Direct Acetylation of the Estrogen Receptor α Hinge Region by p300 Regulates Transactivation and Hormone Sensitivity

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Regulation of nuclear receptor gene expression involves dynamic and coordinated interactions with histone acetyl transferase (HAT) and deacetylase complexes. The estrogen receptor (ERα) contains two transactivation domains regulating ligand-independent and -dependent gene transcription (AF-1 and AF-2 (activation functions 1 and 2)). ERα-regulated gene expression involves interactions with cointegrators (e.g. p300/CBP, P/CAF) that have the capacity to modify core histone acetyl groups. Here we show that the ERα is acetylated in vivo. p300, but not P/CAF, selectively and directly acetylated the ERα at lysine residues within the ERα hinge/ligand binding domain. Substitution of these residues with charged or polar residues dramatically enhanced ERα hormone sensitivity without affecting induction by MAPK signaling, suggesting that direct ERα acetylation normally suppresses ligand sensitivity. These ERα lysine residues also regulated transcriptional activation by histone deacetylase inhibitors and p300. The conservation of the ERα acetylation motif in a phylogenetic subset of nuclear receptors suggests that direct acetylation of nuclear receptors may contribute to additional signaling pathways involved in metabolism and development.

Nuclear receptors coordinate diverse physiological roles in metabolism and development through ligand-dependent and -independent mechanisms (1). Nuclear receptors form multiprotein complexes with coactivator and corepressor proteins to orchestrate dynamic transcriptional events in response to ligand. In the absence of ligand, nuclear receptors repress transcription through a dominant association with corepressor complexes with histone deacetylase activity (2). Conformational changes induced upon nuclear receptor ligand binding release corepressors, with subsequent transient association of coactivator proteins (2–4). Estrogen binds the estrogen receptor (ERα), thereby regulating important functions in development and reproduction and in human diseases including breast cancer, cardiovascular disease, osteoporosis, and Alzheimer’s disease. The ERα contains domains conserved with other members of the “classical” receptor subclass (termed A—F) and two activation domains, AF (activation function)-1 and AF-2.

The two activation domains of ERα contribute synergistically to transcription of target genes. The AF-1 function is both constitutive and induced by mitogen-activated protein kinases (MAPKs) induced by growth factors or oncoproteins (5). p300 (6) and a p300/CBP-binding protein, p68 RNA helicase A (7), also induce AF-1 activity. Thus, p300 binds AF-1 in the absence of ligand (6, 8) inducing ERα activity 2–3-fold in either reporter or in vivo transcription assays (6, 8). p300/CBP binding to ERα is also detectable in MCF7 cells in the absence of ligand (4). The ligand-dependent transactivation function (AF-2) domain of ERα consists of a conserved carboxyl-terminal helix. The AF-2 domain contributes to ligand-induced activity through further recruitment of coactivator proteins including the p160 family, (SRC-1, TIF2/GRIP1, AIB1/ACTR), the cointegrators (CBP, p300), and p300/CBP-associated factor (P/CAF) (2, 8, 9). The role of p300 as an ERα cointegrator is complex; p300 contributes to ERα induction through several separable subdomains including the histone acetyltransferase (HAT) and the bromodomain (4, 8, 10), which make separate contacts to distinct domains of the ERα.

The enhancement of transcriptional activity by p300/CBP involves several different functions. The cointegrators provide a bridging function, which associates transcription factors with the basal transcription apparatus (11). Second, p300/CBP pro-

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1 The abbreviations used are: Erα, estrogen receptor α; AF, activation function; MAPK, mitogen-activated protein kinase; MEK, MAPK/ERK (extracellular signal-related kinase) kinase; CBP, CREB (cAMP-response element-binding protein)-binding protein; IP, immunoprecipitation; HAT, histone acetyl transferase; HPLC, high pressure liquid chromatography; GST, glutathione S-transferase; TSA, trichostatin A; MALDI-TOF, matrix-assisted laser desorption/ionization time-of-flight; E2, estradiol; P/CAF, p300/CBP-associated factor; EKLF, erythroid Kruppel-like factor; ERE, estrogen response element.
vides a scaffold, interacting with numerous transcription factors through dedicated domains to assemble high molecular weight “enhancesomes” (reviewed in Ref. 12). Third, the HAT activity of p300/CBP, which may be either intrinsic or mediated through the recruitment of associated proteins such as P/CAF, contributes to the transcriptional coactivator function. Transcriptional activation in chromatin-containing systems has correlated transcriptional activity with acetylation of specific lysines in the NH₂ termini of histones (13, 14). Histone acetylation is thought to facilitate binding of transcription factors to specific target DNA sequences by destabilizing nucleosomes bound to the promoter region of a target gene (15). In addition, p300/CBP and P/CAF directly acetylate non-histone proteins including a subset of transcription factors and coactivators (p53, EKLF, HMG1(Y), GATA-1, E2F-1, and ACTR (16–20). Transcription factor acetylation by coactivators has divergent effects. p300/CBP-dependent acetylation enhanced the activity of the tumor suppressor p53 (21), the Kruppel-like factor (EKLF) (19), and the erythroid cell differentiation factor, GATA-1 (22) (reviewed in Ref. 23). In contrast, CBP repressed the transcriptional activity of T cell factor (24), and direct acetylation of the coactivator ACTR by p300 contributed to an inhibition of hormone-induced nuclear receptor signaling (4). Together these studies are consistent with a model in which coactivator proteins, through their acetylation function, are engaged in a dynamic interplay to coordinate both the induction and repression of gene expression.

Although transcription factors can serve as substrates for HATs, no direct role for such molecules in hormone signaling had been identified (25). Intrinsinc HAT activity for histone lysines is shared redundantly by ERα transcriptional regulatory proteins, which include p300, CBP, P/CAF, SRC1, and ACTR (26, 27). Redundancy of the HAT function among coactivators raises the fundamental question of whether alternate substrates to histones may be involved in hormonal signaling. In the current studies we show that the ERα is acetylated in vivo and is directly and selectively acetylated by p300, but not by P/CAF, within the ERα hinge region at conserved lysines in vitro. Substitution mutation established an important role for these acetylated residues in both ligand-dependent and -independent functions, suggesting local conformational changes may regulate interactions between the two activation domains of the ERα. Conservation of the ERα motif acetylated in vitro between a subset of nuclear receptors raises the possibility that direct acetylation may regulate diverse functions of phylogenetically related nuclear receptors.

**MATERIALS AND METHODS**

**Reporter Genes, Expression Vectors, and Luciferase Assays**—The ERα luciferase reporter gene ERE, TK81 pαLUC (28), the Flag-tagged P/CAF mutants (29), the ERα fusion proteins (30), pCDNAS-HP300 (31), the constitutively active MEK1 plasmids, pCMV-NAV (32), pCMV-RAF (JN3-S218E-S229D) (28), and the catalytically inactive mutant MEK1 (K97M) (32, 33) were described previously. The ERα mutants were derived by polymerase chain reaction-directed amplification using sequence-specific primers. Both the wild type ERα and ERα mutants were cloned into pCI-neo (Promega, Madison, WI). The integrity of all constructs was confirmed by sequence analysis.

Cell culture, DNA transfection, and luciferase assays were performed as previously described (30, 34). Cells were incubated in media containing 10% charcoal-stripped fetal bovine serum prior to experimentation using estradiol and transfected by calcium phosphate precipitation or Superfect transfection reagent (Qiagen, Valencia, CA). The medium was changed after 6 h and luciferase activity was determined after 24 h. Luciferase activity was normalized for transfection using β-galactosidase or Renilla luciferase as an internal control exactly as described previously (20).

**Protein Expression and Western Blots**—The antibodies used in Western blot analysis were anti-M2 Flag (Sigma), anti-guanine nucleotide dissociation inhibitor (55), anti-acetyl lysine (16), and GST (B-14) and ERα (H-184) antibodies from Santa Cruz Biotechnology (Santa Cruz, CA).

*In vitro* [3H]methylene-labeled proteins were prepared by coupled transcription-translation with a Promega TNT®-coupled reticulocyte lysate kit (Promega), using 1 μg of plasmid DNA in a total of 50 μL. Flag-tagged P/CAF proteins were expressed in SF9 cells by infecting with recombinant baculovirus and purified using an anti-Flag antibody (Sigma, M2) (36). Full-length recombinant baculovirus ERα was obtained from Affinity Bioreagents, Inc. (Golden, CO).

**Immunoprecipitation**

**Histone Acetyltransferase Assays**—Immunoprecipitation histone acetyl transferase (IP-HAT) assays were performed using p300 as described previously (16, 37). For immunoprecipitation, the protein concentration was adjusted to 1 μg/ml in 500 μL. The relevant antibodies from Santa Cruz Biotechnology (p300, N15) were added (2 μg/500 μg of extract) and incubated at 4 °C for 2 h. A standard HAT assay was performed containing 5 μg of substrate and enzyme, either 200 ng of purified histone acetyl transferase (purified baculovirus p300 or P/CAF) or immunoprecipitated p300 from cultured cells (16, 37). The mixture was incubated at 30 °C for 1 h. 90 pmol of [3H]Acetyl-CoA reaction was electrophoresed on a SDS-polyacrylamide gel and viewed following autoradiography of the gel. [3H]Acetyl incorporation into the substrates was also determined by liquid scintillation counting or filter assays.

**In Vitro Protein-Protein Interactions and Mapping the ERα Acetylation Sites**—The interactions between the two activation domains of the ERα were mapped by using p300 as described previously (38). The in vitro translated protein (15 μL of ERα), 1 μg of rabbit anti-ERα polyclonal antibody (H184, Santa Cruz Biotechnology), and 5 μg of purified Flag-tagged baculovirus-expressed P/CAF were incubated in 300 μL of binding buffer.

**In vitro acetylation assays were performed as described previously (17).** Synthetic peptide corresponding to the ERα (ER1, residues 293–310, NH2-PSPPLMKRRKSSLSL-OH, and ER2, residues 353–370, NH2-ELVHMINWAKRVPFGVFDL-OH) were synthesized by BioSynthesis (Lewisville, TX) and purified to 95% purity by HPLC. The peptides were acetylated in vitro by incubation with 5 mM acetyl-CoA and baculovirus-purified Flag-p300 or P/CAF at 30 °C for 2 h. After incubation, acetylated peptides were separated from contaminating p300 by passage through a micro filter (Amicon Inc., Beverly, MA) and further purified by analytical reversed phase HPLC. The reaction products were analyzed with a PE-Biosystems DE-STR MALDI-TOF mass spectrometer. Further analysis by Edman degradation was performed on a PE-Biosystems Procise sequencer. Phenylthiohydantoin-acetyl-lysine was measured by absorbance at 259 nm.

**RESULTS**

**The ERα Is Acetylated by p300 in Vitro and in Vivo**—The p300/CBP coactivator proteins have been shown to regulate several promoters in a manner dependent upon their histone acetylation activity (25), and p300 can both bind and stimulate the activity of the ERα (4, 8, 10). In addition, p300/CBP and P/CAF have been shown to acetylate non-core histone-related transcription factors directly through a conserved motif. We assessed whether p300 could acetylate recombinant ERα in vitro. Recombinant p300 acetylated recombinant ERα but did not acetylate GST (Fig. 1A). In contrast, recombinant baculovirus-expressed P/CAF did not acetylate ERα, although it was capable of acetylating histone H3 and itself (Fig. 1B) as shown previously (39).

**The ERα Is an Efficient and Selective Substrate for p300 Acetylation in Vivo**—Two fundamental types of questions raised by these studies are, first, the relative efficiency of ERα acetylation and, second, whether the failure of P/CAP to acetylate the ERα is due to failed binding or substrate selectivity. To assess the relative efficiency with which p300 acetylates the ERα, a direct comparison was made between equimolar amounts of ERα and histone H3. The products acetylated by increasing amounts of p300 were electrophoresed in a SDS-polyacrylamide gel and the incorporation of [3H]Acetyl-CoA assessed (Fig. 2A). The efficiency of incorporation on an equimolar basis was ~3-fold greater for histone H3 (16 kDa) than ERα (66 kDa) (Fig. 2B), suggesting ERα is acetylated with substantially efficiency. Thus the ERα is efficiently and selectively acetylated by p300 in vitro.
P/CAF has been reported to associate with ERα in vitro (40). We examined whether the recombinant P/CAF used in the HAT assays bound to the ERα. As shown in Fig. 2C, recombinant P/CAF bound with high affinity to ERα, and binding required the HAT domain. Thus, although P/CAF acetylates histone H3 and H4, the failure of P/CAF to acetylate ERα is not due to...
IP-HAT assays were performed using (Fig. 3D).p300 acetylated the ER1 polypeptide but did not acet-

tylylated ER2 (Fig. 3D). An additional lysine, residue 362, was identified that had been conserved between species and is homol-
gous to the acetylation motif of the mu-
rine GATA-1 and human p53 proteins. The ER-(293–310) peptide was selectively acetylated by p300.

Identification of the ERa Acetylation Sites—To identify the residues required for ERa acetylation in vitro, recombinant

GST-ERa fusion fragments were expressed, their integrity was confirmed by Western blotting using a GST antibody, and equal amounts of proteins were assayed in HAT assays using recombinant p300 as a source of HAT activity and the previously described filter assay (16). As shown in Fig. 3, B and C, the ERa from residues 282–337 was sufficient to function as a substrate for acetylation by p300.

Peptides were synthesized to encompass the two lysine-containing motifs identified within the region of the ERa acetylated in vitro (Fig. 3D). We identified residues resembling an acetylation motif found in the p53 and GATA-1 transcription factors, which were conserved between species (Fig. 3D). An additional lysine, residue 362, was identified that had been implicated previously in ligand-regulated ERa function (41). Polyptides were synthesized therefore to include residues encoding the consensus acetylation motif ERa-(293–310) (ER1) and a second polyptide including lysine 366 (ERa-(353–370)) (ER2). HAT assays were performed using recombinant p300 or P/CAF. p300 acetylated the ER1 polypeptide but did not acetylate ER2 (Fig. 3D). Recombinant P/CAF failed to acetylate either ER polypeptides.

Mass analysis of the acetylated ER1 peptide confirmed the presence of two major ions differing by 42 mass units, with the smaller molecular weight product corresponding to the unmodified ER1 peptide and the higher molecular weight component corresponding to the acetylated ER1 product (Fig. 4A). Following in vitro acetylation of the ER1 peptide, Edman degradation assays were performed. As only monoaacetylated lysine-containing peptides were detected in the samples by MALDI-TOF mass spectrometry, the product analyzed by Edman degrada-
tion was a heterogeneous population of polypeptides, each acetylated at a single site (Fig. 4A). These studies demonstrated that lysines 302 and 303 of the ERa were preferentially acetylated by p300 with an additional acetylation site at lysine 299 (Fig. 4B).

The ERa Acetylated Residues Regulate Basal Activation of the ERa by TSA—To examine the role of histone acetylases in the regulation of ERa activity, an estrogen-responsive lucifer-

ase reporter gene was assessed in ERa-deficient cells (MDA MB231). Inhibitors of histone deacetylase(s) trichostatin A (TSA) and sodium butyrate were added to transfected cells for 24 h. TSA induced the ER-LUC reporter (ERE1 TKpA LUC) 4–6-fold (Fig. 5A). Similarly, sodium butyrate (1 mm) induced ER reporter activity 2-fold (Fig. 5B). To examine the functional consequence of lysines 302 and 303 in ERa function, point mutation of the ERa acetylation sites was performed. The ER-responsive reporter was assessed in ERa-deficient cells (MDA MB231 and HeLa). Activity was assessed through normalization to the internal standard β-galactosidase reporter. The 2-fold induction of wild type ERs by sodium butyrate was abolished by the ERa K302A/K303A mutant (Fig. 5C). The abundance of the ERaK302A/K303A mutant was similar to ERa wild type in cultured cells (Fig. 5D). HeLa cells were transfected with either wild type ERs or mutants of the acetylation site and assessed for ERa activity. The wild type ERa was induced 2-fold by the addition of TSA in a dose-dependent manner (Fig. 5E). Both the alanine and threonine substitutions failed to respond to TSA (Fig. 5E). Together these findings suggest that direct ERa acetylation contributes to induction by histone deacetylase inhibitors.

MAPK-induced ERa Functions Independently of the ERa Acetylation Site—To investigate further the in vivo conse-
quence of the ERa acetylation site, point mutation substitu-
tions were introduced into the wild type ERs at the lysine residues acetylated in vitro. It was reasoned that the acetyl-

failed binding. These findings are consistent with the observation that p300 and P/CAF have distinguishable substrate speci-

ficiencies (21).
tion of a lysine, a positively charged, hydrophobic residue, is thought to both reduce its charge and increase its polarity. If acetylation augments activity through increasing the polarity or reducing the charge, a mutation of the two ERα lysines to polar residues, ER(K302Q/K303Q), may function as an activating mutant. The introduction of a large positively charged amino acid with a significant side chain (ER(K302Z/K303Z)) might be anticipated to mimic acetylation if increasing polarity is of greater importance. Substitution of lysine to alanine, (ER(K302A/V303A), or another small hydrophobic threonine residue (ER(K302T/K303T)) was anticipated to result in a loss of function. If the post-translational modification of acetylation itself were important in regulating ERα activity, the substitution of the lysine residues with any of these other residues would be expected to have a similar effect. The results of these studies are shown in Fig. 6. The mutant ERα proteins were expressed equally in transfected cells (data not shown). HeLa cells were transfected with either wild type ERα or mutants of the acetylation site and assessed for their ability to regulate the activity of a synthetic ERE in the absence of ligand.

Assessment was made of the AF-1 function mediated by MAPK signaling. Growth factors induce ligand-independent activity of the ERα through activation of MAPK (5) and the p160 coactivator AIB1 (also named RAC3, ACTR, or SRC3) (42). p160 proteins bind p300 (43) and contact both the AF-1 and AF-2 of the ERα (44, 45). To determine whether the lysine substitutions within the ERα hinge regulated MAPK-dependent ERα activity, constitutively activated MEK1 (ΔN3, ΔN3-S218E-S222D) were coexpressed with the ERα mutants (Fig. 6A). The wild type ERα was induced 3.5-fold by activated MEK1 but was not significantly induced by the catalytically defective MEK1 (K97 m). The basal activity of the ERα(C302A/K303A) mutant was reduced 2.5-fold; however, the magnitude of induction by activated MEK1 was not significantly changed for any of the mutants (Fig. 6A). The finding that the ERα acetylation mutants are not altered in their responsiveness to MAPK activation suggests the mechanisms governing ligand-induced ERα activity through the ERα acetylation site are distinct from those governed by ACTR.

The ERα Acetylation Site Governs Ligand Sensitivity—in previous studies of ERα activity in HeLa cells using a similar reporter assay, estradiol (10⁻⁸ M) induced ERE-dependent luciferase activity 2-fold (41). In our studies the wild type ERα gave a similar 2-fold induction upon the addition of estradiol (10⁻⁸ M) (Fig. 6B). This ERE₂TK81LUC reporter is not induced by 10⁻¹⁰ M E2 in HeLa cells with the wild type ERα; however, both the glutamine and arginine substitutions were induced by 2–3-fold, suggesting the positive charge of these residues may contribute to ligand sensitivity (Fig. 6B). The hinge domain mutants were compared with the wild type ERα for ligand-dependent transactivation using increasing concentrations of E2. Enhanced E2-dependent activity was observed for each of the ERα mutations of the hinge region lysine residues. Thus, uncharged, polar, or hydrophobic substitutions of the ERα enhanced ligand sensitivity. As each of the ERα mutants exhibited similar levels of expression to wild type ERα, and the wild type ERα functioned in the same manner as the ERα wild type in other studies in this cell type (41), these findings suggest that the wild type lysine residues within the ERα hinge region may play a role in normally repressing ligand-dependent ERα activity.

We next assessed the role of the hinge domain lysine residues in p300-dependent regulation of ERα function. The modest induction of wild type ERα activity by p300 in the absence of ligand (Fig. 6C) is consistent with studies by others. Binding of p300 to the ERα in the absence of ligand and a 2–3-fold induction of ERα activity in the absence of ligand were observed both in reporter assays (6) and in in vitro transcription assays (8). Conformational changes induced by the addition of estradiol recruits p160 coactivators to a hydrophobic fold in the ERα with the p300 cointegrator (9). Because mutation of the lysine residues of the ERα enhanced ligand sensitivity, we hypothesized that substitutions of these lysines may also enhance p300-dependent transactivation of the ERα in the presence of E2. In keeping with this model each of the ERα acetylation mutants demonstrated enhanced activation by p300 in the presence of hormone (Fig. 6C). These findings raise the possibility that this region of the ERα may serve to dock repressor proteins or that direct acetylation of the ERα may play a role in ligand-dependent transcriptional attenuation, as was recently described for the direct acetylation of ACTR by p300 (4). Crystal structural analyses showed the LXXLL motif of the coactivator GRIP1 forms the core of a short amphipathic α helix that contacts helices 3, 5, 6, 11, and 12 of the ERα; however, the exact proximity of the ERα(C302A/K303A) residues

![Fig. 4](http://www.jbc.org/figs/4_4.png)  
Fig. 4. A conserved acetylation motif in the ERα. A parallel reaction to that used in Fig. 3D using unlabeled acetyl Co-A was analyzed by MALDI-TOF mass spectrometry (A) and sequenced by Edman degradation (B). In A, the resulting ER-(293–310) peptide mass spectrum is shown with mass/charge expressed in atomic mass units (amu). The major peak labeled X corresponds to the expected mass of the unmodified ERα peptide. The major peak labeled Y, larger by 42 atomic mass units, represents singly acetylated peptide. The minor peaks are methionine oxidation products present in the starting material. In B, the bars represent the amount of phenylthiohydantoin-acetyl-lysine present in the corresponding positions. The major acetylated products correspond to residues 302 and 303.
required for the acetylation of ERα in cultured cells. This would seem unlikely, however, as mutations of the ERα acetylation site, which could not be acetylated in vitro, conveyed enhanced ligand sensitivity in cultured cells. To determine whether ERα is acetylated in vivo, a polyclonal antibody raised against acetylated lysines (16) was used to immunoprecipitate acetylated proteins from MCF7 cells. The IP product was subjected to SDS-polyacrylamide gel electrophoresis and probed with an ERα antibody. Fig. 6D shows that the ERα antibody specifically recognized ERα protein within the anti-acetylated lysine immunoprecipitate (upper panel). Because the coactivator ACTR is acetylated by itself (4), the co-immunoprecipitation of the ERα may potentially be due to cross-reactivity with ACTR. Therefore, a reciprocal immunoprecipitation was performed in which we used the ERα antibody to IP ERα from MCF7 cells, and Western blotting was performed with the anti-acetyl lysine antibody (Fig. 6D, lower panel). The acetyl lysine immunoreactive band corresponding to the molecular weight of the ERα was observed in the ERα IP but not with the control IgG IP. Together these studies indicated that the ERα is acetylated in cultured cells consistent with previous findings that p300 binds and regulates ERα in the absence of ligand in vivo (4, 6, 8).

DISCUSSION

The regulation of estradiol signaling by direct ERα acetylation reveals an unexpected and novel role for histone acetyltransferase in hormone signaling. Nuclear receptors have been shown to form multiprotein complexes with coregulatory proteins that possess either histone acetylase or histone deacetylase activity (4, 47). The evidence that the ERα is a direct substrate for HAT activity and may thereby regulate hormone-dependent transactivation function remained to be demonstrated. Here we have shown that ERα is acetylated in vivo and is a substrate for selective acetylation by p300 in vitro. Although coactivators recruited to ERα share a redundant capacity to acetylate histones, herein the ERα was selectively acetylated by p300. The select enzymatic activities of p300 and P/CAF toward ERα are consistent with their structurally divergent HAT domains (36, 48). Mutagenesis demonstrated a critical role for the ERα acetylation site in regulation by histone deacetylase inhibitors. The finding that mutations with the ERα hinge domain lysine residues enhanced hormone sensitivity suggests these residues may be involved in ligand-dependent transcriptional repression or transcriptional attenuation. The finding that the lysine residues within the ERα that are substrates for the HAT activity of p300 may function in transcriptional repression suggests that coactivator proteins acetylate several distinct substrates with distinct effects to coordinate genomic responses.

The mechanisms governing substrate specificity of HATs are not well understood at this time (49). P/CAF did not acetylate ERα but was capable of efficiently acetylating histone H3 and binding ERα. These findings suggest that p300 and P/CAF, although both capable of binding ERα, convey select enzymatic activities, consistent with the lack of sequence similarity within their HAT domains (36, 48). From previous studies of TAFⅭ50 it is known that the bromodomain modules form selective interactions with multiple acetylated histone H4 peptides (50). To understand the mechanisms responsible for the failure of P/CAF to acetylate ERα, we performed an analysis of P/CAF domain mutants to identify the sites of interaction with the ERα lysine motif peptide. These studies revealed the surprising result that the P/CAF bromodomain was dispensable and that the HAT domain was required for binding to ERα. It is possible that the interaction surfaces may determine subsequent acetylation activity. Alternatively, the acetylation motif of the substrate may be critical. The ERα acetylation motif re-

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Footnote: 2. G. Greene, personal communication.
sembles the GATA-1 and p53 acetylation sites. GATA-1, EKLF, and ACTR are selectively acetylated by p300/CBP (4, 19, 22). By contrast, P/CAF preferentially acetylates E2F-1 and MyoD in vitro (20, 51). p53 contains two acetylation sites differentially acetylated by either p300 (16) or P/CAF (21). Although the determinants of the histone acetylase substrate preference are poorly understood, this substrate specificity may form the biochemical basis for functional synergy and promoter selectivity.

In the current studies, mutation of the ERα in vitro acetylation site enhanced ligand sensitivity. The 2-fold induction of the synthetic estrogen-responsive enhancer reporter gene ERαS218ES222D and ΔN3 is shown compared with either vector or the catalytically inactive mutant MEK1 (K97M). Results are shown on the left as the mean ± S.E. for the luciferase activity. B, E2-induced transactivation of the ERE-LUC reporter was determined for the wild type and ERα mutants; the mean -fold induction is shown at each of the E2 concentrations used. The data are the mean of six separate experiments. The S.E. was <3% for the data points. The ERα mutants were increased significantly in ligand-induced activity at each ligand concentration compared with wild type (wt) ERα (p < 0.05). C, the effect of p300 on wild type and ERα mutant activity was determined in the presence and absence of ligand. Data are the mean ± S.E. with significant differences shown (*, p < 0.05) compared with wild type ERα. D, upper panel, MCF7 cells were subjected to IP with polyclonal anti-acetylated lysine antibody (New England Biolabs, Beverly, MA), and the IP product was subjected to Western blotting with the ERα antibody. Lower panel, MCF7 cells were immunoprecipitated with an anti-ERα antibody or control IgG and the electrophoresed product was subjected to Western blotting with an anti-acetyl-lysine antibody (16). The immunoreactive band detected with the anti-acetyl lysine antibody is of identical molecular weight to the ERα.

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transcription assays (8) or in reporter assays (6), together suggest that p300 conveys important ligand-dependent and -independent functions. Estradiol treatment of MCF7 for 24 h cells does not change the abundance of p300, histone deacetylase-1, or ERα (4), and the induction of histone H4 acetylation at target promoters in response to ligand is quite transient (4). Conformational changes induced by the addition of estradiol are known to recruit p160 coactivators to a hydrophobic fold in the ERα with the p300 cointegrator (9). As noted above, the LXXLL motif of the coactivator GRIP1 forms the core of a short amphipathic α helix that contacts helices 3, 5/6, 11, and 12 of the ERα; however, the exact proximity of the ERα(K302A/K303A) residues to the ERα hydrophobic fold remain unknown (46). Future studies will discern whether the increased ligand sensitivity of these ERα acetylation mutants is due to enhanced recruitment of coactivators within the local promoter context or to loss of binding to transcriptional repressors.

These studies raise several important new types of question regarding the direct acetylation of the ERα affects interactions with other coactivators and corepressors, DNA binding within native chromatin at estrogen-responsive promoters of target genes, the function of the ERα in \textit{in vitro} transcription assays, and the effect of these mutations on selectivity of estrogen signaling pathways. In the current studies, mutational analysis of the ERα acetylation site demonstrated dissociable effects of histone deacetylase inhibitors (TSA) and the addition of ligand on ERα activity. The induction of ERα activity by the histone deacetylase inhibitors TSA and sodium butyrate was abolished upon substitution of the acetylated lysine residues with small hydrophobic residues, either alanine or threonine, suggests that basal ERα activity is under constitutive repression by histone deacetylase-containing complexes and that the lysine residues may contribute to a surface recruiting such complexes. In the absence of ligand, nuclear receptors have been shown to exist in multiprotein complexes containing N-CoR (nuclear receptor corepressor) or related proteins (54) together with histone deacetylases and homologues of the yeast corepressor Sin3, which repress gene transcription (47, 55, 56).

As estrogen is mitogenic in mammary epithelial cells, the enhancement of ligand-dependent transactivation induced by mutation of these ERα target lysines may be predicted to confer a growth advantage. The same mutant that we demonstrated as conveying enhanced ligand sensitivity for transactivation (ERα(K302R)) was recently shown to occur in 34% of premalignant human breast lesions, suggesting that these acetylated residues play an important role in ERα function and biology (57). The ERα acetylation motif is conserved between species and between phylogenetically related nuclear receptors (58) (Fig. 7). Mutations of the conserved lysine motif have been identified in the ERα in breast cancer as has the androgen receptor in prostate cancer. Because nuclear receptors that contain the candidate acetylation motif contribute to diverse roles in the regulation of growth, development, and homeostasis (1), these studies may have possible implications in understanding regulation and function of many nuclear receptors.
Direct Acetylation of the Estrogen Receptor α Hinge Region by p300 Regulates Transactivation and Hormone Sensitivity

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