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The Catalytic Mechanism of Histone Acetyltransferase p300: From the Proton Transfer to Acetylation Reaction

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Abstract

The transcriptional coactivator and histone acetyltransferase (HAT) p300 acetylates the four core histones and other transcription factors to regulate a plethora of fundamental biological processes including cell growth, development, oncogenesis and apoptosis. Recent structural and biochemical studies on the p300 HAT domain revealed a Theorell-Chance, or "hit-and-run", catalytic mechanism. Nonetheless, the chemical mechanism of the entire reaction process including the proton transfer (PT) scheme and consequent acetylation reaction route remains unclear. In this study, a combined computational strategy consisting of molecular modeling, molecular dynamic (MD) simulation and quantum mechanics/molecular mechanics (QM/MM) simulation was applied to elucidate these important issues. An initial p300/H3/Ac-CoA complex structure was modeled and optimized using a 100 ns MD simulation. Residues that play important roles in substrate binding and the acetylation reaction were comprehensively investigated. For the first time, these studies reveal a plausible PT scheme consisting of Y1394, D1507 and a conserved crystallographic water molecule, with all components of the scheme being stable during the MD simulation and the energy barrier low for PT to occur. The two-dimensional potential energy surface for the nucleophilic attack process was also calculated. The comparison of potential energies for two possible elimination half-reaction mechanisms revealed that Y1467 reprotonates

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Supporting Information

Supporting Information Available: The model selection description of L1 loop in the structure of the p300/H3/Ac-CoA, Figure S1–9 and Table S1 including H3 peptide structural alignment during the MD simulation, structure mapping of the PT processes in three HAT subfamilies, RMSFs of MD simulation on different models, the interaction between L1 Loop and neighboring residues, and the predicted pKa for ionizable residues of the active site and PT scheme with different internal dielectric constant. This material is available free of charge via the Internet at http://pubs.acs.org.

the coenzyme-A leaving group to form product. This study provides new insights into the detailed catalytic mechanism of p300 and has important implications for the discovery of novel small molecule regulators for p300.

Keywords

Epigenetics; p300; Proton Transfer; Histone Acetylation; QM/MM

Introduction

Histone modifications such as methylation, acetylation, ADP-ribosylation, ubiquitination, citrullination, and phosphorylation regulate specific and distinct functional outputs of genomes at the transcription level^{1, 2}. Among all these modifications, histone acetylation mediated by histone acetyltransferases (HATs), which occurs both on the unstructured tails and the globular core domains of histones, controls many aspects of chromatin structure and function^{3–6}. In addition, HATs also exert various influences on the genome by catalyzing the lysine acetylation of many other cellular proteins including themselves^{7–9}. Different HAT members show varying degrees of substrate specificity, a property consistent with a role in maintaining a dynamic, acetylation-based epigenetic code^{3, 10}. Moreover, recent evidence raises an interesting possibility that an acetylation-based code may operate through both mitosis and meiosis, providing a possible mechanism for germ-line transmission of epigenetic changes^{10–13}.

P300, one of the major HAT proteins as it controls the expression of numerous genes in a cell-type-specific and signal-dependent manner. Accordingly, p300 has been shown to play an important role in cellular proliferation, apoptosis, and embryogenesis ^{14–16}. More recently, p300 has been shown to play a role in regulating embryonic stem (ES) cell pluripotency by modulating the expression of the pluripotency marker Nanog ¹⁷, and to modulate the specification of liver progenitors over pancreas progenitors ¹⁸. Additional data demonstrates that abnormal p300 function is associated with deregulated target gene expression, and is implicated in inflammation, cancer, cardiac hypertrophy, and genetic disorders such as the Rubinstein-Taybi syndrome, making p300 an attractive therapeutic target ^{19–21}. Because of these associations, small molecular p300 modulators may apply to determining the direction of stem cell differentiation ¹⁸, be useful as probes to study p300 function ^{22–25} for developing new chemical entities for therapeutic use ^{26–28}. Rational design of small molecules to interfere with p300 function on the basis of the catalytic mechanism and transition state structures may represent a feasible and efficient way to achieve these goals²⁹.

Most HATs utilize two distinct mechanisms to catalyze the acetyl-transfer reaction. One is ternary complex mechanism, in which the cofactor acetyl coenzyme A (Ac-CoA) and the substrate bind to the enzyme simultaneously followed by the acetyl transfer reaction in the HAT/Ac-CoA/substrate ternary complex³⁰. The other mechanism is a ping-pong mechanism, in which the acetyl group is covalently attached to the enzyme before being transferred to the substrate³¹. One major HAT family, GCN5, was shown to use the ternary

complex mechanism^{32–34}, whereas another major HAT family, MYST, was shown to employ the ping-pong mechanism³⁵ or the ternary complex mechanism³⁶. Intriguingly, it was reported that neither of these mechanisms applies to p300. Instead, p300 was shown to use a Theorell-Chance, or "hit-and-run", catalytic mechanism³⁷. As a special case of the ternary mechanism, ternary complex is also generated in Theorell-Chance mechanism. While distinct from the classic ternary mechanism, Theorell-Chance is characterized that the ternary complex never accumulates and the steady-state concentrations of the ternary complex is kinetically insignificant. Following the association of Ac-CoA, the protein substrate binds transiently to the p300 surface, allowing the lysine residue to snake through the enzyme active site to receive the acetyl group, followed by rapid protein dissociation. However, answers to several important questions regarding the detailed molecular mechanism of catalysis remain unanswered: 1)how does the lysine substrate bind to the enzyme; 2) how is the e-amino group of the substrate lysine deprotonated prior to the reaction; 3) what is the chemical pathway for the entire catalytic process and which structural elements of p300 are critical for this process.

In this study, a p300/H3/Ac-CoA ternary structure model was constructed on the basis of the crystal structures of the p300/Lys-CoA and GCN5/H3/Ac-CoA complexes. Then a 100 ns MD simulation was performed on the complex model to obtain a suitable structural configuration for further investigations. The recognition network of substrate H3 was analyzed and compared with the experimental data. Meanwhile, two possible proton transfer (PT) schemes were proposed based on structural analysis, and the scheme consisting of Y1394, D1507 and a crystallographic water molecule important for the PT was validated by MD simulations and QM/MM calculations. Subsequently, the entire acetylation reaction pathway proceeding through a ternary complex was fully investigated by QM/MM calculations. Two residues in the active site that were previously shown to be important for catalysis, Y1467 and W1436, were also investigated. In summary, these computational studies provide a deeper understanding of Ac-CoA cofactor and lysine substrate binding, the PT pathway and the acetylation reaction mechanism of p300 and its related HAT proteins. The snapshots of the transition states of the acetyl-transfer reaction offers important insights for the design of small molecule modulators for regulating p300 function.

Experimental section

Preparation of the simulation system

To date, the only crystal structure containing p300 is that of the p300/Lys-CoA complex. No p300/Ac-CoA/peptide complex structure is available. Therefore, we modeled a p300/H3/Ac-CoA complex based on the crystal structures of p300/Lys-CoA (PDB code: 3BIY)³⁷ and GCN5/H3/Ac-CoA³⁸ (PDB code: 1PU9) complexes. First, the lysine portion of the Lys-CoA was modified to an acetyl group to generate a p300/Ac-CoA structure. Second, the crystal structure of the GCN5 HAT bound to a 15-residue H3 peptide was aligned to p300/Ac-CoA by structural alignment with PyMOL plugin CEAlign³⁹, then the H3 peptide was extracted to construct the p300/H3/Ac-CoA complex model. Since the H3 peptide is involved in the catalytic reaction of p300. and whether it locates on the top of the L1 loop or bound to the bottom of the binding site is still a question, therefore, we performed a series

computational and structural analysis to select the H3 peptide conformation (see in the section of "The model selection of H3 conformation in the structure of the p300/H3/Ac-CoA model" in supporting information). Briefly, considering the structural and biophysical factors, the H3 substrate peptide binding to a shallow pocket on the top L1 loop will be modeled for further analysis.

The autoacetylation loop, which consists of L1520–L1580, was genetically deleted during crystallization and the resulting p300 HAT domain showed comparable acetyltransferase activity with the intact enzyme, thereby we didn't include it in our model $^{40, 41, 37}$. This model was then minimized using the AMBER force field 42 with the following parameters: a distance-dependent dielectric function, a non-bonded cutoff of 8 Å, amber charges for the enzyme and H3 peptide, and Gastieger-Hückel charges for Ac-CoA and water molecules. The structures were minimized by the simplex method, followed by the Powell method to an energy gradient of < 0.05 kcal/(mol·Å). All procedures were performed using the Sybyl software package (Tripos, St. Louis, MO).

MD simulations

A 100 ns MD simulation was performed on the p300/H3/Ac-CoA system. Before simulations, this system was put into a suitably sized box with a 10 Å buffer distance between the solvent box wall and the nearest solute atoms. Then the box was solvated with the SPC (Simple Point Charge) water model. Each solvated system was submitted to energy minimization. Afterwards, counterions were added to the system to provide a neutral simulation system. The whole system was subsequently minimized again. The charges of the atoms of Ac-CoA were calculated by using the RESP method⁴³ encoded in the AMBER suite program⁴⁴ at the level of RHF/6-31G*. Covalent and non-bonded parameters for the ligand atoms were assigned, by analogy or though interpolation, from those already present in the AMBER force field.

MD simulations were carried out using the AMBER package (Version 10.0) with constant temperature, constant pressure (NPT), and periodic boundary conditions. The AMBER Parm99 force field⁴² was applied for the enzyme and H3 peptide. The Particle Mesh Ewald method⁴⁵ was used to calculate the long-range electrostatics interactions. The non-bonded cutoff was set to 10.0 Å, and the non-bonded pairs were updated every 25 steps. The SHAKE method⁴⁶ was applied to constrain all covalent bonds involving hydrogen atoms. Each simulation was coupled to a 300 K thermal bath at 1.0 atm of pressure (1 atm=101.3 kPa) by applying the algorithm of Berendsen et al⁴⁷. The temperature and pressure coupling parameters were set as 1 ps. An integration step of 2 fs was set up for the MD simulations.

QM/MM calculations

The hybrid QM/MM mechanical approaches have been widely employed to enzymatic reaction studies^{48, 49}. In this research, QM/MM calculations were performed by using a two-layered ONIOM method⁹ encoded in the Gaussian 03 program. In this approach, the molecular system under study is divided into an inner layer and an outer layer. The inner layer consists of the most critical elements of the system, and the rest of the system comprises the outer layer, the full system is called "real" and is treated with a low level of

theory. The inner layer is termed "model" and is treated with both a low and a high level of theory. The total ONIOM energy EONIOM is given as following:

where E(high, model) is the energy of the inner layer (plus the link atoms) at the high level of the theory, E(low, real) is the energy of the entire system at the low level of theory, and E(low, model) is the energy of the model system at the low level of theory. Thus, the ONIOM method allows one to perform a high-level calculation on just a small, critical part of the molecular system and incorporate the effects of the surrounding elements at a lower level of the theory to yield a consistent energy expression on the full system.

The minimized p300/H3/Ac-CoA complex system optimized using the AMBER Parm99 force field was further optimized at the ONIOM (B3LYP/6-31G*:Amber) level to investigate the PT and acetylation mechanism. The quantum mechanics (QM) region for the PT consists of the methylene amonium CH2-CH2-NH3+ group of H3K14, the water molecule in the transfer wire (Wat1861 in the crystal structure), the side chain of Y1394 and D1507 (the proton shuttling residues), and the backbone of I1395 and W1436 (h-bonding with the proton shutting residues). The QM region for the acetylation reaction consists of the deprotonated side chain (-CH2-CH2-NH2) of H3K14, the -CH2-S-COCH3 group of Ac-CoA, the main chain of W1436, and side chain of Tyr1467. Each QM region was calculated by using the density functional theory with the B3LYP exchange-correlation functional and 6-31G* basis set. The remainder of the system (MM region) was treated with the AMBER Parm99 force field. A total of 5,512 atoms were included for the QM/MM calculations by using the ONIOM module as implemented in Gaussian 03. The electrostatic interactions between the QM and MM regions were calculated by using the electronic embedding method, which treats the polarization of the QM region by the MM region with scaled partial atomic charges of MM atoms, and the response of the QM region with the Merz-Singh-Kollman scheme for charge fitting so as to produce the changing partial charges of the QM atoms.

Results and discussions

The overall structure of the p300/H3/Ac-CoA model

Initially, the p300/H3/Ac-CoA model was constructed as described in materials and methods

The overall structure of p300 can be divided into two main parts according to the structural features of all HAT families, as shown in Figure 1: 1) the central core region, consisting of the $\alpha 3$ helix and $\beta 2$ – $\beta 5$ strands that are structurally conserved in all HAT enzymes. This region participates in Ac-CoA binding and in templating the lysine side chain for acetyl-transfer $^{50,\,51}$. 2) The flanking N- and C-terminal regions, which are structurally divergent between the different HAT subfamilies. These more variable regions have been shown to participate in histone substrate binding by forming a binding cleft that flanks the central core region $^{50,\,51}$. Structural comparison reveals that this cleft in p300 is highly acidic and shallower than the corresponding clefts in the Gcn5/PCAF and MYST HAT subfamilies. H-bond interactions and hydrophobic interactions between H3/p300 complex are also shown in

Figure 1A and Figure 1B, respectively. p300 harbors an additional L1 loop in its N-terminal region, which is unique to this subfamily of HATs and participates in both Ac-CoA and lysine substrate binding. The autoacetylation loop, which consists of L1520–L1580, was genetically deleted during crystallization and the resulting p300 HAT domain showed comparable acetyltransferase activity with the intact enzyme, thereby we didn't include it in our model^{37, 52, 53}.

To investigate the stability of the active site cavity, a 100-ns MD simulation was performed on the p300/H3/Ac-CoA complex. Firstly, the protonation states of key residues in active site and involved in PT were determined computationally by H++ program⁵⁴. As shown in Table S1, the basic residues H3K14, Y1446 and Y1467 around the active site, and the PT shuttling residue Y1394 were largely protonated. While the depronation state was the dominate for D1444 and D1507, which are considered as the proton acceptor residues in PT process. Therefore, the calculation results were applied to the model used in MD simulation and the QM/MM study.

During the MD simulation, the root-mean-square deviation (RMSD) was monitored (from t= Ops), using the backbone coordinates of the model structure as the reference (Figure 2A). In addition, to represent the stability of residues, the root-mean-square fluctuation (RMSF) of p300 in the complex model was also calculated. The B-factor (thermal factor) of p300 in the crystal structure of p300/Lys-CoA, was used as a measure for any type of displacement of an atom from its mean position, and was extracted and transformed to RMSF using the formula $B=8\pi^2*RMSF^2/3$ (Figure 2B). The RMSD tends to be flat after a 20 ns' simulation, which indicates that structures from 20 to 100 ns are more reasonable. Notably, the RMSD of H3 is apparently higher than other components of the complex. Consistently, the average RMSD of apo-p300 is only about 1.8 Å while that of the complex is above 2.2 Å, likely due to the flexibility of the H3 peptide. This result is in good agreement with the Theorell-Chance mechanism, where the protein substrate is proposed to bind p300 for acetylation and leave rapidly before forming a stable ternary complex. The consistency between the theoretical and experimental RMSF shown in Figure 2B validates the quality of our model. We propose that the peaks of the RMSF curve represent the different loop regions of the protein. Interestingly, the, 23-resisue L1 loop (residues 1436–1459) is fairly stable during the simulation, displaying a lower RMSF value than other regions of p300 with defined secondary structure. This is consistent with the extensive interactions that the L1 loop makes with Ac-CoA in the p300/Lys-CoA crystal structure.

H-bond and hydrophobic interaction analysis for the p300/H3/Ac-CoA model

To probe the molecular basis for H3 binding, both the H-bond and hydrophobic interactions between p300 and H3 were analyzed. First, the reaction residue H3K14 h-bonds strongly with the main chain oxygen atom of W1436. Its adjacent residue A315 h-bonds with P1439 and P1440 on the L1 loop, which contributes to further stabilize the H3K14 interaction. Second, two h-bond networks located at both ends of H3 together display important roles for anchoring the peptide chain. For one of these h-bonding networks, the electropositive side chains of terminal A7, R8 and K9 form an interaction network with the electronegative side chains of E1505, D1625 and D1628 from p300. This interaction is consistent with previous

in vitro studies showing that the D1625R/D1628R mutant shows significantly decreased p300 HAT activity, further supporting the importance of this h-bond network for peptide substrate binding⁷. This interaction network provides a rationale for the observed preference for additional Lysine or Arginine residues two to three residues downstream or upstream of the primary acetylation site. For the other h-bonding network, R17, Q19 and A21 of H3 interact with E1442, K1456, I1457, K1459 and R1462 of p300, with most of these residues coming from the L1 loop. This same anchoring pattern (one h-bond network at each end of the histone) was also observed in the Gcn5 HAT subfamily^{38, 55}. R8 and Q19 in the tGCN5/H3/Ac-CoA system and K166 and Q176 in the hGCN5/H3/Ac-CoA system h-bond with the enzyme at both ends of the H3 peptide, just like the corresponding K9 and Q19 residues in the p300/H3/Ac-CoA system. Third, Y1397 in the middle of the substrate binding cleft h-bonds with the main chain carbonyls of T11 and G12 next to the substrate lysine, which acts like a bridge to connect the substrate lysine and positively charged region mentioned above to help maintain the configuration of the H3 peptide.

Hydrophobic interactions between p300 and the H3 peptide are not only conducive to H3 binding, but also provide a favorable environment for nucleophilic attack of the deprotonated lysine side chain. It has been reported that in tGCN5 and hGCN5, there is a G-K-X-P recognition sequence in the middle of the histone peptide, which is responsible for most of the hydrophobic interactions between H3 and the enzyme^{38, 55}. In our simulation, most hydrophobic contacts are located from G12 to P16 of the H3 peptide, which includes a G-K-A-P region. In the reaction center, H3K14 is surrounded by the hydrophobic side chains of Y1397, W1436, P1439, P1440 and Y1446. This highly hydrophobic environment could greatly facilitate the deprotonation of H3K14 and subsequent nucleophilic attack.

It is worth noting the structure alignment of different snapshots at a 20 ns interval also shows the fluctuations of H3 (Figure S6). Interestingly, the binding cleft is relatively shallow and polar comparing to GCN5 and Esa1 HATs. As required by the Theorell-Chance mechanism, the protein substrates associate weakly with the enzyme so that a rapid disassociation would occur and the ternary complex never accumulates during catalytic turnover. Taken together, the unstable and flexibility characteristic of the H3 peptide bound to p300 strongly supports the Theorell-Chance mechanism.

Identification of rational PT scheme

As the positive charged state of reaction residue H3K14 was the dominating form under physiological environment (Table S1), the first step of the acetylation reaction should involves the deprotonation of the lysine side chain. For the Esa1 MYST HAT, the lysine proton is directly delivered to a conserved glutamate residue (E338)³⁵. In contrast to Esa1, in the reaction center of p300 there is no electronegative residue within 5 Å of the ε-N atom of the target lysine, which leads to a hypothesis for the existence of a long-range mechanism in the PT process. In this case, the proton acceptor may be far from the lysine, the proton donor, and the long-range PT is usually mediated by waters or residues containing a hydroxyl group^{56, 57}. We also note that in the GCN5 HAT, the proton is delivered to a conserved glutamate (E173 in yGCN5³³, E122 in tGCN5³⁸ and E80 in hGCN5⁵⁵). In particular, for tGCN5, a conserved water molecule is ideally located to shuttle a proton from

the reactive lysine to E122, representing a long-range transfer scheme. A similar mechanism may be employed in p300. Analysis of the p300/Lys-CoA crystal structure (PDB code: 3BIY) indicates that the nearest electronegative residues that could be the proton acceptors around the reaction center are D1444 and D1507. Three crystal water molecules, Wat1777, Wat1861 and Wat1878, and two residues with a hydroxyl group, Y1394 and Y1446, may also serve as the intermediate proton carriers. Based on this comparison and analysis, we proposed two possible PT modes showed in scheme 1.

MD simulations were incompatible with scheme (b) since the positions of Wat1777 and Wat1878 were not stable enough to facilitate PT. In contrast, Wat1861 in scheme (a) was quite stable during the entire 100 ns simulation time. The distances between the heavy atoms involved in scheme (a) were all steady and close enough for the PT (Figure 3A). Detailed MD simulation analysis suggested that the high stability of scheme (a) is induced by a strong h-bond interaction between Wat1861 and the backbone carbonyl oxygen of 11395, which retains the location of the water molecule. In addition, the strong h-bond interaction between H3K14 and W1436 may help align the departing proton toward the water molecule, which is also apparently facilitated by the hydrophobic interactions between the same two residues (Figure 3B). Moreover, a hydrophobic local environment contributes to increase the pKa of the D1507 side chain, allowing it to function as a general base. An analogous favorable environment was not present in scheme (b), which instead showed remarkable fluctuation. The importance of residues Y1394 and D1507 in the PT mechanism of scheme a is supported by mutational studies. Specifically, Y1394F, D1507N and D1507A mutations significantly impair p300 acetylation activity³⁷. In summary, due to the instability of the two water molecules in scheme (b), and the significantly more favorable MD simulation results with scheme (a) along with the supportive mutational studies, we propose that scheme (a) is used for PT lysine deprotonation for the first step of the p300 acetylation reaction.

QM/MM calculations were employed to validate PT scheme (a) and further investigate the subsequent chemical mechanism. The QM region included Wat1861, the side chain of H3K14, Y1394 and D1507, the backbone of I1395 and W1436, where I1395 h-bonds with Wat1861 to keep its position, W1436 h-bonds with H3K14 to orient the -NH3+ group, and W1436 contributes to the hydrophobic environment that facilitates lysine deprotonation as mentioned above. When systematically lengthening (at 0.05 Å intervals) the N-H bond of the target lysine toward Wat1861 with each step subjected to a geometry optimization, the protons in Wat1861 and the hydroxyl group of Y1394 migrated in a concerted manner. As a consequence, the hydroxyl hydrogen was bound to D1507, while the other two hydrogen atoms were abstracted by oxygen atoms, On and O, of Y1394 and water, respectively, generating a protonated D1507, a neutral Y1394, and a deprotonated H3K14. A potential energy profile captured the concerted nature of the PT reaction (Figure 4): the proton movement has a barrier of 0.65 kcal/mol, while the product (P) is 10.15 kcal/mol lower in energy than the reactant (R). This energy profile displays a notably low PT barrier and a dramatic energy decrease in the immediate products, which leads us to conclude that such a single-direction proton shuttle is energetically inescapable, or even intrinsic.

The PT schemes in three major HAT subfamilies (Gcn5/PCAF, MYST and p300) were compared by structure alignment. Interestingly, the PT processes in all three HAT

subfamilies are common in transfer direction, but differ in molecular transfer scheme (Figure S7). The locations of E338 of Esa1, E122 of tGCN5 and Y1394 of p300 mapped very well. However, the conserved proton acceptor residues among these enzymes belong to different secondary structure elements. E338 is located on the end of the β 10 strand of the C-terminal region of Esa1, and E122 in tGCN5 is located on the β 4 strand of the core region. For p300, D1507 is found on the short loop before the α 6 helix of the N-terminal region. It is also notable that the complexity of the three schemes is different. In Esa1, the proton is directly delivered to E338. In tGCN5, the proton is transferred to E122 by a water molecule. While in p300, the proton is shuttled to Y1394 on the β 4 strand by a water molecule, and then the hydrogen atom from the hydroxyl group of Y1394 is delivered to D1507. In summary, the PT process in Esa1 is direct from substrate to enzyme, tGCN5 uses a water-mediated pathway, and p300 employs the most complex scheme, in which both a water molecule and a shuttle residue Y1394 are involved.

The catalytic mechanism of the acetylation reaction

The acetylation reaction can be divided into two steps: an addition half-reaction and an elimination half-reaction. In the addition half-reaction, nitrogen atom (N3) of H3K14 nucleophilically attacks the carbonyl carbon atom (C4) of Ac-CoA, promoting this carbon atom to change hybridization state from sp2 to sp3. Subsequently, the N3-C4 bond forms and the hydrogen (H2) of H3K14 is transferred to the oxygen atom (O5) of the acetyl portion of Ac-CoA, forming an intermediate. In the elimination half-reaction, normally it is the hydroxyl group of the intermediate that protonates CoAS- to generate the CoASH product, as found for N-acetyltransferases^{58, 59}. However, the side chain hydroxyl group of Y1467 can also function as the general base to protonate the CoAS- since it forms a direct hydrogen bond to the sulfur atom of Ac-CoA (Scheme 2). The two steps and the two possibilities of the elimination half-reaction were investigated by QM/MM simulation. The reactant system for this simulation of the acetylation reaction was constructed based on the optimization of the product structure of the PT process.

The QM region included the side chain of the deprotonated H3K14, the -CH₂-S-COCH₃ group of Ac-CoA, the backbone of W1436, and the side chain of Y1467, considering their important roles during the reaction process. W1436 strongly h-bonds with the ε-nitrogen of the target lysine, which may contribute to orient the lysine amino group. Moreover, the aromatic side chain of W1436 together with the phenol group of Y1446 forms a hydrophobic environment that facilitating lysine deprotonation to trigger the PT process and acetylation reaction. Notably, this h-bond interaction may activate the reaction by making the reactive nitrogen atom partially negative and by stabilizing the intermediate. Indeed, this activation and stabilization mechanism is widespread in HAT proteins. For tGCN5, the backbone amide group of L126 forms a hydrogen bond with the carbonyl oxygen of the acetyl group of Ac-CoA so that the thioester carbonyl group can be polarized to achieve these activation and stabilization functions⁶⁰. Y1467 coordinates with the sulfur atom of Ac-CoA, which may help position the carbonyl carbon. More importantly, Y1467 may also help activate the chemical bond between the sulfur atom and the carbonyl carbon. Finally, after the nucleophilic attack, Y1467 may also function as a general acid to protonate the CoASH leaving group as shown in scheme 2 (a).

Along the nucleophilic attack reaction, the energies of the reactant (R), transition state (TS), and intermediate (IntMed) were determined by a two-dimensional QM/MM potential energy surface by defining the distances of R(N3-C4) and R(N3-H2) as the reaction coordinates (Figure 5). In the optimized reactant, R(N3-C4)=3.32 Å and R(N3-H2)=1.03 Å. In the optimized intermediate structure, the N-C bond is formed (R(N3-C4=1.43 Å)), the N3-H2 bond is broken, and C4=O5 of Ac-CoA changes to C4-O5H2. The calculated potential energy barrier of this step is 20.14 kcal/mol. The structure of the TS1 was determined by adiabatic mapping at the QM/MM level. In the TS, R(N3-C4)=1.60 Å and R(N3-H2)=1.30 Å. This structure illustrates that atom N3, H2, C4 and O5 were tuned to be in a plane to form a tetrahedron ring. The N3-H2 bond was partially broken, the N3-C4 bond was partially formed, and the C4-S6 bond was also partially broken simultaneously. Furthermore, the bond order of C4=O5 was transformed from double to single gradually during the whole process. Consistently, we also noticed that the two h-bond pairs (H3K14-W1436 and Ac-CoA-Y1467) mentioned above remained stable. This two h-bonds contributes to stabilizing the reactive components so that a energetically favored IntMed could properly formed. The distance between H3K14 and W1436 became even closer. This observation confirms the role of W1436 and Y1467 in stabilizing the IntMed.

As aforementioned, there are two possible mechanisms to protonate the leaving group after the intermediate is formed. To determine which scheme is more reasonable, we scanned the coordinates of S6-H2 and S6-H7 bond formation in both schemes (Figure 6). The energy barrier of scheme (a) is 12.08 kcal/mol, while for scheme (b) the energy barrier is 15.75 kcal/mol. Although the energy difference between these two schemes is not large, the lower energy barrier of scheme (a) indicates that the Y1467-meidated elimination half-reaction is more likely to occur. In addition, Y1467F showed a 430-fold reduction in V/K^{3737} also suggesting an important role of this residue in the reaction. Based on all these analyses, we consider scheme (a), in which Y1467 serves as the major general acid to protonate the leaving group, as the preferred reaction scheme. In TS2(a), the sulfur atom, the carbon atom of the acetyl group, the hydroxyl group of the IntMed, and the phenolic hydroxyl group of Y1467 forms a hexahedron ring together with two hydrogen bonds. In addition, the hydrogen bond between W1436 and the deprotonated lysine amino nitrogen is probably even stronger as the distance (1.77Å) between them is shorter than in the intermediate (1.83Å). Since a Y1467F mutant displays significantly reduced enzyme activity, it is mostly likely that, the acetylation reaction adopts scheme (a) to proceed. Altogether, the elimination half-reaction is mainly triggered by Y1467 and may be conducted directly by the IntMed when Y1467 is mutated.

Conclusion

As the founding member of one of the major HAT subfamilies, p300 plays important roles in a plethora of biological processes, including cell growth, differentiation and transformation, oncogenesis and apoptosis. The catalytic mechanism of the p300 HAT has started to unfold through recent structural and biochemical studies. However, a complete delineation of the entire chemical pathway of the acetylation reaction remained unclear. Knowledge was lacking for some of the most critical components of the pathway, including

the structure of the PT wire and the chemical mechanism of the acetyl transfer reaction. No study had been reported to date to address these important issues.

In this study, molecular modeling was applied to construct the p300/H3/Ac-CoA system. A 100ns MD simulation was carried out on this model to obtain the most suitable configurations for further study. The h-bond interactions between p300 and the H3 peptide were mainly formed by the electronegative residues E1505, D1625, D1628 of the substrate binding cleft and E1442, K1456, I1457, K1459 and R1462 of the L1 loop. In addition to residues G12 to P16 of the H3 peptide, which include a conserved G-K-A-P sequence in HAT substrates, hydrophobic interactions are mediated by non-polar part of Y1397, W1436, P1439, P1440 and Y1446 of the L1 loop. Structure analysis reveals two possible "protonwires" for the deprotonation process of substrate H3K14. Consistent with previous mutation data, our MD simulation indicates that the proton wire consisting of H3K14, Wat1861, Y1394 and D1507 is highly likely. Detailed analysis suggests that a strong h-bond interaction between Wat1861 and the backbone carbonyl oxygen of I1395 might be the major stabilization factor for this wire by fixing the location of the water molecule. The extremely low potential energy in the QM/MM calculation provides additional support for this PT wire. Finally, QM/MM calculations were also performed to investigate the acetylation mechanism. The two-dimensional potential energy barrier of the addition halfreaction and structures of two TSs and IntMed were obtained. In particular, for the elimination half-reaction both potential energy barrier calculations and previous experimental data support the model whereby the protonation of the CoAS- group by Y1467 is more energetically favorable than protonation by the reaction IntMed.

In summary, our research elucidated several essential aspects of the acetylation reaction catalyzed by p300. The rational PT scheme was proposed and validated for the first time. Detailed knowledge of the configurations of the two TSs and IntMed do not only shed light on the mechanism research of other HAT families, but also opens up a novel avenue to discover a new family of small molecule effectors for p300. These small molecules can be designed specifically focused on specifically blocking the PT wire, interfering with the addition half-reaction or affecting the protonation of the leaving group on the basis of the TS structures.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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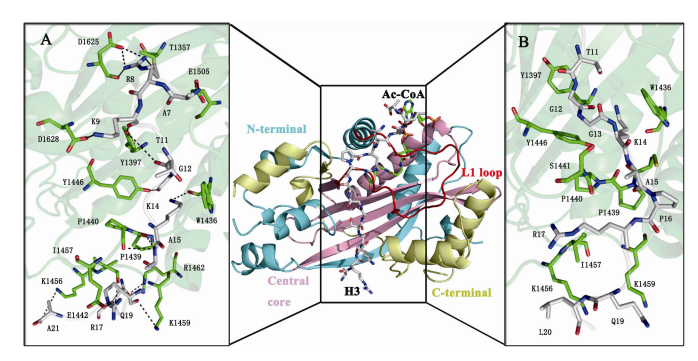


Figure 1. The overall structure of the p300/H3/Ac-CoA complex model. Cyan: N-terminal region; Magnet: central core region; Yellow: C-terminal region; Red: L1 loop. (A) H-bond interactions and (B) hydrophobic interactions between H3 and p300. Green sticks: p300; White sticks: H3 peptide.

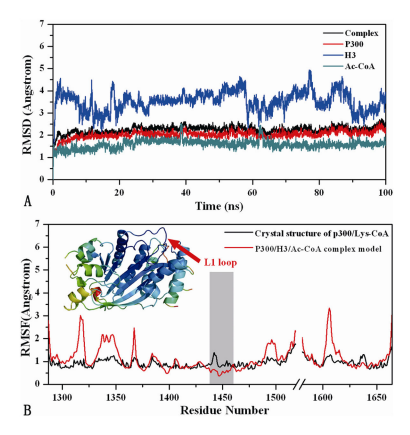


Figure 2.

(A) The time dependencies of the root-mean-square deviations (RMSDs) for the back bone atoms of the p300/H3/Ac-CoA complex, p300, H3 and Ac-CoA from their initial positions during the 100 ns simulation. (B) Residue fluctuations obtained by average residual fluctuations over 100 ns simulation and experimental result calculated from B factors of p300 (PDB code: 3BIY). The residue number of genetically deleted autoacetylation loop was represented by a gap. The color of the cartoon structure ranges from blue to red with increasing RMSF values.

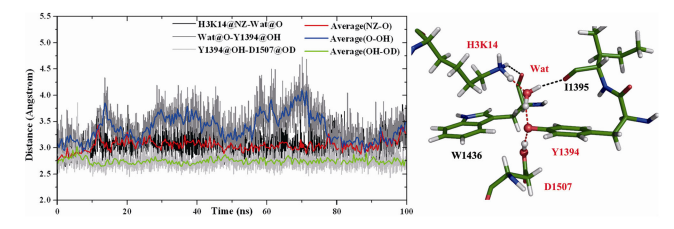


Figure 3.(A) the time dependencies of the distances between heavy atoms composing the PT wire of scheme (a) during the 100 ns simulation. (B) The interaction network that stable the configuration in scheme (a). The atoms directly involved in the transfer wire are shown as spheres.

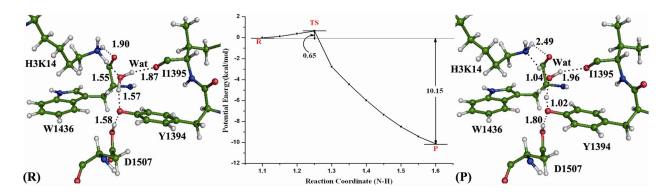


Figure 4.The potential energy barrier for the PT process. The structures of reactant (R) and product (P) are displayed.

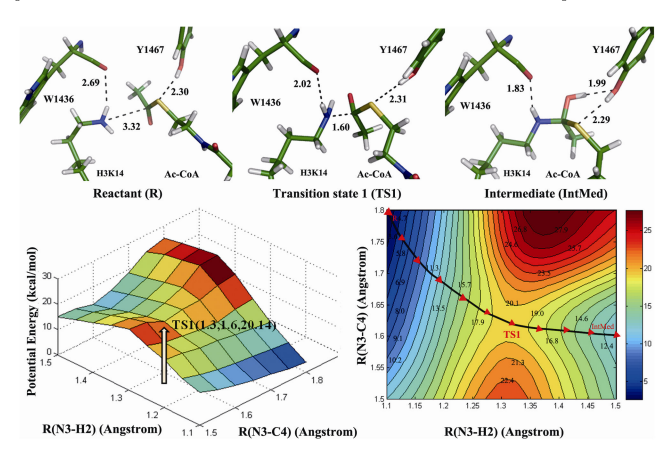


Figure 5.

The potential energy surface (PES) and contour plot of the PES of the addition half-reaction pathway. Energy barriers of transition state 1 (TS1) is marked and structures of reactant (R), intermediate (IntMed) and product (P) are displayed.

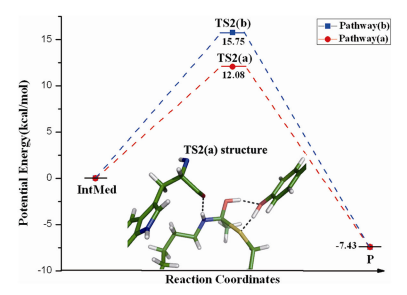


Figure 6.The potential energy barrier comparison of the two elimination half-reaction mechanisms. The structure of TS2(a) which is more favorable is displayed.

Scheme 1. Two proposed PT schemes.

Scheme 2.
Two possible acetylation schemes