Leuprorelin acetate affects ERK1/2
tivity in prostate cancer cells

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Abstract. The mechanisms by which a GnRH analogue, leuprorelin acetate (LA), antagonizes the mitogenic effect of dihydrotestosterone (DHT) or epidermal growth factor (EGF) in prostate cancer cells is poorly understood. The mitogen-activated protein kinase system has a central role in growth regulation and, for this reason, we investigated the involvement of the extracellular signal-regulated kinase (ERK1/2) pathway in the response of both androgen-sensitive (LNCaP) and -insensitive (PC-3) prostate cancer cells to LA alone or combined with EGF or DHT. The evaluation of ERK activation was performed by using Western blot analysis and immunocytochemistry. EGF specifically induced ERK1/2 activity in both models and this effect was counteracted by an inhibitor of EGF-receptor phosphorylation. The addition of LA produced an appreciable reduction of ERK phosphorylation promoted by EGF in LNCaP cells, while it generally determined an increase in ERK activity in androgen-unresponsive PC-3 cells. The slight ERK activation induced by DHT in LNCaP cells was counteracted by LA and this effect was evident only by immunocytochemistry. Our findings suggest that the antiproliferative effect of LA in prostate cancer cells stimulated by hormones and growth factors may be, at least in part, mediated by the reduction of ERK1/2 activation in LNCaP cells and linked to the unexpected increase of this activity produced by the analogue in PC-3 cells.

Introduction

The hypothalamic decapptide gonadotropin-releasing hormone (GnRH) is a key regulator of the mammalian reproductive system, triggering the synthesis and release of follicle-stimulating hormone (FSH) and luteinizing hormone (LH) in the pituitary (1).

In the past, hundreds of agonists and antagonists of GnRH have been developed and various applications in different fields of medicine have been established for these analogues. In particular, they have been investigated for their therapeutic advantages in hormone-related tumors. GnRH agonist administration results in hormone ablation (chemical castration) which can effectively replace surgical castration (2). The slowing-down of neoplastic cell growth is based on the reduction of sex steroid supply (3). In addition, it is widely accepted that GnRH analogues may act directly on tumor cells via specific receptors expressed on cell membrane. Several reports have provided evidence that GnRH analogues negatively control the proliferation of breast, ovarian, endometrial and prostate cancer cells (2). Nevertheless, the mechanisms that underlie this direct action are poorly understood.

In our experience, the GnRH agonist, leuprorelin acetate (LA), which is ineffective when used alone, counteracts or even suppresses the androgen-induced growth of androgen-sensitive prostate cancer cells, LNCaP, and reduces the mitogenic effect of epidermal growth factor (EGF) in androgen-insensitive PC-3 cells. Moreover, in both models, LA reduces the expression of the PSA gene, evaluated by reverse transcriptase-polymerase chain reaction, and inhibits the gene expression induced by dihydrotestosterone (DHT) in LNCaP cells and EGF in PC-3 cells (4,5).

More recently, we have shown that LA significantly reduces the DHT-promoted expression of the antiapoptotic gene, bcl-2, evaluated at both the mRNA and protein level, without inducing apoptosis in the LNCaP cells (6).

The extracellular signal-regulated kinase, ERK1/2, belongs to the larger mitogen-activated protein (MAP) kinase family and is generally involved in cell division and differentiation (7). In the pathogenesis and development of prostate cancer, most growth and oncogenic signals have been implicated and lead to tyrosine phosphorylation which, in turn, activates MAP kinase cascade (8,9).

The present study deals with the role played by ERK1/2 in determining the antiproliferative effect of LA observed by our group in prostate cancer cells only in the presence of DHT or EGF. Further experiments were performed regarding the effect of EGF, alone or combined with LA, on both LNCaP and PC-3 cells. ERK1/2 expression has been evaluated by both Western blotting and immunocytochemistry in LNCaP and PC-3 cells exposed to different treatments.
Materials and methods

Compounds. Leuprolerin acetate (LA), D-Leu6-(des-Gly10-NH2) LHRH ethylamide was kindly donated by Takeda Italia Farmaceutici SpA, Rome, Italy. It was dissolved in saline solution and stored at 4°C.

5α-dihydrotestosterone (DHT), epidermal growth factor (EGF) and tyrphostin (AG1478) were purchased from Sigma (St. Louis, MO, USA). Stock solutions were made by dissolving DHT and AG1478 in absolute ethanol, and EGF in a 10 mM acetic acid buffer containing 0.1% BSA and stored at 4°C.

Antibodies. Anti-phospho-p44/42 MAP kinase, Thr202/Tyr204 (pERK1/2), anti-phosphorylation state independent p44/42 MAP kinase (ERK1/2), rabbit polyclonal antibodies and the secondary antibody, horseshadish peroxidase (HRP)-labeled goat anti-rabbit IgG, were from Cell Signalling Technology, Inc. (Newcastle, USA). The secondary biotinylated goat anti-rabbit antibody was from Vector Laboratories (Burlingame, CA, USA).

Cell cultures. The hormone-sensitive LNCaP cell line was used between passages 33 and 54 and cultured in RPMI-1640 medium (Eurobio, Les Ulis Cedex B, France), supplemented with 10% (v/v) foetal bovine serum (FBS, ICN Biomedicals, Costa Mesa, CA, USA), 10 mM N-2-hydroxyethylpiperazine-N'-2-ethane sulfonic acid (Hepes buffer, Eurobio) and antibiotics.

Hormone-insensitive PC-3 cells were used between passages 76 and 92 and were routinely cultured in Dulbecco's modified Eagle's medium (DMEM, Eurobio), supplemented with 5% FBS, Hepes buffer and antibiotics.

Both cell lines were sub-cultured weekly. They were maintained in a humidified air:CO2 atmosphere (95%:5%) at 37°C.

Cell proliferation assays. The mitogenic effects of EGF on both LNCaP and PC-3 cells were investigated. No agreement in the literature about the culture conditions used to investigate the proliferative effects of EGF on LNCaP cells (10-12). In the present study, LNCaP cells were seeded into 60-mm plastic Petri dishes at two different densities, i.e. 25,000 and 50,000 cells/ml culture medium, and allowed to adhere to the culture plate. After 48 h, cells were fed with fresh medium supplemented with 5% charcoal-treated FBS (CH-FBS) and EGF (0.1-50 ng/ml of culture medium).

PC-3 cells were plated at a density of 25,000 cells/ml of standard culture medium in 60-mm plastic plates. Cells were allowed to adhere and, 24 h after plating, the seeding medium was replaced with fresh DMEM supplemented with 5% CH-FBS and the above-mentioned EGF concentrations. The fresh medium containing EGF was renewed 2 days after the first change.

In another series of experiments, both LNCaP and PC-3 cells were exposed to 10 ng/ml EGF in combination with LA at concentrations from 10-11 to 10-5 M. Medium containing EGF was changed 2 days after the first medium renewal and LA was added every day. Cell counts were performed with a hemocytometer after 2 and 4 days.

Finally, proliferation experiments were also performed in the same conditions as those used for Western blot analysis. Cells were counted after 2 days of treatment with the different compounds.

Triplicate cultures were set up for each drug concentration and control dishes containing untreated cells were run in parallel with the same amount of vehicle. Cell viability was assessed by Trypan blue exclusion test.

Western blot analysis

Cell treatment. Cells were seeded and cultured in 100-mm plates to approximately 60% confluence in the presence of serum-containing medium. They were then cultured in low serum medium (0.5% CH-FBS) for 48 h to reduce basal levels of phosphorylated MAP kinases. Prior to treatment with hormones and growth factors, the medium was renewed again to abolish the MAP kinase phosphorylation due to factors secreted by cells. Then, LNCaP cells were treated for 5, 10, 30, 60 and 240 min with various concentrations of LA (10-11, 10-6, 10-5 M), DHT (10-9-10-7, 10-5 M) and EGF (10 ng/ml) in serum-free medium, while PC-3 were exposed to LA (10-11 and 10-5 M) and EGF (10 ng/ml) for the same time periods.

In a series of experiments, cells were treated with LA and EGF simultaneously or they were pre-treated with LA 15 min prior to the exposure to 10 ng/ml EGF.

A number of plates from each experiment were treated with 300 nM tyrphostin (AG1478), a selective inhibitor of the tyrosine kinase of the EGF receptor, 30 min prior to treatment with EGF.

Following exposure to different compounds for the various time periods, cells were washed twice with ice-cold phosphate buffered saline (PBS) without Ca2+ and Mg2+, pH 7.6, collected by mechanical scraping and lysed in lysis buffer [50 mM Tris-HCl (pH 8.0), 150 mM NaCl, sodium dodecyl sulfate 0.1% (SDS), 1% Nonidet P-40, 0.5% sodium deoxycholate, 100 μg/ml phenylmethylsulfonylfluoride, 1 mM sodium vanadate and 30 μg/ml aprotinin] on ice for 30 min. Cell debris was removed by centrifugation (12,000 x g per 10 min at 4°C) and the protein content of the supernatants was determined using a modified version of Lowry's method (13).

Immunoblot of phosphorylated-ERK1 and ERK2 (pERK1 and pERK2). Samples (40 μg of proteins) and control protein (nonphosphorylated and fully phosphorylated ERK2) were resolved on denaturing polyacrylamide gels 12% (SDS/PAGE) and transferred to polyvinylidene difluoride (PVDF; Immobilon P, Millipore Corp., Bedford, MA, USA) membrane by electroblotting. Membranes were blocked with TBS-T (Tris-HCl buffer saline with 0.1% Tween-20) containing 5% non-fat dry milk on ice for 1 h. After washing for 3x5 min, membranes were probed for 1 h at room temperature with an HRP-conjugated anti-rabbit antibody (1:2000) in TBS-T containing 5% non-fat dry milk. After washing for 3 x 5 min, membranes were probed for 1 h at room temperature with an HRP-conjugated anti-rabbit antibody (1:2000) in TBS-T containing 5% non-fat dry milk. Specific proteins were visualised by enhanced chemiluminescence reagents using protocols described by the supplier (Phototope-HRP Western detection kit; Cell Signalling Technology) and visualised on Hyperfilm ECL (Amersham, Buckinghamshire, UK). The signals were quantitated by densitometric scanning.
magnification lens. Images from the cells were captured using a Zeiss Axiophot light microscope and a x40 objective. Evaluation of immunoreactivity of LNCaP cells. Slides were prepared by fixing the cells with 3% paraformaldehyde in PBS and washed with 50 mM Tris-HCl (pH 7.4), 150 mM NaCl solution (TBS) containing 0.1% Triton X-100 (TBST). Subsequently, non-specific binding was blocked with 5.5% normal goat serum in TBST for 1 h. Incubation with primary antibody anti-pERK1/2 diluted (1:500 in TBST containing 3% BSA) for 60 min at room temperature. Immunostaining was performed by the avidin-biotinylated peroxidase complex (Vectastain Elite ABC Kit; Vector Laboratories) as chromogen. 3,3'-diaminobenzidine containing nickel chloride (DAB substrate kit for peroxidase; Vector Laboratories). Antigen-antibody complex was visualized by DAB reaction product and counterstained with hematoxylin. The immunostaining was quantitated in a semiquantitative fashion incorporating both the intensity and distribution of specific staining. Scoring was generated as follows: [3 x (% of strongly stained cells)] + [2 x (% of moderately stained cells)] + [1 x (% of weakly stained cells)] + [0 x (% of unstained cells)]. The range of staining intensity was scored as [0 = absent; 1 = weak; 2 = moderate; 3 = strong] staining. Densitometric units of the protein pERK1 and pERK2 were calculated for the densitometric units of total ERK1 and ERK2. The ratio pERK/total ERK from each treated sample was then divided by the value determined under control or EGF conditions to obtain the fold enhancement or reduction of the activated protein.

Immunocytochemistry. In immunocytochemical experiments, LNCaP cells (density: 25,000 cells/ml) and PC-3 cells (density: 13,000 cells/ml) were seeded in standard medium on sterile circular glass coverslips (Ø 18 mm) placed on the bottom of tissue culture plates (353043, Multiwell™ 12 well; Becton-Dickinson and Company, Franklin Lakes, NJ, USA). The procedure for cell culture and treatment was the same as described for the Western blot experiments. However, before treatment, the cells were maintained in serum-free medium for 2 h.

At the end of treatment, the culture medium was removed from the tissue culture plates and cells were washed with PBS. Then they were fixed with 3% paraformaldehyde in PBS and washed with 50 mM Tris-HCl (pH 7.4), 150 mM NaCl solution (TBS) containing 0.1% Triton X-100 (TBST). Subsequently, non-specific binding was blocked with 5.5% normal goat serum in TBST for 60 min at room temperature. Incubation with primary antibody anti-pERK1/2 diluted (1:250) in TBST containing 3% BSA was performed for 24 h at 4°C. After rinsing, cells were incubated with biotinylated goat anti-rabbit secondary antibody (Vector Laboratories) (1:500 in TBST containing 3% BSA) for 60 min at room temperature. Endogenous peroxidase activity was inhibited with 0.3% hydrogen peroxide in TBS for 30 min at room temperature. Immunostaining was performed by the avidin-biotinylated complex (Vectastain Elite ABC Kit; Vector Laboratories). Antigen-antibody complex was visualized by 3,3'-diaminobenzidine containing nickel chloride (DAB substrate kit for peroxidase; Vector Laboratories) as chromogen. After staining, coverslips were dehydrated in ascending ethanol, immersed in xylene and mounted on glass microscope slides.

Densitometry was performed by omitting the primary antibody. 3T3-Swiss albino cells treated with the phorbol ester 12-O-tetradecanoylphorbol-13-acetate (TPA) 160 mM were used as positive control for pERK1/2.

Evaluation of immunoreactivity of LNCaP cells. Slides were examined using a Zeiss Axiophot light microscope and a x40 magnification lens. Images from the cells were captured using a Sony XC77 CCD camera connected to a Macintosh IV computer (Apple Computer, Cupertino, CA, USA) by means of a Scion LG3 frame grabber (Scion Corporation, Frederick, MA, USA). The intensity of staining reactivity was evaluated on a grey scale ranging from 0 (white) to 255 (black). In 100 cells from at least 5-6 random fields per slide. The sampled cell was outlined by hand on the computer screen and the mean grey level was measured. The background grey value was determined in at least four cell-lacking areas near the sampled cell. The specific mean grey value was calculated by deducting the background grey value. Measurements were performed on slides under constant lighting conditions.

Evaluation of immunoreactivity of PC-3 cells. Owing to the peculiar morphology of the attached PC-3 cells as well as the numerous spherical grape-like aggregates arising from them, an evaluation as described for LNCaP cells was impossible. Therefore, PC-3 cells were counted with a Zeiss Axiophot light microscope and the staining intensity of each MAP kinase was graded as absent, weak, moderate or strong. Two independent observers counted at least 500 cells in 5-10 random fields per slide and the percentage of stained cells at each intensity was determined.

The immunohistochemical analysis was scored in a semi-quantitative fashion incorporating both the intensity and distribution of specific staining. Scoring was generated as follows: [3 x (% of strongly stained cells)] + [2 x (% of moderately stained cells)] + [1 x (% of weakly stained cells)] + [0 x (% of unstained cells)]. This gave a possible range of 0-300. Scoring was performed on high-power fields (x400).

Data analysis. For each dose-response curve, at least two proliferation experiments in triplicate were performed. Data of cell growth were expressed as mean ± SE. All Western blot analyses were conducted at least twice on different occasions, and the most representative results are shown.

Proliferation, Western blotting and PC-3-immunocytochemistry data were analyzed using the Student's t-test to determine the significance of differences between two populations. Immunocytochemical data in LNCaP cells were analyzed by one-way ANOVA followed by Tukey's multiple comparison test. A value of p<0.05 was always considered statistically significant.

Results

Cell proliferation. We previously reported that LA, added daily to the culture medium, did not induce any variation in the growth of either LNCaP or PC-3 cells, even at the highest concentration tested (10−5 M). Nevertheless, in LNCaP cells at doses of 10−1-10−4 M, it inhibited the DHT-stimulated growth, which was suppressed by the highest concentrations (10−3-10−2 M) of the analogue used (6).

In the present study, we focused our attention on the mitogenic effects of EGF (0.1-50 ng/ml) alone or associated with LA (10−11-10−8 M ) in both models. In LNCaP cells, plated at the lower density, a 2-day treatment with EGF at concentrations of 25 and 50 ng/ml determined a statistically significant increase in cell numbers of ~20% with respect to control (not shown). After 4 days,
the stimulation of cell growth started from the dose of 5 ng/ml EGF and reached ~25% with respect to control at the concentration of 50 ng/ml (Fig. 1A). The results were similar to those obtained when cells were plated at the higher density (not shown).

In PC-3 cells, seeded at an initial density of 25,000 cells/ml of culture medium, no response to EGF was observed after 2 days (not shown), while a statistically significant stimulation of ~15% was seen at EGF concentrations ranging from 5 to 50 ng/ml of culture medium after 4 days. No difference in the effectiveness seemed to exist among the concentrations tested (Fig. 1B).

On the basis of these results, we used the concentration of 10 ng/ml EGF, which stimulated cell growth in both cell lines, to investigate the effect of the combination with LA.

When LNCaP cells were seeded at the lower density, the EGF stimulation was reduced by 50% (p<0.05) with 10^-11 M LA and by 80% with 10^-5 M LA (p<0.01) (Fig. 1C). When cells were seeded at 50,000 cells/ml, it was completely suppressed by concentrations of LA from 10^-11 to 10^-5 M (p<0.001) (not shown). In PC-3 cells, LA at both low and high concentrations produced a high reduction of EGF-mitogenic activity (77-80%) (Fig. 1D).

LA was also able to significantly counteract or abolish the DHT or EGF-stimulated proliferation of both LNCaP and PC-3 cells in the conditions in which ERK activity was evaluated (see Materials and methods), as shown in Fig. 2, which refers to the second day of treatment. Moreover, in LNCaP cells, stimulated with EGF, high doses of LA determined a diminution in cell number which decreased below the controls (Fig. 2, middle panel). In all experiments, cell viability was higher than 90%.

Expression of phosphorylation status independent and phosphorylated p44/42 in untreated LNCaP and PC-3 cells.

To determine the presence of both phosphorylated and total ERK1/2, LNCaP and PC-3 cells were grown in standard medium until they reached sub-confluence, then they were placed in a low-serum medium which was subsequently changed with serum-free medium to abolish ERK activation due to factors contained in the serum or produced by the cells.

To assess the presence of total ERK1/2 in both androgen-sensitive and -insensitive prostate cancer cells, a phosphorylation status independent p44/42 MAP kinase antibody was used. Western blot analysis showed that these proteins are highly expressed in both LNCaP and PC-3 cells at the correct molecular weight. No activation of ERK1 and 2 was observed in LNCaP cells, while ERK1 and 2 were constitutively activated and equally expressed in PC-3 cells (Fig. 3, top panel).

Phosphorylated ERK1/2 expression was also evaluated by immunocytochemistry to confirm our findings from Western blot analysis and to determine their intracellular localization. It is important to note that immunocytochemistry could not discriminate between ERK1 and 2 or indicate the rate of activated enzymes with respect to total ERK.

In contrast with data from Western blotting, LNCaP cells displayed a heterogeneous pattern of pERK1/2 staining and, even if most of the immunoreactivity was localized in the
nucleus, it was also observed in the cytoplasm in many cells, as depicted in Fig. 3 (bottom panel). The same pattern of immunoreactivity was observed in PC-3 cells (Fig. 3, bottom panel). In the negative controls, no staining was present (Fig. 3, bottom panel).

ERK activity in LNCaP cells treated with hormones and EGF

Western blots indicated that the addition of DHT or LA to LNCaP cells did not promote ERK activation (data not shown). Otherwise, some modifications were found by using immunocytochemical analysis. In fact, after 10 min of treatment with \(10^{-9}\) M DHT, an increase of 16.7% (\(p<0.001\)) in the intensity of the reaction compared to the control was observed (Fig. 4). Treatment with \(10^{-11}\) or \(10^{-6}\) M LA alone did not change the pERK1/2 expression (not shown).

On the contrary, the combination of \(10^{-11}\) or \(10^{-6}\) M LA with DHT, determined a slight but statistically significant (\(p<0.001\)) reduction (13.3% and 11.9% respectively), in the expression of ERK1/2 obtained with the androgen alone (Fig. 4). The stimulation induced by DHT was substantially suppressed. Similar results were seen when cells were pre-treated with the analogue and then exposed to androgen. No relevant modification of staining was observed at the other times examined (data not shown).

There was a substantial coincidence of the results obtained by both Western blotting and immunocytochemistry when cells were exposed to EGF alone or in combination with LA.

The time course of ERK activation was examined by Western blotting in LNCaP cells stimulated with EGF for multiple time points, ranging from 5 min to 4 h. ERK activation was detected early (5 min after the initiation of EGF treatment),
ERK2 being more expressed than ERK1. The activity peaked by 60 min, and decreased over the next 4 h. Fig. 5A refers to data from three separate curves and a representative blot is shown (Fig. 5B). To provide evidence that EGF receptor phosphorylation was required for ERK activation, cells were 30-min pre-treated with the specific inhibitor of the tyrosine kinase of the EGF receptor (300 nM AG1478). After 5 min, a 90% inhibition of EGF-induced ERK activity was observed which gradually diminished until 30 min or 1 h. The AG1478 effect disappeared after 4 h (Fig. 5C), when ERK activation was higher than that observed in cells treated with EGF alone.

Figure 4. Immunocytochemical staining for pERK1/2 in LNCaP cells using anti-phospho-p44/42 MAP kinase, (Thr202/Tyr204) polyclonal antibody. Immunocytochemistry was performed after 10 min of treatment with 10^{-6} M DHT, 10^{-6} M DHT plus 10^{-11} M LA, 10^{-6} M DHT plus 10^{-6} M LA or in untreated cells. Original magnification x400.

Figure 5. Time course (from 5 to 240 min) of ERK activation in LNCaP cells treated with 10 ng/ml EGF. (A) Total ERK and pERK levels were analyzed by immunoblot assay. pERK levels were quantified by densitometry and standardized against the levels of total ERK per sample. The resulting values were normalized against the values from EGF-treated cells set as 1. Values are the mean ± SE of two individual experiments. Immunoblot from a representative experiment is shown below. Untreated cells (lane 1), cells pre-exposed to the inhibitor, AG1478, and then treated with 10 ng/ml EGF (lane 2), 10 ng/ml EGF (lane 3), 10 ng/ml EGF plus 10^{-11} M LA (lane 4), 10 ng/ml EGF plus 10^{-5} M LA (lane 5), 10 ng/ml EGF plus 10^{-6} M LA (lane 6). **p<0.01, Student's t test versus EGF-treated cells.

Figure 6. Western blot analysis of pERK and total ERK expression in LNCaP cells exposed to different treatments for 5 min (top panel) or for 10 min (bottom panel). Total ERK and pERK levels were analyzed by immunoblot assay. pERK levels were quantified by densitometry and standardized against the levels of total ERK per sample. The resulting values were normalized against the values from EGF-treated cells set as 1. Values are the mean ± SE of two individual experiments. Immunoblot from a representative experiment is shown below. Untreated cells (lane 1), cells pre-exposed to the inhibitor, AG1478, and then treated with 10 ng/ml EGF (lane 2), 10 ng/ml EGF (lane 3), 10 ng/ml EGF plus 10^{-11} M LA (lane 4), 10 ng/ml EGF plus 10^{-5} M LA (lane 5), 10 ng/ml EGF plus 10^{-6} M LA (lane 6). **p<0.01, Student's t test versus EGF-treated cells.
To assess the role of ERK activation in the LA reduction of cell proliferation induced by EGF, LNCaP cells were treated with LA plus EGF or were pre-incubated for 15 min with the analogue prior to EGF exposure. When LA at doses of $10^{-11}$ M and $10^{-6}$ M was combined with EGF, a reduction in ERK activation was seen. This effect was evident with both of the LA concentrations used after 5 min, but the lower dose of the analogue was more effective; 30% inhibition for ERK2 and 40% for ERK1 (Fig. 6, top panel). The analogue at the lower dose continued to determine, after 10 min, a higher reduction in ERK activation (38-60%), (Fig. 6, bottom panel). The reduction produced by the analogue in ERK activation persisted after 30 and 60 min (data not shown). It is worth mentioning that the highest dose of LA employed ($10^{-5}$ M) did not affect ERK activity.

Of note, the signal given by Western blotting referring to cells exposed to EGF was almost undetectable after 4 h, but it seemed to be nullified by the analogue (not shown).

If LNCaP cells were pre-incubated for 15 min with the analogue and then exposed to EGF, a trend similar to that produced by AG1478 was observed. In fact, LA was able to counteract the EGF-induced ERK phosphorylation within 5-30 min (~30% inhibition). The effect of the analogue disappeared after prolonged treatment (60 min) and, after 4 h, ERK phosphorylation exceeded that seen with EGF alone. In this condition, $10^{-3}$ M LA produced a reduction in ERK activation with a behaviour similar to that displayed by the other concentrations after 5-30 min of treatment (Fig. 7).

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Similar results were obtained with immunocytochemistry. Fig. 8 shows a representative experiment after 30 min of exposure to different compounds.

**ERK activity in PC-3 cells untreated or treated with LA and EGF.** In PC-3 cells, which express basal levels of pERK, the analogue generally determined an increase in this activity. The enhancement in phosphorylation status, induced by both the concentrations used, peaked after 4 h of exposure (>100%) (Fig. 9).

In PC-3 cells, the addition of 10 ng/ml EGF resulted in a rapid stimulation of ERK phosphorylation. The enhancement was evident after 5 min, peaked after 60 min, and persisted until 4 h (Fig. 10A and B).

As with LNCaP cells, pre-exposure of PC-3 cells to AG1478 counteracted the EGF-stimulated activity. This was evident after 5 min (>70% inhibition), persisted until 60 min (40-60%), and was lost after 4 h (Fig. 10C).

When LA and EGF were combined, a more pronounced increase in ERK activity was generally observed, with respect to that observed with EGF or LA alone, using both Western blot analysis and immunocytochemistry. The enhancement was observed in both modalities of treatment (simultaneous or in sequence) and, in some cases, ERK phosphorylation was >5-fold with respect to control after 1 h of treatment. Fig. 11 refers to Western blotting (top panel) and immunocytochemistry (bottom panel) results, after 60 min of treatment.

Of note, there was no change in the localization of immunostaining in both LNCaP and PC-3 cells due to the different treatments.

**Discussion**

The present study demonstrates that LNCaP and PC-3 cells respond to EGF with an increase in cell proliferation, as measured by cell counting. The stimulation that we obtained in our experiments concerning LNCaP cells is lower than that observed in the majority of studies, which show that the increase in cell number was generally >25% with respect to control (10,12,15,16).

The evidence in the literature concerning EGF activity in PC-3 cells remains contradictory and no clear effect emerges. Some authors described an increase (4,10,17-19) in cell growth after EGF treatment while, in certain cases, no effect on proliferation was observed (17,20,21). The divergent responses to EGF in PC-3 cells and the differences in the mitogenic effect in LNCaP cells may reflect the differences in cell passages, cell density, culture conditions or EGF concentrations used.

The increase in cell number found in PC-3 cells after EGF exposure is similar to that which we reported in our previous manuscript (4).

To establish the role of the modulation of kinase phosphorylation in the ability of LA to counteract the stimulation of cell growth produced by DHT or EGF, ERK activity was evaluated in LNCaP and PC-3 cells untreated or after different treatments.

Our data from Western blotting clearly demonstrate that untreated androgen-sensitive LNCaP cells in serum-free conditions is lower than that observed in the majority of studies, which show that the increase in cell number was generally >25% with respect to control (10,12,15,16).

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Consistent with other data in the literature in which other analogues or other models have been used (22,23), Leuprorelin acetate inhibited DHT- and EGF-induced proliferation in both LNCaP and PC-3 cell lines, also when very low concentrations are used (10⁻¹¹ M) and in all conditions tested; i.e. presence of 5% or 0.5% CH-FBS.

Many reports indicate that the MAP kinase pathway may be involved in the mitogenic action of hormones and growth factors in prostate cancer as well as in other kinds of tumour.
We assessed that LA alone did not determine ERK activation in LNCaP cells; on the other hand, the analogue did not modify cell growth. The slight increase in pERK expression that we observed by immunocytochemistry after DHT treatment could not be confirmed by Western blotting, hence it is not possible to draw any conclusions in agreement with data in the literature. In fact, ERK activation following androgen treatment is a point of controversy among various authors, some of whom found an increase (9,25,32) while others reported no variation (10,27,31).

Figure 11. Top panel, Western blot analysis of pERK and total ERK expression in PC-3 cells exposed to different treatments for 60 min. Total ERK and pERK levels were analyzed by immunoblot assay. pERK levels were quantified by densitometry and standardized against the levels of total ERK per sample. The resulting values were normalized against the values from untreated cells set as 1. Values are the mean ± SE of two individual experiments. Immunoblotting from a representative experiment is shown below. Untreated cells (lane 1), cells pre-exposed to inhibitor (AG1478) and then treated with 10 ng/ml EGF (lane 2), 10 ng/ml EGF (lane 3), 10 ng/ml EGF plus 10^{-11} M LA (lane 4), 10 ng/ml EGF plus 10^{-9} M LA (lane 5), cells pre-exposed to 10^{-6} M LA (lane 6) or 10^{-4} M LA (lane 7), and then treated with 10 ng/ml EGF, 10^{-11} M LA (lane 8), 10^{-9} M LA (lane 9). *p<0.05, †p<0.01; Student’s t-test versus control cells. ‡p<0.05, ††p<0.01, Student’s t-test versus EGF treated cells. §p<0.05, †††p<0.01, Student’s t-test versus EGF treated cells. Bottom panel, immunocytochemical staining for pERK1/2 in PC-3 cells using anti-phospho-p44/42 MAP kinase, Thr202/Tyr204 polyclonal antibody. Immunocytochemistry was performed after 60 min with the treatments reported above. Negative control was obtained in the absence of primary antibody. Original magnification x400.
On the other hand, EGF determined an increase in ERK activation with both Western blotting and immunocytochemistry as well as mitogenic activity in LNCaP cells and our data are in agreement with those reported in the literature (10,15,24,27,30,31,33).

Consistent with results obtained by other authors, the decrease in ERK activation following AG1478 treatment supports the idea that EGF receptor tyrosine kinase activity may be an intermediate in rapid EGF-induced ERK activation (10).

More interestingly, we found that LA, particularly at a low concentration, induced a reduction in ERK activation produced by EGF after a brief treatment. However, the effect was more evident concerning ERK1, as revealed by Western blot analysis. A reduction of ERK activity induced by DHT was seen by immunocytochemistry.

Our findings are in complete agreement with a report by Emons et al (23), who showed that the 5-fold increase in ERK activity induced by EGF in ovarian and endometrial cancer cell lines was nullified by pre-exposure to triptorelin, another GnRH agonist. It is worth mentioning that EGF produced an increase in cell numbers that was <30% in 2 of the models used.

As far as PC-3 cells are concerned, EGF determined an increase in ERK activity, as expected, in accordance with other authors (10,24). In spite of the analogue inefficacy on cell proliferation, an induction of ERK activity occurred after 60 and 240 min of LA treatment. Still more surprising was the pERK increase produced by the LA/EGF combination, whereas the analogue showed antagonizing properties on EGF mitogenic activity. It may be hypothesized that both these events may reflect a protracted ERK activation after LA treatment. Transient activation of MAP kinase is thought to be associated with cellular proliferation, whereas prolonged activation seems to be involved in differentiation. In this context, Reiss et al (34) observed that, in pituitary αT-3-1 cells, EGF, TPA and triptorelin induced ERK activation. Maximal activity after triptorelin and TPA treatment persisted for 60 min and dropped to near-basal levels only after 2-4 h of incubation whereas activation by EGF was shorter and dropped to basal levels within 20 min. Recently, Kraus et al (35) have shown that triptorelin induced ERK activation and, at the same time, apoptosis in a human androgen unresponsive prostate cancer cell line (DU-145). Moreover, Kimura et al (36) demonstrated that LA, which had antiproliferative activity in a human ovarian cancer cell line, Caov-3, activated ERKs and its effect was sustained until 24 h. Finally, it has been reported that other agents, such as phenethyl isothiocyanate, which has apoptotic activity, induce prolonged activation of ERK1/2 in PC-3 cells (26).

There would seem to be a difference concerning the involvement of ERK activation in the effect of LA on cell proliferation of LNCaP and PC-3 cells stimulated by mitogens. This might be due to the different expression of pERK1 and 2 in these cell lines. In fact, PC-3 cells expressed constitutive ERK1/2 activation while LNCaP cells did not. Recently, Unni et al (9) reported that a constitutively active kinase pathway in LNCaP cells is associated with their transition to androgen independence. The presence of this constitutively active kinase pathway is related to unresponsiveness to manipulation with androgen receptor agonists or antagonists. Of note, in our experience, pERK2 was more expressed than pERK1 in LNCaP cells while, in PC-3 cells, both proteins were expressed at the same level. The role of the two ERK isoforms remains to be clarified and few papers have been dedicated to this topic. Reiss et al (34) showed that a GnRH analogue preferentially activated the ERK1 isoform, whereas other compounds (i.e. doxorubicin) were reported to especially activate ERK2 (37).

In some cases, ERK1 might be involved in the regulation of differentiated cellular functions while ERK2 might mainly regulate cell proliferation (35) but, in other cases, ERK1 activation seems to be an important determinant in cell transformation (38). Moreover, ERK2 activation may result in growth arrest and apoptosis (37) or in increased proliferation (39).

In conclusion, the findings presented in this manuscript indicate that the inhibitory effect of GnRH analogues in the presence of mitogens may be linked to interference with growth factor signal transduction. The ERK pathway seems to be involved in this phenomenon, the activation being dependent on the cell characteristics (androgen-responsiveness or androgen-unresponsiveness). The possibility of obtaining the mentioned effects in vitro with a very low concentration of LA suggests that these effects might also be obtained in vivo, which seems interesting in view of the potential use of the analogue in hormone-refractory prostate cancer.

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References


