Abstract. For estrogen-responsive B-1F cells, established from estrogen-responsive mouse Leydig cell tumor, it has been reported that the 5-lipoxygenase (5-LOX) metabolic pathway appears to be associated with cell growth. The addition of 5-LOX inhibitor 2-(12-hydroxydodeca-5,10-diyl)-3,5,6-trimethyl-1,4-benzoquinone (AA861) to the medium resulted in a dose-dependent increase in cell yield as described previously. When the growth of the palpable tumors was measured, AA861 had stimulated in vivo tumor growth in adult male mouse inoculated B-1F cells. The effects of AA861 and 17ß-estradiol (E2) on the contents of various arachidonic acid metabolites in B-1F cells and their conditioned medium were examined. Although AA861 and E2 decreased the contents of leukotrienes (LTs), the two did not significantly change those of prostaglandins, thromboxan, prostacyclin, 12-hydroxyeicosatetraenoic acid (HETE) and 15-HETE. In immunohistochemical study B-1F cells show positive staining for 5-LOX in the E2-depleted condition, while E2 decreased the expression of 5-LOX. The decrease of the intensities of 79-kDa 5-LOX protein and 403-bp RT-PCR product bands was observed. The growth of Morpholino-anti oligo delivered B-1F cells was higher than that of Standard control oligo delivered cells. The delivery of Morpholino-anti oligo into B-1F cells caused the decrease of contents of LTs and 5-HETE in the cells and medium, and the reduction of 5-LOX activity. When LTD4 was added in the culture medium, the increasing concentrations of LTD4 resulted in a significant inhibition of cell yields of E2-treated B-1F cells. Morphological changes such as nuclear condensation and fragmentation, and DNA ladder pattern were demonstrated in E2-stimulated B-1F cells treated with LTD4 as well as in control cells cultured in the basal medium. These results implicate that 5-LOX at least plays an important role in the growth of B-1F cells and LTD4 induces the apoptosis of B-1F cells.

Introduction

Tumor cells are derived from cells which constitute the body and retain their original features to various degrees. In a physiological condition, in the target cells of hormones, increase or decrease of cell population is caused by hormonal stimulation. When the stimulation disappears, the cell population returns to basal levels. Endometrial and mammary glands are good examples of this. In these tissues, the growth of the target cells is regulated in response to menstrual cycle and hormonal state in pregnancy. In the growth of tumor cells which originate from these target cells, hormones play important roles and thus they are called hormone-dependent tumors. Mammary and endometrial cancers are representative examples of the female sex hormone-dependent tumors. The representative of the androgen-dependent tumor is prostate cancer. The cell line B-1F was established from estrogen-responsive mouse Leydig cell tumor (T124958-R) in our laboratory in order to study the mechanism of steroid hormone-dependent cancer growth (1). 17ß-Estradiol (E2) stimulated the growth of B-1F cells in a dose-dependent manner, the maximum effect being at a range of 10^{-8}~10^{-7} M (1). When the growth of the palpable tumors was measured to delineate the in vivo proliferative behavior of B-1F cells inoculated s.c. into adult male mice, E2 had significantly stimulated the tumor growth (1).

There are three major metabolic pathways of arachidonic acid, the cyclooxygenase (COX) pathway, lipoxygenase (LOX) pathway and cytochrome p450 pathway (2,3). It is known that NSAIDs (nonsteroidal anti-inflammatory drugs) affect the development of colorectal cancers (4-6) and familial
adenomatous polyposis (5,6). The major target and mechanism of NSAIDs' effects on the cancers were thought to be cyclooxygenase-2 (COX-2). However, the involvement of other molecular targets besides COX-2 has now been shown (7-12). For estrogen-responsive B-1F cells, we have reported that the 5-lipoxigenase (5-LOX) metabolic pathway appears to be associated with cell growth (13-19). Moreover, there are reports of a relationship between 5-LOX metabolites and the growth of prostate and pancreas cancers (20,21). Hormone-dependent cancers are characteristic in the control of their growth by hormones. It was expected that the knowledge obtained from the study of hormone-dependent cancers would be useful for the development of treatment and prevention of not only hormone-dependent cancers but also other cancers. Therefore, additional studies were conducted on the role of 5-LOX in the estrogen-responsive growth of B-1F cells.

Materials and methods

Cell culture. B-1F cells were maintained in Eagle’s minimum essential medium-Ham’s F12 (1:1, vol/vol) containing 0.1% (wt/vol) bovine serum albumin (BSA) (HMB medium, basal medium) supplemented with 10⁻⁸ M E2 (HMBE medium) in a humidified atmosphere of 5% CO₂ at 37°C. Cells were grown to confluence and passed with the use of trypsin (0.01%, wt/vol)-EDTA (0.02%, wt/vol) in phosphate-buffered saline (PBS).

Cell growth experiment. The effects of appropriate treatments on the cell yield or DNA synthesis were examined with slight modification as previously published (15,18,19). Briefly, cells were plated into four replicate 35-mm dishes at an initial density of 1x10⁴ cells/dish in 2 ml of HMB medium. On the following day (day 0), the medium was replaced with 2 ml of HMB medium with or without the appropriate treatments. Test compounds were dissolved as directed in the manual on day 6, unless otherwise specified. [Methyl-1',2',3'H]-thymidine incorporation assay was also performed as described previously (13,18). B-1F cells were plated onto a 96-well plate at an initial density of 4x10⁴ cells/well in 0.15 ml of HMB medium, and on the next day (day 0) the medium was replaced with 0.15 ml of HMB medium with or without test compounds. On day 3, the cells were pulsed with [³H]-thymidine (0.15 μCi/0.15 ml per well) for 2 h at 37°C in a humidified incubator. The incorporation of radioactivity into the cells that were briefly trypsinized and collected with a Micro96 Harvester (Molecular devices, USA) was measured.

In vivo growth. The male Balb/c mice (seven-week-old) were obtained from SLC Co. (Shizuoka, Japan), housed 5/cage in a temperature-controlled (25°C) room with a 12-h light/dark cycle, and given ad libitum access to water and powdered diets. After being harvested and washed, B-1F cells (5x10⁵) dispersed in MEM were injected subcutaneously (s.c.) into castrated male Balb/c mice. On the following day (day 0) mice were subcutaneously implanted with a fused pellet of 5% E2 in cholesterol or started on diets supplemented with or without 0.615% (w/w) 2-(12-hydroxydodeca-5,10-diyl)-3,5,6-trimethyl-1,4-benzooquinone (5-LOX selective inhibitor, AA861) (22). Palpable tumor was measured with a caliper at the indicated times. The tumor volume (v) was then calculated by the formula: v=4/3π(a/2xb/2)³, where a and b are the largest and the smallest diameter of the tumor, respectively.

Immunohistochemistry for 5-LOX. Immunohistochemistry for 5-LOX was performed using a TSA™ (Tyramide Signal Amplification) Plus DNP system as directed in the manual from the supplier.

B-1F cells were grown in four-well chamber slides (Falcon) at an initial cell density of 2x10⁴ cells/well in HMB medium. On the following day (day 0), the medium was changed to HMB medium with or without E2. On day 4, the medium was aspirated, and cells were fixed in 4% paraformaldehyde in 0.1 M phosphate buffer at 4°C for 30 min, and washed in three changes of PBS, followed by freezing-thawing. The chamber slides were incubated for 15 min in 0.3% hydrogen peroxide in methanol to quench endogenous peroxidase activity and washed three times in PBS containing 0.01% (v/v) Tween-20 (PBS-T) for 5 min each. Following rinses, for antigen retrieval before the immunostaining, the slides were immersed in a container with 0.01 M citrate buffer, pH 6.0, and microwaved for 10 min (600 W, 5 min, twice). The container with slides was then cooled for approximately 20 min at room temperature. After rinsing three times in TMB buffer (0.1 M Tris/HCl, pH 7.5, 0.15 M NaCl, 0.05% Tween-20) for 5 min each, the slides were incubated for 30 min in TMB buffer (0.1 M Tris/ HCl, pH 7.5, 0.15 M NaCl, 0.5% blocking reagent supplied in kit). Then, the slides were incubated with primary antibody (final 1000-fold dilution) against mouse 5-LOX for 60 min at room temperature. Primary antibody was incubated for 60 min at 37°C with Envision+/ Peroxidase™ (mouse) in order to form a primary-secondary antibody complex prior to use. After washing three times for 5 min each with TMB buffer the slides were incubated with dinitrophenyl (DNP)-labeled amplification reagent for 5 min at room temperature followed by washing three times for 5 min each with TMB buffer. The slides were incubated with anti-DNP-HRP (horseradish peroxidase) in TMB buffer for 30 min at room temperature, washed three times for 5 min each with TMB buffer and then incubated with diaminobenzidine tetrahydrochloride (DAB) for 5 min at room temperature until the desired stain intensity developed. The slides were counterstained in hematoxylin for 5 sec, washed with tap water, dehydrated through alcohols and xylene, mounted, and observed by light microscopy. For the negative control, the primary antibody was replaced with isotype-matched negative control antibody (Dako), which showed no specific reactivity.

Western blotting. B-1F cells were plated on culture dishes at a density of 5x10⁴ cells/100-mm dish in HMB. On the following day (day 0), the medium was changed to HMB or HMBE. On day 4, B-1F cells were scrapped into lysis buffer (PBS, 2 mM sodium vanadate 1.0 mM sodium fluoride, 1% NP40, 0.5% sodium deoxycholate, 0.1% SDS, 100 μg/ml
Ethoxylated polyethylenimine (EPEI) with Morpholino/DNA into the cytosol/nuclear compartment of the cells using the supplier. The special delivery system delivers M-oligos into B-1F cells was performed as directed in the manual from the supplier. The supernatants were recovered and their protein concentrations were measured using Bio-Rad protein assay reagent. The supernatants (~100-500 μg protein) were immunoprecipitated with anti 5-LOX mouse monoclonal antibody. For immunoblotting, equivalent amounts of immunoprecipitates were separated by SDS-PAGE and transferred to PVDF membranes by electoblotting using a Bio-Rad semidy requirement transfer blotting apparatus (Nippon Eido Co.) The membranes were subsequently blocked in PBS-T with 5% nonfat milk overnight at 4˚C and then incubated with anti 5-LOX mouse monoclonal antibody diluted 1:100 in PBS-T for 1 h at room temperature. After incubation the membranes were washed in PBS-T and then incubated with horseradish peroxidase-conjugated anti-mouse secondary antibodies [Envision+/Peroxidase™ (mouse)] at a dilution of 1:1000 for 1 h at room temperature. The membranes were washed, detected by chemiluminescence (ECL reagent kit) and exposed to X-ray film for 1-5 min.

**RT-PCR analysis.** B-1F cells were plated on culture dishes at a density of 2x10^6 cells/100-mm dish in HMB. On the following day (day 0), the medium was changed to HMB or HMBE. On day 4, total RNA was isolated from B-1F cells using the TRIzol reagent (Invitrogen), according to the manufacturer's protocol. RT-PCR was performed as directed in the manual from the supplier (Qiagen OneStep RT-PCR kit) using specific primers: 5-LOX, 5'-ATGGATGGAGTG AACCCCCG-3' (upstream) and 5'- CTGTAACCTCCTGTTCAAAACT-3' (downstream); and GAPDH (32 cycles), 5'- ATCTTACAGGAGGGACAGCCCA-3' (upstream) and 5'-TCC ACAAAGTTGTCATG-3' (downstream). The RT profile for 5-LOX was 50˚C for 30 min and 95˚C for 15 min. The PCR profile for 5-LOX was 94˚C for 45 sec, 46˚C for 45 sec, and 72˚C for 1 min 30 sec for 25 cycles. The PCR profile for 5-LOX was performed twice. The RT profile for GAPDH was 42˚C for 30 min and 95˚C for 10 min. The RT-PCR profile for GAPDH was 94˚C for 45 sec, 55˚C for 45 sec, and 72˚C for 2 min for 35 cycles.

The final PCR products were separated on a 2% agarose gel with ethidium bromide and visualized under UV light. Negative control for 5-LOX RT-PCR that contained no reverse transcriptase showed no PCR products (data not shown).

**Delivery of Morpholino oligonucleotide into B-1F cells.** Morpholino antisense oligonucleotides (M-anti oligos) and standard control oligos (Std. Ctl. oligos) fluorescent at the 3’ end were purchased from Gene Tools, LLC (Philomath, OR, USA). The sequence of M-anti oligos against mouse 5-LOX mRNA, located at the translation initiation site is 5'-CCACGGTGACAGTGAGGGAGCAT-3' and the sequence of standard control oligos (Std. Ctl. oligos) is 5'-CCTCTTACCTCAGGATAATTATA-3'. M oligos were dissolved in water at a concentration of 0.5 mM (stock solution). The delivery of M-anti or standard control oligos into B-1F cells was performed as directed in the manual from the supplier. The special delivery system delivers M-oligos into the cytosol/nuclear compartment of the cells using Ethoxylated polyethyleneimine (EPEI) with Morpholino/DNA duplexes. After incubation with the complete delivery solution containing M-anti or standard control oligos for 3 h, 1x10^6 or 2x10^6 cells/dish of B-1F were plated in 35- or 60-mm dishes, respectively for cell growth or LT assay.

**Assay of arachidonic acid metabolites.** The contents of various arachidonic acid metabolites in B-1F cells and their conditioned medium were examined as previously described (13,15). Briefly, B-1F cells were seeded at a cell density of 1x10^6 cells/60-mm dish in 3 ml of HMB. On the following day (day 0), the various test compounds were added to the medium. After 2 days, the medium was replaced with fresh medium containing the same compound. After incubation for a further 7 h, the medium and cells were collected separately, and the viable cells were counted with a hemocytometer using the trypan blue dye exclusion method. The cells were washed in fresh medium and homogenized in 10 ml of ice-cold PBS. In this condition the cell yield per dish was not significantly different between cells irrespective of the presence of test compounds. Various metabolites were separated by reverse-phase HPLC. LTs present in the cells and medium were measured as described previously (13,15). In our system, the extraction efficiency for LTC4 was 60±4% (data not shown). The radioimmunoassay for LTs was performed as directed in the manual from the supplier, using LTB4 or LTC4 kits. The peak areas were estimated for other metabolites of arachidonic acid.

**Lipoxygenase assay.** The activities of lipooxygenases (5-LOX and 12-LOX) were measured by incubation of the cytosol with [^3H]-arachidonic acid. In these assays, B-1F cells were precultured in HMB for 1 day and then in fresh HMB containing the test compounds for 2 days. On the day of the assay further stimulation with the test compounds for 7 h was performed. Then, B-1F cells were twice homogenized in 7 vol of 20 mM Tris-HCl (pH 7.8 at 20˚C)-5 mM dithiothreitol with Polytron PT-10 at setting 5 for 10 sec and 10-sec cooling followed each burst. The homogenate was centrifuged at 105,000 x g for 60 min. For the assay of the 5-LOX activity, the aliquots of the cytosol were incubated with 10 μM [^3H]-arachidonic acid (2-3x10^6 dpm) in 50 mM potassium phosphate buffer at pH 7.4, 2 mM CaCl₂, 1 mM glutathione and 2 mM ATP at 30˚C for 5 min in a final volume of 0.2 ml. For the assay of the 12-LOX activity, the mixture contained 50 mM Tris-HCl buffer at pH 7.4, 25 μM [^3H]-arachidonic acid (2-3x10^6 dpm), and the aliquots of the cytosol in a final volume of 0.2 ml. Reactions were performed at 30˚C for 5 min. Hydroxyeicosatetraenoic acids (HETEs) that formed were separated using HPLC and the radioactivity in fractions for 5-HETE and 12-HETE was measured as described previously (13,15). In our system, the activity of 5-LOX was estimated as arachidonic acid release from B-1F cells. Arachidonic acid release from B-1F cells was measured as previously published (15).

**Detection of apoptosis.** B-1F cells were plated on culture dishes at a density of 5x10^6 cells/100-mm dish in HMB. On the following day (day 0), the medium was changed to HMBE with or without 10⁻⁸ M LTD₄. As a control, the medium was changed to HMB. The medium was changed again on day 3. On day 4, floating and attached cells in all dishes of the same
group were collected together and suspended in PBS. Aliquots of suspension were used to extract fragmented DNA as directed in the manual from the supplier of the ApopLadder EX™. Fragmented DNA extracted from approximately 1x10⁶ cells was electrophoresed on a 2% NuSieve (3:1) agarose gel (12x13 cm) for 2 h at 65 V. After electrophoresis, the gel was stained with ethidium bromide (0.5 μg/ml) for 10 min at room temperature, washed with distilled water and photographed.

For morphological examination B-1F cells were fixed in 2% glutaraldehyde in 0.1 M phosphate buffer at 4˚C for 2 h, stained with Hoechst 33342 (final concentration 0.167 mM) and examined by fluorescence microscopy.

Materials. Materials were obtained from the following sources: M-oligos from Gene Tools, LLC (Philomath); LTD4 from Cayman Chemical Co. (Ann Arbor, MI); BSA (crystallized and lyophilized), trypsin from Sigma Chemical Co. (St. Louis, MO); Eagle’s minimum essential medium, Ham’s F12 from Nissui Pharmaceutical Co. (Tokyo, Japan); [3H]-arachidonic acid, LTB4 and LTC4 [3H] RIA kits from DuPont/NEN Research Products (Wilmington, DE); the materials for HPLC from Waters Associates (Bedford, MA); anti-5-LOX monoclonal mouse antibody from Transduction Lab.; Envision+™ polymer (peroxidase, mouse), normal goat serum, mouse IgG1 from Dako; DAB from Nitirei Co.; TRizol reagent from Gibco; RNA OneStep RT-PCR kit from Quagen; NuPage Bis-Tris gel from Invitrogen; ECL Western Blotting Kit, [methyl-1'2'-3H]-thymidine from Amersham Biosciences Co. (NJ, USA); Bio-Rad Protein Assay reagent from Bio-Rad; TSA™ Plus DNP (HRP) System from NEN Life Science Products. The other reagents used were of analytical grade.

Statistics. The data were expressed as mean ± standard error (SE). Data were analyzed by ANOVA with Dunnett or Bonferroni's corrections for multiple comparisons, as appropriate. Calculated p values of less than 0.05 were considered to be significant.

Results

Possible linkage between E2-responsive growth and arachidonic acid metabolism. Arachidonic acid metabolism inhibitors were incubated with B-1F cells in the absence of E2. As previously described (13,14) and shown in Fig. 1, an addition of AA861 to the HMB medium resulted in a dose-dependent increase in cell yield at a range of 10⁻⁸ to 10⁻⁶ M. The phospholipase A2 inhibitor, p-bromophenacyl bromide (BPB) was examined for its growth-stimulatory activity. While BPB was less potent than AA861, it showed growth-stimulatory activity. The cyclooxygenase inhibitor, Indomethacin, did not elicit a significant effect on B-1F cell growth at a range of 10⁻⁸ to 10⁻⁶ M. To study the in vivo effect of AA861 on the B-1F cell growth, the cells were inoculated s.c. into intact adult male mice (Fig. 2). As positive control, mice were treated with E2 pellet. The incidence of tumor growth was not different between AA861-treated mice and non-treated mice (6 to 10 for AA861-treated mice, 6 to 10 for non-treated mice). When the growth of the palpable tumors was measured, AA861 stimulated the tumor growth. Although statistical significance was not shown in this experiment (Fig. 2), the other 2 separate trials also gave similar results.

Arachidonic acid metabolites in B-1F cells and their conditioned medium. The effects of AA861 and E2 on the contents of various arachidonic acid metabolites in B-1F cells and their conditioned medium were examined. As shown in
Table I and our previous report (13), the contents of LTs decreased to ~20% in the cells and 24% in the conditioned medium by 10^{-6} M AA861. E2 (10^{-8} M) suppressed the contents of LTs to ~27% in the cells and 26% in the conditioned medium. Although AA861 and E2 decreased the contents of LTs in B-1F cells, the two did not significantly change those of prostaglandins, thromboxan, prostacyclin, 12-HETE and 15-HETE in B-1F cells and their conditioned medium (Table I).

The expression of 5-LOX in B-1F cells. Immunohistochemical study for 5-LOX was carried out in E2-treated and E2-untreated B-1F cells. Since the activity of 5-LOX in B-1F cells is very low compared with stimulated leukocytes (data not shown), the detection of 5-LOX was difficult by a usual immunohistochemical detection method. Therefore, the TSA method was used in this experiment. B-1F cells show positive stain for 5-LOX under the E2-depleted condition, while E2 decreased the expression of 5-LOX (Fig. 3). Moreover, this E2-responsive decrease of the expression of 5-LOX was confirmed at the levels of protein and mRNA by Western blot analysis and RT-PCR for 5-LOX. Fig. 3 shows the significant decrease of the intensities of 79-kDa 5-LOX protein and 403-bp RT-PCR product bands.

Delivery of Morpholino oligonucleotide into B-1F cells. Gene function and control can be studied with the appropriate antisense oligo. M-anti oligo is a novel antisense structural type that belongs to RNAase-H independent oligonucleotides and exhibits higher antisense activity, higher water solubility, and more resistance to nucleases in target cells than the mostly used phosphorothioate oligonucleotides. These type of antisense oligonucleotides against 5-LOX mRNA were used to examine the role of 5-LOX in B-1F cell growth. The

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Table I. Arachidonic acid metabolites produced by B-1 cells after stimulation in the presence of various concentrations of Estrogen (E2), 10^{-6} M AA861 or 10^{-6} M Indomethacin (INDO) were measured. The results are expressed as % relative to control, unstimulated cells. ND, no detectable values. The data represent the average of at least two experiments, each performed in five replicates.
delivery of M-anti oligos into B-1F cells was confirmed by the intracellular fluorescence (data not shown). The growth of M-anti oligo delivered B-1F cells was statistically higher than Standard control oligo delivered B-1F cells (Fig. 4), or higher than B-1F cells incubated with only delivery solution. The delivery of M-anti oligo into B-1F cells caused the decrease of contents of LTs and 5-HETE in the cells and medium (Fig. 5). The activity of 5-LOX was lower in M-anti oligo delivered B-1F cells than in Standard control oligo delivered B-1F cells. The activity of PLA2, estimated by arachidonic acid release, was not significantly affected (Fig. 5).

Inhibition of B-1F cell growth and induction of apoptosis by LTD4. In order to confirm the role of LT in estrogen-responsive growth of B-1F cells, various concentrations of LTD4 were added in the culture medium and cell growth was examined after 6 days. As shown in Fig. 6A, the increasing concentrations of LTD4 resulted in a significant inhibition of cell yields of E2-treated B-1F cells. These results were similar to our previous report (15). The cells were stained with Hoechst 33342 and examined by fluorescence microscopy. Morphological changes, such as nuclear condensation and fragmentation, were observed in B-1F cells treated with LTD4. DNA fragmentation analysis was also carried out in B-1F cells cultured in the basal medium (Fig. 6C). These showed LTD4 induced apoptosis in B-1F cells even if in the presence of estrogen.

Discussion

E2 stimulated the growth of B-1F cells in a dose-dependent manner in vitro. B-1F cells inoculated s.c. into intact adult male mice showed similar estrogen-responsive behavior in vivo. The incidence of tumor growth was 50-60%. Histological analysis of the tumor demonstrated characteristics of malignant tumor such as many mitotic figures, and atypical cell and nuclear features. Our present results, together with our previous results,
implicate the important role of 5-LOX in estrogen-responsive growth enhancement of B-1F cells. The exogenous addition of LTs in the cultured medium led to a reduction of the proliferation and induced apoptosis of B-1F cells. It was considered that the inhibition of LT production by estrogen resulted in an enhancement of B-1F cell growth. One possible explanation for the inhibition of LT production by estrogen was the reduction of 5-LOX itself as shown in this experiment. As an additional possibility, catechol estrogens, the metabolites of estrogens, might modulate LT synthesis in B-1F cells, although the metabolism of E2 in B-1F cells was not examined in this study. In A23187-stimulated human whole blood, catechol estrogens are reported to be potent 5-LOX inhibitors (23). The reduction of the catalytically active ferric enzyme to the catalytically inactive ferrous form is shown as a possible mechanism for the regulation of lipoxygenase activity by catechol estrogens (23,24). Moreover, interestingly, a bioactive form of E2, such as estradiol-17β, is suggested to interact with the LTD4 receptor to induce granulocytic differentiation in myeloblasts (25).

Tumor growth is enhanced by the suppression of negative growth control and the stimulation of positive growth control. In B-1F cells, estrogen appears to demonstrate its growth promotion ability in part by the suppression of negative growth control. Probably, this mechanism is not the only one for estrogen's effects on B-1F cell growth, and there might be other pathways in which estrogen stimulates positive growth control.

Many reports on the role of LOX in tumor growth have shown the growth stimulatory potency of LOX products (26-29). Moreover, possible correlations are shown of cysteinyl leukotriene (1) receptor [CysLT(1)R] to COX-2, 5-LOX and Bcl-x(L) (30). An increased localization of CysLT(1)R in the nucleus of colorectal adenocarcinoma cells has been reported (31). The existence of constitutive CysLT(1)R signalling that mediates both survival and proliferation is revealed in intestinal cells, proposing that tumor-derived intestinal cells are resistant to CysLT(1)R antagonist-induced apoptosis, a phenomena that could be explained by nuclear CysLT(1)R (32). In other reports, LOX metabolites of arachidonic acid inhibited the growth (33-35). In addition to LOX metabolism of arachidonic acids, it has been reported that LOX metabolism of linoleic acids plays important roles in carcinogenesis (10). Reasons for these controversial results, including our results, are unknown. Since the estrogen-responsive element in the upstream of the mouse 5-LOX gene has not been recognized, the mechanism of the decrease of 5-LOX expression by E2 in B-1F cells is not clear. Further study might help to clarify these issues.

Acknowledgements

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References


