Downregulation of 15-lipoxygenase 2 by glucocorticoid receptor in prostate cancer cells

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Abstract. 15-lipoxygenase 2 (15-LOX-2) is lost or significantly reduced in prostate cancer. However, the regulation of 15-LOX-2 remains unclear. In this study, we independently cloned the 5' upstream promoter fragments of 15-LOX-2 gene. Target DNA fragments each were cloned into an expression vector containing luciferase reporter gene, which were called LF1(-1533/+87), LF2(-628/+87), LF3(-253/+87), LF4(-157/+87), LF5(-33/+87), LF6(-253/+1), and LF7(-157/+1). Each of these individual promoter fragments was transfected into primary prostate epithelial cells and prostate cancer LNCaP cells. The promoter activity gradually decreased with progressive deletions from LF2 to LF4. A significant drop was noted in the LF5. LF6 and LF7 that did not contain the 87-bp region downstream the transcription start site (TSS) have significant luciferase activities similar to those of corresponding fragments (LF3 and LF4) that contain 87-bp region downstream the TSS. This suggests that the 125-bp region (-157 to -33) of LF4 is critical for the promoter activity of 15-LOX-2 in the primary prostate epithelial cells PrEC and cancer cells LNCaP. Moreover, we discovered a specific glucocorticoid receptor (GR) responsive element (GRE) in this key region. The luciferase activities of the LF4 and LF7 were decreased in the LNCaP cells co-transfected with GR (hGR or hGRβ) expression vectors. This inhibitory effect is reversed after treatments with dexamethasone or two specific GR inhibitors (siRNAs of GR and RU486). Results from this study suggest a 125-bp region (-157 to -33) is critical for the 15-LOX-2 promoter activity in prostate epithelial cells and cancer cells, which was significantly downregulated by GR via the GRE in this region.

Introduction

15-lipoxygenase 2 (15-LOX-2) is a 76-kDa enzyme and consists of 662 amino acids and two domains (www.ncbi.nlm.nih.gov). The domain one called Plat_LOX domain contains 110 amino acids located in residual number 2 through 111. The domain two called lipoxygenase domain contains 518 amino acids located in residual number 137 through 654. 15-LOX-2 oxygenates carbon-15 in arachidonic acid (AA) and is one of important lipid peroxidizing enzymes that have been linked to carcinogenesis including prostate cancer. Unlike 15-lipoxygenase 1 (15-LOX-1), 15-LOX-2 has limited tissue distribution and significant substrate preference (1). 15-LOX-2 is mainly expressed in epithelia from prostate (2), esophagus (3), head and neck (4), breast, bladder, lung, and skin (1,5). 15-LOX-2 preferentially converts arachidonic acid to 15-S-hydroxyeicosatetraenoic acid (15-S-HETE) and metabolizes linoleic acid poorly.

15-LOX-2 has been considered to be a tumor suppressor. First, 15-LOX-2 is significantly reduced or lost in cancer of prostate (2), esophagus (3), head and neck (3) breast, bladder, lung, and skin (5). Second, immunohistochemical studies further showed that the reduced expression of 15-LOX-2 was inversely correlated with the degree of tumor differentiation and pathologic grade (Gleason scores) in prostate cancer (6). 15-LOX-2 expression was decreased in high-grade prostatic intraepithelial neoplasm (HGPIN) compared with benign tissues, suggesting that the loss of 15-LOX-2 is an early event in prostate carcinogenesis (6). Third, results from us (4) and others (2,3) showed that restoration of 15-LOX-2 resulted in significant inhibition of cancer growth proliferation. Moreover, 15-LOX-2 is found to be a negative cell cycle regulator in normal prostate epithelial cells (7) and may be associated with the ability to induce cell senescence (8). All together, 15-LOX-2 plays an anti-carcinogenic role in many epithelia-derived carcinomas including prostate cancer.

It remains unclear why tumor suppressor 15-LOX-2 is lost or significantly reduced in epithelia-derived cancer including prostate cancer. The previous study (7) revealed that loss of 15-LOX-2 expression in prostate cancer cells is not caused by promoter (gene) hypermethylation or histone

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Abbreviations: COX, cyclooxygenase; EMSA, electrophoretic mobility shift assay; GR, glucocorticoid receptor; HRPC, hormone refractory prostate cancer; 15-LOX-2, 15-lipoxygenase 2; PC, prostate cancer

Key words: 15-lipoxygenase 2, glucocorticoid receptor, prostate cancer, promoter
deacetylation. Recently there were two other studies focused on the promoter regulation of 15-LOX-2 gene. Tang et al. (9) reported that Sp1 positively and Sp3 negatively regulate and androgen does not regulate 15-LOX-2 gene expression in normal human prostate epithelial cells. Subbarayan et al. (5) reported that 15-LOX-2 can be negatively regulated by its product 15(S)-hydroxyicosatetraenoic acid and PPARγ through a PPAR half-site present in the 15-LOX-2 promoter region (-560 to -596). The above studies might partly explain the gene regulation of 15-LOX-2 in prostate cancer, but additional studies are clearly required to reveal molecular mechanisms for the gene regulation of this functional tumor suppressor 15-LOX-2 in prostate cancer. This information would be important leading to the identification of a molecular target(s) for prevention, early diagnosis and treatment of prostate cancer.

In this study, we independently cloned 5′ upstream promoter fragments of the human 15-LOX-2 gene from human fetal skin DNA through progressive deletions. Sequences of these fragments were verified. These promoter fragments each were cloned into the expression vector containing luciferase reporter gene. The luciferase activities of these promoter fragments each were examined in the primary normal prostate epithelial cells and the prostate cancer cell line LNCaP. We determined a 125-bp fragment (-157 to -33) as a key region for the promoter activity of 15-LOX-2 and identified a specific glucocorticoid receptor (GR) responsive element (GRE) in this key region through a computer search analysis and a gel shifting experiment. Furthermore, we have determined that 15-LOX-2 promoter was downregulated by over-expression of GR in prostate cancer cells. This inhibitory effect by GR was reversed when the cells were treated with either steroid dexamethasone, or a specific GR inhibitor (siRNA of GR or RU486).

Materials and methods

Cloning of immediate upstream 15-LOX2 promoter region. Primer design: the human 15-LOX-2 gene was found to be located in chromosome 17 (10). The primer sequences (Table I) for cloning were designed based on the genomic sequences and cDNA sequence of 15-LOX-2 deposited in the GenBank (Accession numbers: NT_086857, AJ305028-AJ305031).

Cloning of 15-LOX2: Fetal skin genomic DNA was purchased from the American Type Culture Collection (ATCC, Manassas, VA). Expand long template PCR system and Tag DNA polymerases were obtained from Roche Diagnostics Corporation, Roche Applied Science (Indianapolis, IN). TA cloning kit was purchased from Invitrogen (Carlsbad, CA). The PCR for target DNA fragments was performed using Bio-Rad iCycler (Bio-Rad, Hercules, CA). These PCR products were gel purified using QIAquick gel extraction kit (Qiagen, Valencia, CA), and cloned respectively into the T/A vector pCRII and transformed into OneShot cells (Invitrogen) following the manufacturer’s protocol. These cells were cultured in LB medium containing 100 μg/ml ampicillin. Plasmids were purified with Concert™ Rapid Plasmid Miniprep System (Gibco-BRL, Rockville, MD). The target DNA fragments were further excised, gel purified and cloned into pGAW expression vector (gifted from Dr Fei Xiao, University of Stanford, CA). The orientation and sequence of each insert were confirmed by restriction digestion and sequencing. A dye terminator cycle sequencing was performed to determine sequence of each of DNA fragments using an ABI PRISM™ 310 Genet Analyzer. The other primers that were used for PCR and sequencing, were T7, SP6 primers, pGAW-For-primer, luc-Rev-primer designed according to the sequence of the parental pGAW vector (Table I). A computer analysis was performed on the nucleotide sequence of each clone using the software from the National Center for Biotechnology Information BLAST network service.

Cell cultures. The prostate cancer cell line LNCaP was obtained from ATCC. The LNCaP cells were cultured in RPMI-1640 medium supplemented with 10% heat-inactivated fetal bovine serum (FBS) plus 1X antibiotic-antimycotic solutions (Gibco, Grand Island, NY). A normal primary prostate epithelial cell line PrEC was purchased from Clonetics and maintained in PrEBM media (Cambrex or Clonetics, Charles City, IA). Cultures were maintained in a humidified incubator with 5% CO2, 95% O2 air at 37°C, and the media was changed twice a week.

Luciferase reporter assays. Luciferase reporter assays were performed in the cells. The cells grown in six-well culture plates (5x10^5 cells/well) were transiently transfected, in triplicate, with either promoterless parental vector pGAW or various luc constructs (2 μg plasmid/well) together with the β-gal plasmid (0.125 μg plasmid/well) to normalize the transfection efficiency. To determine effect of human glucocorticoid receptor (GR) on promoter activities of 15-LOX-2, a LF fragment was co-transfected with a GR expression vector into the LNCaP cells. The human GR expression vectors (pcDNA3-hGRα containing α isoform of GR and pcDNA3-hGRβ containing β isoform of GR) were kindly provided by Dr Julian R.E. Davis, Manchester University, UK. Forty eight hours after the transfection, the cells were harvested by scraping into 1X lysis buffer (Promega, Madison, WI) for a subsequent measurement of luciferase activity.

Lysates containing equal amounts of protein were assayed for luciferase activity using luciferase assay kit (Promega). β-galactosidase activity was measured using Tropix Galacto-Light Plus assay system. After subtracting the baseline values, the relative luciferase unit (RLU) activity was determined as the ratio of luciferase/β-gal activities. The relative promoter activities were expressed as relative values (i.e. %).

Electrophoretic mobility supershift assays (EMSA). Immuno-precipitation (IP) of GR protein was performed using LANCaP cells. Briefly the cells were transfected with pcDNA3.1-hGRα. After the selection in the media containing G418 for 2 weeks, the surviving tumor cells were collected and the nuclear lysates were generated using a lysis buffer (50 mM Tris-HCl, pH 8.0, 150 mM NaCl, 1% NP-40). The nuclear lysates were then incubated overnight at 4°C with Sepharose-G bind beads (Santa Cruz Biotechnology, Inc., Santa Cruz, CA) to which rabbit anti-GR antibody (Santa Cruz Biotechnology, Inc.) was bound. These beads were then washed, collected and subjected to EMSA on 6% polyacrylamide gels.

One 26-bp double-stranded oligonucleotide probe containing putative GR binding element and a double-stranded oligonucleotide probe containing mutations in the core of
putative GR binding element were designed and synthesized (Table I). The probes were designed based on the putative sequence CTCCTGTTCC of GRE, among which the sequence TGTTCC is critically required (11). These probes were labeled with or without biotin using the Biotin 3' End DNA Labeling kit (Pierce, Rockford, IL). The gel shift reaction was accomplished by using the LightShift Chemiluminescent EMSA kit (Pierce) according to the manufacturer's instructions. For each gel shift reaction, an aliquot of labeled oligonucleotide (20 fmoles) was incubated with 3 μg of nuclear extract from the cells for 20 min at room temperature in a 10-μl mixture containing 1X binding buffer, 2.5% glycerol, 10 mmol/l MgCl₂, 50 mg/l poly (deoxyinosinic-deoxyctydilic acid), and 0.05% Nonidet P-40. Unlabeled probes with either 200- or 400-fold molar excess were used in order to competitively block the binding of GR to labeled probes. The unlabeled probes were first preincubated for 2 min at room temperature with the GR IP extracts before the addition of the labeled probe. Samples were run on a non-denaturing 6% polyacrylamide gel in 0.5X Tris-borate EDTA buffer at 100 V for 70 min at 4°C. The electrophoresed binding reactions were transferred to positively charged nylon membrane in a Mini Trans-Blot Cell (Bio-Rad) at 380 mA for 40 min, and crosslink was performed for 90 sec in an XL-1500 UV Crosslinker chamber (Fisher Scientific, IL). Detection of biotin-labeled DNA was performed using stabilized streptavidin/horseradish peroxidase conjugate (Pierce) according to the manufacturer’s procedures.

**Table I. Primers and oligonucleotides for cloning, sequencing, electrophoretic mobility supershift assay (EMSA) and site-directed mutagenesis (putative and mutant GRE are highlighted).**

<table>
<thead>
<tr>
<th>Name</th>
<th>Sequence</th>
<th>Restricted enzyme</th>
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<tr>
<td>Cloning primers</td>
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<tr>
<td>LOX2-1R</td>
<td>5'-ATCGGTATCCGCCACGCTAAGTCCAG-3'</td>
<td>KpnI</td>
</tr>
<tr>
<td>LOX2-2F</td>
<td>5'-ACTTCTAGAGGGCACTTGTGCTTC-3'</td>
<td>XhoI</td>
</tr>
<tr>
<td>LOX2-3F</td>
<td>5'-AGCCCTAGAGGCCTGTGAATCCAGC-3'</td>
<td>XhoI</td>
</tr>
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<tr>
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<td>5'-AGCAAGCTTCCATCGGTATTCTCTAGTC-3'</td>
<td>HindIII</td>
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<td>Sequencing primers</td>
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<td>pGAW-For-primer</td>
<td>5'-CTAGCAAAAATAGGTGTCCTCC-3'</td>
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<tr>
<td>Luc-Rev-primer</td>
<td>5'-TCCACGGGATAGAATGGGCG-3'</td>
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<td>Gel shift oligonucleotides</td>
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<td>GRGS-1</td>
<td>5'-CTTAGTCCCTCCCGCTGCAGTTTG-3'</td>
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<td>GRGS-2</td>
<td>5'-GCTGGAGAGAGGAGGGACTAAAGAATA-3'</td>
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<tr>
<td>GRGSM-3</td>
<td>5'-CTTAGTCCCTACTGACCCAGCGCGTG-3'</td>
<td></td>
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<tr>
<td>GRGSM-4</td>
<td>5'-GCTGGAGAGGAGGGACTAAGAATA-3'</td>
<td></td>
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<tr>
<td>Primers for site-directed mutagenesis</td>
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<tr>
<td>GRGSM-5</td>
<td>5'-CTGGGGTGCAGTGGACTAAGAATACCGTCCG-3'</td>
<td></td>
</tr>
<tr>
<td>Amp rev</td>
<td>5'-GTCAAGAAGTGGGATTTGCGATATCGT-3'</td>
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USA). The GRGSM-5 and amp rev primers were used for the substitutions (Table I). The mutations in the primer GRGSM-5 were the same as in GRGSM-4 which was used in EMSA. The primer GRGSM-5 phosphorylated at their 5' end was synthesized by Invitrogen. Briefly, 5 nmol of each primer, 50 ng pGAWLF7, 0.8 μl Change-IT enzyme were mixed together with 20 μl DMSO. The thermal cycler was programmed as follows: initial denaturation at 95°C for 2 min; 30 cycles at 95°C for 30 sec, 55°C for 30 sec and 68°C for 10 min; post-incubation at 72°C for 10 min. Two microliters of the final sample was used to transform competent E. coli cells DH5α (Invitrogen). The purified plasmids were sequenced to confirm the mutant substitution successfully. The luciferase activity assay was obtained by transforming mutant LF7 fragment into LNCaP cells with or without Dex or RU486 treatment as above described.

Western blot analysis. The above LNCaP cells were lysed in a buffer containing protease inhibitors. The protein concentration was determined using a bicinchoninic acid (BCA) assay kit (Pierce). Bovine serum albumin was used as a standard. SDS-PAGE was carried out using Bio-Rad gel system. Equal aliquots of total cell lysates (50 μg per sample) were loaded per lane in Laemmli SDS-PAGE sample loading buffer under denaturing conditions. The Western blot was carried out according to the protocol of Santa Cruz Biotechnology, Inc. GAPDH from Santa Cruz Biotechnology was used as an internal control. The blots were quantitatively analyzed using Kodak Electrophoresis Documentation and Analysis System (EDAS) 290 (Eastman KODAK Company, Rochester, NY). Statistical analysis was done by paired Student’s t-test and P-values were determined to evaluate the statistical significance (P<0.05) of the changes observed.

Results

Characteristics of 5' upper stream structure of 15-LOX-2. A total of 7 fragments were amplified from 5' untranslated structure of 15-LOX-2 gene and were cloned through progressive deletions using human fetal skin DNA as a template. These seven fragments were LF1 (-1533/+87, 1620 bp), LF2 (-628/+87, 715 bp), LF3 (-253/+87, 340 bp), LF4 (-157/+87, 244 bp), LF5 (-33/+87, 120 bp), LF6 (-253/+1, 254 bp), and LF7 (-157/+1, 158 bp), respectively (Fig. 1A). The LF6 and LF7 are the same to the LF3 and LF4, respectively; except that the 87-bp sequence downstream the transcription start site (TSS) (+1 in Fig. 1A) have significant luciferase activities contain the 87-bp region downstream the transcription start (TSS) (+1 in Fig. 1A) have significant luciferase activities similar to those of corresponding fragments (LF3 and LF4) that contain 87-bp region downstream of the TSS (+1). Similar results were obtained in primary normal prostate epithelial cells PrEC (Fig. 2B). Results from the above studies suggest that the 125-bp region (-157 to -33) is critical for the promoter activity of 15-LOX-2 gene in the primary prostate epithelial cells and prostate cancer cells.

Existence of a specific glucocorticoid receptor responsive element (GRE) in the 15-LOX-2 promoter fragment. Based on the above discovery that the 125-bp region (-157 to -33) is critical for promoter activity of 15-LOX-2 in the prostate cancer cells, we hypothesized that an important transcription factor response element(s) may exist in this region. We have performed an extensive computer analysis to search for transcription binding elements (www.gene-regulation.com). Five potential binding elements of transcription factors GR, Egr-1, Krox-20, GLI3 and ETF were identified in this region, but not in other regions of 15-LOX-2 gene promoter structure. Among these 5 potential binding elements, a putative glucocorticoid receptor (GR) responsive element (GRE) was identified in this region, but not in other regions of 15-LOX-2 gene promoter structure.
through its binding to glucocorticoid receptor (GR) (14-17),
activation of GR results in the suppression of prostate cancer (18),
and GR was not significantly expressed in many prostate
cancer cell lines including LNCaP (18).

To examine whether or not this partially matched GRE in
the key region of LF4 was indeed bound by GR, we transfected
hGRα expression vector into the LNCaP cells. The surviving
tumor cells containing hGRα were obtained after the selection
with G418. Immunoprecipitation of GR protein were then
performed using the nuclear lysates from the tumor cells
containing hGRα. Finally we performed electrophoretic
mobility shift assays (EMSA) to determine the binding of GR
to GRE. Results from the EMSA showed that the wild-type
double strand oligonucleotide probe containing a putative
GR binding element was bound by the immunoprecipitated
protein containing highly expressed GR (lane 1, Fig. 3).
However, this binding was competitively blocked by a high
excess of unlabeled probe (Lanes 2 and 3, Fig. 3), but not
blocked by the mutant oligonucleotides (lane 4). Fig. 3 is a
representative picture. These results indicated that this partially
matched GRE is able to be bound by the glucocorticoid
receptor.

**Downregulation of 15-LOX-2 promoter activity by the gluco-
corticoid receptor (GR) was reversed by steroid dexamethasone**

(DEX). To address whether or not GR is able to regulate
the 15-LOX-2 promoter through the above GRE, we performed
functional assays to determine the effect on the luciferase
activities of LF4 and LF7 that contain the GRE in the
LNCaPs cells co-transfected with a GR (hGRα or hGRβ)
expression vectors. In addition, we evaluated the effect of
dexamethasone (DEX) on the luciferase activities of LF4 and
LF7 in the cells co-transfected with a GR expression vector.
This is because DEX is known to downregulate GR expression
in LNCaP cells (19,20). First, we examined the luciferase
activities of LF4 in the LNCaP cells, which was co-transfected
with either hGRα or hGRβ, with or without dexamethasone
(DEX) treatment. LF5 that does not contain the GRE was
used as a control. The luciferase activity of LF4 significantly
decreased to 59.58±2.74 when LF4 and hGRα were co-
transfected into the cells. However, this downregulation of
LF4 activity by hGRα was reversed (99.56%±5.89) when the
cells were treated with DEX at a concentration of 0.1 μM
(Fig. 4A). We further evaluated whether or not this effect by
DEX was dose-dependent. Interestingly, no significant
changes were observed in the luciferase activity of LF4 when
much higher concentrations of DEX (1 or 2 μM) were utilized
(Fig. 4A). Similarly, the luciferase activity of LF4 was
decreased to 68.75±3.28 when LF4 and hGRβ expression
vector were co-transfected into the cells. This downregulation
of LF4 activity by GRβ was slightly reversed by the DEX
treatment, but without statistical significance (Fig. 4B). The
luciferase activities of LF4 was found to increase slightly
(119.72±3.97) with DEX 1 μM (Fig. 4E, lane 2), however,
the effect by DEX was not dose-dependent since different
concentrations of DEX treatment (0.1 μM, or 1 or 2 μM) produced
no significant difference in the luciferase activities
of the cells co-transfected with LF4 and GRβ expression
vectors (Fig. 4B). Similar results were obtained when the
luciferase activities of LF7 were evaluated in the LNCaP
cells that were co-transfected with hGRα or hGRβ
and were treated with or without DEX treatment (Fig. 4C and D,
Fig. 4C). LF5 that does not contain the GRE was used as a
control. The above results indicate that 15-LOX-2 promoter
was downregulated by GR, however this inhibitory effect by
GR was reversed by steroid DEX treatment.

The inhibitory effect on 15-LOX-2 promoter activity by GR
was reversed by a specific GR inhibitor (siRNAs of GR or
RU486). To make sure that the above luciferase activities of
15-LOX-2 promoter were specifically downregulated by expressed GR, we further evaluated the effect of two specific GR inhibitors (siRNAs of GR and RU486) on the luciferase activities of LF4 and LF7 in the LANCaP cells co-transfected with a GR expression vector compared with controls. The siRNAs of GR was previously shown to specifically abolish expression of GR (21) and RU486 was shown to inhibit the function of 15-LOX-2 (22,23). The luciferase assays showed that the treatment with siRNAs of GR was able to reverse the inhibitory effect of GR on promoter activities of LF4 and LF7. GR reduced luciferase activities of LF4 and LF7 to 60.73%±2.09 and 56.11%±2.94, respectively, but these were up to 86.17%±3.60 and 138.57%±3.03, respectively after the treatment with siRNAs of GR. More effectively, the treatment with RU486 was able to reverse the inhibitory effect of GR on luciferase activities of LF4 from 60.73%±2.09 to 172%±7.78 and on the activities of LF7 from 56.11%±2.94 to 290.18%±9.79, respectively. The results are summarized in Fig. 5A and B. The inhibitor RU486 slightly increased the luciferase activities of LF4 and LF7 (Fig. 4E, 5C, lane3). In this study, we also performed Western blot analysis of GR in the LANCaP cells, which showed the treatment with the siRNAs of GR completely abolished the GR exogenous or endogenetic expression (Fig. 5D). However, the treatment with RU486 did not affect the expression level of GR (Fig. 5D). This finding was consistent with the previous studies that RU486 block the GR function through high affinity for the glucorticoid receptors, but does not change the expression level of GR (24). The results from the above experiments further confirmed that GR expression downregulated 15-LOX-2 promoter through the GRE in the key region of 15-LOX-2.
To further examine whether or not the above GRE is functional, we performed site-directed mutagenesis to create a mutant GRE containing three mismatch mutations. This GRE mutant was called LF7M. This was generated using Change-IT Multiple Mutation Site Directed Mutagenesis kit following by the manufacturer's instruction. The mutations in the primer GRGSM-5 were the same as in GRGSM-4, which was used in EMSA (Table I). The luciferase activity of LF7M was found to dramatically decrease to 11.93±0.61 after its normalization to 100. The treatment with DEX (1 μM) did not result in significant increase in the luciferase activity of LF7M (15.23±1.018), and the treatment with RUM486 (1 μM) resulted in increase in the luciferase activity of LF7M (33.54±1.47), but not to the level of LF7 (Fig. 5C).

Collectively, the above findings suggested the GRE exists and is functional in this 125-bp key region (-157 to -33) of the 15-LOX-2 promoter.

Discussion

Through progressive dissection and analysis of 5' untranslated structure of 15-LOX-2 gene, we have determined that the 125-bp region (-157 to -33) of the LF4 fragment is critical for the promoter regulation of 15-LOX-2 gene in prostate cancer cells LNCaP. This is because the luciferase activity (111±6.48) of LF4 was similar (slightly reduced) to the dissected upstream fragments LF1-LF3, and activity of downstream fragment LF5 was dramatically reduced (1.94±0.07) almost to the level of the control background. This was further demonstrated using the fragments LF6 and LF7 that share the same sequence with LF3 and LF4, but do not contain the 87-bp region downstream the transcription start site (TSS). This suggests that the sequence downstream of the TSS in 15-LOX-2 promoter might not contain a special binding element(s) that significantly influences the gene expression in the prostate epithelial cancer cells LNCaP. Compared with the previous two studies on the promoter dissection and analysis for 15-LOX-2 gene, we found this 125-bp region (-157 to -33) is within the most active fragments identified in their studies (5,9). In this study, we have advanced the 15-LOX-2 promoter dissection into many smaller fragments and have determined the 125-bp key region (-157 to -33) for the promoter regulation of 15-LOX-2 gene. Therefore, we believe that the results from our study have advanced our research in revealing key transcriptional factors that regulate the 15-LOX-2 promoter in cancer cells.

Through the extensive computer analysis and powerful electrophoretic mobility shift assay (EMSA) (Fig. 3) and site-directed mutagenesis (Fig. 5C), we have determined existence of functional GRE in the 125-bp key region of 15-LOX-2 gene. Furthermore, we have determined that dexamethasone (DEX), which is known to downregulate GR in prostate cancer...
cells (19), have reversed the inhibitory effect on the 15-LOX-2 promoter activity (Fig. 4). This discovery is very important in revealing the molecular mechanism on the tumor suppressing function of 15-LOX-2 gene. This is because glucocorticoid hormones are known to regulate proliferative, inflammatory and immune responses via its receptors GR (14-17). Clinically, glucocorticoid hormones, combined with or without chemotherapy, have been extensively used for the treatment of hormone refractory prostate cancer (HRPC). Glucocorticoids were even used as the ‘standard’ therapy arm in several randomized phase II–III clinical trials for the combination therapy of HRPC (14,18,25). However, it still remains to be determined whether or not glucocorticoid hormones suppress prostate cancer growth through the activation of tumor suppressor gene 15-LOX-2 in vivo.

In addition, we have further confirmed the functional regulation of 15-LOX-2 promoter through the use of two specific GR inhibitors (siRNAs of GR and RU486) (Fig. 5). A siRNAs of GR is a short RNA sequence specific to GR that completely abolishes the expression of GR (21). In our study, we demonstrated that addition of this siRNA almost abolished expression of both endogenous and exogenous GR. RU486 is another well-known GR inhibitor, which specially binds glucocorticoid receptor, estrogen receptor and progesterone receptor and inhibits their functions, but does not affect their protein expression (24). In our study, we indeed see no significant change in the level of GR protein in the cells treated with RU486 (Fig. 5D). Of great interest is that the degree of inhibitory effect by RU486 is much greater than the siRNAs of GR. The reason for this difference is unknown. This may be due to inhibition of other proteins including estrogen receptor and progesterone receptor by RU486, which may be responsible for additional inhibitory effect by RU486.

The glucocorticoid receptor GR is a well-known transcription factor (26,27). To our understanding, two major mechanisms on the gene regulation by GR: direct or indirect transcriptional regulation were previously reported (28,29). The direct transcriptional regulation (transactivation) occurs via binding of the GR homodimer to palindromic promoter DNA sequences called glucocorticoid-response elements (GRE). The indirect regulation is mediated via cross-talk with other transcription factors (TFs) (28,29). Most of such GR-TF interactions repress the activity of partner TFs and their target genes (transpression) (18). In this study, we have demonstrated an existence of a specific and functional GRE in the key region (-157 to -33) of 15-LOX-2 promoter (Fig. 3). However, it still remains unclear whether or not GR regulates 15-LOX-2 promoter through indirect transcriptional regulation mechanism. This certainly points to our future research direction.

GR expression was strongly decreased or absent in 70-85% of PC. Similar to primary prostate cancer, some prostate cancer cell lines including LNCaP, also lack GR expression (18). The GR expression is found to decrease at the high-grade prostatic intraepithelial neoplasia (HGPIN), early stage of prostate carcinogenesis. The early loss of GR expression in prostate tumorigenesis resembles changes reported for estrogen receptor β, an inhibitor of prostate growth (30). In contrast, the expression of other steroid hormone receptors either remains stable, like AR, or is increased, like estrogen receptor α and progesterone receptor (30,31). Remarkably, the loss of GR was specific only for the epithelial compartment of PCs. In benign and malignant prostate specimens alike, the stromal cells showed predominant nuclear localization of GR (32), suggesting an important role of GR specifically in prostate epithelium (18). 15-LOX-2 is normally expressed in apical epithelial cells of prostate epithelium, but either significantly reduced or lost in majority of prostate cancer (2). Decrease of 15-LOX-2 expression also starts at the HGPIN. In this study, we have demonstrated that the 15-LOX-2 promoter can be negatively regulated by overexpression of GR in prostate cancer cells in vitro. However, we still do know how 15-LOX-2 and GR interact in primary prostate cancer. Many questions still remain. For example, is 15-LOX-2 inversely related to GR in terms of their expression pattern and function in an individual cell? Additional mechanistic studies are certainly required.

In summary, we have discovered that a 125-bp region (-157 to -33) of 15-LOX-2 promoter is critical for the promoter regulation of 15-LOX-2. In addition, we have demonstrated the existence of a specific and functional GR-responsive element (GRE) in this key region through computer analysis, electrophoretic mobility shift assay and site-directed mutagenesis. Overexpression of GR was shown to downregulate the promoter activity of 15-LOX-2 gene. This inhibitory effect was partially or completely reversed by steroid dexamethasone (DEX) and specific GR inhibitors (siRNAs of GR and RU486). Results from our study have demonstrated that promoter activity of 15-LOX-2 was inhibited by overexpression of GR possibly through a direct binding mechanism.

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References


