COX-2 silencing enhances tamoxifen antitumor activity in breast cancer in vivo and in vitro

YE DU1, AIPING SHI1, BING HAN1, SIJIE LI1, DI WU1, HONGYAO JIA1, CHAO ZHENG1, LIQUN REN2* and ZHIMIN FAN1*

1Department of Breast Surgery, The First Hospital, Jilin University; 2Department of Experimental Pharmacology and Toxicology, School of Pharmacy, Jilin University, Changchun 130021, Jilin, P.R. China

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Abstract. Tamoxifen (Tam), a selective estrogen receptor modulator, is in wide clinical use for the treatment and prevention of breast cancer. However, extended TAM administration for breast cancer induces increased VEGF levels in patients, promoting new blood vessel formation and thereby limiting its efficacy and highlighting the need for improved therapeutic strategies. Cyclooxygenase-2 (COX-2) silencing via a replication-incompetent lentivirus (LV-COX-2) induce cancer apoptosis and suppresses VEGF gene expression. In this study, the effect of LV-COX-2 infection, either alone or in combination with TAM, was analyzed in a breast cell lines for suppressing VEGF expression and simultaneously reducing doses of TAM. Cell proliferation, apoptosis, angiogenesis, metastasis, cell cycle distribution, an receptor signaling were determined after LV-COX-2 combination with TAM treatment. In addition, tumor growth ability in nude mice was detected to define the combination treatment effect in tumorigenesis in vivo. It is found that LV-COX-2 combination with TAM treatment in breast cancer cell significantly suppressed the proliferation and metastasis, and induced tumor apoptosis in vitro, and tumor growth also was suppressed in vivo. In addition, we also found that LV-COX-2 combination with TAM treatment could inhibit angiogenesis and VEGF expression. Taken together, our experimental results indicate that LV-COX-2 combination with TAM has promising outcome in anti-metastatic and apoptotic studies. Furthermore, these results showed that LV-COX-2 combination with TAM is a potential drug candidate for treatment of breast tumors expressing high levels of VEGF.

Introduction

Breast cancer is the most common female cancer and ~70-75% of cases express oestrogen receptor α (ERα) (1). Tamoxifen (TAM), a selective estrogen receptor (ER) modulator, has been used extensively in the clinical management of primary and advanced breast cancer and is also widely employed as a preventive agent after surgery for breast cancer (2). A larger number of clinical studies over the past 30 years have shown that TAM can reduce the incidence and regression of breast carcinoma among women worldwide (3). Despite the relative safety and significant anti-neoplastic activities of tamoxifen, most initially responsive breast tumors develop resistance to its (4). In addition, TAM is also known to increase the expression of vascular endothelial growth factor (VEGF), which is an undesirable effect in breast cancer treatment (5,6). Extensive studies have shown that VEGF, a receptor tyrosine kinase, plays an important role in tumorigenesis, and blocking the VEGF signaling pathway can reduce tumor-associated angiogenesis and blood vessel-dependent metastasis (7,8). Therefore, combination therapies of tamoxifen with an anti-VEGF signaling agent aimed at inhibition of both ER-mediated signaling and VEGF stimulated stromal activation, may be a potential means of delaying the arrival of resistance.

Cyclooxygenase-2 (COX-2) is an inducible gene, whose expression is undetectable in most normal tissues. Accumulating evidence shows that overexpression of COX-2 was commonly related to different types of carcinomas, including breast carcinoma (9-12). COX-2 has been implicated in breast tumorigenesis based on its increased expression in a significant fraction of breast carcinomas and the protective effects of nonsteroidal anti-inflammatory drugs (NSAIDs) against breast cancer (13). Hoeben et al (14) have indicated that COX-2 overexpression is correlated with induction of VEGF expression and therefore tumor angiogenesis in human breast cancers. Inhibition of COX-2 by non-steroidal anti-inflammatory drugs (NSAD) leads to restricted angiogenesis and downregulates production of VEGF (15). Although accumulating evidence suggests that NSAIDs, such as celecoxib sensitize cancer cells to ATM-induced apoptosis, it remains...
unclear whether it is COX-2 dependent or independent (16). Furthermore, NSAIDs could result in liver injury, which would limit their clinical applications (17,18).

Silencing of COX-2 by RNAi would avoid the side effects induced by NSAIDs. Accumulating evidence shows that down-regulation of COX-2 expression by RNAi inhibited tumor cell proliferation and colony formation in vitro in different types of cancer cells (12,19,20). COX-2 silencing abolished the metabolic potential of highly malignant breast cancer cells (10). In addition, Lin et al documented that combining gene therapy with various drugs enhances transduction efficiency resulting in enhanced tumor cell killing (21). Therefore, in this study, we knocked out the COX-2 gene via a replication-incompetent lentivirus (LV-COX-2), then LV-COX-2 combination with TAM for suppressing VEGF expression and simultaneously reducing doses of ATM.

The objective of the present study was to evaluate the potency of LV-COX-2 in combination with TAM in inhibiting breast cancer cell growth, proliferation, and angiogenesis in vitro and reveal the underlying molecular mechanisms involved in TAM-induced apoptosis. In addition, tumor growth ability in nude mice was detected to define the combination treatment effect in tumorigenesis in vivo.

Materials and methods

Cell culture. Human breast cancer cell lines MCF-7 and HEK293 (human embryonic kidney cells) were bought from the Shanghai Cell Collection (Shanghai, China). Cells were cultured in Dulbecco's modified Eagle's medium (DMEM; Gibco-BRL, Grand Island, NY, USA) supplemented with 10% heat-inactivated fetal bovine serum (FBS; Gibco-BRL) at 37°C in a 5% CO₂ atmosphere and at 95% humidity.

Reagents. Stock solutions of 10 mM TAM (Sigma-Aldrich, St. Louis, MO, USA) were dissolved in dimethylsulfoxide (DMSO, Sigma-Aldrich, St. Louis, MO, USA) at 20°C, and diluted in fresh medium just before use. TRIZol reagent kit and Coomasie Blue R-250 from Gibco-BRL, Invitrogen Corp. (Carlsbad, CA, USA); Nonidet P-40 lysis buffer, chemiluminescent peroxidase substrate, propidium iodide (PI), 4',6-diamidino-2-phenylindole (DAPI), 3-(4,6-diamidino-2-yl)-2,5 diphenyltetrazolium bromide (MTT), and sense and antisense VEGFR2 oligo primers from Sigma-Aldrich; and pyrogallol and H₂O₂ from Merck (Whitehouse Station, NJ, USA). Stock solutions of PI, DAPI, and MTT were prepared by dissolving 1 mg of each compound in 1 ml of phosphate buffered saline (PBS). The solution was protected from light, stored at 4°C, and used within 1 month.

Animals. Female BALB/c nude mice aged 4-5 weeks were obtained from the Experimental Animal Center of the Jilin University (Changchun, China). All procedures were performed according to institutional guidelines and conformed to the National Institutes of Health guidelines on the ethical use of animals.

Generation of plasmids and recombinant lentivirus. To inhibit the expression of COX-2, a short hairpin RNA (shRNA) targeting the COX-2 transcript was designed. The synthesized oligonucleotides containing specific target sequence, a loop, the reverse complement of the target sequence, a stop codon for U6 promoter and two sticky ends were cloned into pGCSIL-GFP lentivirus vector according to the instructions (Shanghai Gene Chem Co. Ltd., China). The targeting sequences corresponding to the siRNAs for COX-2 (GeneBank accession no. NM_000963.2) was as follows: bases on 290-310 (sense 5’-AAACTGCTCAACACCCGAATT-3’). Sequences for the scrambled control (NC) for siRNA are AATTCTCCGAAAC TGTCAACGT (sense). This sequence does not target any gene product and have no significant sequence similarity to human gene sequences, being essential for determining the effects of siRNA delivery.

To produce the lentivirus, the HEK293 cells were transfected with 20 µg of LV-COX-2, and LV-NC plasmid together with 15 µg of pHelper-1.0 and 10 µg of pHelper-2.0 packaging plasmids, respectively (22). The culture medium was collected within 48 h after transfection, concentrated by ultracentrifugation, aliquoted, and stored at -80°C until used. The titer of lentivirus was determined by hole-by-dilution titer assay as described (23). Four days after a single exposure of HEK293 cells to the lentivirus, strong green fluorescence was observed in >90% of cells, indicating a high and stable transduction of TAM.

Materials and methods

RNA isolation and real-time RT-PCR. Total RNA was extracted from cultured cells using TRIZol reagent (Invitrogen) according to the manufacturer’s instructions. RNA was reverse-transcribed into cDNA by a Primerscript™ RT reagent kit according to the manufacturer’s protocols (Takara, Japan). Quantitative real-time polymerase chain reaction (RT-PCR) assays were carried out using SYBR Green Real-Time PCR Master Mix (Toyobo, Osaka, Japan) and RT-PCR amplification equipment using specific primers: COX-2, sense strand 5’-CCTTTGGGT TGTTTAAATTAA-3’, antisense strand 5’-AAACTGATGTC GTGAATTCTG-3’, β-actin, sense strand 5’-GGAGAGCACAGACCTCGCCTTTG-3’, antisense strand 5’-GATGGCCGGTGCTCAGTTA-3’. The PCR conditions were as follows: a pre-denaturing at 95°C for 3 min, followed by 40 cycles of denaturation at 95°C for 10 sec, annealing/extension at 58°C for 20 sec, final extension of 72°C for 10 min. The amplification specificity was checked by melting curve analysis. The 2-ΔΔCT method was used to calculate the relative abundance of target gene expression generated by Rotor-Gene Real-Time Analysis Software 6.1.81. The expression of interest genes were determined by normalization of the threshold cycle (Ct) of these genes to that of the control β-actin.

Western blot analysis. Cultured cells were washed twice with phosphate-buffered saline (PBS, pH 7.2), then cells were lysed with Triton X-100 in HEPES buffer (150 mM NaCl, 50 mM HEPES, 1.5 mM MgCl₂, 1% Triton X-100, 0.1% SDS, protease inhibitor cocktail (Sigma), 100 mM NaF and 100 mM Na₂VO₃) for 30 min. Cell lysates were clarified by centrifugation (10,000 g, 15 min), and protein concentrations were determined using the Bradford reagent (Sigma, Germany). Protein samples were separated on an 8-15% SDS-polyacrylamide gel (SDS-Page) and transferred onto nitrocellulose membranes (Santa Cruz Biotechnology, Inc., Santa Cruz, CA, USA),
followed by blocking in TBS containing 5% skim milk and 0.1% Tween-20 for 2 h at room temperature, and then immunoblotted with specific primary antibodies and incubated with corresponding horseradish peroxidase-conjugated secondary antibody. The other primary antibodies used in the western blot analyses were: anti-VEGF, anti-VEGFR, anti-COX-2 (all Cell Signaling Technology, Beverly, MA, USA), mouse monoclonal anti-β-actin (Sigma-Aldrich), and mouse monoclonal anti-caspase-3, mouse monoclonal anti-caspase-8 (Santa Cruz Biotechnology). Secondary Abs used for immunodetection were: HRP-conjugated goat anti-mouse IgG and goat anti-rabbit IgG (Amersham Biosciences, Uppsala, Sweden). All immunoblots were visualized by enhanced chemiluminescence (Pierce).

Proliferation assays. To measure the effect of LV-COX-2 and ATM alone or both on cell proliferation, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay was used. MCF-7 cells grown in monolayers were harvested and dispersed in 96-well culture plates in 100 µl of DMEM at a concentration of 5x10^4 cells per well. After 24 h, cells without any treatment, cells treated only with LV-NC, and cells transfected with LV-COX-2, cells treated only with ATM, cells treated with ATM combination with LV-COX-2 were seeded in 96-well plates and left at 37˚C for 48 h. The cells were then washed with PBS and incubated in 50 µl of 0.5 mg/ml MTT in culture medium at 37˚C for 4 h. Following the addition of 100 µl of isopropanol, the absorbance was read at 595 nm in an ELISA plate reader (Molecular Devices Corp., Sunnyvale, CA, USA). The mean proliferation of cells without any treatment was expressed as 100%.

Detection of apoptosis. MCF-7 cells were cultured in 6-well plates in DMEM with 10% FBS and medium were treated with LV-NC, LV-COX-2, ATM and ATM combination with LV-COX-2, respectively, for 48 h. The cover slips were washed three times with phosphate-buffered saline (PBS) and single cell suspensions were fixed in 1% PBS. Cells were stained with 100 µg/ml acridine orange (AO) and 100 µg/ml ethidium bromide (EB) for 1 min. Then cells were observed under a fluorescence microscope. At least 200 cells were counted and the percentage of apoptotic cells was determined. In addition, we also detected caspase-3 and -8 protein expression by western blotting as an additional indicator of apoptosis.

Cell cycle analysis. To determine the cell cycle distribution, 5x10^5 MCF-7 cells were plated in 60-mm dishes and treated with LV-NC, LV-COX-2, ATM and ATM combination with LV-COX-2, respectively, for 48 h. After treatment, the cells were collected by trypsinization, fixed in 70% ethanol, and kept at -20˚C overnight for fixation. Cells were washed in PBS, resuspended in 1 ml of PBS containing 100 µg/ml RNase and 40 µg/ml PI and incubated in the dark for 30 min at room temperature. The distribution of cells in the cell cycle phases were analyzed from the DNA histogram with a FACSCaliber flow cytometer (Becton-Dickinson, San Jose, CA, USA) and CellQuest software (CA, USA).

Cell migration assay. The migration assay was performed using a 12-well Boyden chamber (Neuro Probe) with an 8-µm pore size. Approximately 1x10^5 cells were seeded into upper wells of the Boyden chamber and incubated for 6 h at 37˚C in medium containing 1% FBS. Medium containing 10% FBS was used as a chemoattractant in the bottom wells. Cells that did not migrate through the pores of the Boyden chamber were manually removed with a rubber swab. Cells that migrated to the lower side of the membrane were stained with hematoxylin and eosin and photographed using an inverted microscope.

Wound-healing assay. To assess the effect of TAM and LV-COX-2 on cell migration, wound-healing assay was performed. MCF-7 cells (1x10^5) were plated in 12-well plates in complete growth medium. After 24 h of growth, a scratch was made through the confluent cell monolayer, and then the cells were treated with LV-COX-2, LV-NC, TAM, TAM combination with LV-COX-2, respectively, in 3 ml of complete medium. At 48 h post-treatment, cells were stained with hematoxylin and eosin (HE). Cells invading the wound line were observed under an inverted phase-contrast microscope using 20x, Leica DMR, Germany.

Cell invasion assay. Cell invasion was determined using transwell chambers made from polycarbonate membrane filters with a pore size of 8 µm. Transwell filters in 6-well plates were coated with Matrigel, hydrated for ~2 h in the tissue culture incubator with 500 µl serum-free culture media in the bottom and 500 µl in the top of the chamber. After hydration of the Matrigel, 5x10^5 MCF-7 cells were plated in 500 µl serum-free medium on top of chamber, while 2 ml medium 10% FCS were placed in the lower chambers. TAM, LV-NC, LV-COX-2, LV-COX-2 combination with ATM were added to the upper chambers, respectively. Cells without any drug were used as control. After 48 h of incubation, the filters were removed, washed twice in PBS and fixed in 10% formalin for 15 min. After fixing at room temperature, the chambers are rinsed in PBS and stained with 0.2% crystal violet staining solution for 30 min. After washing the chambers by PBS, the cells at the top of the Matrigel membrane were carefully removed by cotton swabs. At this time all cells that remain had invaded to the bottom side of the membrane. Cell invasion was observed with an immunofluorescence microscope by counting the cells that had invaded into the bottom of the Cell Culture Insert. All experiments were performed in triplicate.

Measurement of prostaglandin-E2 (PGE2) production. PGE2 synthesis was determined by competitive enzyme-linked immunosorbent assay (ELISA) as previously described (24). In brief, MCF-7 cells were treated with LV-NC, LV-COX-2, ATM and ATM combination with LV-COX-2 respectively for 48 h in 12-well plates, and then these culture media were centrifuged to remove cell debris. Cell-free culture media were collected at indicated time, and then PGE2 levels were determined by competitive ELISA as described using the kit manufacturer (Cayman Chemical, Ann Arbor, MI, USA).

Measurement of VEGF levels. To measure VEGF levels, 5x10^5 MCF-7 cells were plated in 6-well plates and incubated under culture conditions overnight, and the medium was replaced by serum-free culture conditioned medium. LV-NC,
LV-COX-2, ATM and ATM combination with LV-COX-2, respectively, were added to the culture, and the medium was collected at 72 h. VEGF levels were measured using a VEGF enzyme linked immunosorbent assay (ELISA) kit (DVE00, R&D Systems, Minneapolis, MN, USA) according to the manufacturer's instructions. The optical density at 570 nm of each well was measured using an ELISA reader (µQuant; Biotek Instruments, Inc., Winoski, VT, USA). In addition, we also detected VEGF and VEGFR protein expression level by western blotting.

Tumor xenograft in nude mice. Exponentially growing MCF-7 cells were harvested and a tumorigenic dose of 2x10^6 cells was injected intraperitoneally into 4-5-week-old female BALB mice. When tumors reached 100-200 mm^3, mice were divided randomly into five groups (10 mice per group). The control group received 1% polysorbate resuspended in deionized water. The other four groups were treated with LV-NC (1x10^8 PFU/dose), ATM (2 mg/kg body weight), LV-COX-2 (1x10^8 PFU/dose), or ATM plus LV-COX-2 (ATM, 1 mg/kg body weight; LV-COX-2, 1x10^8 PFU/dose respectively) intraperitoneally on alternative days for 3 weeks. The tumor size was measured using caliper before the treatment injections were given, and on day 7, 14 and 21 of treatment. On day 21, the animals were euthanized using chloroform and their spleen tissues were collected and cultured for a splenocyte surveillance study.

Assay of splenocyte proliferation. Spleens from treated mice were collected, and single-cell spleen suspensions were pooled in serum-free RPMI-1640 by filtering the suspension through a sieve mesh with the aid of a glass homogenizer to exert gentle pressure on the spleen fragments. The detail assay of splenocyte proliferation was previously described (16).

Statistical analysis. Data from at least three independent experiments are expressed as mean ± SD. Statistical comparison of more than two groups was performed using one-way ANOVA followed by a Tukey post hoc test. Statistical analyses were undertaken using GraphPad Prism version 5.01 (GraphPad Software, San Diego, CA, USA) and the SPSS® statistical package, version 16.0 (SPSS Inc., Chicago, IL, USA) for Windows®. P<0.05 was considered statistically significant.

Results

Downregulation of COX-2 expression by LV-COX-2. To silence COX-2 expression, we constructed a recombinant lentiviral carrying a based shRNA against COX-2, LV-COX-2, to inhibit COX-2 expression in breast carcinoma cell lines. To evaluate the silencing capacity of LV-COX-2, MCF-7 were treated with LV-COX-2, LV-NC as negative controls. Real-time RT-PCR and western blotting were performed to detect the COX-2 mRNA expression and protein expression levels at 3 days posttransfection. Our results showed no significant inhibition in COX-2 mRNA expression in LV-NC group, there is no significantly difference in LV-NC group and control group (PBS group) (P>0.05). On the other hand, COX-2 mRNA expression in LV-COX-2 group was significantly decreased compared to LV-NC group and control group after injection (Fig. 1A, p<0.01). On protein level, there was no significant inhibition in COX-2 protein expression found in LV-NC group and control group (P>0.05), while the band density decreased dramatically in the LV-COX-2 group as compared with the LV-NC and control group (P<0.01) (Fig. 1B). These results demonstrated that LV-COX-2 significantly decreased COX-2 protein expression in the breast cancer cell line (P<0.01).

Effects of ATM and LV-COX-2 alone or combination on MCF-7 cell proliferation and the cell cycle. To evaluate the effect of LV-COX-2 and ATM alone and both on the cell viability of breast cancer cells in vitro, MTT assay was performed for 48 h when MCF-7 were treated with LV-COX-2 and ATM alone or combination. It was found that the inhibitory rates of LV-COX-2 and ATM alone or combination treatment were higher than control group and LV-NC treatment group (P<0.01). There is no significance different between LV-NC group and control group (P>0.05). In addition, the inhibitory rates of LV-COX-2 in combination with ATM group were higher than LV-COX-2 group and ATM treatment group (Fig. 2A).

The effects of LV-COX-2 and ATM on the cell cycles of MCF-7 cells were then analyzed by flow cytometry. MCF-7 cells treated with LV-COX-2 or ATM had an increased...
percentage of arrest at the G0/G1 phase compared with untreated cells and LV-NC control (Fig. 2B). The LV-COX-2 combination with ATM resulted in an even greater percentage of arrest at the G0/G1 phase than the higher doses of either drug alone (P<0.01).

Effects of ATM and LV-COX-2 alone or combination on MCF-7 cell apoptosis. To investigate whether the LV-COX-2 and ATM alone or combination could induce of apoptosis, we analyzed apoptosis after treatment with LV-COX-2 and ATM. It was found that MCF-7 cells treated with LV-COX-2 or ATM could significantly induce cell apoptosis compared with untreated cells and LV-NC control (Fig. 3A). Treatment with combination of LV-COX-2 and ATM led to a dramatic increase in apoptotic cells compare to single LV-COX-2 group and control group (P<0.01) (Fig. 3A).

To explore the possible mechanism of induction of cell apoptosis of combination with LV-COX-2 and ATM, expression patterns of caspase-3 and -8 were determined by western blotting. The results showed that combination with LV-COX-2 and ATM could significantly decrease the expression of apoptosis inhibiting genes, caspase-3 and -8, in MCF-7 cells, compared to LV-COX-2 or ATM alone (P<0.01) (Fig. 3B).

Effects of ATM and LV-COX-2 on MCF-7 cell migration and cell invasion. To ascertain the inhibitory effect of TAM and LV-COX-2 as a single or combined treatment on breast cancer migration, wound-healing assay was performed to investigate their effects on the migration potential of MCF-7 cells. After 48-h treatment, cells in the control group and LV-NC group efficiently spread into the wound area to such an extent that the wound boundary was not apparent, while only some cells

Figure 2. Effect of ATM and LV-COX-2 alone and in combination on cell proliferation and cell cycle of MCF-7 cells. Cell proliferation (A) and cell cycle (B) of MCF-7 cell was determined 48 h after treatment with ATM and LV-COX-2 alone or combination. *P<0.05 versus control, #P<0.05 versus ATM alone.

Figure 3. Effect of ATM and LV-COX-2 alone and in combination on cell apoptosis in MCF-7 cells. (A) Cell apoptosis of MCF-7 cell was determined 48 h after treatment with ATM and LV-COX-2 alone or combination; (B) Expression of caspase-3 and -8 in MCF-7 cell was determined by western blot analysis. Data are expressed as the means ± SD. *P<0.01 versus control, #P<0.01 versus ATM alone.
in TAM or LV-COX-2 treated group spread forward in MCF-7 cells. The cell migration in combination group was less than either drug alone (Fig. 4A).

The ability of TAM and LV-COX-2 to reduce the invasiveness of MCF7 cells was further investigated by the transwell system assay. It was found that invasion was also decreased...
significantly with TAM or LV-COX-2 treatment compared to control and LV-NC (P<0.01) (Fig. 4B). Compared with the results with either agent alone, the combination of TAM and LV-COX-2 greatly inhibited MCF-7 cell invasion.

**Effects of LV-COX-2 and ATM on PGE2 production and VEGF expression level in MCF-7 cells.** To examine the effect of ATM and LV-COX-2 alone or combination on PGE2 production in MCF-7 cells, ELISA was performed. As shown in Fig. 5A, LV-COX-2 single or combination with ATM inhibited PGE2 production. However, ATM increased PGE2 production compared to the other groups.

We also investigated the role of TAM and LV-COX-2 in the inhibition of secretory VEGF, a pro-angiogenic factor responsible for the migration and invasion of breast cancer cells. VEGF secretion in serum-free culture conditioned medium was assessed in MCF-7 cells by ELISA 24 h post-treatment. The results showed that TAM alone considerably upregulated VEGF secretion and the LV-COX-2 or LV-COX-2 combination TAM significantly decreased VEGF secretion compared with no treatment and LV-NC treatment (Fig. 5B).

To further study the possible mechanism of induction VEGF expression with LV-COX-2 and ATM, VEGF and VEGFR expression were determined by western blotting. The results showed that combination with LV-COX-2 and ATM or single LV-COX-2 could significantly decrease the expression of VEGF, VEGFR expression compared to control group and LV-NC group (P<0.01) (Fig. 5C and D). On the other hand, ATM dramatically increased the expression of VEGF, VEGFR expression compared to control group and LV-NC group (P<0.01) (Fig. 5C and D). These results implied that combination with LV-COX-2 and ATM could inhibit PGE2 production and VEGF secretion by inhibition VEGF and VEGFR expression.

**Antitumor activity of TAM and LV-COX-2 in nude mice bearing MCF-7 tumors.** We assessed the in vivo therapeutic efficacy of TAM and LV-COX-2 in female BALB mice bearing MCF-7 breast tumors cells. Mice were sacrificed and tumor tissue harvested 21 days after treatment. Then tumor weight of the animals was measured. Tumor weight of LV-COX-2 and ATM alone or combination group was lower than those of control group and LV-NC group (Fig. 6A). Compared with the results with either agent alone, the combination of TAM and LV-COX-2 greatly inhibited tumor growth. In addition, we found that tumor volume after treatment with LV-COX-2 and ATM was significantly slower for MCF-7 tumor cells compared with control group and LV-NC group (Fig. 6B). Treatment with combination of LV-COX-2 and ATM led to a dramatic inhibition of tumor growth compared to single LV-COX-2 group and control group (P<0.01) (Fig. 6B). These results indicate that LV-COX-2 combination with ATM treatment in breast cancer cells markedly suppresses their tumorigenicity in mice.

We also assess the efficacy of LV-COX-2 combination with ATM in modulating splenocyte proliferation using MTT assay. As shown in Fig. 6C, the inhibitory rates of LV-COX-2 and ATM alone or combination significantly increased compared to control group and LV-NC group (P<0.01). The inhibitory rates of LV-COX-2 combination with ATM treatment group were higher than single drug groups, which demonstrated that combination treatment could inhibit MCF-7 cell proliferation.

**Discussion**

Estrogen can cause a general upregulation of genes regulating cell proliferation and survival and the downregulation of genes with anti-proliferative or pro-apoptotic activity resulting in growth stimulation and apoptosis suppression (25). Therefore, anti-estrogens are able to decrease cancer cell proliferation and induce cell death signaling pathways (26). Consequently, tamoxifen (TAM), a selective estrogen receptor (ER) modulator, has been described as ‘the most important drug developed in the history of breast cancer’ (27) and can induce cell cycle arrest leads to an accumulation of cancer cells in G0/G1 phase of the cell cycle (28) and induce apoptosis of breast cancer cells (29). However, a large number of initial clinical studies of TAM displayed its anti-angiogenic and VEGF reducing ability in various tumor models (30-33). In addition, an undesirable response leading to enhanced metastasis and angiogenesis and resulting in inferior outcomes, when prolonged administration of TAM...
causes intracellular VEGF levels to rise in patients (34). In this context, we made an attempt to decrease intracellular VEGF levels by decreasing ATM dose in breast cancer cells. Furthermore, recent studies showed that siRNA-mediated knockdown of COX-2 expression inhibited VEGF expression (35). Therefore, in this study, we knocked out COX-2 gene via a replication-incompetent lentivirus (LV-COX-2), then LV-COX-2 combination with TAM for reducing doses of ATM. The result showed that this novel combination inhibited COX-2 expression, downregulated the level of PGE2 and decreased the production of vascular endothelial growth factor (VEGF) in tumors.

A recent study showed that the combination of TAM and CXB at nontoxic levels exerts potent anti-angiogenic effects by decrease in VEGF expression (16), which was in agreement with our results. However, the celecoxib analogs also faces the gastrointestinal side effects (36) and tolerance as observed in a rat model of inflammatory pain (37). Furthermore, celecoxib could result in liver injury (17,18). These side effects would limit their clinical applications. In this study, we selected LV-COX-2 combination with ATM avoiding the side effects induced by celecoxib.

Extensive studies showed that shRNA and transgene carried by lentiviral vector can be transcribed into precursor mRNA where shRNA is embedded in the 3'UTR or in the intron of transgene (38-40). It has been shown that RNA combination anticancer drug could achieve better antitumor activity (21). Luni et al demonstrated that when melanoma differentiation associated gene-7 (mda-7) via a replication-incompetent adenovirus (Ad.mda-7) and gefitinib are used in combination they might provide an effective therapeutic approach for NSCLC since these agents target multiple cell survival pathways and are equally effective against NSCLC cells (41). In the present study, we used a novel strategy of combining a gene therapy approach, LV-COX-2, with ATM, and evaluated its antitumor effects against breast cancer cells. The result showed that LV-COX-2 combination with TAM in breast cancer cells significantly suppressed the proliferation and metastasis, and induced tumor apoptosis in vitro, and tumor growth also was suppressed in vivo.

In conclusion, our in vitro studies demonstrate that when LV-COX-2 and ATM are used in combination they might provide an effective therapeutic approach for breast cancer since these agents target induction breast cell apoptosis and are equally effective against breast cancer cells. Additionally, further studies with in vivo tumor models also confirmed that LV-COX-2 combination with ATM could suppress breast tumor growth. Therefore, it may be worthwhile to consider the combination treatment as novel therapeutic strategy for further evaluation in clinical trials.

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