Acetylation of conserved lysines in bovine papillomavirus E2 by p300

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Quinlan, Edward J.; Culleton, Sara P.; Wu, Shwu-Yuan; Chiagn, Cheng-Ming; and Androphy, Elliot J., "Acetylation of conserved lysines in bovine papillomavirus E2 by p300" (2013). *University of Massachusetts Medical School Faculty Publications*. 197. 
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The p300, CBP, and pCAF lysine acetyltransferase (KAT) proteins have been reported to physically interact with bovine (BPV) and human (HPV) papillomavirus E2 proteins. While overexpression of these KAT proteins enhances E2-dependent transcription, the mechanism has not been determined. Using RNA interference (RNAi) to deplete these factors, we demonstrated that E2 transcriptional activity requires physiological levels of p300, CBP, and pCAF. Each protein appears to have a unique function in E2-dependent transcription, since overexpression of one KAT failed to compensate for RNAi knockdown of another KAT. Using an in vitro acetylation assay, we identified highly conserved lysines that are targeted by p300 for acetylation. The conservative changes of lysines at positions 111 and 112 to arginine were of particular interest. The K111R and the K111R/K112R mutants showed reduced transcriptional activity that was not responsive to p300 overexpression, while the K112R mutant retained activity. p300 and CBP were detected at the viral promoter; however, pCAF was not. We propose a model by which E2 transcriptional activity is controlled by p300-mediated acetylation of lysine 111. This model represents a novel mechanism regulating papillomavirus gene expression.

The papillomavirus E2 protein controls transcription of viral early gene products by binding to specific DNA motifs (ACCCN6GGT) critically placed within the viral genome (1). While viral gene expression is controlled by a variety of cellular transcription factors, including TFIID, Sp1, Oct1, and AP1 (2–6), expression of E2 results in increased transcriptional activation from the bovine papillomavirus (BPV) promoter and enhancer elements containing E2 binding sites (7,8). DNA binding activity is conferred by the C-terminal DNA binding domain (DBD); however, the N-terminal transactivation domain (TAD) of E2 is also necessary for specific activity (9–12). The ability of the TAD to activate transcription in the absence of the DBD (13) indicates that its function is at least partly mediated through complex formation with cellular factors (10,12).

Prior studies have identified E2-interacting proteins that facilitate E2-dependent transcription; however, the precise mechanisms through which these factors contribute to E2 activation of the viral early promoter remain unclear. Several reports have demonstrated that general transcription factors TFIIB and TFIID, including the TATA binding protein (TBP) and several TBP-associated factors, interact with E2 (14–18) and that TFIIB and TBP can enhance transcription by E2 (3,16,18). Cellular coactivators such as Brd4, hBrm, Gps2 (also known as AMF-1), and Tax1BP1 interact with E2 and enhance E2-dependent transcriptional activation (19–24). Brd4 and Tax1BP1 stabilize E2 protein, which partially explains how these factors increase E2 transcriptional activity (20,24–26). Association with the Brm ATP-dependent chromatin remodeling complex enhances E2-dependent transcriptional activity specifically on episomally maintained templates (19). E2-dependent coactivation by either Tax1BP1 or Gps2 is enhanced further through complex formation with the cellular acetyltransferase p300 (22,24).

Histone acetyltransferases (HATs), also known as lysine (K) acetyltransferases (KATs), catalyze the transfer of an acetyl group from acetyl coenzyme A (acyetyl-CoA) molecules to the ε-amino group of lysine side chains. Each KAT is thought to recognize a distinct molecular surface area of the target lysine and not necessarily a specific consensus peptide sequence (27,28). Further specificity is conferred through the formation of higher-order complexes containing one or more KATs and a variety of subunits (28–30). Subunits commonly encode chromatin binding domains such as bromodomains or chromodomains that recognize acetylated (31–33) and methylated (34,35) lysine residues, respectively. Through recognition of various combined modifications on histone tails, acetyltransferase complexes are recruited to specific promoter or enhancer regions where activity is required (36–40).

Acetylation of nonhistone proteins, such as transcriptional cofactors, can affect substrate stability, subcellular localization, and interactions with other proteins or DNA (41).

Published data reveal that p300, CBP, and the p300/CBP-associated factor (pCAF) interact directly with E2 proteins of BPV type 1 (BPV-1) and human papillomavirus type 16 (HPV-16), HPV-18, HPV-11, and HPV-6b (22,42–44). These KAT proteins have also been demonstrated to enhance E2-dependent transcription (22,42–45). Further, p300 coexpressed with small amounts of HPV-16 E2 enhances transcriptional activity from the HPV-16 promoter, which is generally repressed at higher levels of E2 (45). The E2 coactivator Gps2, which also directly binds p300, was proposed to facilitate enhancement of BPV-1 E2 activity by p300 (22). The KIX domain of CBP, which is also the CREB binding region, interacts with the N-terminal
transactivation domain of HPV-18 E2. Heterologous expression of the CBP KIX domain fused to the VP16 activation domain enhanced E2-dependent transcription (43). The N-terminal 390 amino acids of pCAF are necessary for interaction with HPV-6b, -11, -16, and -18 E2 (42). Enhancement of E2-dependent transcription by pCAF was further stimulated by coexpression of CBP (42), suggesting that these KAT proteins act through distinct pathways. These studies also showed that the acetyltransferase domains of CBP and pCAF were required for enhancement and that acetyltransferase-defective mutants were unable to enhance E2-dependent transcription (42, 43).

The functional interaction of KATs p300, CBP, and pCAF with E2 suggests a role for acetylation in the regulation of E2-dependent transcription. This could conceptually occur through histone alteration and subsequent chromatin remodeling, similar to the proposed role for hBRM, and/or through posttranslational modification of lysines within E2. We have examined the role of p300, CBP, and pCAF in E2-dependent transcription using selective RNA interference (RNAi) depletion and explored the possibility that E2 is a substrate for acetylation. The results shown here are consistent with distinct roles for each KAT protein in regulating E2 activity. We demonstrate that E2 is acetylated by p300 in vitro and that the mutation of conserved N-terminal lysines restricts E2 activity and uncouples E2-dependent transcription from the effects of p300 overexpression. These data provide the first evidence for the direct acetylation of E2 as a mechanism to regulate its transcriptional activity.

**MATERIALS AND METHODS**

**Cells and transfections.** RPE-1 cells were cultured in a 1:1 mix of Dulbecco’s modified Eagle’s medium (DMEM) and Ham’s F-12 medium (Invitrogen). C33a, RKO, C127, ID13, and C127-A3 cells were cultured in DMEM (Invitrogen). ID13 and C127-A3 cells are BPV-1-transformed cell lines, BPV genomes in C127-A3 cells contain three mutations within the E2 reading frame leading to increased E2 protein levels (46). All cell culture media were supplemented with penicillin-streptomycin solution (Invitrogen) as well as 10% fetal bovine serum (Atlas Biologicals and Sigma-Aldrich). Transfections of all cells except C33a were performed using Lipofectamine 2000 transfection reagent (Invitrogen) according to the manufacturer’s specifications. All DNA, small interfering RNA (siRNA), and transfection reagents for these experiments were diluted in Opti-MEM serum free medium (Invitrogen) in the absence of antibiotics. The calcium phosphate transfection method was used for experiments with C33a cells.

**Plasmids and siRNAs.** pGL2-E24BS contains 4 high-affinity E2 binding sites upstream of a simian virus 40 (SV40) promoter which drives luciferase expression (47). Expression plasmids used include pCG-E2, pCMVβ-p300, pRSV-CBP, and pCI-pCAF. All E2 mutations from lysine to arginine (R) were generated with the Quikchange II site-directed mutagenesis system according to the manufacturer’s specifications. All DNA, small interfering RNA (siRNA), and transfection reagents for these experiments were diluted in Opti-MEM serum free medium (Invitrogen) in the absence of antibiotics. The calcium phosphate transfection method was used for experiments with C33a cells.

**ChIP.** A total of 10^5 C127, ID13, or C127-A3 cells were plated onto 10-cm tissue culture dishes. Cells were lysed and sonified prior to chromatin immunoprecipitation (ChIP) as described previously (24). Cells to be probed for endogenous protein were harvested 24 h after plating; otherwise, cells were transfected the day after plating and harvested 24 h later. Immune complexes were captured using Dynabeads (Invitrogen), and DNA was then purified using a QIAquick PCR purification kit (Qiagen) according to the manufacturer’s specifications. Purified samples were amplified by PCR using the following BPV-1 LCR primers: sense, AAAAGTTCATGGCCTGTG; antisense, GCTTTATAGTTAGCCTGTATT.

Purified ΔNcsl BPV-1 DNA was quantified with an Eppendorf Mastercycler using the BPV-1 LCR primers listed above. A four-point, 10-fold dilution series was prepared from input DNA of each individual lysate for a reference curve. Threshold cycle (CT) values for each immunoprecipitated DNA sample were compared to the appropriate reference curves, and the amount of immunoprecipitated DNA was calculated as a percentage of input DNA. Two-way analysis of variance (ANOVA) was performed with Bonferroni’s post hoc analysis using GraphPad Prism, version 5.01.

**Immunofluorescence and immunoblotting.** A total of 3.5 × 10^5 RKO cells or 2.5 × 10^5 RPE-1 cells were plated on collagen-coated coverslips (BD Biosciences) and transfected the following day with expression plasmids. Twenty-four hours later, cells were fixed for 30 min with 3.7% paraformaldehyde (Electron Microscopy Sciences), permeabilized for 10 min with 0.2% Triton X-100, and then incubated in blocking buffer (5% normal goat serum, 1% bovine serum albumin [BSA], and 0.05% Triton X-100 in phosphate-buffered saline [PBS]) overnight. Primary and secondary antibodies were diluted in blocking buffer and incubated on the coverslips for 1 h at approximately 20°C. Coverslips were washed with 0.05% Triton X-100 in PBS and then mounted on glass slides using Pro-Long Gold antifade reagent with 4′,6-diamidino-2-phenylindole (DAPI) (Invitrogen). Cells were visualized and images collected using a Leica TSC SP2 AOBS confocal microscope with Leica imaging software. For quantification, 50 E2-positive cells were counted per coverslip and a minimum of 4 coverslips were counted for wild-type E2 and each mutant. Each coverslip represents an individual experiment. Counted cells were sorted into one of four categories, and each category was presented as a percentage of total cells counted.

RKO and RPE-1 cells for immunoblot analysis were plated onto six-well dishes at the above-stated densities and transfected the following day with expression plasmid alone or cotransfected with siRNA. Twenty-four hours posttransfection or 48 h after siRNA transfection, cells were lysed using 2% SDS in 50 mM Tris-HCl (pH 8.0) and 1 mM dithiothreitol (DTT). Following protein estimation using bicinchoninic acid (BCA) protein assay reagent (Thermo Scientific), equal amounts of protein were loaded and separated by SDS-polyacrylamide gel electrophoresis (SDS-PAGE). Gels were transferred to polyvinylidene fluoride (PVDF) membranes (Millipore), blocked, and then probed with specific antibodies. Signal was detected using Super Signal enhanced chemiluminescence substrates (Thermo Scientific).

Antibodies used for both immunofluorescence and immunoblot experiments included BPV-1 E2 B201; Santa Cruz p300 N-15 (sc-583), CBP A-22 (sc-369), and pCAF E-8 (sc-13124); and Sigma-Aldrich actin (A-2668). Secondary antibodies included Alexafluor 488 and 555 (Invitrogen) for immunofluorescence and horseradish peroxidase conjugates (Jackson Laboratory) for immunoblotting.

**In vitro acetylation assay.** Two hundred nanograms of protein substrate was mixed with 50 ng of acetyltransferase in a 10-μM reaction mixture containing 40 mM Tris-HCl (pH 8.0), 75 mM potassium chloride (KCl), and 10 μM acetyl-CoA (Sigma-Aldrich). Each reaction was carried out at 30°C for 1 h and terminated by adding SDS-PAGE sample buffer and heating the samples to 95°C for 5 min. Proteins were resolved by SDS-PAGE and transferred to a PVDF membrane, which was probed for acetylated substrates using an anti-acetyl lysine antibody (Cell Signaling Technology; catalog number 9441). Duplicate samples were separated by SDS-PAGE, and protein input was visualized using Coomassie brilliant blue R250. Substrates included BPV-1 E2 (49), p53, and histone protein (50), which were incubated with p300, pCAF, and Gcn5 (50) acetyltransferases. All proteins were purified from bacteria or baculovirus-infected Sf9 insect cells except for histone protein, which was purified from HeLa cells.
Luciferase reporter assay. A total of 3.5 × 10^5 RKO cells or 2.5 × 10^5 RPE-1 cells were plated onto six-well dishes and transfected in triplicate the following day with pGL2-E24BS, expression plasmids, and 10 nM siRNA as described previously (47). Cells were lysed 24 h posttransfection or 48 h after siRNA transfection using reporter lysis buffer (Promega), and luciferase activity was detected on an EnVision multilabel plate reader (PerkinElmer) after addition of luciferase assay reagent or Steady-Glo luciferase assay reagent (Promega) according to the manufacturer’s specifications. Means were averaged from at least four independent experiments, and error bars in figures represent standard errors of the means (SEM). Wild-type E2 transcriptional activation in the presence or absence of control siRNA was set to 100%, and each experimental value was calculated as a percentage of the wild-type activity with control siRNA. Student’s t test was performed comparing knockdown to control. *, P < 0.05; **, P < 0.01; ***, P < 0.001.

RESULTS

E2 transcriptional activity is dependent on physiological levels of p300, CBP, and pCAF. Transfection of expression vectors for p300, CBP, and pCAF has been reported to increase E2-dependent transcription (22, 42–44). How this occurs is not known. To first investigate dependence on each KAT, we used RNAi to examine the consequences of selective KAT depletion on activation of an E2-responsive luciferase reporter. BPV-1 E2 was cotransfected with a reporter plasmid and either control siRNA or siRNA targeting p300, CBP, or pCAF. E2 stimulated reporter expression nearly 80-fold, which is consistent with previous reports (47). Individual siRNA knockdown of p300, CBP, or pCAF resulted in approximately 40 to 50% reduction of wild-type E2 activity compared to that in samples transfected with the scrambled siRNA (Fig. 1A). A minimum of 50% reduction in KAT protein levels was confirmed by immunoblot analysis.

Luciferase reporter assay. A total of 3.5 × 10^5 RKO cells or 2.5 × 10^5 RPE-1 cells were plated onto six-well dishes and transfected in triplicate the following day with pGL2-E24BS, expression plasmids, and 10 nM siRNA as described previously (47). Cells were lysed 24 h posttransfection or 48 h after siRNA transfection using reporter lysis buffer (Promega), and luciferase activity was detected on an EnVision multilabel plate reader (PerkinElmer) after addition of luciferase assay reagent or Steady-Glo luciferase assay reagent (Promega) according to the manufacturer’s specifications. Means were averaged from at least four independent experiments, and error bars in figures represent standard errors of the means (SEM). Wild-type E2 transcriptional activation in the presence or absence of control siRNA was set to 100%, and each experimental value was calculated as a percentage of the wild-type activity with control siRNA. Student’s t test was performed comparing knockdown to control. *, P < 0.05; **, P < 0.01; ***, P < 0.001.

Mass spectrometry and data analysis. Following acetylation with p300, BPV-1 E2 protein was excised from a polyacrylamide gel stained with Coomassie brilliant blue R250 (Fisher). The gel band was destained and digested with trypsin, chymotrypsin, or GluC overnight at 30°C. Peptides from each sample were injected onto a Symmetry C18 trapping cartridge (Waters, Inc.) using a NanoAquity autosampler (Waters, Inc.) and separated by in-line gradient elution onto a 75-μm (internal diameter) by 10-cm column packed with a BEH 130 stationary phase (Waters). Samples were separated using a linear gradient from 3% to 90% of solvent B where solvent A was 2% acetonitrile and solvent B was 98% acetonitrile, both solvents containing 0.1% formic acid and 0.01% trifluoroacetic acid. During the gradient elution, data-dependent scans were performed with 8 scan events per cycle consisting of one full mass spectra (MS) from m/z 400 to 2,000 followed by product ion scans (collision-induced dissociation = 35%) on the 10 most intense ions in the full scan. Precursor ions used for product ion scans were dynamically excluded for 30 s. Proteins were identified from the product ion spectra using SEQUEST (Thermo Scientific) and X!Tandem search engines across the entire Swiss-Prot database (51); the results from each search were combined using Scaffold, version 3.00 (Proteome Software, Portland, OR).

RESULTS

E2 transcriptional activity is dependent on physiological levels of p300, CBP, and pCAF. Transfection of expression vectors for p300, CBP, and pCAF has been reported to increase E2-dependent transcription (22, 42–44). How this occurs is not known. To first investigate dependence on each KAT, we used RNAi to examine the consequences of selective KAT depletion on activation of an E2-responsive luciferase reporter. BPV-1 E2 was cotransfected with a reporter plasmid and either control siRNA or siRNA targeting p300, CBP, or pCAF. E2 stimulated reporter expression nearly 80-fold, which is consistent with previous reports (47). Individual siRNA knockdown of p300, CBP, or pCAF resulted in approximately 40 to 50% reduction of wild-type E2 activity compared to that in samples transfected with the scrambled siRNA (Fig. 1A). A minimum of 50% reduction in KAT protein levels was confirmed by immunoblot analysis.
Depletion of p300, CBP, and pCAF had no significant effect on E2 protein levels (Fig. 1B).

These data imply that each KAT protein has a unique function that cannot be compensated for by the remaining KAT proteins. This was unexpected, particularly for p300 and CBP, which are highly homologous and similar in function (52, 53). The residual E2-dependent transcriptional activity following depletion of each KAT might be due to the reduced but persistent levels of the targeted KAT. While we considered using p300 or CBP null mouse embryonic fibroblasts (MEFs), unrecognized compensatory changes might make results difficult to interpret. We decided to explore potential redundancy among p300, CBP, and pCAF by measuring the ability of each KAT to restore E2-dependent transcriptional activity following depletion of another KAT protein. In these experiments, we observed that overexpression of pCAF was unable to restore E2 transcriptional activity following p300 depletion (Fig. 1C); however, overexpression of CBP resulted in a small but significant increase in activity. E2-dependent transcriptional activity following siRNA silencing of CBP or pCAF was not functionally replaced through overexpression of another KAT protein (Fig. 1C). There was an increase in E2 activity following expression of CBP in pCAF-depleted cells; however, this activity was not significantly different from that in pCAF-depleted cells without overexpression and was 20% lower than E2 activity in the presence of control siRNA.

p300 and CBP interact with the BPV-1 genome. A previous report demonstrated that the presence of E2 increased pCAF-mediated acetylation of histone H3 using a reporter plasmid system (42). While this report indicates that E2 may recruit KATs, specifically pCAF, to DNA containing E2 binding sites, it did not directly examine the presence of the KAT protein at the promoter. We investigated the ability of endogenous KAT proteins to interact with the BPV-1 LCR through ChIP assay. As expected, we readily detected E2 in complex with the BPV-1 genomes present in ID13 cells and C127-A3 cells but not in parental C127 cells that do not contain BPV DNA (Fig. 2A). ChIP assay using p300 and CBP antibodies revealed their interaction with the viral genomes, specifically with the LCR, in both of the BPV-transformed cell lines (Fig. 2A). Interestingly, pCAF antibody failed to coimmunoprecipitate BPV-1 DNA in cross-linked lysates of ID13 or C127-A3 cells (Fig. 2A).

While it is difficult to evaluate quantitative differences between samples using traditional PCR methods, ChIP of the viral genome with CBP antibodies appeared to be greater with the C127-A3 cells, in which E2 protein levels are higher (Fig. 2A). To address this observation, the ability of E2 to recruit KATs to the LCR was investigated using a modified ChIP assay. C127 cells were transfected with mutant BPV-1 genomes that do not express E2, and the viral DNA in complex with the KAT was quantified by real-time PCR. Following transfection of E2, the amount of viral DNA coprecipitated by E2 antibodies increased 8-fold over background (Fig. 2A). There was a 2-fold increase in the amount of genome coprecipitated with CBP antibodies in E2-transfected samples (Fig. 2B). However, no such increases were observed following immunoprecipitation with p300 or pCAF antibodies. The amount of pCAF associated with the genome was comparable to background levels detected following control immunoprecipitation (Fig. 2B).

E2 is acetylated by p300. Previous reports investigating the functional interactions between p300, CBP, and pCAF with BPV and HPV E2 proteins have focused on the effects of KAT overexpression on E2-dependent transcriptional activation (22, 42–45). While these reports provide evidence that these proteins affect E2-dependent transcription, little is known about their mechanism of action. Despite evidence that acetyltransferase activity is necessary for enhancement by CBP and pCAF (42, 43), acetylation of E2 has not been reported. Previous attempts to identify acetylated BPV-1 E2 in vitro and in cultured cells were inconsistent or unsuccessful (data not shown) (22). These results were likely hampered by low levels of E2 proteins being immunoblotted with insensitive anti-acetyl lysine antibodies. We performed in vitro acetylation reactions using p300, pCAF, and Gcn5 (a pCAF paralog) proteins that were bacterially expressed or purified from baculovirus-infected Sf9 cells. Purified KAT and E2 proteins were combined in the presence of acetyl-CoA, and resulting acetylated protein was detected by immunoblot analysis with an acetyl lysine antibody. Purified histones and p53 were included as positive controls for acetylation. Robust acetylation was detected on histone substrates by each KAT (Fig. 3A). p300 mediated strong acetylation of p53, while pCAF and Gcn5 were much less active (Fig. 3A). This may reflect the fact that p300 acetylates p53 on several more lysines than does pCAF or Gcn5 (54, 55) or an epitope preference for the acetyl lysine antibody. After incubation of BPV-1 E2 with p300, a strong band corresponding to acetylated E2 was observed. Acetylation of E2 was not detected in reaction mixtures containing either pCAF or Gcn5 (Fig. 3A). Autoacetylation of the short isoform of Gcn5 used in
this experiment is detectable in these blots at a migration slightly slower than those of E2 and p53 (50).

Lysines in BPV-1 E2 that were targeted by p300 were identified using a proteomic approach. BPV-1 E2 was acetylated in vitro and digested, and the resulting peptides were analyzed by mass spectrometry. Each peptide was compared to the entire Swiss-Prot database and assembled using both the SEQUEST and X!Tandem search engines. These two utilities directly compare uninterpreted tandem mass spectra to protein databases using different algorithms, resulting in protein identification. Acetylated peptides of BPV-1 E2 were not identified following trypsin digestion. This is likely due to the inability of trypsin to cleave after acetylated lysines. Analysis of BPV-1 E2 digested with chymotrypsin yielded 97% total sequence coverage and identified 11 acetylated lysine residues (Fig. 3B). Analysis of GluC digestion yielded 89% sequence coverage and nine acetylated lysines, of which three were unique to GluC digestion (Fig. 3B). Nine acetylated lysines were found in the N-terminal TAD, five in the C-terminal DBD, and one in the central hinge region. Acetylated lysines were discovered in an average of 22% of each unique E2 peptide identified. The frequency of lysine acetylation in each unique peptide ranged from 3% to 60%. The low average frequency of acetylation could explain the difficulty detecting acetylated E2 in vivo. A summary of data compiled from chymotrypsin and GluC digests, including SEQUEST and X!Tandem correlation scores, is presented in Table 1.

E2 lysine mutations display transcriptional defects. We next sought to characterize the functional significance of E2 acetylation by p300 in vivo. Six lysines identified in the proteomic analysis were selected for mutation on the bases of sequence conservation among BPV-1 and several HPV types, frequency with which the modified lysine was identified in proteomic analysis, and SEQUEST and X!Tandem correlation scores. Lysines 107, 111, and 112 are part of a putative BPV-1 E2 N-terminal nuclear localization signal (NLS) (56). Lysines 111 and 112 are conserved throughout eight HPV types, including high-risk 16, 18, and 31 (Fig. 4A), and lysine 339 is a critical residue that mediates interaction with E2 DNA binding sites (57). These lysines were conservatively mutated to arginine to avoid effects related to variations of residue charge. Each mutant was screened for its ability to activate transcription from an E2-responsive reporter and for protein expression levels.

Mutation of BPV-1 E2 lysines (K) 70 and 391 to arginine (R) resulted in transcriptional activity equivalent to that observed with wild-type E2 in RPE-1 cells (Fig. 4B) despite protein levels approximately 30% lower than that of the wild type (Fig. 4C). The K107R E2 mutant protein was present at ~85% of the wild type, with a proportional decrease in its transcriptional activity (Fig. 4B and C). K112R E2 was present at levels similar to those of K107R E2 and displayed comparable transcriptional activity. This activity is not significantly different from that of wild-type E2 in RPE-1 cells; however, when K112R is transfected into C33a cells, the mean transcriptional activity is slightly lower (64.5% for RPE-1 cells and 58.1% in C33a cells), resulting in a significant difference (Fig. 4B and D). The K339R E2 mutant was impaired in its ability to activate transcription in RPE-1 cells (Fig. 4B). This may be partially attributable to protein levels at 45% of that of wild-type E2; however, transfection of increasing amounts of the K339R mutant resulted in a peak of transcriptional activity at less than 50% of that observed for wild-type E2 (data not shown). K111R and double-mutant K111R/K112R proteins were expressed at lower levels (Fig. 4C).

Table 1: Proteomic analysis of acetylated E2

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<th>Highest X!Tandem Score</th>
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Table 2: E2 Acetylation and Transcription

E2 lysine mutations display transcriptional defects. We next sought to characterize the functional significance of E2 acetylation by p300 in vivo. Six lysines identified in the proteomic analysis were selected for mutation on the bases of sequence conservation among BPV-1 and several HPV types, frequency with which the modified lysine was identified in proteomic analysis, and SEQUEST and X!Tandem correlation scores. Lysines 107, 111, and 112 are part of a putative BPV-1 E2 N-terminal nuclear localization signal (NLS) (56). Lysines 111 and 112 are conserved throughout eight HPV types, including high-risk 16, 18, and 31 (Fig. 4A), and lysine 339 is a critical residue that mediates interaction with E2 DNA binding sites (57). These lysines were conservatively mutated to arginine to avoid effects related to variations of residue charge. Each mutant was screened for its ability to activate transcription from an E2-responsive reporter and for protein expression levels.

Mutation of BPV-1 E2 lysines (K) 70 and 391 to arginine (R) resulted in transcriptional activity equivalent to that observed with wild-type E2 in RPE-1 cells (Fig. 4B) despite protein levels approximately 30% lower than that of the wild type (Fig. 4C). The K107R E2 mutant protein was present at ~85% of the wild type, with a proportional decrease in its transcriptional activity (Fig. 4B and C). K112R E2 was present at levels similar to those of K107R E2 and displayed comparable transcriptional activity. This activity is not significantly different from that of wild-type E2 in RPE-1 cells; however, when K112R is transfected into C33a cells, the mean transcriptional activity is slightly lower (64.5% for RPE-1 cells and 58.1% in C33a cells), resulting in a significant difference (Fig. 4B and D). The K339R E2 mutant was impaired in its ability to activate transcription in RPE-1 cells (Fig. 4B). This may be partially attributable to protein levels at 45% of that of wild-type E2; however, transfection of increasing amounts of the K339R mutant resulted in a peak of transcriptional activity at less than 50% of that observed for wild-type E2 (data not shown). K111R and double-mutant K111R/K112R proteins were expressed at lower levels (Fig. 4C).
While all other E2 mutants tested precipitated the BPV genome at levels comparable to those of wild-type E2, precipitation of the K111R and K111R/K112R mutants with the genome was reduced (data not shown). The K111R and K111R/K112R mutants were both severely impaired in their ability to activate transcription in RPE-1 and C33a cells (Fig. 4B and D). The transcriptional defects of the K111R and K111R/K112R mutants were not due to reduced protein levels; transfection of increasing amounts of either construct had no effect on transcriptional activity (Fig. 4E).

**E2 lysines 111 and 112 are important for nuclear retention.** A region that is rich in basic amino acids in the N-terminal transcriptional domain (TAD) of BPV-1 E2 has been reported to function as a NLS in the absence of the highly conserved C-terminal NLS (56). Four of the six lysines mutated in this study are within these two NLS sequences and prompted examination of mutant E2 localization. The subcellular distributions of four mutants, the K70R, K107R, K339R, and K391R mutants, were nearly indistinguishable from that of wild-type E2; nuclear localization was observed in nearly all cases (Fig. 5A). The three remaining K111R, K112R, and K111R/K112R mutant proteins deviated from wild-type localization and exhibit a spectrum of subcellular distributions ranging from exclusively cytosolic to exclusively nuclear (Fig. 5A). The severity of mislocalization, which was characterized as the extent of observed nuclear exclusion, was different for each mutant protein. The K112R mutant was observed to be more nuclear than the K111R/K112R double mutant, which was predominantly cytosolic. The K111R mutant displayed an intermediate phenotype, although its increased cytosolic localization more closely resembled that observed for the K111R/K112R mutant than for the K112R mutant (Fig. 5A).

Quantifying the extent of mislocalization was necessary to fully characterize these mutants due to the spectrum of localization patterns observed for the K111R, K112R, and K111R/K112R mutants. The localization spectrum was divided into four discrete categories. E2 mutant-expressing cells were visually inspected, and each cell was assigned to a category according to predetermined selection criteria. All three mutant proteins were considerably mislocalized to the cytosol (Fig. 5B). The K111R and K111R/K112R mutants were most commonly found to be evenly distributed throughout both the nuclear and cytosolic compartments (Fig. 5B). While the localization profile of the K111R/K112R mutant was similar to that of the K111R mutant, it was found to be completely excluded from the nucleus more often than either of the other mutants (Fig. 5B; compare second and fourth graphs). While expression of the K112R mutant resulted in cytosolic accumulation, E2 staining was more prevalent in the nuclear compartment (Fig. 5B). The increase in cytosolic localization for each mutant is not due to Crm-1–mediated nuclear export; treatment of mutant-transfected cells with leptomycin B (LMB) had no effect on the distribution of E2 mutant proteins (data not shown).

**p300 cannot enhance the transcriptional activity of E2 K111 mutants.** The mechanism by which p300 enhances E2-dependent transcription is unclear (22, 44, 45). The K111R and K111R/K112R E2 mutants retain their overall charge but cannot be acetylated and were severely defective transcriptionally. The relationship between K111 and p300 was further addressed in RKO cells that do not express endogenous p300 (58). Coexpression of p300 with wild-type E2 enhanced activation of E2-dependent transcription in RKO cells an average of 2-fold (Fig. 6). The K111R and K111R/K112R E2 mutants showed 10 to 20% of the transcriptional activity of wild-type E2, and this activity was not enhanced by coexpression of p300 (Fig. 6).

The ability of wild-type E2 to activate transcription in RKO cells suggests that p300-mediated acetylation of lysine 111 is not necessary for this function. We also considered that in these cells, the p300 paralog CBP may be compensating for the absence of p300. Interestingly, RNAi depletion of CBP in RKO cells dramatically reduced the activity of wild-type E2, comparable to levels observed with the K111R and K111R/K112R E2 mutants (Fig. 6).
p300 and CBP promote E2 nuclear localization. While we demonstrate that E2 is acetylated by p300 on K111 and K112, the role of acetylation in the mislocalization of the E2 mutants is unclear. The contribution of p300 to E2 localization was investigated in RKO cells. Expression of wild-type E2 in these p300-deficient cells resulted in accumulation of E2 protein in the cytosol (Fig. 7A), resembling the pattern observed for the K112R mutant in RPE-1 cells (Fig. 5A). Transfection of p300 restored nuclear localization of E2 (Fig. 7A). Introduction of p300 had no effect on the cytosolic mislocalization of the K111R, K112R, and K111R/K112R E2 mutants in RKO cells (data not shown). The reduction of E2 transcriptional activity in RKO cells following depletion of CBP prompted investigation into a compensatory role for this protein in E2 nuclear localization. Following depletion of CBP, wild-type E2 was present diffusely throughout the cytosol and nucleus (Fig. 7B). This phenotype was similar to that of the K111R mutant in RPE-1 cells (Fig. 5A).

DISCUSSION

The lysine acetyltransferases p300, CBP, and pCAF have been reported to interact with the E2 proteins from several papillomaviruses (22, 42, 43). In each instance, overexpression of the KAT protein was found to increase E2-dependent transcription. In contrast, we used RNAi to specifically deplete p300, CBP, and pCAF under conditions in which the other KAT proteins remained present at endogenous levels. Interestingly, knockdown of each KAT reduced E2-dependent transcriptional activation. This demonstrates that the functions of the
FIG 6 p300 does not enhance the transcriptional activity of E2 K111R mutants. RKO cells were transfected with an E2-responsive luciferase reporter, wild-type or mutant E2, and a p300 expression construct or empty vector. The sample transfected with wild-type E2 was cotransfected with control siRNA for comparison to the siCBP-transfected sample. Results are presented as a percentage of wild-type E2 activity. One-way ANOVA was performed with Bonferroni’s post hoc analysis comparing all means. ***, P < 0.001. ns, not significant.

silenced KATs were not being replaced by the other nontargeted KAT proteins and implies that each KAT mediates nonoverlapping activities. The requisite KAT activities were then investigated. We demonstrated that p300 and CBP are present at the BPV-1 LCR by chromatin immunoprecipitation assay and that CBP was enriched at the LCR in the presence of E2 about 2-fold. We suggest that E2 recruitment of CBP to the viral promoter may serve to acetylate histones and lead to increased chromatin access for the RNA polymerase complex.

Lysine acetylation of a transcription factor leading to its activation is well documented (41); however, this has not been reported for any papillomavirus E2 protein. Because we were unable to isolate a sufficient amount of BPV E2 protein for microsequencing from mammalian cells with replicating BPV-1 genomes, we performed in vitro reactions using purified E2 and KAT proteins. Several E2 lysine residues were found to be acetylated. To test their biological relevance, each lysine was replaced with arginine to maintain charge at each position. Mutations of K111 alone and in combination with K112 resulted in substantial decreases in transcriptional activation.

Several reports have described mutation of E2 at lysines 111 and 112, which are highly conserved (59–62). Replacement of these lysines with alanines in BPV-1 E2 results in proteins that are transcriptionally defective in several cell types (59, 61). Additionally, the same mutations resulted in E2 mislocalization to the perinuclear region (59), indicating a possible deficiency in nuclear import. The conservative substitution of BPV-1 E2 lysine 111 to arginine reportedly results in severely impaired transcriptional activity, and K112R mutation was shown to activate transcription to near wild-type levels (60). These results are consistent with our data for the K111R mutant, and while we demonstrate that K112R mutant activity is lower than was previously shown, it is not significantly different from that of wild-type E2 in RPE-1 cells. The activity for the K112R mutant in C33a cells, however, is slightly lower, resulting in a significant difference that may also be attributable to variations in activity between cell types, which we suspect may relate to different KAT protein availability.

K111R and K111R/K112R mutant proteins were expressed at reduced levels. This likely reflects their lack of transcriptional activity; we routinely observe that E2 stimulates its own transcription in transient assays (63, 64). However, titration of increasing amounts of the K111R mutant did not result in increased activation, indicating that the deficiency is not due to a dosage effect (Fig. 4E). The reduction of transcriptional activity with K111 mutation to arginine suggests that acetylation of this residue is necessary for transcriptional activation.

The subcellular distribution of E2 may also be modulated through acetylation of lysines 111 and 112. This is supported by our finding that conservative mutation of these two residues to arginine resulted in E2 accumulation in the cytosol. Similar cytosolic accumulation was observed with wild-type E2 in cells in which p300 and CBP were absent. Our results for the K111R and K112R mutants are consistent with previous studies examining the localization of E2 proteins with deletions that disrupted the N-terminal NLS including lysines 111 and 112 (56) and for mutants with changes of lysine to alanine at both of these positions (59). Incomplete inclusion or exclusion of the mutants suggests that either inefficient nuclear import due to a weakened NLS or passive diffusion may be responsible. A defect in Crm-1-dependent nuclear export may be excluded given that LMB had no effect on E2 mutant localization. Another possibility is that K111 acetylation favors E2 retention within the nucleus. Notably, it was recently proposed that pCAF-mediated acetylation of lysines within the NLS of the retinoblastoma proteins promotes its nuclear retention (65). This scenario can be envisioned by the entry of E2 into a macromolecular KAT complex that prevents its loss to the cytosol.

How E2 acetylation on K111 stimulates its transcriptional activity requires further study. One possibility is that this creates an
additional site for docking to the bromodomain of Brd4, which facilitates E2-dependent transcriptional activation (20, 23, 66, 67). While bromodomains have affinity for most acetylated lysines, they do exhibit binding preference (68, 69). The bromodomain interaction preference on histone H3 and H4 peptides is for di-acetylated lysines (70, 71). This would indicate that acetylation of a combination of K111, K112, and possibly K107 would result in a higher-affinity interaction. A co-crystal of the N-terminal region of E2 with a short C-terminal peptide of Brd4 revealed the site of their interaction to span the three alpha helices and not the K111-containing peptide, which is at the N terminus of the beta sheet region (66). However, HPV-11 E2 has been demonstrated to interact with a region containing the second bromodomain of Brd4 in addition to the CTD (12). This supports the possibility that a second region of E2 may also participate in this interaction. Although the structures of several functional domains on Brd4, including both bromodomains and the C-terminal domain that interacts with E2, have been solved, the full-length protein structure has not been reported.

We previously reported that the transcriptional coactivator Gps2 is necessary for E2 transcription and directly binds to both p300 and E2 (22). E2 association with p300 using bacterially expressed proteins was weak and barely above background (22). Gps2 binding was mapped to residues 134 to 216 on the BPV-1 E2 TAD (47). We hypothesize that the interaction between E2 and Gps2 facilitates complex formation with p300, and this leads to acetylation of lysine 111. This, in turn, may stabilize the association of Brd4 in the transcription complex.

Here, we have outlined potential roles for the cellular acetyltransferases p300 and CBP in E2-dependent transcription. A specific role for pCAF in E2-dependent transcriptional control was not identified in this study. Acetylation of E2 by pCAF or Gcn5 may occur in vivo despite our inability to detect it using the acetyl lysine antisera in in vitro reactions. It is possible that modification of E2 by these KATs occurs only under specific conditions. While data indicate that p300, CBP, and pCAF may function independently, there does appear to be some biochemical redundancy. The reduction in transcriptional activation in RKO cells following CBP depletion suggests that CBP may be actively compensating for the loss of p300 in these cells. This is also consistent with a small but significant increase in activity following overexpression of CBP in RPE-1 cells that are depleted of p300.

It has been reported that p300 levels are low in basal keratinocytes and may increase with differentiation (44). As these epithelial cells transit into the suprabasal layers, increased expression of p300 favors acetylation of p53 and induction of p21Waf1/CIP1, mediated cell cycle arrest, further promoting commitment to keratinocyte differentiation (72). Our results suggest that elevated p300 expression in differentiating cells could also induce K111 acetylation, thereby stimulating E2-dependent transcription. The resulting increase in early gene protein levels, especially E1, could induce viral DNA amplification, linking it to the differentiated status of the infected cell. The role of K111 acetylation by p300 in regulation of the viral replicative program is currently under active investigation.

ACKNOWLEDGMENTS

This work was supported by NIH grants R01CA58376 to Elliot J. Androphy and R01CA103867 and R01CA124760 to Cheng-Ming Chiang. Sara P. Culleton was supported by the NIH (T32 AI 060519).

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February 2013 Volume 87 Number 3 jvi.asm.org 1505


