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TITLE: A Unique Breast Cancer Cell Model for Studying Reported Functions of Membrane-Localized Estrogen Receptor (Alpha)

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We previously developed a cell line system in which exogenous expression of estrogen receptor alpha (ERα) in an ERα-negative cell line resulted in ERα-mediated signaling and proliferation. We previously reported generation of a cell lines that expressed ERα only in the cytoplasm (cERα) to characterize the putative cytoplasmic (non-genomic) function of ERα. However, while we found that cERα was not able to stimulate genomic ER action, and found interesting differences in estrogen-mediated downregulation of cERα, we were unable to show that this receptor could activate short-term non-genomic signaling. Since last year we have now started studying a membrane-targeted ERα (rhodopsin fused to ERα). We have generated stable cells expressing this receptor and show that this form of ERα is exclusively localized to the plasma membrane, and also estrogen is able to rapidly activate ERK1/2 in these cells. We have now also generated MCF-7 or MCF-7/HER2 cells overexpressing either cERα or rho-ERα and are currently examining the effect of this receptor on hormone response in these cells. We will finish the study using a one-year no cost extension.
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INTRODUCTION

Since the discovery of membrane estrogen receptor α (ERα) more than 20 years ago, reports on this form of ERα signaling have continued to be documented and have recently received increasing attention. However, this field remains very controversial with nuclear ERα action being studied in much greater detail and becoming much better understood. The IDEA of this proposal is to create a novel and unique model of breast cancer cells that express only cytoplasmic or membrane estrogen receptor (and not nuclear ERα) and then compare and contrast ERα action to cells that express no ERα or wild-type ERα. We are in a unique situation to perform this, as we have recently shown for the first time that ER can be functionally expressed and regulate proliferation in an ERα-ve breast cancer cell line (C4-12).

BODY

Summary

This progress report is for year 3 of the project. This was supposed to be a final report for the project, however, due to the generation of many exciting models at the end of the project period, we have requested and had approved a one-year no cost extension. In the first year of the project we made significant progress, having generated stable transfectants that express ERα in the cytoplasm (cERα) and having performed a preliminary characterization of these cells. However, as described in the body of the report in the second year, we were unable to identify any increase in rapid estrogen signaling in these cells that expressed the cERα (although we did generate interesting data regarding the ability of estrogen to cause downregulation of cERα, but the inability of ICI182780 to do this). This suggests that either these cells are not suitable for studying rapid estrogen effects or that perhaps the ERα needed to be in or near the plasma membrane to signal in this manner. In the second year we also struggled to create an ER variant that would reside in the plasma membrane. We expect that the small localization tag (myr or CAAX), placed on the N or C-terminus of ERα is folded within the protein and inaccessible for attachment to the plasma membrane. However, in this third year we were able to target the ER to the plasma membrane by using a fusion protein consisting of rhodopsin and ERα. We have stably expressed this construct in C4-12 cells and find that it is expressed exclusively in the plasma membrane. Furthermore, we have proven our hypothesis by showing that this receptor can now respond to short-term (minutes) estrogen by increased ERK1/2 phosphorylation. We have also generated stable transfectants of MCF-7 or MCF-7 cells overexpressing HER2 (MCF-7/HER2), and will use these to study the effect of plasma membrane ER on hormone action and anti-estrogen resistance.

Task 1: To create and characterize ERα-negative MCF-7 cells (C4-12) that stably express GFP tagged membrane ERα (mERα), cytoplasmic ERα (cERα), or wild type (wtERα) (Months 1-12): i) Stably transfect ERα-negative MCF-7 (C4-12) cells with GFP, GFP-wtERα, GFP-mERα and GFP-cERα, and select cell lines that have low and high levels of the receptor (Months 1-6).

We have stably transfected C4-12 cells with GFP, GFP-wtERα, and GFP-cERα. We have isolated multiple clones and using immunofluorescence microscopy shown that the cERα is indeed expressed in the cytoplasm (compared to wt ERα which is mainly nuclear) (Figure 1). GFP alone is expressed all over the cell.

Figure 1: Stable expression of GFP-cERα in C4-12 cells. cERα was generated by deletion of the nls
(245-270aa) in ERα. Stable clones of GFP, GFP-wtERα, and GFP-cERα were obtained and visualized by fluorescence microscopy (top panels) or by phase contrast (lower panels). GFP was widely distributed over the cell. In contrast, wt-ERα was exclusively nuclear; however, cERα was again widely distributed over the cell and did not show nuclear localization.

We have confirmed that GFP-cERα does not show nuclear localization by biochemical fractionation (data not shown) and confocal microscopy (Figure 2).

A setback came however, when we tried to express mERα, which was not targeted to the membrane (data not shown). Tagging of ERα with either C or N-terminal membrane signals does not send ERα to the membrane. This is probably due to folding of the protein making the tag inaccessible. We have therefore entered into collaboration with Dr Wang from Johns Hopkins University. He generated an ERα construct that consists of rhodopsin linked to ERα. Rhodopsin is membrane bound and so directs ERα to the plasma membrane (Xu Y Mol Endocrinol 2004 Jan, 18:86-96.) We found the rho-ERα to be in the endoplasmic reticulum following transient transfection with high concentrations of DNA (data not shown), however, lower amounts of DNA caused the rho-ERα to give only a plasma membrane signal (Figure 3).

In this year (year 3) we have generated several stable transfectants that express rho-ERα (Figure 4). Figure 4 shows immunofluorescence for ERα (red) or a nuclear stain (DAPI – blue) on MCF-7 cells or
C4-12 cells stably expressing rho-ERα. This figure clearly shows that MCF-7 cells express mainly nuclear ER with overlay of both red and blue signal. In contrast C4-12 cells show no nuclear ER (nucleus is only blue) but the ER is now exclusively localized to the plasma membrane. Figure 4B shows an immunoblot of C4-12 cells (left panel) and MCF-7 cells that were screened for expression of various forms of ER (this is an example, we screened over 400 colonies and generated more than 4 clones per construct – GFP-wtER, GFP-cER, and rho-ERα). The highlighted lanes with numbers represents 1) C4-12 rho ERα, 2) MCF-7 wGFP-wtER, 3) MCF-7 GFP-cER, 4) MCF-7HER2 GFP-cER, 5) MCF-7 rho-ERα, 6) MCF-7 rho-ERα. The tagging of GFP or rho makes it appear with a higher molecular weight. The rho-ER often appears as multiple bands probably due to it insertion in the plasma membrane and resulting posttranslational modifications of the rhodopsin protein. This immunoblot is only showed as an example of the tools we now have to use.

**Figure 4: Stable expression of rho-ER in C4-12, MCF-7 and MCF-7/HER2 cells.**

A) Immunofluoresence using antibodies against ER (red) and a DAPI as a nuclear stain (blue). Note the nuclear ER in MCF-7 (left) but the lack of ER in the nucleus of C4-12-rhoER and the presence of a strong plasma membrane localization. B) Representative immunoblot (from 400 screened clones) for various forms of ER. Immunoblot is with an ER antibody. Lane 21 shows C4-12 cells positive for rho-ER (increased molecular weight due to fusion with rhodopsin). 2 is wt-GFP-ER. 3 and 4 are GFP-cER (with an unknown lower species). 5 and 6 are rho-ER. Note the multiple isoforms of rho-ER which we believe maybe due to the posttranslational (probably glycosylation) of rhodopsin.

**ii) Use biochemical fractionation and confocal microscopy to determine whether mERα and cERα are expressed only in the membrane and cytoplasm respectively, and test whether mERα and cERα are capable of binding estradiol (E2) and tamoxifen (Tam) (Months 4-8).**

Figure 1 confirms that cERα is only expressed in the cytoplasm, while wt-ERα is mainly nuclear. Figure 3 shows that rho-ERα is mainly membranous by transient transfection, but is exclusively associated with the plasma-membrane upon stable transfection (Figure 4).

We have not tested directly whether cERα can bind E2 or tam, however an indirect measure is the ability of E2 to downregulate the receptor (which occurs after E2 binding). We found that cERα is degraded following E2 stimulation, suggesting in an indirect way that this receptor can bind E2. in one preliminary experiment we have found that rho-ERα is not degraded by estrogen. This highlights the interesting ability of cERα to be degraded by estrogen. This also argues against the literature suggesting a link between ER transcriptional activity and degradation – something which has been refuted by multiple recent publications.

**iii) Examine whether mERα or cERα associate with membrane or cytoplasmic structures (e.g. clathrin-coated pits) (Months 7-12).**

We have not yet performed these assays. This will now be done on the rho ERα stable transfectants.

**Task 2: To compare and contrast the effects of estrogen in C4-12-cERα, mERα and wtERα cells (Months 12-24):**
i) Analyze the effect of short (mins) and long-term (hours) E2 stimulation on localization, movement, and degradation of the different GFP-ERα variants (Months 12-16).

We have not examined localization and movement, but we have found that the cERα can be degraded by E2 (Figure 5).

![Figure 5: cERα is completely resistant to antiestrogen ICI 182780-mediated degradation. C4-12wt-ERα and cERα stable transfectants were starved in serum-free overnight and then treated for 8 hours with increasing concentrations of estradiol (E2) or ICI182780 (ICI). As expected, wtERα protein levels were reduced upon exposure to both E2 and ICI. This effect was blocked with the proteasome inhibitor (lactacystin 10uM) (data not shown). Of note, ICI required a 10-fold higher excess, which has previously been noted by others. In contrast to wtERα, cERα proteins levels decreased upon exposure to E2, but were not affected by any concentration of ICI.](image)

This is an important result given that a number of groups have proposed that E2-mediated degradation of ERα is linked to transcription. cERα is a variant ER that can't activate transcription, thus the degradation of cER by E2 represents a new paradigm for E2-mediated degradation of ER. Interestingly, while wt- ERα is degraded by antiestrogens such as ICI182780, the cERα is not degraded significantly by ICI182780 suggesting that this is a nuclear mediated event and that E2 and ICI degradation mechanisms are distinct. We have found that membrane localization of rho-ERα also inhibits degradation.

ii) Examine whether ER-responsive genes (e.g. TGFα, PgR, cathepsin D, pS2, IRS-1, cyclin D1) are induced by E2 and inhibited by Tam (Months 15-20) by the different GFP-ERα variants.

We have found that cERα is incapable of inducing expression of genes such as IRS-1, IGF-IR, and cyclin D1 (Figure 6). This is consistent with it not being able to activate gene transcription in an ERE-luc reporter assay (data not shown).

![Figure 6: cERα doesn't confer estrogen-induction of ER-responsive genes. MCF-7, C4-12cERα and C4-12wtERα cells were starved in serum-free medium overnight and then treated with either antiestrogen (ICI, 10-9M) or estrogen (E2, 10-9M) for 24 hrs. Cells were lysed in 5% SDS and immunoblotted with antibodies to insulin receptor substrate-1 (IRS-1), insulin-like growth factor receptor 1 (IGF-IR) or cyclin D1. As expected, all 3 genes were induced by estrogen in MCF-7 cells and also in C4-12 cells expressing wtERα. In contrast, two stable clones of C4-12ERα did not show estrogen regulation of IRS-1, IGF-IR or cyclin D1.](image)

In a preliminary result with the C4-12rho ERα we have found that this receptor is also unable to confer estrogen induction of IRS-1. This is not surprising given its attachment to the plasma membrane.
iii) Determine whether E2 stimulation results in an increase in S-phase and cell proliferation in C4-12-cERα and mERα compared to C4-12wtERα (Months 18-24).

We have found that E2 stimulation is able to increase S-phase in wt- ERα cells, but is unable to have an effect in cERα cells (Figure 7), consistent with this variant not inducing gene transcription (Figure 5). This is despite the fact that the cERα can clearly bind E2 and be degraded.

![Figure 7: cERα does not confer estrogen stimulated S-phase entry. MCF-7, C4-12, and C4-12 cells expressing cERα were starved in serum-free medium overnight, and then stimulated with estradiol (1nm) or ICI (100nM) or the combination for 16hrs. Cells were then fixed in alcohol, stained with propidium iodide and FACS analysis performed. MCF-7 cells stimulated with estradiol showed an induction in S-phase fraction (red) and also an increase in cells entering G2/M (blue). These changes were completely blocked by ICI. In contrast, ERα negative C4-12 cells, or the cERα expressing cells showed no changes in response to E2 or ICI. We will now be performing these experiments in the C4-12rho ERα to see whether the transient induction of ERK1/2 phosphorylation that we see with estrogen stimulation can confer a proliferative advantage.

Task 3: To determine whether previously reported short-term (minutes) E2-mediated effects are observed in C4-12-cERα or mERα cells (Months 24-36):

i) Perform coimmunoprecipitation and colocalization to determine if mERα and cERα can bind p85 and activate PI3K (Months 24-30).

Despite preliminary evidence that cERα was able to associate with p85, subsequent experiments failed to confirm an association with either p85 or IGF-IR. In addition we have been unable to show that cER can enhance short-term mediated activation of ERK1/2 or Akt by estradiol (Figure 8).

![Figure 8: cERα does not allow short-term estrogen signaling in C4-12 cells. MCF-7, C4-12 and C4-12-wtERα were starved in serum-free medium overnight, and then stimulated with estradiol (1nm) or ICI (100nM) or the combination for 16hrs. Cells were then fixed in alcohol, stained with propidium iodide and FACS analysis performed. MCF-7 cells stimulated with estradiol showed an induction in S-phase fraction (red) and also an increase in cells entering G2/M (blue). These changes were completely blocked by ICI. In contrast, ERα negative C4-12 cells, or the cERα expressing cells showed no changes in response to E2 or ICI. We will now be performing these experiments in the C4-12rho ERα to see whether the transient induction of ERK1/2 phosphorylation that we see with estrogen stimulation can confer a proliferative advantage.

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12cERα cells were starved in SFM overnight and then stimulated with estradiol (1nm, 15 or 30mins) or IGF-I (10nM, 15mins). Cells were lysed and immunoblotted for pIGF-IR, pERK1/2 and pAkt. MCF-7 cells did not show a response to estradiol at 15 or 30mins. Similar results were also seen in both C4-12 cells. As a positive control, all 3 cell lines responded to IGF-I.

We do not know whether this result simply shows that cERα is not able to activate these pathways in these cells, or whether the specific system we chose is not suitable for detection of short-term estrogen effects. However, we were able for the first time to detect rapid estrogen signaling in the stable transfectants of C4-12 cells expressing rho-ER (Figure 9). In this experiments (repeated three times) we were able to show that 15mins of estrogen exposure caused an increase in p-ERK1/2 that was equivalent to the positive control (15 mins exposure to IGF-I). However the induction was always transient and decreased after 30 mins (similar to that seen with growth factors).

Figure 9: Rapid phosphorylation of ERK1/2 in response to short-term estrogen in C4-12-rho-ER. C4-12 and C4-12rho-ER were starved overnight in serum free medium and then stimulated with estradiol (1nM) for 15 or 30 minutes. As a positive control for ERK1/2 activation, cells were stimulated with IGF-I (10nM) for 15 mins). Cells were lysed and immunoblotted with an antibody to pERK1/2. Total ERK1/2 levels were unchanged (data not shown).

We would like to now examine the interaction with Her2 signaling, which has been shown by our collaborators Drs Schiff and Osborne to be important for rapid ER signaling in MCF-7 cells (Figure 10).

Figure 10: Rapid estrogen signaling in MCF-7 cells that overexpress HER-2 receptor. MCF-7 and MCF-7 cells stably overexpressing HER-2. Cells were starved overnight in serum-free medium and then stimulated for 15 minutes with estradiol (1nM), epidermal growth factor (EGF, 100nM), heregulin (HRG, 100nM) or tamoxifen (Tam, 100nM). These incubations were performed in the presence or absence of the EGFR kinase inhibitor Iressa (gefitinib). Cells were then lysed and immunoblotted for phospho-EGFR, HER-2, Akt and ERK1/2 (MAPK). The left panel shows that estradiol or tam is unable to stimulate short-term signaling in MCF-7 cells, which can only be seen with heregulin (red box). In stark contrast, estradiol and tamoxifen were able to activate and phosphorylate all signaling intermediates in MCF-7/HER2 cells, and this was completely inhibited by EGFR blockade. We will therefore use these cells to better understand the role of cERα and rhoERα in this response. Figure kindly provided by Drs Osborne and Schiff, Breast Center, Baylor College of Medicine.

We have therefore stably expressed GFP-wtER, GFP-cER, and rho ERα in MCF-7 and MCF-7/HER2 cells and are examining rapid estrogen signaling and anti-estrogen resistance in collaboration with Drs Rachel Schiff and C. Kent Osborne. A first preliminary experiment showed little effect of the receptor on ERK1/2 signaling, however we couldn’t repeat previous results from Dr Schiff in parental cells that weren’t transfected (Fig 9). Therefore we have given the cells to her lab and she is repeating the experiment using their conditions.
ii) Examine whether E2-stimulation of C4-12-mERα and cERα cells results in mobilization of intracellular Ca²⁺ and activation of PKC (Months 30-34).
Not yet performed

iii) Determine whether E2 can induce an anti-apoptotic response in C4-12-mERα and cERα cells (Months 32-36).
Not yet performed

KEY RESEARCH ACCOMPLISHMENTS

- Generation of C4-12 cells that express ERα only in the cytoplasm (C4-12-cERα).
- Evidence that cERα is degraded by estrogen but not by anti-estrogen.
- Generation of C4-12 cells expressing rho-ERα.
- Evidence that rho-ERα is able to respond to short-term estrogen stimulation by enhancing ERK1/2 phosphorylation.
- Generation of MCF-7 or MCF-7/HER2 cells stably expressing GFP-wtER, GFP-cER, or rho-ER.

REPORTABLE OUTCOMES

Development of stable cell lines:
C4-12-GFP
C4-12-GFP-wtER
C4-12-GFP-cERα
C4-12-rhoER
MCF-7-GFP-wtER
MCF-7-GFP-cER
MCF-7-rhoER
MCF-7/HER2-GFP-wtER
MCF-7/HER2-GFP-cER
MCF-7/HER2-rhoER

CONCLUSIONS

This project will use a unique cell line model (C4-12) to test if cytoplasmic (cERα) or membrane targeted ERα (mERα) can perform signaling and mediated proliferation. This research is critical, as several recent studies have suggested that cERα or mERα is important, and pathologists only analyze nuclear ERα, which might misclassify a number of breast cancer patients. We have generated cells that express ERα only in the cytoplasm. We find that this receptor can't activate gene transcription or proliferation, despite the fact that the receptor is degraded by E2 and thus presumably can bind E2. However, this receptor is unable to stimulate short term estrogen events. We have also expressed ER in the plasma membrane and find that this receptor can respond rapidly to estrogen by enhancing ERK1/2 phosphorylation. We are currently examining the effect of this receptor in MCF-7 or MCF-7/HER2 cells. Understanding any potential role of cERα or mERα is critical for the complete understanding of estrogen action and targeting in breast cancer.

PRESENTATION

2004 “Expression of non-nuclear ER in ER-negative breast cancer cells doesn’t confer estrogen-stimulated growth”. 14th Annual Breast Cancer Think Tank Meeting, St Kitts.