Estrogen receptor-α interaction with the CREB binding protein coactivator is regulated by the cellular environment

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Abstract

The p160 coactivators, steroid receptor coactivator-1 (SRC-1), transcriptional intermediary factor-2 (TIF2) and receptor-associated coactivator-3 (RAC3), as well as the coactivator/integrator CBP, mediate estrogen receptor-α (ERα)-dependent gene expression. Although these coactivators are widely expressed, ERα transcriptional activity is cell-type dependent. In this study, we investigated ERα interaction with p160 coactivators and CBP in HeLa and HepG2 cell lines. Basal and estradiol (E2)-dependent interactions between the ERα ligand-binding domain (LBD) and SRC-1, TIF2 or RAC3 were observed in HeLa and HepG2 cells. The extents of hormone-dependent interactions were similar and interactions between each of the p160 coactivators and the ERα LBD were not enhanced by 4-hydroxytamoxifen in either cell type. In contrast to the situation for p160 coactivators, E2-dependent interaction of the ERα LBD with CBP or p300 was detected in HeLa but not HepG2 cells by mammalian two-hybrid and coimmunoprecipitation assays, indicating that the cellular environment modulates ERα-CBP/p300 interaction. Furthermore, interactions between CBP and p160 coactivators are much more robust in HeLa than HepG2 cells suggesting that poor CBP-p160 interactions are insufficient to support ERα–CBP–p160 ternary complexes important for nuclear receptor–CBP interactions. Alterations in p160 coactivators or CBP expression between these two cell types did not account for differences in ERα–p160–CBP interactions. Taken together, these data revealed the influence of cellular environment on ERα–CBP/p300 interactions, as well as CBP-p160 coactivator binding, and suggest that these differences may contribute to the cell specificity of ERα-dependent gene expression.


Introduction

Estrogen receptor-α (ERα) is a ligand-regulated transcription factor, which plays an important role in the growth, development, proliferation and maintenance of reproductive and various other tissues. Based on sequence comparisons, the 595 amino acid length of human ERα is divided into 6 regions named A through F (reviewed in Carson-Jurica et al. 1990, Green & Chambon 1991, Tsai & O’Malley 1994). The N-terminal A/B domain which contains a ligand-independent activation function (AF-1), exhibits little conservation among nuclear receptor superfamiliy members. The centrally located DNA binding domain (DBD), which is highly conserved across members of the receptor superfamily, corresponds to region C and enables the receptor to bind to estrogen response elements located in the regulatory regions of estrogen-responsive target genes. The D and E regions encode the hinge and ligand-binding domain (LBD), respectively, and are important for nuclear localization, receptor dimerization and transcriptional activity. The LBD is composed of 12 α-helices arranged as a 3-layered antiparallel α-helical sandwich which forms the hydrophobic pocket to which ligands bind (Brzozowski et al. 1997, Shiau et al. 1998). In addition to ligand binding, region E contains a second, ligand-dependent, activation function (AF-2). The function of the most C-terminal domain, region F, is not well understood, but appears to play a role in...
modulating antiestrogen activity (Montano et al. 1995, Nichols et al. 1998). The relative contribution of each activation function to ERα transcriptional activity varies depending on the cell and promoter context in which receptor function is assessed (McDonnell et al. 1995, Tzukerman et al. 1994). However, in most contexts, AF-1 and AF-2 synergize to maximally induce ERα-dependent gene expression (Lees et al. 1989, McInerney et al. 1996, Tora et al. 1989).

Coactivators are required for efficient regulation of transcription by nuclear receptors (Glass & Rosenfeld 2000, Robyr et al. 2000) and many ER-binding cofactors have been identified. These include members of the p160 steroid receptor coactivator (SRC) family of coactivators, SRC-1 [NCoA-1; (Onate et al. 1995)], transcrip

ational intermediary factor-2 [(TIF2) GRIP1/NCoA-2/SRC-2; (Hong et al. 1996, Voegel et al. 1996)], and receptor-associated coactivator-3 [(RAC3) p/CIP/ACTR/AIB1/TRAM-1/SRC-3; (Anzick et al. 1997, Chen et al. 1997, Li et al. 1997, Takeshita et al. 1997, Torchia et al. 1997)] as well as other coactivators such as E6-AP (Nawaz et al. 1999), CREB binding protein (CBP)/p300 (Kamei et al. 1996, Smith et al. 1996), DRIP205 (Burakov et al. 2000) and the p68/p72 RNA binding proteins (Endoh et al. 1999, Watanabe et al. 2001). Upon ligand binding, the ER undergoes a conformation change in its ligand binding domain which stabilizes a coactivator interaction site comprising a hydrophobic groove formed by residues in helices 3, 4, 5 and 12 (Beekman et al. 1993, Brzozowski et al. 1997, McDonnell et al. 1995, Shiau et al. 1998). Moreover, recruitment of p160s to ER is dependent on the integrity of helix 12 (Mak et al. 1999). The p160 coactivators, through a highly conserved signature motif termed the nuclear receptor (NR) box which encompasses the core consensus sequence LXXLL (where L is leucine and X is any amino acid) are recruited to nuclear hormone receptors (Heery et al. 1997, McInerney et al. 1998). There are multiple NR boxes within CBP and SRC family coactivators, and the amino acid sequences surrounding the core LXXLL motifs play an important role in regulating receptor–coactivator interactions (Chang et al. 1999, McInerney et al. 1998). Recruitment of coactivators to ERα enables the resulting complex to bridge the receptor to the general transcription machinery as well as remodel chromatin structure and thereby facilitate gene expression (Glass & Rosenfeld 2000, Jacq et al. 1994, Kraus & Kadonaga 1998).

It has been well established that ERs exert their effects in a cell- and tissue-specific manner (Katzenellenbogen et al. 1996). This has been attributed to differences in the relative strength of the AF-1 and AF-2 domains of ERα in various cell types which in turn are dependent on the cell-specific expression, activity or accessibility of cellular factors with which ERα must interact to efficiently stimulate gene expression. These cellular factors could include other transcription factors (Porter et al. 1997, Webb et al. 1995), molecules that regulate ERα interaction with DNA (Verrier et al. 1997, Zhang et al. 1999) and coregulatory proteins (e.g. coactivators and corepressors) (Glass & Rosenfeld 2000, McKenna et al. 1999, Robyr et al. 2000). The interactions of p160 coactivators and CBP with ERα have been studied extensively by in vitro approaches that have defined the regions of ERα and coactivators critical for protein-protein interactions (Demarest et al. 2002, Heery et al. 1997, Henttu et al. 1997, Hong et al. 1996, Kalkhoven et al. 1998, Kamei et al. 1996, Mak et al. 1999, Norris et al. 1998, Torchia et al. 1997, Voegel et al. 1996). In comparison there are fewer studies of the in vivo interactions between ERα and coactivators, although these support the general conclusion that ERα binds to coactivators in an estrogen-dependent manner (Harnish et al. 2000, Llopis et al. 2000, Shang et al. 2000, Stenoien et al. 2000, 2001a, 2001b, Tikkanen et al. 2000). However, intracellular signaling pathways influence p160 coactivator and CBP/p300 activity (Font de Mora & Brown 2000, Lopez et al. 2001, Rowan et al. 2000b, See et al. 2001), and ERα and ERβ phosphorylation has been shown to influence their interactions with coactivators (Dutertre & Smith 2003, Tremblay et al. 1999) demonstrating that the cellular environment can influence ER–coactivator activity and/or interaction.

In this study we investigated the contribution of cellular environment to the ability of the ERα LBD to interact with the p160 family of coactivators as well as CBP using mammalian two-hybrid and coimmunoprecipitation assays. These studies were performed in two cell lines chosen for their relative differences in AF-1 and AF-2 contributions to ERα transcriptional activity (Smith et al. 1997, Tzukerman et al. 1994). HeLa (human cervical
cancer) cells utilize the receptor’s AF-2 activity predominantly while in HepG2 (human hepatoma) cells the receptor’s AF-1 domain is required for efficient activation of target gene expression (Tzukerman et al. 1994). Furthermore, these cell lines differ in their ability to support activation of ERα by 4-hydroxytamoxifen (4HT), a selective estrogen receptor modulator (SERM), further highlighting the influence of cellular environment on ERα transcriptional activity. We demonstrate that while the ERα LBD binds well to p160 coactivators in both cell types, its interaction with CBP is cell-type dependent. Intriguingly, the extent of interactions between CBP and p160 are also cell-type dependent. The differences between HeLa and HepG2 cells with respect to CBP interactions with ERα and other coactivators is consistent with the hypothesis that cellular environment influences coactivator function, and is thereby an important determinant of cell-specific ERα function.

Materials and methods

Chemicals

17β-Estradiol was obtained from Sigma Chemical Company (St Louis, MO, USA). The partial antiestrogen 4-hydroxytamoxifen and the pure antiestrogen ICI 182 780 were gifts of D. Salin-Drouin (Laboratoires Besins Iscovesco, Paris, France) and A. Wakeling (Zeneca Pharmaceuticals, Macclesfield, UK), respectively.

Plasmids

The pBIND vector which encodes the GAL4-DNA binding domain (amino acids 1–147) and pACT plasmid which drives the expression of herpes virus VP16 activation domain (amino acids 411–456) were obtained from Promega Corporation (Madison, WI) as was the pG5-Luc target gene, which contains 5 DNA binding sites for the GAL4 DBD and a TATA promoter upstream of a luciferase reporter gene. The expression vectors for GAL-TIF2, GAL-RAC3 and GAL-CBP have been previously described (Lonard et al. 2000) as have those for VP16-CBP and VP16-SRC-1a (Rowan et al. 2000a). The GAL-p300 expression vector was obtained from Dr Tso-Pang Yao (Duke University). The GAL-SRC-1e expression plasmid was constructed by substituting the BstZ17I-XbaI fragment of pBIND-hSRC-1a (Lonard et al. 2000) with a 270 nucleotide BstZ17I-SpeI fragment of the SRC-1e cDNA which was generated by reverse transcribing total RNA isolated from HeLa cells with SuperScript II reverse transcriptase (Life Technologies, Grand Island, NY, USA), followed by PCR amplification using 5’-TGTGTTTCAG TCAAGCTGTCC-3’ and 5’-GAGCATTCCAC TAGTCTGTAG-3’ as primers.

The VP16-ERα LBD chimera expression vector was generated in pACT as follows: 882 bp of the ERα LBD cDNA corresponding to amino acids 302–595 were PCR amplified from pCMV-hERα (LeGoff et al. 1994) using 5’-GGGATCCGTAA GAAGAACAGCTGGCCTTGTTCC-3’ and 5’-TCTAGAGACTGTGGCAGGGAAACCCTCTG CC-3’ as primers. The resulting PCR fragments were subcloned into pCR3·1 using the TA Cloning kit (Invitrogen, Carlsbad, CA, USA), and then re-isolated as a BamHI–XbaI fragment and subcloned into the corresponding sites of pACT to generate pACT-LBD. All constructs were sequenced to verify that mutations did not occur during PCR amplifications.

Cell culture and transfection

HeLa and HepG2 cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS). Twenty-four hours prior to transfections, cells were plated in six-well culture dishes at a density of 3 × 10^5 or 6 × 10^5 cells per well for HeLa and HepG2 cells, respectively, in phenol-red free DMEM containing 5% charcoal-stripped fetal bovine serum (sFBS). DNAs were introduced into cells in the indicated amounts using Lipofectamine (Life Technologies) following the manufacturer’s recommendations. Five hours later, serum-free medium was replaced with phenol red-free DMEM and 5% sFBS, and 18–20 h thereafter, cells were treated with various hormones as indicated. Twenty to twenty-four hours later, cells were harvested and cellular extracts were prepared and assayed for luciferase activity using the Luciferase Assay System kit (Promega) and a Monolight 2010 Luminometer (Analytical Luminescence Laboratory, San Diego, CA, USA); values were normalized to protein content measured with Bio-Rad Protein Assay reagent (BioRad, Hercules, CA, USA).
Mammalian two-hybrid assays

In most mammalian two-hybrid assays, HeLa or HepG2 cells were transfected with 100 ng of the expression vectors for GAL-coactivator (pBIND-SRC-1e, pBIND-TIF2, pBIND-RAC3 or pBIND-CBP) along with 1000 ng of pACT-hERα-LBD and 1000 ng of pG5-Luc reporter plasmid. CBP–p160 interaction assays were performed with 100 ng pBIND-coactivator, 1000 ng of pACT-coactivator and 1000 ng pG5-Luc. Plasmid amounts employed in the GAL–CBP dose response curves are given in the legend to Fig. 3. Control experiments employed equivalent amounts of the pACT and pBIND empty vectors. Cells were treated with either 0·1% ethanolic vehicle, 10⁻⁸ M E₂ or 10⁻⁷ M 4HT for 20–24 h prior to cell harvest and assayed as described above.

Northern analysis

Total RNA was extracted from HeLa and HepG2 cells using TRIZOL reagent (Life Technologies) according to manufacturer’s protocol. Prior to RNA extraction, HeLa and HepG2 cells were treated with either vehicle, E₂, 4HT or the pure antiestrogen ICI 182 780. Forty micrograms per lane of total RNA were size fractionated on a 1·2% agarose/3-[N-morpholino]propanesulfonic acid/formaldehyde gel then transferred by capillary action to a GeneScreen Plus nylon membrane (NEN Life Sciences, Boston, MA, USA). Membranes were air-dried and UV-cross linked with a model FB-UVXL-1000 crosslinker (Fisher Scientific, Houston, TX, USA). Probes were prepared from 25 ng of cDNA fragments for SRC-1 (nucleotides 829–1896), TIF2 (nucleotides 4204–4815), RAC3 (nucleotides 119–1062) and mouse cyclophilin (Hasel and Sutcliffe 1990) which were labeled with [³²P]dCTP (3000 Ci/mmol; ICN Biochemical Inc., Irvine, CA, USA) using the RadPrime DNA Labeling System (Life Technologies). The blots were probed overnight under high stringency conditions [65 °C in hybridization buffer consisting of 0·5% SDS, 6 × SSC (0·9 M sodium chloride, 0·09 M sodium citrate, pH 7·0), 5 × Denhardt’s solution (Sambrook et al. 1989) and 100 µg/ml of salmon sperm DNA]. After stringent washing, radiolabeled blots were subjected to autoradiography at −80 °C using Kodak Biomax MS films, and band intensities were quantified by scanning laser densitometry (Personal Densitometer SI, Molecular Dynamics, Sunnyvale, CA, USA). Blots were subsequently stripped and reprobed with cyclophilin for standardization.

Western analysis

For measurement of CBP levels, HeLa and HepG2 cells were lysed in modified RIPA buffer [50 mM Tris–HCl (pH 7·4) containing 1% NP-40, 0·25% sodium deoxycholate, 150 mM NaCl, 1 mM EGTA, 1 mM PMSF, 1 µg/ml aprotonin, 1 µg/ml leupeptin, 1 µM pepstatin, 1 mM Na₃VO₄, 1 mM NaF]. After mixing with SDS-PAGE loading buffer, protein was resolved by 7·5% SDS-PAGE and electrophoretically transferred onto a nitrocellulose membrane (Osmonics Inc., Westborough, MA). CBP protein was detected immunologically using CBP antibodies from Upstate Biotechnology (Lake Placid, NY, USA; CBP-CT), Affinity Bioreagents (Golden, Co) or from Santa Cruz Biotechnology (Santa Cruz, CA, USA; A-22). Donkey anti-rabbit conjugated horseradish peroxidase (HRP) antibody (Amersham, Piscataway, NJ, USA) was used for detection. For Western analysis of p160 expression levels in HeLa and HepG2 cells 50 µg of total proteins extracted with lysis buffer [50 mM HEPES (pH 7.5), 100 mM KCl, 0·2 mM EDTA and 0·1% NP40 supplemented with Complete Mini-Tablets protease inhibitor tablets (Roche Applied Sciences, Indianapolis, IN, USA)] were separated by 6·5% SDS-PAGE. The blots were probed with either anti-SRC-1 (GeneTex, San Antonio, TX, USA), anti-TIF2 or anti-AIB1 antibodies (the latter two from BD Biosciences, San Diego, CA, USA). Anti-mouse conjugated to HRP (Amersham Biosciences) was used as secondary antibody. Visualization of specifically bound proteins was accomplished with ECL+PLUS reagent from Amersham, according to the manufacturer’s protocol using XL-1 Blue film (Kodak, Rochester, NY, USA). Band intensities were quantified by scanning laser densitometry.

Coimmunoprecipitation

For coimmunoprecipitation, four plates containing 5 × 10⁶ of either HeLa or HepG2 cells were harvested and incubated for 30–60 min on ice using lysis buffer similar to the one used above for
p160 Western analysis. Following that, the cell lysate was centrifuged for 5 min at 21,000 × g, and the protein content of the lysate was quantified using Bio-Rad Protein Assay reagent. Prior to immunocomplex preparation, 100 ng of recombinant ERα (Panvera, Madison, WI, USA) was incubated with either ethanol or 270 ng of estradiol for 30 min in 30 µl of buffer [50 mM Tris–HCl (pH 7·5), 10 mM MgCl₂, 1 mM EGTA, and 2 mM dithiothreitol]. The immunocomplex was prepared in a total volume of 1 ml that contained 60 µl of prewashed protein G+ agarose beads, 0·5 mg cell lysate, 100 ng of prebound ERα, and either 1 µg of anti-CBP antibody (C-1; Santa Cruz Biotechnology) or 1 µg of anti-SRC-1 antibody (GeneTex, San Antonio, TX, USA). The immunocomplex was rotated at 4 °C for 1·5 h then centrifuged and washed three times with lysis buffer. Subsequently, the immunocomplex was boiled for 5 min in 50 µl of 2 × Laemmli solution and resolved by 7·5% SDS-PAGE and transferred to nitrocellulose membrane and probed with rat anti-human ERα antibody (H222) followed by goat anti-rat HRP.

Results

Interaction of p160 coactivators with the ERα LBD

To examine the influence of cellular environment on interactions between p160 coactivators and the ligand binding domain of ERα, a mammalian two-hybrid assay was established. HeLa or HepG2 cells were transfected with expression vectors for the GAL4 DBD (GAL) or each of the chimeras of GAL fused to the amino termini of full length SRC-1, TIF2 or RAC3 in the presence of the VP16 activation domain alone (VP16) or VP16 fused to the amino-terminus of the ERα LBD (VP16–LBD). In contrast to other studies (Harnish et al. 2000, Mak et al. 1999), the GAL–coactivator constructs employed in this report encompassed the full-length forms of coactivators to ensure that the influence of regions apart from the previously defined primary interaction sites was taken into account (Heery et al. 1997). Therefore, when GAL–coactivator expression vectors were introduced into cells in the absence of ERα, the intrinsic transcriptional activity of SRC-1, TIF2 and RAC3 was apparent in comparison to the GAL4 DBD alone (Fig. 1A and B). Cotransfection of VP16–LBD with either GAL–SRC-1, GAL–TIF2 or GAL–RAC3 into HeLa cells further increased luciferase activity 3·5- to 6-fold indicating an interaction between receptor and each coactivator in the absence of hormonal stimulation (Fig. 1A). This is in consistent with a previous study from our lab showing that p160/SRC coactivators and CBP can interact with ERα in a ligand independent manner and that this interaction can be regulated by phosphorylation (Dutertre & Smith 2003, Stenoien et al. 2001a). Control luciferase values obtained for cells transfected with expression vectors for the GAL4 DBD in the presence of VP16 or VP16–LBD, or GAL–coactivator with VP16 were very low. Estradiol treatment significantly increased luciferase activity indicating a ligand-dependent enhancement of the ERα LBD interaction with SRC-1, TIF2 and RAC3. Treatment with the partial antiestrogen 4HT reduced luciferase expression below vehicle controls consistent with a reduction in basal interaction between coactivators and the ERα LBD. Simultaneous treatment of cells with 10 nM E2 and increasing concentrations of 4HT (0·1 → 100 nM) or 100 nM of ICI 182 780 inhibited interaction between receptor and each of the coactivators tested in a dose-dependent fashion (data not shown) further illustrating the ability of ligands to modulate this interaction. Similar results were obtained for HepG2 cells (Fig. 1B). Taken together, these results indicate that p160 coactivators interact with the ERα LBD in both HeLa and HepG2 cells.

Cell-specific interactions between CBP and the ERα LBD

CBP also contributes to steroid receptor transcriptional activity, and we therefore tested CBP’s ability to interact with the ERα LBD in HeLa and HepG2 cells cotransfected with GAL–CBP and the VP16–LBD chimera described above in cells treated with vehicle, 10 nM E2 or 100 nM 4HT. Similar to the results obtained for SRC family coactivators, an E2-dependent increase in LBD–CBP interaction was observed in HeLa cells (Fig. 2A). Surprisingly, however, luciferase activity in HepG2 cells transfected with GAL–CBP and VP16 was similar to that measured for cells transfected with GAL–CBP and VP16–LBD, even after E2 treatment, indicating that LBD–CBP
Figure 1  Mammalian two-hybrid assay of SRC family coactivator interactions with the ERα LBD. (A) HeLa or (B) HepG2 cells were transfected with expression vectors for the GAL4 DBD alone (GAL) or GAL–SRC–1 (SRC-1), GAL–TIF2 (TIF2) or GAL–RAC3 (RAC3) chimeras in the presence of an expression vector for VP16 alone (VP16) or a chimera consisting of the VP16 activation domain fused to the amino-terminus of the ERα LBD (VP16–LBD) along with the pG5-Luc reporter plasmid as indicated in the Experimental Procedures. Cells were treated with 0·1% ethanol (vehicle), 10 nM E2 or 100 nM 4HT for 20–24 h. Cells were subsequently harvested and assayed for luciferase activity as described in the Materials and Methods. Values are normalized to those obtained for GAL–SRC–1 and VP16 in the presence of vehicle, which were defined as ‘1’. Bars represent the average±S.E.M. of triplicate samples and are representative of at least 3 experiments.
interactions are very weak to nondetectable (Fig. 2B). In HepG2 cells, the apparent intrinsic transcriptional activity of GAL–CBP (e.g. in the presence of VP16 alone) was ~100–150-fold greater in comparison to that measured for the GAL4 DBD alone, while the differences in these

Figure 2  Cellular environment influences interactions between CBP and the ERα LBD. (A) HeLa or (B) HepG2 cells were transfected with expression vectors for the GAL4 DBD alone (GAL) or a GAL-CBP chimera in the presence of an expression vector for VP16 alone or VP16–LBD along with the pG5-Luc reporter plasmid. Cells were treated with 0.1% ethanol (vehicle), 10 nM E2 or 100 nM 4HT for 20–24 h, and subsequently harvested and assayed for luciferase activity. Values are normalized to those obtained for GAL–CBP and VP16 in the presence of vehicle which was defined as ‘1’. Bars represent the average±S.E.M. of triplicate samples and are representative of at least 3 experiments.
values in HeLa cells were much less pronounced; this was also the case for GAL-p160 versus GAL activity (see Fig. 1). Therefore, to ensure that the apparent lack of ER–CBP interaction in HepG2 cells was not due to CBP’s intrinsic transcriptional activity masking the activity of VP16 alone, we compared the transcriptional activity of GAL–VP16 to that of GAL–CBP in a dose–response experiment in both HeLa and HepG2 cells. In both cases, VP16 activity was greater than CBP (data not shown). Both of these findings argue against the possibility that CBP recruitment of VP16–LBD would be undetectable in our HepG2 cell two-hybrid assay due to relatively poor VP16 transcriptional activity. Thus, while the interactions between p160 coactivators and the ERα LBD are similar in HeLa and HepG2 cells, robust CBP-LBD interaction is only observed for HeLa cells.

As a further control to ensure that the apparent lack of CBP-ERα interaction in HepG2 cells was not due to saturation of the transcriptional machinery by high levels of intrinsic GAL–CBP activity in the cells, mammalian two-hybrid assays were performed using lower levels of GAL–CBP than used in the previous assay (10, 25, 50 and 100 ng) along with a fixed amount of VP16–LBD and pG5-Luc reporter. As shown for HeLa cells (Fig. 3A), there was a dose-dependent increase in luciferase activity measured in cells transfected with increasing amounts of GAL–CBP and VP16 in the presence of vehicle or estrogen. At each GAL–CBP dose, transfection of VP16–LBD resulted in an increase in luciferase activity in the absence of ligand that was further increased by estrogen treatment, confirming the results obtained in Fig. 2A. In contrast, curves obtained for HepG2 cells transfected with GAL–CBP and either VP16 or VP16–LBD were overlapping, regardless of estrogen treatment (Fig. 3B). This was not due to GAL–CBP saturation of reporter gene expression since transfection of increasing amounts of GAL–CBP expression vector yielded a dose-dependent increase in luciferase activity (Fig. 3B, inset).

To further support the mammalian two-hybrid results, we examined CBP interactions with full-length ERα by coimmunoprecipitation. Equal amounts of full-length recombinant ERα were added to either HeLa or HepG2 cell lysates. Following CBP coimmunoprecipitation, Western blotting for ERα demonstrated that E2 induces interaction between ERα and CBP in HeLa cells (Fig. 4A, top panel). Moreover, this experiment confirmed that the CBP-ERα interaction in HepG2 cells is weak and not hormone regulated.
contrast to ERα-CBP binding, ERα-SRC-1 levels of interaction were comparable; in both cell types E2 enhances ER-SRC-1 interaction (Fig. 4A bottom panel). These results are consistent with our mammalian two-hybrid data indicating that CBP interaction with the ERα LBD differs between HeLa and HepG2 cells. In order to determine whether differences in endogenous expression levels of CBP might account for the very weak LBD–CBP interaction observed for HepG2 cells, CBP expression levels were assessed in HeLa and HepG2 cells by Western blot analysis. In comparison to actin levels, which were used as an internal control, CBP expression levels were similar in both cell types. Moreover, scanning densitometry of Western blots using several CBP antibodies...
showed almost equal CBP expression levels in HepG2 cells compared with HeLa cells (Fig. 4B). Mammalian two-hybrid assays demonstrate that the interaction of the p300 coactivator, which is closely related to CBP, also interacts with ERα in a cell-specific manner with little to no binding between these proteins in HepG2 cells (Fig. 5). Taken together, these data demonstrate that CBP and p300 interactions with the ERα LBD or full length ERα are readily apparent only in HeLa cells indicating that the cellular environment influences the binding of these coactivators to ERα.

Expression of SRC family coactivators are similar in HeLa and HepG2 cells

Since p160 coactivators bind to CBP and enhance CBP physical and functional interactions with nuclear receptors (Demarest et al. 2002, Kamei et al. 1996, Li et al. 2000, Torchia et al. 1997, Voegel et al. 1998), Northern blot experiments were performed to determine whether gross differences in SRC family coactivator expression could account for differences in CBP–ER binding properties. Analyses were performed using total RNA extracted from cells exposed to either vehicle, E2, 4HT, or the pure antiestrogen ICI 182 780 to ensure that coactivator expression was not influenced by hormonal treatment. As shown in Fig. 6 (panels A and B), hormonal treatment had no gross effects on expression levels of SRC-1, TIF2 and RAC3 mRNAs in either HeLa or HepG2 cells. SRC-1 expression was similar in both cell lines (Fig. 6C). Taken together, these data indicates that cellular environment affects CBP–ER binding properties. Analyses were performed using total RNA extracted from cells exposed to either vehicle, E2, 4HT or the pure antiestrogen ICI 182 780 to ensure that coactivator expression was not influenced by hormonal treatment. As shown in Fig. 6 (panels A and B), hormone treatment had no gross effects on expression levels of SRC-1, TIF2 and RAC3 mRNAs in either HeLa or HepG2 cells. SRC-1 expression was similar in both cell lines and less than 2-fold differences in mRNA levels for TIF2 and RAC3 were found between HeLa and HepG2 cells. Endogenous protein levels of p160 coactivators were also examined in HeLa and HepG2 cells and were found to be similar in the two cell lines (Fig. 6C). Taken together, there appear to be insufficient differences in expression of either CBP/p300 or p160 coactivators between HeLa and HepG2 cells to account for the lack of ERα LBD interaction with CBP.

Interactions between p160s and CBP are stronger in HeLa than HepG2 cells

Since p160s and CBP bind to nuclear receptors as a ternary complex (Kamei et al. 1996, Li & Chen 1998, Xu et al. 2000) poor receptor–p160 coactivator or CBP–p160 coactivator interactions may compromise the ability of CBP to bind to ERα. Since the data in Fig. 1 demonstrate good ERα interactions with all three p160s in both HeLa and HepG2 cells, we investigated whether the lack of CBP–ERα LBD binding in HepG2 cells may represent a failure of p160s to interact sufficiently well with CBP to support an ERα–CBP–p160 ternary complex. HeLa and HepG2 cells were transfected with each of the GAL-SRC-1, GAL-TIF2, and GAL-RAC3 expression vectors along with VP16–CBP. As shown in Fig. 7A, SRC-1, TIF2 and RAC3 interactions with CBP were very strong in HeLa cells and yielded luciferase activity 35–70 times greater than the appropriate controls. In contrast, GAL–p160 interactions with VP16–CBP in HepG2 were present, but much weaker, with luciferase values only 4–6-fold higher than for GAL–p160 and VP16 alone (Fig. 7B). This finding was confirmed in a reverse two-hybrid assay between GAL–CBP and VP16–SRC–1 in which CBP–SRC-1 interactions were ~4-fold stronger in HeLa than HepG2 cells (Fig. 7C and D). Although it was noted that the magnitude of differences in interaction between cell types varied between the two-hybrid contexts, the result in both cases is the same; CBP interactions with p160s are more robust in HeLa than in HepG2 cells. Taken together, this data indicates that cellular environment affects CBP’s interactions with the ERα LBD as well as with p160 coactivators, even though the same methodology reveals equivalent ERα–p160 interactions in both HeLa and HepG2 cells, indicating that the lack of CBP–ERα binding corresponds to poor CBP–p160 interactions.

Discussion

The interaction between coactivators and ERα is critical for receptor-dependent activation of target gene expression. Extensive in vitro work has characterized the basic structural requirements for receptor–coactivator binding and the role of agonistic ligands in promoting recruitment of coactivator to the LBD (Brzozowski et al. 1997, Heery et al. 1997, Henttu et al. 1997, Kalkhoven et al. 1998, Mak et al. 1999, Shiau et al. 1998). In this report we demonstrate that the interaction between the ligand binding domain of ERα and CBP/p300 varies between two cell lines in which the ER activation domains differ in their contribution to receptor activity (Smith et al. 1997, Tora et al. 2000).
Cellular environment influences interactions between p300 and the ERα LBD. (A) HeLa or (B) HepG2 cells were transfected with expression vectors for the GAL4 DBD alone (GAL) or a GAL–p300 chimera in the presence of an expression vector for VP16 alone or VP16–LBD along with the pG5-Luc reporter plasmid. Cells were treated with 0·1% ethanol (vehicle), 10 nM E2 or 100 nM 4HT for 20–24 h, and subsequently harvested and assayed for luciferase activity. Values are normalized to those obtained for GAL–p300 and VP16 in the presence of vehicle which was defined as ‘1’. Bars represent the average±S.E.M. of triplicate samples and are representative of at least 3 experiments.

**Figure 5** Cellular environment influences interactions between p300 and the ERα LBD. (A) HeLa or (B) HepG2 cells were transfected with expression vectors for the GAL4 DBD alone (GAL) or a GAL–p300 chimera in the presence of an expression vector for VP16 alone or VP16–LBD along with the pG5-Luc reporter plasmid. Cells were treated with 0·1% ethanol (vehicle), 10 nM E2 or 100 nM 4HT for 20–24 h, and subsequently harvested and assayed for luciferase activity. Values are normalized to those obtained for GAL–p300 and VP16 in the presence of vehicle which was defined as ‘1’. Bars represent the average±S.E.M. of triplicate samples and are representative of at least 3 experiments.
et al. 1989, Tzukerman et al. 1994). In vivo interactions between SRC-1, TIF2 or RAC3 and the wild-type ERα LBD are hormone-dependent in both HeLa and HepG2 cells, supporting a role for these coactivators in ERα action in both cell types. However, very little interaction between CBP or p300 and the ERα LBD was observed in HepG2 cells in vivo and in vitro, suggesting that ERα transcriptional activity in this cell type may be less dependent on CBP/p300 than in HeLa cells.

Figure 6 Endogenous expression levels of SRC family coactivators and CBP are similar in HeLa and HepG2 cells. (A) Total RNA isolated from HeLa (left) and HepG2 (right) cells which had been treated with ethanol (vehicle) or 10 nM E2, 100 nM 4HT or 100 nM ICI 182 780 (ICI) for 24 h were subjected to Northern blot analyses using radiolabeled cDNA probes against SRC-1 (top), TIF2 (middle) or RAC3 (bottom). Blots were subsequently stripped and reprobed with a radiolabeled cyclophilin (CYC) cDNA fragment to measure cyclophilin mRNA levels, which were used as an internal control. (B) Relative normalized expression levels of SRC-1, TIF2 and RAC3 mRNAs. Laser scanning densitometer measurements of coactivator mRNAs were normalized to values obtained for cyclophilin, and the expression of each coactivator in vehicle-treated HeLa cells was arbitrarily set to ‘1’. (C) Fifty micrograms of HeLa and HepG2 cell lysate were subjected to Western analysis to examine endogenous levels of SRC-1, TIF2 and RAC3. Western analysis of actin levels were used to ensure equal loading.
Although we and others had hypothesized several years ago that differences in coactivator expression could be important for influencing the nature of ER-coactivator interactions (Jackson et al. 1997, Lavinsky et al. 1998, Smith et al. 1997), here we have shown differences in coactivator–ER interactions in cells in which SRC-1, TIF2, RAC3 and CBP expression levels are comparable. Thus, cellular factors other than the ligand and the expression of the coactivators themselves such as post-translational modifications (positive or negative), or the expression of other modulatory coregulators, are important determinants of receptor–coactivator interactions.

Estradiol promotes interactions of coactivators with ERα by inducing a conformational change in the structure of the receptor’s LBD. Recent crystallographic analyses of the LBDs of ERα and/or ERβ complexed with E2 (Brzozowski et al. 1997), diethylstilbestrol (Shiau et al. 1998), raloxifene (Brzozowski et al. 1997, Pike et al. 1999), 4HT (Shiau et al. 1998), genistein (Pike et al. 1999)
or ICI 164 384 (Pike et al. 2001) provides a structural framework within which LBD–coactivator interactions can be examined. Upon binding to an ERα agonist such as E2 or diethylstilbestrol, helix 12 of the LBD is oriented over the ligand-binding cavity. Helices 3, 5 and 12 in turn generate a functional AF-2 domain consisting of a hydrophobic groove on the LBD surface (Brzozowski et al. 1997, Shiau et al. 1998), and the co-crystal structure of amino acids 686 to 698 of the NR box II peptide of glucocorticoid receptor-interacting protein-1 (GRIP1) (mouse homologue of TIF2) and the LBD confirm that this receptor surface is a coactivator-binding site (Shiau et al. 1998). As anticipated, our data indicate that p160 coactivator interaction with the ERα LBD was significantly enhanced by E2 in both HeLa and HepG2 cells, consistent with ligand-induced changes in receptor structure and formation of a coactivator-binding site. However, receptor-coactivator interaction in the absence of ligand was also observed in both cell types, and 4HT reduced this interaction below levels observed under basal conditions. In this regard, ERα is similar to the constitutive androstane receptor-β (CAR-β) that can also interact with SRC-1 in the apparent absence of ligand (Forman et al. 1998). Recent work has demonstrated that the extent of ligand-independent interactions between ERα and co-activators is influenced by ERα phosphorylation (Dutertre & Smith 2003).

Although CBP/p300 can bind to various members of the nuclear receptor superfamily (Hanstein et al. 1996, Kamei et al. 1996, Kobayashi et al. 2000), its relative importance to the transcriptional activity of steroid receptors versus its role as an integrator/coactivator for numerous types of transcription factors is not clear. This contrasts with SRC family coactivators whose actions appear to be more restricted to nuclear receptors and for which evidence of the importance of these factors to steroid receptor function is abundant (Goodman & Smolik 2000). Nevertheless, CBP increases ERα-dependent gene expression and/or binds to ERα (Kamei et al. 1996, Smith et al. 1996), and ERα-CBP functional interactions therefore appear to be significant. Our results indicate that the cellular environment plays an important role in regulating the magnitude of CBP–LBD and CBP–p160 interactions. Both are much stronger in HeLa than HepG2 cells, suggesting that cellular factors that regulate each of these interactions may be related. This is consistent with reports indicating that CBP/p300–receptor interactions, in the absence of p160s, are poor. For instance, p300 binds weakly to thyroid hormone receptor on its own, but the addition of SRC-1 greatly enhances p300 interaction with this receptor (Li et al. 2000). It also has been shown that the p160-binding region of CBP is required for this coactivator to stimulate retinoic acid receptor transcriptional activity (McInerney et al. 1998). Those results are in agreement with the finding that CBP, p160s and nuclear receptors binding to one another as a ternary complex (Kamei et al. 1996, Li and Chen 1998, Xu et al. 2000). Our data therefore are consistent with a model in which the inability of CBP to interact well with the ERα LBD in HepG2 cells is due to the relatively poor CBP–p160 interactions observed for this cell type. Whether this is due to a negative acting factor in HepG2 cells, or lack of a positive influence in HeLa cells is currently under investigation. It is, however, unlikely to be due to failure of p160–ERα LBD interactions as they were quite strong in both cell types. It is important to note that CBP may still alter ERα transcriptional activity in HepG2 cells by indirect competition with other transcription factors or coactivators, and/or through interactions with the AF-1 domain (Kobayashi et al. 2000). Indeed, another investigator has demonstrated indirect effects of ERα on NF-κB activity that can be partially overcome by CBP overexpression (Goff et al. 1994), suggesting that ERα–CBP interactions may still be functionally important. Taken together, our data demonstrate that the molecular nature of ER interactions with CBP and SRC family coactivators are distinct.

Although the role of ligand in promoting interaction between nuclear receptors and different coactivators is widely recognized, there are few examples of the ability of cellular environment to alter coregulator function in a cell-type specific manner. For instance, the histone deacetylase inhibitor trichostatin A stimulates thyroid hormone receptor transcriptional activity in GH3 but not CV1 cells (Lizcano et al. 2001), while GRIP1 and RAC3 have preferential effects on stimulating vitamin D receptor activity in CV1 and P19 cells, respectively (Issa et al. 2001). Our studies provide evidence that extend these observations and demonstrate that coactivator binding to ERα or
other coactivators is not simply a reflection of the relative abundance of potential interacting proteins in a given cellular environment, but is also regulated by other factors within the cell. Taken together with the complexity contributed by the organizing influence of the promoter regions of ERα target genes, the ability of ERα to stimulate gene expression can be viewed in terms of multiple possible coactivator partners for ERα and the ability of ERα to bind to these coregulators in a cell-dependent fashion. As the selectivity of SERMs is thought to depend on differences in ERα interactions with coregulators between different cell and tissue types, these results also have important implications for understanding the molecular basis of SERM action.

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