Expression of melanocortin receptors in human prostate cancer cell lines: MC2R activation by ACTH increases prostate cancer cell proliferation

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Abstract. The melanocortin receptors (MCRs 1-5) are G protein coupled-receptors (GPCRs) that regulate food intake, inflammation, skin pigmentation, sexual function and steroidogenesis. Their peptide ligands, the melanocortins, are α-, β- and γ-melanocyte-stimulating hormone and adrenocorticotropic hormone (ACTH) all of which are secreted from the anterior pituitary gland under hypothalamic control. MC2R binds ACTH but has no affinity for the other melanocortins and is, thereby, pharmacologically different from MCRs that bind those ligands. Evidence suggests that elevated GPCRs transactivate the androgen receptor (AR), the critical mediator of prostate cell growth, and consequently promote prostate cancer cell proliferation. It may be that reduced central melanocortin signaling is coincidental with reversal of prostate cancer cachexia, but no data are available on the expression of, or the role for, MCRs in prostate cancer. Here, we show that MCR (1-5) mRNAs are expressed in androgen-dependent LNCaP and androgen-independent PC3 and DU-145 human prostate cancer cell lines. Further, MC2R, the specific target of ACTH, is expressed in LNCaP, PC3 and DU-145 cells. Among the several synthetic MCR peptide ligands that we used, only ACTH promoted concentration-dependent cell proliferation in the three cell lines as shown by MTT cell proliferation assay. In LNCaP cells, the effect was additive with testosterone stimulation and was partially blunted with SHU9119, a non-selective MCR antagonist. In the same cells, ACTH induced cAMP production and increased AR nuclear labeling in immunocytochemical assays. Our observations suggest that MC2R is involved in prostate carcinogenesis and that targeting MC2R signaling may provide a novel avenue in prostate carcinoma treatment.

Introduction

Prostate cancer (PCa) is a leading cause of cancer mortality among men in the United States and is the second leading cause of cancer mortality in men worldwide (1). Current treatments include hormonal therapy, targeted radiation, and radical prostatectomy. The only alternative is watchful waiting. The limitations and side-effects associated with these therapies, however, warrant the development of new therapies (2).

Testosterone (T) regulates normal cell growth in the prostate gland and is secreted by testicular Leydig cells. The action of T is mediated by the androgen receptor (AR), a nuclear receptor expressed in the stromal and luminal prostate gland epithelial cells. Androgen-dependent and independent neoplastic PCa cells depend on T, its more potent 5 α-reductase metabolite, 5 α-dihydrotestosterone (DHT), and AR (3). Huggins (4) demonstrated PCa dependency on T and, based on his early findings that showed reduced prostate growth following removal of androgen, local and metastatic cancers are currently treated with endocrine therapies aimed at reducing androgen production and/or blocking AR with anti-androgens. These therapies limit tumor growth and reduce tumor mass but the cancer may recur as androgen-insensitive (AI) disease (5).

Multiple direct and indirect mechanisms are thought to cause the AI state through changes in AR activity (6-8). These
changes include AR signaling modulation by a kinase receptor such as human epidermal growth factor receptor 2 (HER-2/ neu) tyrosine kinase (9); overexpression of, or mutations in, the AR gene (10); increased AR sensitivity (11); or activation of the AR by growth factors such as insulin-like growth factor 1, epidermal growth factor and keratinocyte growth factor (6,12). Additional indirect and equally important mechanisms by which PCa cell growth circumvent androgen-dependency involve overexpression of GPCRs (13,14). An effective cure for AI is currently unavailable despite the intense research in this area (15).

Since mutations in the AR are responsible for a small number of prostate cancers, indirect mechanisms are likely to be involved in the majority of androgen-refractory cases. One such mechanism is activation and overexpression of GPCRs (8). Several studies showed that GPCRs are overexpressed in various cancer types and contribute to tumor growth when activated by circulating or locally produced peptide ligands (13). These ligands induce cell proliferation and prevent apoptosis of hormone-independent PCa cells, which suggest a critical role in PCa progression (14,16-20). Normal GPCR functions are altered in malignant cells leading to autonomous cell proliferation (13). Hence, GPCR-targeted therapy for the treatment of PCa is actively pursued (21).

GPCR actions in PCa involve cell growth enhancement via ERK6, a mitogen-activated protein kinase-related serine/threonine kinase (MAPK)(22). ERK can phosphorylate AR at several sites thereby upregulating AR target genes (13). GPCRs that signal through stimulatory G proteins α (Gsa) can also transactivate AR through cyclic adenosine monophosphate (cAMP) production (23,24) and protein kinase A (PKA)(25). In addition, the activated Gsa subunit can directly activate AR and act synergistically with low androgen concentrations to increase AR activity in PCa cells (24).

Several studies suggest a role for at least some of the MCRs in cancer development. An association between MC1R and PCa risk was reported in human (26) and central MCR blockade with SHU9119-reversed PCa anorexia in Wistar rats (27). Additionally, MC5R is widely expressed in normal rodent exocrine glands including the prostate (28).

MC2R, activated by ACTH, is involved in androgen and steriodogenic enzyme synthesis. It is normally expressed in the adrenal gland cortex and epidermis as well as in basal cell carcinoma tissue and cultured melanocytes. As with other GPCRs, MC2R binding ACTH activates adenyl cyclase with subsequent production of cAMP. Cyclic AMP can then directly activate or interact with various proteins such as ion-channels, transcription factors, guanine nucleotide exchange factors, and protein kinases such as PKA. PKA and cAMP play critical roles in the progression and differentiation of prostate carcinogenesis (25). Activation of MC2R can also result in phosphorylation of MAPK and the MC2R accessory protein isofrom β (MRAP-β) in cells stably expressing MC2R (29). Similarly, ectopic production of ACTH, the putative ligand for MC2R, has been reported in cases of prostate carcinoma associated with Cushing's syndrome (30).

Currently, the role of MCRs in PCa is unclear. The objective of this study was to investigate the MCRs expression in human PCa cell lines and their effect on cell proliferation, with particular emphasis on MC2R.

**Materials and methods**

**Reagents.** ACTH and DHT were obtained from Sigma (St. Louis, MO). Melanocortin peptides, NDP-MSH (a potent α-MCR analogue for all MCRs except MC2R) and THIQ (highly selective MC4R agonist), D-Trp<sup>5</sup>-γ-MSH (highly selective MC3R agonist), and SHU9119 (a universal MCR blocker) were obtained from Tocris Bioscience, Ellisville, MO. A selective MC4R inverse agonist (Ipsen 5i) was synthesized by Enzo Life Sciences International Inc., Plymouth Meeting, PA. RPMI-1640 media with and without phenol red, Dulbecco's phosphate buffered saline (DPBS), without calcium and magnesium, Forskolin (cAMP inducer), and penicillin streptomycin antibiotics were purchased from Lonza (Portsmouth, NH). Heat-inactivated and charcoal-dextran-treated fetal calf serum (FCS) was obtained from Atlanta Biological (Atlanta, GA). Cell culture flasks and other disposable cell culture supplies were purchased from VWR International, LCC (Atlanta, GA). MTT cell viability assay kits were purchased from the American Type Culture Collection (ATCC). Primary antibodies and control peptides for MC2R (SA-639), and AR (Sc-816) were obtained from Enzo Life Science (Farmingdale, NY) and Santa Cruz Biotechnology (Santa Cruz, CA), respectively. Dylite 488 conjugated goat anti-rabbit IgG (Jackson Immunoresearch Laboratories, PA) was used for the detection of primary binding in immunocytochemistry assays.

**Cell culture.** The human prostate normal epithelial cells, RWPE-1, and human prostate adenocarcinoma cell lines, LNCaP, PC3, and DU-145, were obtained from the American Type Culture Collection (Manassas, VA). The RWPE-1 cells were propagated in base medium containing bovine pituitary extract and human recombinant epidermal growth factor (Invitrogen, Carlsbad, CA). PCa cell lines were maintained in RPMI-1640 media supplemented with 1% (v/v) streptomycin-penicillin antibiotics (Invitrogen), and 10% (v/v) heat-inactivated FCS. Cells were grown in T-75 vented filter cap tissue culture flasks until they reached approximately 75-90% confluency at 37°C in a humidified incubator under a 5% CO<sub>2</sub> atmosphere. Detachment of the cells from the T-75 flasks was accomplished with trypsin after the cells were washed with DPBS to remove residual serum. Cells were then counted using an automated coulter cell counter and cultured at the desired concentration. For culture cell proliferation/viability experiments, ligands were diluted in PBS-BSA and added in 20 µl volumes containing various concentrations. Twenty-four hours before treatments, cells were depleted of androgen by replacing growth media with phenol red-free RPMI-1640 supplemented with 5% charcoal-dextran-treated FCS. Treatment peptides were replenished every 24 h for 72 h. Cells were seeded at a density of 2x10<sup>5</sup> cells per well when grown in 6-well culture plates (for protein and RNA isolation), and at 2x10<sup>4</sup> cells per well when grown in flat-bottom 96-well plates for MTT cell viability assay. Passage number was recorded with each split and cells were used between passages 5 and 38.

**MTT cell viability assay.** Cells in log-phase growth were harvested via trypsinization and counted using a coulter counter. The effect of test drugs on cell viability/
proliferation was determined using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) cell viability assay ATCC kit according to the manufacturer's instructions. This colorimetric assay measures the reduction of a tetrazolium component (MTT) into an insoluble formazan product by the mitochondria of metabolically active cells. Forty-eight to 72 h after test drugs were added (eight replicates per treatment), 10 µl MTT reagent was added per well for 2 h at 37°C. Next, 100 µl detergent solution was added to lyse the cells and solubilize colored crystals. Plates were then incubated in the dark for 6 h at room temperature (RT). Optical density (OD) was obtained using Spectramax Plus microtiter plate reader at 570 nm wavelength. The amount of purple color produced in this test is directly proportional to the number of viable and metabolically active cells. The effect on cell proliferation/viability was calculated based on three independent experiments.

Conventional and real-time PCR. Total RNA was isolated using TRIzol reagent (Invitrogen-Life Technologies Inc., Carlsbad, CA, USA), according to the manufacturer's protocol and as described in our previous study (31). RNA concentrations were determined at 260 nm wavelength and the ratio of 260/280 was obtained using UV spectrophotometry (DU640, Beckman Coulter, Fullerton, CA, USA). Samples with 260/280 ratio of ≥1.8 were used. First strand synthesis was performed with SABiosciences first strand kit C-03 (SABiosciences, Frederic, MD, USA). Conventional PCR was used initially to determine the presence of MCR (1-5) genes in LNCaP, PC-3, and DU-145 using validated human primer sets obtained from SABiosciences (Table I). RT-PCR was performed using ReactionReady™ HotStart ‘Sweet’ PCR master mix (SABiosciences). PCR products were analyzed in parallel with GAPDH housekeeping gene. Final end PCR products were fractionated and visualized on 2% ethedium bromide-stained agarose gel. Next, real-time PCR was used to determine the expression level of the MCRs (1-5) in LNCaP, PC-3 and DU-145 cancer cells relative to RWPE 1 normal prostate epithelial cells. The assay was carried out as we previously described (31). Briefly, cell lysates were prepared by homogenization in RIPA buffer (150 mM sodium chloride, 50 mM Tris-HCl, pH 7.4, 1 mM ethylenediaminetetraacetic acid, 1 mM phenylmethylsulfonyl fluoride, 1% Triton X-100, 1% sodium deoxycholic acid, 0.1% sodium dodecylsulfate) containing 5 µg/ml of aprotinin, and 5 µg/ml of leupeptin. Cell debris was removed by centrifugation. Protein concentration was determined with the Bio-Rad protein assay. Cell lysates containing 5 µg/ml of aprotinin, and 5 µg/ml of leupeptin. Cell lysates containing 5 µg/ml of aprotinin, and 5 µg/ml of leupeptin. Cell lysis was performed by equilibration in PBS (3x for 3 min each). Before adding the primary antibody, cells were incubated in appropriate blocker (5% normal goat serum, 2% BSA, in PBS, pH 7.3). Rabbit polyclonal anti-MC2R (extracellular) and anti-AR (nuclear) were incubated (1:100) in blocking solution overnight at RT. Secondary, fluorescence-labeled goat anti-rabbit, IgG antibody was incubated at RT for 1 h. Slides were mounted with Vectashield mounting medium and the preparations were sealed with clear nail polish. Epifluorescence and DIC images were acquired with a Nikon E600 microscope equipped with a Retiga EX CCD digital camera (Q Imaging, Burnaby, Canada). Images were processed and saved with NIS-Elements AR 3.2 software. Exposure times were identical among the treatment groups. Confocal images were captured with a Nikon TE2000E confocal microscope (Nikon, Mississauga, Canada) operated with EZ-C1 3.9i software.

Cyclic AMP. ACTH-induced cAMP was determined in three PCa cell lines. Briefly, PC3, DU-145, and LNCaP cells were plated in 6-well plates and allowed to attach for 16 h. The following day, the cells were washed twice with warm Waymouth/BSA. Then fresh Waymouth/BSA containing 0.5 mM isobutylmethylxanthine (IBMX) (a cyclic AMP promoter

<table>
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<th>Gene symbol</th>
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<th>RefSeq accession no.</th>
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<tr>
<td>MC5R</td>
<td>90</td>
<td>NM_005913.1</td>
<td>103-121</td>
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Western blotting (WB). WB was performed as previously described by us (31). Briefly, cell lysates were prepared by homogenization in RIPA buffer (150 mM sodium chloride, 50 mM Tris-HCl, pH 7.4, 1 mM ethylenediaminetetraacetic acid, 1 mM phenylmethylsulfonyl fluoride, 1% Triton X-100, 1% sodium deoxycholic acid, 0.1% sodium dodecylsulfate) containing 5 µg/ml of aprotinin, and 5 µg/ml of leupeptin. Cell lysates were boiled for 5 min in 1X SDS sample buffer (50 mM Tris-HCl pH 6.8, 12.5% glycerol, 1% sodium dodecylsulfate, 0.01% bromophenol blue) containing 5% β-mercaptoethanol. Adrenal cell lysate (positive control for MC2R) was obtained from VWR.

Immunofluorescence microscopy. Cells were grown on Lab-Tek II chamber slides for 24 h. Appropriate treatments were added as described in the figure legends. Cells were then rinsed 3x in PBS and fixed in buffered 4% paraformaldehyde (30 min). Following fixation, cells were rinsed 2x in dH2O and transferred to absolute methanol for 10 min followed by equilibration in PBS (3x for 3 min each). Before adding the primary antibody, cells were incubated in appropriate blocker (5% normal goat serum, 2% BSA, in PBS, pH 7.3). Rabbit polyclonal anti-MC2R (extracellular) and anti-AR (nuclear) were incubated (1:100) in blocking solution overnight at RT. Secondary, fluorescence-labeled goat anti-rabbit, IgG antibody was incubated at RT for 1 h. Slides were mounted with Vectashield mounting medium and the preparations were sealed with clear nail polish. Epifluorescence and DIC images were acquired with a Nikon E600 microscope equipped with a Retiga EX CCD digital camera (Q Imaging, Burnaby, Canada). Images were processed and saved with NIS-Elements AR 3.2 software. Exposure times were identical among the treatment groups. Confocal images were captured with a Nikon TE2000E confocal microscope (Nikon, Mississauga, Canada) operated with EZ-C1 3.9i software.
and a non-specific inhibitor of cAMP and cGMP phosphodiesterases, Sigma) was added to each well. After 15 min of incubation at 37˚C, different concentrations of ACTH or [Nle<sup>4</sup>, D-Phe<sup>7</sup>]-α-melanocyte stimulating hormone (NDP-MSH) were added and the cells were incubated for another hour. Final concentrations of ligands are indicated in Fig. 7. Intracellular cAMP was extracted using 0.5 N perchloric acid containing 180 µg/ml theophylline (Sigma), and measured using radioimmunoassay (32). All determinations were performed in triplicate. The cAMP levels were calculated as fold increase over basal levels of each cell line. Maximal response (Rmax) and EC<sub>50</sub> values were calculated with GraphPad software.

Statistical analysis. Data from cell culture MTT cell viability assay were expressed as the mean ± SD. The data were analyzed by Student’s t-test, one-way ANOVA followed by Dunnett’s post hoc test (Graph Pad Prism 5.0, San Diego, CA). P<0.05 was considered to indicate statistically significant differences. Statistical analysis for the real-time PCR data was performed using a modification of the ΔΔCt method (ΔΔCt) as described in our previous study (33).

Results

MCR (1-5) transcripts are expressed in PCa cell lines. We used conventional and real-time RT-PCR analysis to determine that MCR (1-5) transcripts are expressed in LNCaP and PC3 cells. DU-145 cells expressed the MCR 1-3, and 5 transcripts, but did not express MC4R mRNA (Fig. 1A). Comparative real-time PCR analysis of MCR (1-5) showed higher expression in cancer cells compared with RWPE-1 normal prostate epithelial cells (Fig. 1B). Similar to conventional RT-PCR data, the MC4R level in DU-145 cancer cells was undetectable by the real-time PCR technique. Table I shows the expected PCR product size and the accession numbers of MCR genes in Genbank.

ACTH (MC2R specific ligand) increases cell proliferation in human PCa cells and SHU9119 (a universal MCR blocker) partially blunts this effect. Because increased GPCRs expression in PCa cells was thought to provide cell survival signal in PCa (34). We used several MCR synthetic peptide agonists and antagonists in cell viability/proliferation assays to test the function of the MCRs in PCa cells. These compounds were: NDP-MSH, the potent α-MSH analogue for all MCRs except MC2R; THIQ, the highly selective agonist of MC4R; D-Trp<sup>8</sup>-γ-MSH, the highly selective agonist of MC3R; SHU9119, a universal MCR blocker, and IPSEN i5, MC4R inverse agonist, did not alter cell viability/proliferation. Values are representative of three experiments. Similar results for ACTH were obtained in PC3 and DU-145 (see Fig. 3).
Figure 3. (A) ACTH increases cell proliferation in a dose-dependent manner in androgen-dependent LNCaP, and androgen independent PC3 and DU-145. Cell viability was determined by MTT viability/proliferation assay using four ACTH concentrations. Values are representative of three experiments. (B) Representative MTT assay plate image showing concentration-dependent increase in cell proliferation (indicated by conversion of MTT to purple color) in LNCaP, PC3, and DU-145. Note the color change from light yellow in wells treated with PBS/BSA vehicle to purple in ACTH/BSA-treated wells. (C) Kinetics of PC3 proliferative responses to ACTH stimulation overtime. Cells were treated with $10^{-5}$ ACTH for 2, 4, 6, or 8 days. (D) ACTH-induced cell proliferation was partially reversed by MCR SHU9119 (a universal MCR blocker) in LNCaP cells. DHT is a potent AR activator and was used as positive control for cell proliferation.

Figure 4. (A) Immunohistochemical localization of MC2R in PC3 using rabbit anti-melanocortin receptor 2 (extracellular) antibody captured with fluorescence-labeled goat anti-rabbit secondary antibody. Note specific cell surface expression of MC2R in panel (1). Insert (panel 2) is the magnification of a cell membrane to show punctuated and discrete plasma membrane sub-domains. Little or no staining is present in negative control (minus primary Ab, panel 3). (B) MC2R protein is present in LNCaP lysate as shown by western blotting and was masked by MC2R specific peptide blocker. Lane 1, MC2R protein; lane 2, MC2R band was partially abolished when MC2R antibody was pre-incubated with a control MC2R peptide; lane 3, positive MC2R control (35 µg adrenal cell lysate). (C) MC2R protein in LNCaP, PC3, DU-145 PCa cell lines and RWPE1 normal prostate cells. Lane 1, adrenal lysate (positive control); lane 2, LNCaP; lane 3, PC3; lane 4, DU-145; lane 5, RWPE1.
ACTH treatment for up to 8 days (Fig. 3C). Treatment with the universal MCR blocker SHU9119 inhibited cell proliferation in ACTH treated LNCaP cells by approximately 50% (Fig. 3D).

Indirect immunocytochemistry showed a punctate label pattern on PC3 cell membranes (Fig. 4A). The labeling pattern was similar to that described in transfected HEK293-related cells (293/FRT cell line) (29). Immunoblots incubated with MC2R antibody that was pre-absorbed with commercially available MC2R polypeptide showed no band (Fig. 4B). Routine western blot analysis showed higher MC2R protein expression in LNCaP, PC3, DU-145 cells compared with normal non-tumorigenic RWPE2 prostate cells (Fig. 4C).

**ACTH proliferative effect is additive with DHT stimulation**. Owing to the fact that adrenal steroids, dehydroepiandrosterone (DHEA), dehydroepiandrosterone sulfate (DHEAS), and androstenedione, are induced by ACTH and are linked to PCA recurrence, especially in patients undergoing long-term androgen ablation (6,35), we investigated a possible interaction with ACTH and androgen. ACTH stimulation of LNCaP cells in combination with DHT, a potent AR agonist, induced a 150–200% increase in cell proliferation compared with ACTH or DHT alone (Fig. 5).

**ACTH treatment enhances AR protein nuclear translocation and AR protein expression in LNCaP**. Production of cAMP/PKA via GPCR activation can transactivate AR in prostate carcinoma (25). Therefore, we considered whether ACTH stimulation of LNCaP cells affected the amount of nuclear AR protein expression. Immunocytochemistry showed increased AR nuclear signal intensity following ACTH treatment compared to that seen in vehicle treated cells (Fig. 6A). Similarly, western blot analysis of ACTH treated LNCaP cells confirmed an increase in total AR expression (Fig. 6B and C). Forskolin, a cAMP inducer and AR activator that mimics ACTH action, and DHT, a potent AR agonist, were included as positive controls. The results of these experiments indicate that ACTH treatment enhances AR nuclear translocation and increases AR protein in PCa cells (Fig. 6A-C).

**ACTH treatment increases cAMP concentration**. Cyclic AMP is a critical mediator of cell proliferation and differentiation; it induces cAMP-dependent PKA production in various cell types. As shown in Fig. 7, all three PCa cell lines responded to ACTH stimulation with concentration-dependent cAMP production increases. The mean EC₅₀ from three independent experiments was 1.79 µM for PC-3; 3.20 µM for DU-145; and 3.67 µM for LNCaP. The highest ACTH concentration (10 µM) increased the cAMP level 81-fold in PC3 cells, 111-fold in DU-145 cells, and 52-fold in LNCaP cells. There was no increase in cAMP production following NDP-MSH treatment (n=3, data not shown).

**MRAP-α and -β are expressed in PCa cells**. As coexpression of MC2R accessory protein isoforms α and β (MRAP-α and -β) is required for cAMP signaling via MC2R in some cells such as human embryonic kidney 293 cells (36), we examined whether MRAP-α and MRAP-β mRNA transcripts are expressed in LNCaP, PC3, and DU-145 PCa cells. RT-PCR analysis showed amplification of both transcripts in all three cell lines (Fig. 8). Positive control cDNA from the adrenal gland was amplified in parallel for validation of the PCR products.

**Discussion**

The findings reported here strongly link the ACTH-MC2R signaling pathway to PCA progression. This conclusion is based on several observations. Transcripts for MCRs, as well as MC2R protein, are expressed in LNCaP, PC3 and DU-145 human PCA cells. ACTH, the primary ligand for MC2R, increased cell proliferation/viability in the androgen dependent LNCaP and the androgen-independent PC3 and DU-145 cells by upregulating cAMP. ACTH-MC2R-induced cell proliferation was partly abrogated by SHU9119, a universal MCR blocker. Finally, AR was transactivated and its nuclear translocation increased in LNCaP cells following treatment with exogenous ACTH.

Physiological and sub-nanomolar ACTH concentrations stimulated PCA cell proliferation under serum-free conditions and in the absence of an exogenous cAMP inducer. The mitogenic effect was above 100% that of the control. NDP-MSH, a potent α MSH analogue and agonist of all but MC2R, as well as D-Trpγ-γ-MSH and THIQ, highly selective agonists for MC3R- and MC4R respectively, failed to stimulate cell proliferation. These results were predictable since it is known that all five MCRs respond to ACTH (37) but, among the MCRs, MC2R binds only ACTH and lacks affinity for other melanocortins. The other MCRs are, however, activated by both MSH and ACTH. The additional observation that the universal MCR blocker SHU9119 reduces but does not cancel the mitogenic effect of ACTH suggests that proliferation is at least partially mediated by MC2R.

Such trophic effects of ACTH were reported in human adrenal gland cells and skin melanocytes (38). Several studies documented what was referred to as ‘ectopic ACTH syndrome’ whereby several cancers from endocrine sites such as the pituitary gland (39) and diverse non-endocrine sites including the prostate gland, lung, liver, pancreas, and neuronal tissue, secrete ACTH that drives cell proliferation in these tissues (40–42).

ACTH-driven cell proliferation was likely mediated via cAMP induction since ACTH is known to bind MC2R,
a G protein-coupled receptor, which subsequently activates adenyl cyclase and results in cAMP production (43). The significant cAMP increases at high ACTH concentrations (above 1 µM) reported here indicated that an endogenous ACTH receptor is present in all three PCa cell lines. Since NDP-MSH is a superpotent agonist for other MCRs, except for MC2R, we reasoned that if the PCa cells cannot respond to NDP-MSH, the observed ACTH-dependent cAMP production in our three PCa cell lines should be caused by direct MC2R activation by ACTH. Indeed, there was no cAMP production by NDP-MSH stimulation (data not shown), suggesting that MC2R is expressed in all three cell lines.

The enhanced transactivation of AR via this pathway suggests cross-talk between MC2R and AR and supports the view of a critical downstream AR role in driving PCa cell growth in androgen-dependent and androgen-independent PCa cell growth (5). The increased AR protein expression following ACTH treatment suggests that the ACTH-MC2R signaling transduction pathway contributes to enhanced AR-DNA binding. That increased binding could promote PCa cell growth, an effect that would be particularly important under low androgen conditions. Such a conclusion is consonant with studies showing that several PCa cell GPCRs are indirectly involved in AR transactivation (13). In PCa, several GPCRs that signal through Gαs, such as prostaglandin E2 receptors (EP2 and EP4), and α-adrenergic receptor, can transactivate AR and induce cAMP and PKA activation which, in turn, can enhance AR sensitivity to low androgen levels. Our data indicate that MC2R is another GPCR that contributes to PCa cell proliferation and survival.

Figure 6. (A) ACTH treatment (10^-5) increased the nuclear staining intensity of the AR protein in LNCaP that have functional AR. Panel 1, DIC, differential interference contrast for cells in panel 2. Panel 2, AR expression in LNCaP cells treated with PBS/BSA vehicle. Panel 3, AR nuclear staining in cells treated with ACTH. Note the increased fluorescence intensity of AR nuclear staining in panel 3 (arrows). Panel 4, negative control (minus MC2R primary antibody). (B) ACTH treatment upregulated AR protein expression in LNCaP. Western blot analysis of AR protein expression in untreated LNCaP cells (lane 1), in cells treated with ACTH (lane 2), cAMP inducer Forskolin (lane 3), or DHT (lane 4). β-actin was included as control to monitor gel loading. (C) Quantitative analysis of band intensity in (B) using Unscan software. Data for (C) were obtained from two experiments.

Figure 7. Treatment of LNCaP, PC3, and DU-145 cells with ACTH increased cAMP production in a concentration-dependent manner. Intracellular cAMP levels were measured using radioimmunoassay. Results are expressed as fold increase over basal cAMP level of each cell line.

Figure 8. MRAP-α and MRAP-β (arrowheads) are expressed in LNCaP, PC3, and DU-145 human PCa cell lines. PCR product sizes for MRAP-α and MRAP-β are 100 and 350 bp, respectively. Lane 1, molecular weight DNA markers; lanes 2 and 3, positive controls for MRAP-α and MRAP-β, respectively (adrenal DNA). Lanes 4 and 5, LNCaP; lanes 6 and 7, PC3; lanes 8 and 9, DU-145. Lane 10, molecular weight DNA markers.
The central feature of the study reported here relates to PCA cell proliferation by ACTH-MC2R interaction that was partially abrogated by the universal MCR blocker SHU9119. This strongly suggests a paracrine role for ACTH and its MC2R in regulating PCA cell growth. The fact that there are two approximately equivalent androgen sources acting in the prostate gland, namely T from the testis and androgen derived from the adrenal gland, makes a strong case for including MC2R blockade as part of a combination therapy that involves inhibitors of androgen and other hormones.

Acknowledgements

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