GGAGTACGTGCAGATCTACA-3'</td>
</tr>
<tr>
<td>MDM-2</td>
<td>F: 5'-ATCACGCGAGGAGAGTGT-3'</td>
</tr>
<tr>
<td></td>
<td>R: 5'-CAATTTCTACAGGGAAGGCCA-3'</td>
</tr>
<tr>
<td>Caspase-3</td>
<td>F: 5'-AGAGCTGACTGGTGATTTGGAG-3'</td>
</tr>
<tr>
<td></td>
<td>R: 5'-GAACCATGACCCGTCCCTTG-3'</td>
</tr>
<tr>
<td>Cyclin D1</td>
<td>F: 5'-GTGCACTACACCCGACAACCTCA-3'</td>
</tr>
<tr>
<td></td>
<td>R: 5'-GTCGATCTACACCCGACAACCTCA-3'</td>
</tr>
<tr>
<td>18S</td>
<td>F: 5'-AACGGCTACACGATCACCAG-3'</td>
</tr>
<tr>
<td></td>
<td>R: 5'-CAATTACAGGGCCTCAGAAG-3'</td>
</tr>
</tbody>
</table>
RNA stem-loop structure that was inserted into or grafted onto the 3' end of the lncRNA MEG3 (17). Twelve copies of the MS2 coat-protein binding site were introduced into the region encoding the MEG3 to generate a MEG3-MS2bs expression plasmid. This expression plasmid and a second plasmid encoding a Flag-tagged MS2 coat-protein were transiently introduced into cultured HEK293 cells. In brief, the pcDNA3.1-Flag-MS2 vector, pcDNA3.1-MS2bs vector and pcDNA3.1-MEG3-MS2bs vector were constructed as previously described by us (16). HEK293 cells are cultured and harvested at 80% confluence, followed by co-transfection with pcDNA3.1-Flag-MS2 vector and pcDNA3.1-MEG3-MS2bs using Lipofectamine 2000. Cells co-transfected with pcDNA3.1 and pcDNA3.1-MEG3-MS2bs vector were used as a control. Cells were further incubated for 48 h, then collected for RIP experiments. The protein components were determined by mass spectrometry, and the candidate proteins were confirmed by western blot analysis.

RNA binding protein immunoprecipitation. HEK293 cells were harvested and lysed in RIP lysis buffer with protein inhibitor, RNA inhibitor and DNase I on ice. Cell lysates were preincubated with the antibody. RNAs were immunoprecipitated with the antibody against RBP and protein A/G magnetic beads. The magnetic bead bound complexes were immobilized with a magnet and unbound material was washed off. Then, RNAs were extracted, precipitated and turned into a cDNA library by ImProm-II reverse Transcription system (Promega, USA). The remaining beads were boiled and separated by SDS-PAGE gel for western blot analysis.

Statistical analysis. All the experiments described were performed at least in triplicate. The data are expressed as mean ± SD. Statistical significance was calculated using either the Student's t-test, or one-way analysis of variance (ANOVA) with a post-hoc test of multiple comparisons. A p<0.05 was considered statistically significant.

Results

Downregulation of MEG3 is due to hypermethylation in the MEG3 promoter region. As shown in Fig. 1A and B, the mRNA relative expression level of MEG3 significantly increased after treatment with adenosine or 5-Aza-CdR (Fig. 1A and B). The unmethylated pattern (U) and a methylated pattern (M) were observed in the cells treated with adenosine or 5-Aza-CdR by MSP. The two DNA methyltransferase inhibitors, which reverse the abnormal methylation status of the MEG3 promoter, increased the expression of unmethylated product (120 bp, Fig. 1C). The results indicate that the methylation status of MEG3 promoter may lead to downregulation of MEG3 expression in HepG2 cells.

PcDNA3.1-MEG3 plasmid is constructed and identified. PcDNA3.1-MEG3 plasmid that expressed lncRNA MEG3 was constructed. LncRNA MEG3 gene was inserted into the pcDNA3.1 vector, which was confirmed by XhoI and BamHI restriction enzyme digestion. The particular sequence of MEG3 insertion into vectors was the same as designed and the fragment was not present in the control vector, showing pcDNA3.1-MEG3 plasmid was successfully constructed (Fig. 2).

Ectopic expression of MEG3 and its effects on cell growth inhibition. HepG2 and Huh7 cells were transiently transfected with either pcDNA3.1 (control) or pcDNA3.1-MEG3, and MEG3 mRNA expression was measured by qRT-PCR. Compared to the control group, MEG3 mRNA expression
levels increased by >90-fold or 80-fold in MEG3-transfected HepG2 or Huh7 cells (Fig. 3A), showing the high level mRNA expression of MEG3 was obtained in cells.

The effect of MEG3 on the proliferation of hepatoma cells was measured using the MTT assay. Ectopic expression of MEG3 significantly decreased cell vitality rate by 29.6 and 31.2% after 48 and 72 h in HepG2 cells, respectively, compared with the control group (p<0.01). Ectopic expression of MEG3 significantly decreased cell vitality rate by 31.5 and 45.9% after 48 and 72 h in Huh7 cells, respectively, compared with the control group (Fig. 3B, p<0.01). These results suggested that ectopic expression of MEG3 can inhibit tumor cell proliferation.

MEG3 induces cell apoptosis and cell cycle arrest in vitro. To detect whether apoptosis was a contributing factor to cell growth inhibition, hepatoma cells were stained with Hoechst 33258 after transfection with plasmids. The nuclei of the control group were uniform and without condensation or fragmentation. Compared with the control group (pcDNA3.1), nuclei in cells with ectopic expression of MEG3 became condensed and shrunken, and typical apoptotic bodies appeared. The percentages of apoptosis cells after MEG3 plasmid transfection are 18.1 and 13.3% in HepG2 and Huh7 cells, respectively, which is significantly higher than that of the control group (Fig. 4C, p<0.01).

In order to further investigate whether MEG3-induced apoptosis is related to the cell cycle, the cell cycle distribu-
Figure 4. Effects of ectopic expression of MEG3 on apoptosis and cell cycles in the hepatoma cells. (A) The morphology of apoptotic nuclei was observed by Hoechst 33258 staining and fluorescence microscopy. The percentage of apoptotic nuclei per optical field was scored. (B) Quantitative analysis of the total apoptotic cells by fluorescence microscopy. Each bar corresponds to the mean ± SD of three independent experiments. (C) Cell cycles and apoptosis percentage were analyzed by flow cytometry and the apoptosis cells in sub-G1 phase (blue peak) were determined. (D) Quantitative analysis of the cell cycles. Bar chart represents the percentage of cells in G0/G1, S, sub-G1 or G2/M phase. *p<0.05 vs. control; **p<0.01 vs. control.
tion was detected by flow cytometry. Ectopic expression of MEG3 significantly decreased the cells of G0/G1 and G2/M phase, but increased the percentage of S phase from 58.08 to 69.27% in HepG2 cells. Ectopic expression of MEG3 significantly increased the percentage of G0/G1 phase from 38.50 to 48.89% and decreased the percentage of G2/S phase in Huh7 cells (Fig. 4C and D, p<0.05). Statistical results showed that ectopic expression of MEG3 increased the percentage of apoptotic cells (sub-G1 phase) in both cell types (p<0.05, Fig. 4D), which is consistent with the result of fluorescence microscopy observations (Fig. 4B). Taken together, these results confirm that cell growth inhibition and cell cycle arrest caused by MEG3 is facilitated by the induction of apoptosis.

MEG3-triggered apoptosis is mediated by p53 pathway and cell cycle arrest. To probe the mechanism by which MEG3 initiated the cell cycle arrest, the investigation was focused on the cell cycle checkpoint, the cyclins (cyclin D1), p53 and other relative factors. Ectopic expression of MEG3 resulted in marked reductions in the mRNA and protein expression of cyclin D1 and MDM2. By contrast, the expression of p53 and caspase-3 mRNA and protein levels significantly increased in HepG2 cells (Fig. 5A-C, p<0.05). These data confirm that MEG3 functions as a tumor suppressor gene by activating the p53-mediated apoptosis pathway and causes cell cycle arrest in HepG2 cells.

ILF3 was also investigated by western blot analysis. The result showed that the expression level of ILF3 decreased after transfection with pcDNA3.1-MEG3, compared with the control group (Fig. 5B and C, p<0.05). The data indicated that ILF3 protein expression is negatively regulated by MEG3 in HepG2 cells.

Screening of MEG3-binding protein. MS2-BioTRAP (MS2 biotin tagged RNA affinity purification) experiment was carried out (Fig. 6). Sequence analysis and restriction endonuclease analysis of target genes demonstrated that the position and size of target gene cDNA insertion were consistent with the design (Fig. 6A-E). The pcDNA3.1-Flag-MS2 and pcDNA3.1-MEG3-MS2bs vectors were co-transfected into HEK293 cells for the RIP experiment. The expression intensity of MEG3 mRNA in co-transfected group with pcDNA3.1-Flag-MS2 and pcDNA3.1-MEG3-MS2bs vectors was ~98-fold higher than that of the control group, showing it is an effective method to study RNA-protein interaction.
The protein components were determined by mass spectrometry, and the candidate proteins were confirmed by RNA pulldown and western blot analysis. Of the four candidate proteins selected in the co-expression network by GO analysis, ILF3 protein expression obviously increased, showing ILF3 is a MEG3-binding protein (Fig. 6G).

Discussion

DNA methylation is a major regulatory mechanism of tumor epigenetic control and it has an important role in gene transcription and splicing. It has been demonstrated that DNA promoter hypermethylation is associated with inactivation of several tumor suppressor genes, and these aberrant DNA methylations are always accompanied by a progressive dysregulation of DNMT expression (18). It is reported that the promoter region of IncRNA MEG3 is rich in CpG dinucleotides. The methylation pattern of the 2.1 kb 5-flanking region of MEG3 contains functionally important sequences for gene expression (19). In general conditions, MEG3 is expressed in normal tissues, with high expression in the brain and pituitary gland (10). Many tumor tissues only express low MEG3, or even no MEG3 (9,20). In pituitary tumors and leukemia, MEG3 promoter hypermethylation is also associated with the loss of MEG3 expression (21,22). In our previous study, adenosine that has been confirmed to have cytotoxic effects on tumor cells also showed a demethylation role in HepG2 cells (15). In order to investigate the relationship between methylation status of MEG3 promoter and the expression of MEG3 gene, we treated hepatoma cells with adenosine or another selective DNMT inhibitor the 5-Aza-CdR. Our results showed that both adenosine and 5-Aza-CdR increased the expression of the unmethylated product and MEG3 mRNA expression (Fig. 1), thus revealing that MEG3 is correlated with a typical hypermethylation status in the promoter region in HepG2 cells, and that we are the first to demonstrate that adenosine can trigger MEG3 activation.
In order to further identify that MEG3 has an antitumor role, pcDNA3.1-MEG3 plasmid vectors were constructed and transfected into hepatoma cells. The results showed that ectopic expression of MEG3 inhibited cell growth in a time-dependent manner (Fig. 3B) and increased cell apoptosis in HepG2 and Huh7 cell lines (Fig. 4A and C).

The cell cycle is divided into four distinct phases, G1, S, G2, and M, and is tightly controlled by a series of key components including cyclins, the cyclin-dependent kinases. In this study, MEG3 induced cell cycle arrest significantly at the S phases and increased cell cycle arrest at the G0/G1 phases in HepG2 cells, accompanying by the decreased cyclin D1 expression, showing that cyclin D1 is essential for cell cycle transition of G1 to S phase and the decrease of cyclin D1 may lead to hampering cells from G0/G1 phase (23). However, we also observed that MEG3 caused different cell cycle arrest in two hepatoma cells (Fig. 4) and we considered the difference might be related to different p53 gene types, for HepG2 is a wild-type p53 and Huh7 is mutant p53 cell line.

Among all apoptosis-related genes, p53 is of particular importance (24). Tight regulation of the anti-proliferative activities of p53 is mostly achieved by the MDM2 ubiquitin ligase, which targets p53 for degradation (25). On the one hand, MDM2/p53 forms an autoregulatory feedback loop that may be needed to maintain a critical MDM2/p53 ratio within cells (26). On the other hand, an excess of MDM2 protein can abrogate the transcriptional activation of p53 (25). Factors that differentially regulate ubiquitination of MDM2 and p53 may affect cell fate profoundly as they can change the MDM2/p53 ratio (27). Because wild-type p53 is a tumor suppressive factor while mutant p53 does not have this function, we chose wild-type p53 HepG2 cells for our research. Fig. 5 shows that ectopic expression of MEG3 decreased MDM2, and increased p53, and cleaved-caspase-3 expression in HepG2 cells, suggesting that downregulation of MDM2 contributes to p53 activation (28,29). In addition, MEG3 might induce apoptosis through p53-independent pathways since exogenous MEG3 also inhibits Huh7 cell proliferation.

ILF3 is known to be a transcription factor and has two main isoforms (NF90 and NF110) (30,31). They participate in many aspects of RNA metabolism, including transcription, degradation and translation. In this study, an improved MS2-BioTRAP technique was used to screen the MEG3-binding proteins. Of the four candidate proteins selected in the co-expression network, ILF3 was identified as the binding protein by RIP, RNA pulldown and western blot analysis in HEK293 cells (Fig. 6). Zhu et al demonstrated that MEG3 can interact with DNA domain of p53 protein and regulate p53 target genes (32). Shamanna et al reported that knockdown of ILF3 by RNA interference led to elevated levels of p53 proteins in HPV-derived HeLa and SiHa cells (33). We speculated that MEG3 and ILF3 might form a complex which participates in the regulation of target genes (such as p53) and induce cell apoptosis. In this study, we observed that the expression level of ILF3 decreased after pcDNA3.1-MEG3 transfection in HepG2 cells. The mechanisms of ILF3 downregulation are due to increased ubiquitination and degradation, or other reasons that need to be further investigated.

In conclusion, this study demonstrates that the antitumor effect of adenosine is involved in MEG3 activation and the hypermethylation of MEG3 promoter region may contribute to its low expression. MEG3 negatively regulates hepatoma cell growth by affecting cell cycle progression and inducing cell apoptosis. Moreover, the present study identified that ILF3 is a MEG3 binding protein and might participate in the anticancer regulation of MEG3.

Acknowledgements

The authors are grateful for funding support from the National Nature Science Foundation of China (no. 30972925) and the Guangdong Natural Science Foundation in China (no. 2014A030313470). The authors would like to thank Professor En-min Li; and Dong-yang Huang and Yu-cai Fu for excellent technical support.

References


