Establishment of a progesterone-sensitive cell line from human lung cancer

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Abstract. For development and function of the lung, progesterone (Prog) fulfils important roles. In a recent report, immunolocalization of Prog and estrogen receptors in non-small cell lung carcinomas were examined and it was shown that the Prog receptor might be a potent prognostic factor. In the present study, a cell line with the sensitivity to Prog was established from a human lung cancer and the growth mechanism was analyzed. The proliferation of established SN96-42 cells was sensitive to Prog and antiprogesterone RU38486 inhibited their proliferation stimulated by Prog. Exposure of these cells to Prog resulted in a decreased formation of leukotriene (LT). The 5-lipoxygenase inhibitor (5-LOX), AA861, effectively stimulated SN96-42 cell proliferation and 5-LOX-catalyzed product(s), especially LTC4, inhibited SN96-42 cell proliferation caused by Prog. Prog-sensitive enhancement of SN96-42 cell proliferation is at least partly mediated through an inhibition of LT formation and these data suggest that 5-LOX and LTs play important roles in SN96-42 cell proliferation stimulated by Prog.

Introduction

Hormone-responsive tumors such as breast, endometrial and prostatic cancers are derived from typical hormonal target tissues (breast, uterus and prostate), in which the proliferation and differentiation of target cells are regulated by various factors such as hormones and polypeptide growth factors. Steroid hormones are known to stimulate initiation and/or promotion step of carcinogenesis. Many studies have shown the roles of steroid hormones in carcinogenesis. Epidemiological work and therapeutic efficacy data of hormone therapy also support the role of estrogen in carcinogenesis.

In order to investigate the action mechanism of estrogen in tumor growth, B-1F cells were established from an estrogen-sensitive mouse Leydig cell tumor in our laboratory. Inhibitors for 5-lipoxygenase (5-LOX), a key enzyme of leukotriene (LT) synthesis, enhanced the B-1F cell proliferation whereas inhibitors for cyclooxygenase had no marked effect. Estradiol-17β (E2) stimulated the proliferation and inhibited LT synthesis in B-1F cells and LTs inhibit the proliferation of B-1F cells (1-7). This indicates that the proliferation of B-1F cells is largely regulated in a negative manner by LTs whose production is suppressed by estrogen.

Recently, the number of the patients with lung cancer has increased. The major types of lung cancer are squamous cell carcinoma and adenocarcinoma. Squamous cell carcinoma often arises from a central part of the lung and the generation of adenocarcinoma in the lung has not been sufficiently clarified. In the peripheral part of the lung, where the microvasculature richly exists, various functions including gas exchange take place. There seems to be a relationship between drug metabolism and the generation of lung cancer.

Sex steroid hormones such as estrogen and progesterone (Prog) are implicated in numerous developmental and functional processes. For the development and function of the lung, Prog fulfils important roles. Studies on the ontogenetic pattern of Prog and estrogen receptors support the idea that both steroids are important for prenatal lung maturation and development (8). It is well known that sex steroid hormones contribute to the development of hormone-dependent carcinomas including breast, endometrial and prostate cancers. However, lung cancer is not generally considered a typical target for sex steroid hormones. In a recent report, immunolocalization of Prog and estrogen receptors in non-small cell lung carcinomas (NSCLCs) were examined and it was shown that the Prog receptor (PR) might be a potent...
Materials and methods

Cell culture. Dissected squamous cell carcinoma tissue from a patient with lung cancer was used to establish the cloned cells. The patient was treated for bronchial asthma with various anti-asthmatic drugs. Our research was started after obtaining informed consent from the patient and his family. The fundamental methods for preparing the dispersed cells from the solid tumor and isolating cytokeratin-positive cells were previously reported (11). The cells were subculturied weekly in Eagle's minimum essential medium (MEM) containing FBS (10%) and 10^{-8} M Prog. The cultured cells in their 50th passage were cloned by limiting dilution. More than 25 clones were examined to select the clonal cells that showed Prog-responsive proliferation in serum-free medium: Ham's F12-MEM (1:1, vol./vol.) containing 0.1% (wt/vol.) BSA (HMB). One was termed SN96-42. SN96-42 cells used in the present experiments were found to proliferate well in the medium without any aid from exogenous peptide hormones or attaching factors and kept in a humidified atmosphere in the presence of 5% CO_{2}-95% air. Cells grown to confluence were transferred with the use of trypsin solution: trypsin (0.01%, wt/vol.)/EDTA (0.02%, wt/vol.) in phosphate buffered saline (PBS).

Cell proliferation experiments in culture. The effects of test compounds on the cell yield or DNA synthesis were examined with slight modification as previously published (3,7,12). SN96-42 cells were plated into four replicate 35-mm dishes at an initial density of 1x10^4 cells/dish in 2 ml of HMB medium. On the following day (day 0), the medium was replaced by 2 ml of serum-free HMB with or without test compounds, unless specified otherwise. The medium was changed every other day. When LTs were used as test compounds, the culture media were changed every day due to their instability as previously described (3,7). At the indicated times, cells were harvested with trypsin solution and counted with a hemocytometer using the trypan blue dye exclusion method on day 6, unless otherwise specified. [3H]-thymidine incorporation assay was also performed as previously described. SN96-42 cells were plated onto a 96-well plate at an initial density of 2x10^3 cells/well in 0.15 ml of HMB medium and on the following day (day 0), the medium was replaced by 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/well) for 2 h at 37˚C in a humidified incubator. The radioactivity (DPM) incorporated into the cells, which were briefly trypsinized and collected with a Cell Harvester LM 101 (Lab Science, Tokyo, Japan), was measured.

Binding analysis of progesterone receptor. The whole cell binding assay was used to obtain the binding parameters of PR in SN96-42 cells as previously described (2,5,12-14). The cells were washed four times with MEM containing 5 mM HEPES (4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid) and 0.1% (wt/vol.) BSA before determining the radioactivity specifically associated with the cells. The amount of [3H]-Prog specifically bound to the cells was determined as previously reported (11). The specific binding was analyzed according to the Scatchard method (15). To examine binding specificity, the cells were incubated with 2x10^{-9} M [3H]-Prog in the presence of various concentrations of Prog, estradiol (E2), dihydrotestosterone (DHT) or dexamethasone (Dx) at 37˚C for 60 min and treated as described above.

Immunohistochemistry. For the immunohistochemistry, 3 μm thick sections were cut from the formalin-fixed, paraffin-embedded blocks of the original tumor. Slides were incubated for 5-10 min in 0.3% hydrogen peroxide in methanol to quench endogenous peroxidase activity and washed twice in PBS for 5 min each. For antigen retrieval before the immunostaining, slides were immersed in a container with citrate buffer, pH 6.0,
microwaved for 15 min (600 W, 5 min, three times) and then cooled for about 30 min before rinsing. Slides were incubated for 30 min with 2% normal goat serum in 0.1% Tween-20-PBS. After removing blocking serum, slides were incubated with primary antibody (200-folds dilution) for PRO overnight at 4°C. On the following day, slides were washed three times for 5 min each with 0.1% Tween-20-PBS buffer, incubated with Envision™+Peroxidase (mouse) as directed in the manual from the supplier and observed by a light microscopy. For the negative control, the primary antibody was replaced with an isotype-matched negative control antibody (Dako), which showed no specific reactivity. In another experiment SN96-42 cells were plated on a sterile four well-chamber slide (Falcon) at a density of 2x10^4 cells/well in MEM containing 10% DCC-treated FBS. On the following day (day 0), the medium was changed to HMB medium with or without 10^-8 M Prog. On Day 2 the cells were fixed in 4% paraformaldehyde for 10 min and washed in three changes of PBS. Omitting the step involving treatment of the cells with hydrogen peroxide, immunofluorescence cell staining was performed with Alexa Fluor 448-labeled goat anti-mouse antibody as directed in the manual from the supplier.

Leukotriene assay. The contents of various leukotrienes in SN96-42 cells and their conditioned medium were examined as previously described (7).

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. The p-values of <0.05 and 0.01 were considered to be significant and highly significant, respectively.

Materials. Materials were obtained from the following sources: BSA (crystallized and lyophilized bovine serum albumin), radioinert Prog and trypsin from Sigma Chemical Co. (St. Louis, MO); Eagle’s minimum essential medium and Ham’s F12 from Nissui Pharmaceutical Co., Ltd. (Tokyo, Japan); (5,6,8,9,11,12,14,15-3H) arachidonic acid (60-100 Ci/mmoll, LTB4 or LTC4 [3H]-RIA kits, (1,2,6,7-3H) Prog (80-110 Ci/mmoll) and (methyl-1’,2’-3H) thymidine (40-100 Ci/mmoll) from DuPont/NEN Research Products (Wilmington, DE). The materials for HPLC from Waters Associates (Bedford, MA). Unlabeled LTs were purchased from Cayman Chemical Co., Inc. (Ann Arbor, MI). Monoclonal antibody (PgR636) for PRO and Envision™+Peroxidase (mouse) were purchased from Dako Co. (Carpinteria, CA). Alexa Fluor 448-labeled goat anti-mouse antibody was purchased from Molecular Probes Inc. (Eugene, OR). The other reagents used were of analytical grade.
Results

Effects of Prog or antiprogesterone on the growth of SN96-42 cells. Prog was found to stimulate the growth of SN96-42 cells in a concentration-dependent manner (Fig. 1A), the maximum effect being at a range of $10^{-9}$ to $10^{-8}$ M. In the following studies, Prog at $10^{-8}$ M was used in order to obtain a maximum and reproducible growth-stimulative effect, unless specified otherwise. The addition of RU38486 with $10^{-8}$ M Prog into a serum-free medium resulted in a decreased cell growth (Fig. 1B). RU38486 was used as the antiprogesterone in the following experiments owing to its potent growth-inhibitory ability. E2 and DHT had no significant effect on SN96-42 cell proliferation, while Dx inhibited SN96-42 cell proliferation at a higher concentration (Fig. 1C, D and E).

PR in SN96-42 cells. Scatchard analysis of the specifically bound [$^{3}$H]-Prog to whole cells suggested the presence of high affinity binding sites (Fig. 2A, 2B). The dissociation constant of [$^{3}$H]-Prog binding sites was $0.5-1 \times 10^{-8}$ M, similar to that of authentic PR. To confirm the presence of PR in SN96-42 cells, the steroid specificity of the binding components was then investigated (Fig. 2C). The experiments using [$^{3}$H]-Prog as a radioactive ligand were conducted. PR binding was found to be specific for Prog. Dx competed to a lesser degree than Prog, while unlabeled E2 or DHT did not compete for [$^{3}$H]-Prog binding even at μM concentrations. The presence of PR was also confirmed by immuno-staining. The nuclear PR immunoreactivity (Alexa Fluor 448) was observed in SN96-42 cells (Fig. 2D) cultured in the medium with Prog as well as the original tumor (data not shown) from which SN96-42 cells were derived.

Effects of enzyme inhibitors in arachidonic acid metabolic pathways on SN96-42 cell proliferation. Several enzyme inhibitors in arachidonic acid metabolic pathways were incubated with SN96-42 cells in the absence of Prog. As shown in Fig. 3, an addition of 2-(12-hydroxydodeca-5,10-diynyl)-3,5,6-trimethyl-1,4-benzoquinone (AA861) to the HMB medium resulted in a dose-dependent increase in cell yield at a range of $10^{-9}$ to $10^{-6}$ M. The PLA2 inhibitor, p-bromophenacyl bromide (BPB) was examined for its effect on the proliferation. While BPB was much less potent than AA861, it showed stimulatory activity. The cyclooxygenase inhibitor, indo-methacin did not elicit a significant stimulatory effect on SN96-42 cell proliferation at a range of $10^{-9}$ to $10^{-6}$ M, and it showed a slight inhibitory effect.

Effects of Prog on LT formation. The effects of Prog and AA861 on the contents of various LTs in SN96-42 cells and their conditioned medium were evaluated (Fig. 4). Both AA861 and Prog decreased the contents of LTs in a dose-dependent manner (Fig. 4).

Figure 4. Effects of AA861 or Prog on LT content. SN96-42 cells were plated in an HMB medium at a cell density of $10^{6}$/60-mm dish. After culture for 24 h, cells were treated without or with $10^{-8}$ M Prog or $10^{-6}$ M AA861 for 48 h and then replaced with a fresh medium containing the same compounds. These cells were subjected to further treatments for 7 h. LT content in the medium (A) or cells (B) was determined as described in Materials and methods. The data (mean ± SE) were obtained by five replicate assays. Cells were also treated with various concentrations of AA861 (AA) (C) or Prog (D) and LT content in the medium was determined. The values (percentage of control) represent means of five replicate assays, taking the contents of LTs produced without AA861 or Prog as 100%. The other 2 trials gave similar results.

Figure 5. Inhibitory effect of peptidyl LTs on the proliferation of SN96-42 cells. SN96-42 cells plated into 35-mm dishes were cultured in 2 ml HMB with various concentrations of LTs in the presence of $10^{-8}$ M Prog (P) (A). In another experiment SN96-42 cells were cultured with various concentrations of LTC4/D4/E4 antagonist pranlukast in the presence of both $10^{-8}$ M Prog and $10^{-8}$ M LTC4 (B). The viable cells were counted on day 6. Points, means of 4 determinations; bars, SE; (**) p<0.01; (*) p<0.05. The other 3 trials gave similar results.
Inhibitory effects of LTs on cell proliferation. In order to confirm the role of LTs in Prog-responsive growth of SN96-42 cells, various concentrations of LTs were added to the culture medium and cell proliferation was examined after 6 days. As shown in Fig. 5A, the increasing concentrations of LTC4 resulted in a significant inhibition of cell yields of Prog-treated SN96-42 cells. LTD4 and LTE4 were less potent than LTC4. Inhibition of the cell proliferation with LTC4 was restored after treatment with LTC4/D4/E4 antagonist pranlukast (Fig. 5B).

Discussion

The proliferation of established SN96-42 cells was sensitive to Prog and antiprogestrone RU38486 inhibited their proliferation stimulated by Prog. Exposure of these cells to Prog resulted in a decreased formation of LTs. Clara cell 10-kD protein (cc10-kD), a secretory phospholipaseA2 inhibitor, is suggested to be the human counterpart of rabbit uteroglobin (UG), that is regulated by progesterone and is similar to lipocortin I (16,17). Human cc10-kD is expressed constitutively at a very high level in the human respiratory epithelium (18). In addition to the lung and trachea, cc10-kD-specific transcripts are present in high levels in several nonrespiratory organs and tissues (prostate, thyroid, mammary gland and pituitary), with a distribution pattern very similar, if not identical, to that of UG in the rabbit using the PCR amplification method (18). In the study using an antibody for rabbit UG, a UG-like immuno-reactivity is observed in human endometrium (19), prostate (20) and respiratory tract (21). In our experiments, at first, it was thought that one possible action site of Prog was PLA2. However, Prog did not have a significant effect on the formation of prostaglandins (data not shown) and lowered all four LTs (LTB4, LTC4, LTD4 and LTE4). Five-lipoxygenase inhibitor, AA861, effectively stimulated SN96-42 cell proliferation and 5-lipoxygenase-catalyzed product(s), especially LTC4, inhibited their proliferation. Therefore, Prog-sensitive enhancement of SN96-42 cell proliferation is at least partly mediated through an inhibition of LT formation. Although the mechanism by which Prog suppresses LT formation remains uncertain, the most probable action site of Prog is 5-lipoxygenase. The addition of antiprogestrone to the Prog-stimulated cells restored LT formation to nearly that of the control (data not shown), indicating that Prog-sensitive inhibition of LT synthesis lies in a PR-mediated process.

Our previous reports suggest the important role of 5-LOX in estrogen-responsive growth enhancement of B-1F cells, established from a mouse estrogen-responsive Leydig cell tumor. The exogenous addition of LTs in the cultured medium led to a reduction of the proliferation and induced apoptosis of B-1F cells (7). It is considered that the inhibition of LT production by estrogen resulted in an enhancement of B-1F cell proliferation. Present results together with these reports show that 5-LOX and LTs play an important role in SN96-42 cell proliferation stimulated by Prog as well as in B-1F cell proliferation stimulated by E2. Although the 5-LOX promoter contains multiple regulatory elements (E2F, Pax-4, USF), no progesterone-responsive element (PRE)-related sequence has been identified in or proximal to the 5-LOX promoter. Therefore, mechanism of decrease of the 5-LOX activity by Prog in SN96-42 cells is not clear. In addition, our present results do not fully agree with other reports in which the proliferation of the tumor cells has decreased by the inhibition of 5-LOX (22-24). It is uncertain why there are differences in the growth response. Recently, it has been reported that medroxyprogesterone acetate induces cell proliferation through up-regulation of cyclin D1 expression via phosphatidylinositol 3-kinase/Akt/nuclear factor-κ B cascade in human breast cancer cells (25). Further study is necessary to elucidate the mechanisms of cell growth regulation by Prog.

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References