Histone modifications in *Trypanosoma brucei*

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Abstract

Several biological processes in *Trypanosoma brucei* are affected by chromatin structure, including gene expression, cell cycle regulation, and life-cycle stage differentiation. In *Saccharomyces cerevisiae* and other organisms, chromatin structure is dependent upon posttranslational modifications of histones, which have been mapped in detail. The tails of the four core histones of *T. brucei* are highly diverged from those of mammals and yeasts, so sites of potential modification cannot be reliably inferred, and no cross-species antibodies are available to map the modifications. We therefore undertook an extensive survey to identify posttranslational modifications by Edman degradation and mass spectrometry. Edman analysis showed that the N-terminal alanine of H2A, H2B, and H4 could be monomethylated. We found that the histone H4 N-terminus is heavily modified, while, in contrast to other organisms, the histone H2A and H2B N-termini have relatively few modifications. Histone H3 appears to have a number of modifications at the N-terminus, but we were unable to assign many of these to a specific amino acid. Therefore, we focused our efforts on uncovering modification states of H4. We discuss the potential relevance of these modifications.

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1. Introduction

The basic unit of chromatin is the nucleosome, formed by 146 bp of DNA wrapped around an octamer containing two copies of each of the core histones, H2A, H2B, H3, and H4. The core histones contain many posttranslational modifications (PTMs), including phosphorylation, acetylation, methylation, and ubiquitination. These PTMs affect chromatin structure, which has been shown to have consequences for nucleosome assembly, DNA replication and repair, and, most notably, gene expression [1]. Histone acetylation, for example, is generally associated with transcriptionally permissive chromatin, or euchromatin [2]. N-terminal acetylation is thought to disrupt interactions that stabilize the nucleosome, making the DNA more accessible to transcription-promoting complexes [3]. Furthermore, transcription factors recruit histone acetyltransferases to the promoter region of target genes, and, accordingly, histone deacetylase activity has been detected in transcriptional co-repressor complexes [4,5]. Histone methylation also affects transcriptional activity of chromatin, but these effects are dependent upon the particular lysine that is methylated: H3 K9 and H4 K20 methylation are associated with silent chromatin, or heterochromatin [6]. In *Saccharomyces cerevisiae* and other organisms, histone modifications clearly play a role in determining whether chromatin is in a transcriptionally permissive or restrictive configuration.
Trypanosoma cruzi and its relatives are the causative agents of human sleeping sickness and widespread animal diseases in Africa and, to a lesser extent, other continents. This parasite evades the host immune response by spontaneously switching its variant surface glycoprotein (VSG), in a process known as antigenic variation. It has been shown that chromatin structure at the subtelomeric VSG expression site (ES) may influence its transcriptional activity. The actively transcribed VSG demonstrates greater sensitivity to digestion with DNase I and single-strand-specific endonucleases than untranscribed VSGs, which suggested a more open chromatin conformation at the active ES [7,8]. Chromatin structure was further implicated in VSG silencing when it was shown that transcription was repressed from ES and rRNA promoters that were inserted in the silent ES, and that this effect was more pronounced as the promoters were placed closer to the telomere [9,10]. These studies suggest that VSG silencing is due to a repressive chromatin conformation.

Apart from VSG expression, several biological processes in trypanosomes involve histone PTMs. H3 K76 di- and tri-methylation and the responsible histone methyltransferases, DOT1A and DOT1B, have been studied in detail in T. brucei [11]. Deletion of trypanosome DOT1 genes caused defects in cell cycle regulation and differentiation between life-cycle stages. Also, it was shown that overexpression of a histone deacetylase, T. brucei SIR2 related protein 1, resulted in increased sensitivity to DNA damaging agents [12].

Identifying histone PTMs in trypanosomes is an important first step towards studying the biological processes that are influenced by chromatin structure. Although histones are highly conserved in most organisms, the amino acid sequences of T. brucei histones are very different (Fig. 1), and we cannot infer which modifications will be present, based on sequence alignments. Commercial antibodies that recognize modifications in other organisms cannot be expected to specifically cross-react with trypanosome PTMs, and the most likely ones do not (unpublished data).

Previous studies in Trypanosoma cruzi and procyclic form (PF) T. brucei, the insect (tsetse) midgut life-cycle stage, have identified some histone PTMs [13,14]. We add to this current knowledge by performing a detailed survey, using Edman degradation and mass spectrometry (MS), to identify as many as possible of the PTMs present on histones H2A, H2B, H3, and H4, which we purified from the bloodstream form (BF) of T. brucei strain Lister 427. Edman analysis provided us with quantitative data of PTMs found at the N-termini of H2A, H2B, and H4, demonstrating that H2A and H2B had relatively few modifications, while the H4 N-terminus possessed numerous modifications. Edman analysis also showed that H2A, H2B, and H4 could be monomethylated at A1, which is a histone modification that, to our knowledge, has only been found in T. brucei and T. cruzi [13,14]. Matrix-assisted laser-desorption/ionization time of flight mass spectrometry (MALDI-TOF MS) showed that the H3 and H4 N-terminal peptides are present in multiple modification states. The H4 N-terminus was studied in greater detail by tandem MS. We conclude by comparing our findings to S. cerevisiae and other organisms.

2. Materials and methods

2.1. Histone purification

BF T. brucei cells of the Lister strain 427 clone 221a were cultured in HMI-9 [15]. Approximately 3 × 10⁸ cells were used to infect CD-1 female Swiss white mice. After 3 days, trypanosomes from the infected mice were used to inoculate Sprague–Dawley male rats (1–2 × 10⁷ cells/rat). After a 3-day infection, trypanosomes were collected after exsanguination of the rats as previously described [16]. For purification of histones, 1 × 10¹⁰ cells were centrifuged at 1800 × g for 10 min at 4°C. Cells were resuspended in 20 ml 8% polyvinylpyrrolidone (containing 0.05% octyl glucoside) and lysed with a Polytron as described elsewhere [17]. Lysed cells were centrifuged at 16,000 × g for 5 min at 4°C. The pellet was washed with 10 ml low salt buffer (10 mM Tris, pH 8.0; 75 mM NaCl; 0.05% octyl glucoside) and resuspended in 10 ml high salt buffer (10 mM Tris, pH 8.0; 2.0 M NaCl, 1% octyl glucoside). Octyl glucoside can be replaced by NP-40 for a better yield of histones, but NP-40 detergent contamination interfered with MS analysis. Nuclei were lysed with a Dounce homogenizer and centrifuged at 16,000 × g for 20 min at 4°C in an SS-34 rotor (Sorvall). Histones from the supernatant were extracted in 0.4 M H₂SO₄ overnight at 4°C. All solutions were treated with 1 mM DTT, 50 mM sodium butyrate, 0.5 mg/ml TLCK, 0.174 mg/ml PMSF, and a protease inhibitor cocktail (Sigma).

The main contaminant in this histone preparation was VSG, the most abundant protein in BF T. brucei. VSG was removed from the sample by passing it through a Concanavalin A (ConA) column. The sample was first dialyzed overnight at 4°C in ConA equilibration buffer (20 mM Tris, pH 7.4; 0.5 M NaCl) and applied to a ConA column (Sigma). The flow-through was collected and lyophilized. The sample was fractionated by RP-HPLC on a C8 column using an acetonitrile gradient (0–60%) in 0.1% trifluoroacetic acid. Fractions were collected every minute for 70 min, lyophilized, then resuspended in 50 μl H₂O. Fractions were analyzed by SDS-PAGE and MALDI-TOF.

2.2. Analysis of histones by Edman degradation and mass spectrometry

Edman degradation was performed using a Procise 494-HT with a PTH C18 HPLC column (Applied Biosystems). Modified PTH standards (acetyl or methyl lysines) were made by reacting free amino acids (Sigma) with PITC to create PTC-amino acids. These were then converted to PTH-amino acids by incubation with 25% trifluoroacetic acid. Histones were digested with endoproteinases Glu-C, trypsin, and Asp-N, according to manufacturer’s instructions (Roche). To prevent trypsin cleavage at lysine, some histone preparations were reacted with 1% propionic anhydride (Sigma) in acetonitrile:50 mM NH₄HCO₃ (1:1 v/v) [18]. After propionylation, the products were dried and redissolved in water.

Digested histones were analyzed by a MALDI-TOF MS or ESI-LC–MS/MS. MALDI-TOF was conducted using a DE-STR mass spectrometer operating in delayed and reflector
Fig. 1. Representative sequence alignments of the N-termini of each core histone and the C-terminus of H2A. Identical residues are shaded. Important sites of homology, which are referred to in the text, are indicated by arrows. The sequence ruler, which is included only when the sequences are very concordant, is numbered according to the Homo sapiens sequence position. Residues found to be acetylated in T. brucei histones are indicated with a blue * and methylated residues with a red *. A putative site of ubiquitination at the H2A C-terminus is indicated with a green *.

(A) H2A N-terminus alignment

(C. elegans H2A) SGRKGGKAKTGGKAKSRSRRAGLQFPFPVR
(D. melanogaster H2A) SGRKGGKAKTGGKAKSRSRRAGLQFPFPVR
(H. sapiens H2A) SGRKQGKAKTGGKAKSRSRRAGLQFPFPVR
(M. musculus H2A) SGRKQGKAKTGGKAKSRSRRAGLQFPFPVR
(S. cerevisiae H2A) SGRKGGKAKTGGKAKSRSRRAGLQFPFPVR
(T. brucei H2A) ATPKQAVKASRSRRAGLQFPFPVR

(B) H2A C-terminus alignment

(C. elegans H2A) GVTIAQGGVLPIQAVLPPKRTGKDKE
(D. melanogaster H2A) GVTIAQGGVLPIQAVLPPKRTGKDKE
(H. sapiens H2A) GVTIAQGGVLPIQAVLPPKRTGKDKE
(M. musculus H2A) GVTIAQGGVLPIQAVLPPKRTGKDKE
(S. cerevisiae H2A) GVTIAQGGVLPIQAVLPPKRTGKDKE
(T. brucei H2A) VTMRSRGCVMSPNLKAKTQKSCHAKATPSV

(C) H2B N-terminus alignment

(C. elegans H2B) PKKPSAKKAKAKTVKTQPDKGKK
(D. melanogaster H2B) PKKPSAKKAKAKTVKTQPDKGKK
(H. sapiens H2B) PKKPSAKKAKAKTVKTQPDKGKK
(M. musculus H2B) PKKPSAKKAKAKTVKTQPDKGKK
(S. cerevisiae H2B) PKKPSAKKAKAKTVKTQPDKGKK
(T. brucei H2B) AIPKSTPAKTRKAAKTRKRTWNVYVS

(D) H3 N-terminus alignment

(C. elegans H3) ARTKQTARKSTGGKAPRQKALTKAARKSAKASAG
(D. melanogaster H3) ARTKQTARKSTGGKAPRQKALTKAARKSAKASAG
(H. sapiens H3) ARTKQTARKSTGGKAPRQKALTKAARKSAKASAG
(M. musculus H3) ARTKQTARKSTGGKAPRQKALTKAARKSAKASAG
(S. cerevisiae H3) ARTKQTARKSTGGKAPRQKALTKAARKSAKASAG
(T. brucei H3) ARTKQTARKSTGGKAPRQKALTKAARKSAKASAG

(E) H4 N-terminus alignment

(C. elegans H4) SGRKKGGKLGRKGRKAAKRRHKVRKLRDNQIGITKPAI
(D. melanogaster H4) TGRKKGGKLGRKGRKAAKRRHKVRKLRDNQIGITKPAI
(H. sapiens H4) SGRKKGGKLGRKGRKAAKRRHKVRKLRDNQIGITKPAI
(M. musculus H4) SGRKKGGKLGRKGRKAAKRRHKVRKLRDNQIGITKPAI
(S. cerevisiae H4) SGRKKGGKLGRKGRKAAKRRHKVRKLRDNQIGITKPAI
(T. brucei H4) AKGKSKQGEAKGSKQKRRKVRVLRENGIGITRGSI

Fig. 1. Representative sequence alignments of the N-termini of each core histone and the C-terminus of H2A. Identical residues are shaded. Important sites of homology, which are referred to in the text, are indicated by arrows. The sequence ruler, which is included only when the sequences are very concordant, is numbered according to the Homo sapiens sequence position. Residues found to be acetylated in T. brucei histones are indicated with a blue * and methylated residues with a red *. A putative site of ubiquitination at the H2A C-terminus is indicated with a green *. (A) H2A N-terminus. (B) H2A C-terminus. (C) H2B N-terminus. (D) H3 N-terminus. (E) H4 N-terminus.
modes (Applied Biosystems). For ESI-LC–MS/MS, digests were injected on a 0.3 mm internal diameter × 65 mm long trapping column (PepMap; Dionex) at a flow rate of 20 μl/min (total loading time of 5 min). By switching the stream valve in the SWITCHOS (Dionex), the trapping column is back-flushed with a binary solvent gradient, which is started simultaneously with the injection cycle. The sample is thereby loaded onto a nanoscale RP C18 column (0.75 mm internal diameter by 150 mm long PepMapT column; LC Packings). Peptides were eluted from the stationary phase using a linear gradient from 0 to 75% solvent B (100% ACN in 0.1% formic acid) applied over a period of 45 min. The solvent delivery system was set at a constant flow of 20 μl/min and using a 1/10000 flow splitter, 200 nl/min of solvent was directed through the nanocolumn. The outlet of the nanocolumn was connected in-line with a distal metal-coated fused silica PicoTipT needle (PicoTipT FS360-20-10-D-C7, New Objective, Woburn, MA, USA) that is interfaced with a QSTAR XL QqTOF mass spectrometer (Applied Biosystems).

3. Results

3.1. Histone purification

We first developed a method for purifying histones from BF T. brucei. Histones were previously purified from PF by isolating nuclei, then using acid extraction and RP–HPLC purification [14,17]. This method was attempted in BF trypanosomes, but the yield was extremely low (<1%). The greatest loss of material occurred during nuclei isolation, which we attribute to the increased fragility of the BF nucleus. Consequently, the protocol we developed (see Section 2) did not involve the purification of nuclei as an intermediate step. Analysis of HPLC fractions by Coomassie staining of polyacrylamide gels and by MALDI-TOF MS demonstrated the high purity of the final histone preparations. When NP-40 was used in the protocol, our final yield of histones was approximately 30%. These samples were analyzed by Edman degradation, but NP-40 contamination of the purified histones prevented us from acquiring MS data. Substituting octyl glucoside for NP-40 prevented detergent contamination of the final samples, but resulted in a lower yield (∼10%).

3.2. Edman degradation and MS analysis

A combination of MS techniques, including MALDI-TOF and MS/MS, and Edman degradation, were used to analyze T. brucei histones for PTMs. Edman degradation provided quantitative data for PTMs on the N-terminal 20–30 amino acids. For MS, histones were first digested with trypsin. Histones contain many arginine and lysine residues, resulting in excessive fragmentation by trypsin. In some experiments, therefore, histones were modified with propionic anhydride to prevent trypsin cleavage at lysine residues by propionylation (+56 Da) of unmodified (K) and monomethylated (Kme1) lysines, and the N-terminus. Dimethylated (Kme2), trimethylated (Kme3) and acetylated (Kac) lysines were not propionylated, but are also not cleaved by trypsin. Histone H3 was also digested with endoproteasines Asp-N and Glu-C and H4 was digested with Glu-C.

MALDI-TOF analysis provided the molecular mass of the digested peptides. Mass shifts of 14, 28, and 42 Da may represent peptides with a monomethyl, dimethyl, and trimethyl or acetyl group, respectively. The peptides were then sequenced by tandem MS to assign PTMs to their parent residue. Acetyl (42.0105 Da) and trimethyl (42.0468 Da) groups are difficult to distinguish by tandem MS. We relied on data generated by Edman degradation to help us assign mass shifts of 42 Da. Acetylated lysines also produce a characteristic immonium ion at m/z 126, and trimethylated lysines have a characteristic ion at MH+ -59, both of which were used to make assignments [19].

3.3. H2A modifications

Histones H2A and H3 co-eluted on the HPLC. We did not attempt to separate them because the presence of both histones in a single sample did not interfere with MS or Edman analysis. Only the H2A N-terminus was susceptible to Edman sequencing, because the H3 N-terminal Serine (S1) was blocked, as previously shown in PF [14]. Analysis of the first 22 amino acids of H2A demonstrated that 60% of A1 is monomethylated, and ∼1% of K4 is acetylated (Table 3).

Tandem MS data were acquired for trypsin-digested H2A (Fig. 2A). Peptides that were analyzed are shown in red. Tryptsin digestion produced several spectra representing modified C-terminal peptides, demonstrating that K115, K119, K120, K122, K125, and K128 can be acetylated (Table 1). We assigned these modifications as acetylation rather than trimethylation, because studies of PF histones showed that H2A exists in multiple acetylation states [14]. The presence of the immonium ion for acetylated lysine at m/z 126 supported this assignment. The MH+ -59 ion was not present on these spectra. Multiple modifications of the H2A C-terminus have not been reported in other organisms.

Next, H2A was modified with propionic anhydride prior to trypsin digest (hereafter referred to as PA-trypsin digest) to acquire information about the N-terminus (Fig. 2B). MALDI-TOF analysis of the PA-trypsin-digested H2A demonstrated that the N-terminus is present in three modified states (Fig. 3A). Tandem MS sequencing of these species demonstrated that A1 could be monomethylated (Fig. 3B), acetylated, or unmodified (Table 1). A spectrum representing peptide 107–133 confirmed acetylation of K115, K119, and K120 (Table 1).

In other organisms, the C-termini of H2A and H2B are ubiquitinated [20,21]. Alignment of the H2A C-terminus (Fig. 1) suggests that T. brucei H2A K122 might be the homologue of the ubiquitinated H2A K119 in other organisms. We therefore performed a directed search for ubiquitinated H2A K122. MS can identify sites of ubiquitination by looking for a GG dipeptide (114 Da) to lysine, which results from tryptic cleavage of covalently linked ubiquitin [22]. The tandem MS spectrum of a peptide with m/z 1029.26+ displayed partial y-series (y1–y6, y8, y15) and b-series (b3, b12, b22–24) ions, demonstrating that the spectrum represents propionylated H2A peptide 107–133 (data not shown). The molecular weight of this species and the y- and
b-ions indicate that K115 and K128 are propionylated, and of the remaining four lysines (K119, K120, K122, and K125), one may be ubiquitinated and three are acetylated.

3.4. H2B modifications

Edman degradation of the first 25 amino acids of H2B demonstrated that 60% of A1 is monomethylated, and ~1% of K4 is acetylated (Table 3). Both H2A and H2B begin with the sequence ATPK, and both are modified at A1 and K4 to the same degree. PA-trypsin digestion of H2B provided MS/MS data for the lysine-rich N-terminus (Fig. 2C). Three spectra representing peptide 1–11 showed that A1 can be unmodified, mono- or di-methylated (Table 1). One spectrum was obtained representing two species of peptide 12–18, which showed that K12 or K16 is acetylated (Fig. 4). These PTMs were designated as acetylated lysines because the Kac immonium ion was present at m/z 126. These marks are presumably minor, because they were not observed by Edman analysis.

3.5. H3 modifications

Multiple attempts were made to acquire information on the modification state of H3. Many of the peptides produced by the PA-trypsin digest of H3 were sequenced by tandem MS (Fig. 2D). A spectrum corresponding to the N-terminus, peptide 1–8, showed that S1 can be acetylated (Table 1). Four spectra representing peptide 70–80 showed that K76 can be unmodified, mono-, di- or tri-methylated (Table 1).

Quadrupole–quadrupole-TOF (QqTOF) analysis of the H3 PA-trypsin digest showed multiple modification-states for the lysine-rich peptide 9–36 (Fig. 5A). Sequencing by tandem MS

<table>
<thead>
<tr>
<th>Histone Digest</th>
<th>Modified peptide</th>
</tr>
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<tbody>
<tr>
<td>H2A trypsin</td>
<td>GGVMPSLN(^{115})K(_{ac}) ALAK (107–119)</td>
</tr>
<tr>
<td>H2B PA-trypsin</td>
<td>(^{1})K(_{ac}) EAKKTR (12–18)</td>
</tr>
<tr>
<td>H3 Trypsin</td>
<td>AS(^{23})K(_{ac}) GSDAASGVK (21–32)</td>
</tr>
</tbody>
</table>

Histones, which in some cases were modified with propionic anhydride, were digested with trypsin. Tryptic peptides were sequenced by tandem MS, and peptides that have covalent modifications are listed here. K\(_{ac}\) refers to acetyl lysine; K\(_{me1}\), K\(_{me2}\), and K\(_{me3}\) refer to mono-methyl, di-methyl, and tri-methyl lysine, respectively; A\(_{ac}\), A\(_{me1}\), and A\(_{me2}\) refer to acetyl, mono-methyl, and di-methyl alanine, respectively; S\(_{ac}\) refers to acetyl serine.
Fig. 3. Histone H2A alanine 1 can be unmodified, acetylated, or monomethylated. H2A was propionylated and digested with trypsin. Peptides 1–17 (ATP-KQA VKKASKGGSSR) was subjected to MALDI-TOF analysis and sequencing by tandem MS. (A) MALDI-TOF shows that peptides 1–17 is present in three modified states. M is the molecular mass ion of peptides 1–17 with all lysines propionylated (1924 Da). Tandem MS sequencing of peaks labeled M + 42 Da, M + 56 Da, and M + 70 Da species indicate that H2A A1 can be acetylated, propionylated (meaning it is unmodified), or both propionylated and monomethylated, respectively. Multiple peaks representing each modified peptide show the isotope pattern, with the first peak representing the monoisotopic m/z. (B) A representative tandem MS showing that H2A A1 is monomethylated. All of the b-series ions are shifted +70 Da relative to the unmodified species. Also, the difference between the MH+ of the spectrum (m/z 1995) and y16 provides the mass of the modified A1. y152+ and y162+ represent doubly charged ions [M+2H]/2+.

showed that all eight lysines are propionylated in the species represented as M + 448 Da. Tandem MS of the three remaining peaks showed that they represent modified species of peptide 9–36, but modifications could not be assigned to specific lysines. The presence of these peaks suggests that several lysines at the H3 N-terminus can be modified.

Although the N-terminus of H3 was blocked to analysis by Edman degradation, we attempted to acquire Edman degradation data by separating the peptides from the PA-trypsin digest by HPLC and sequencing peptide 9–36. A weak profile was obtained, but no PTMs were detectable.

Tandem MS sequencing of trypsin-digested H3 uncovered two modifications at the N-terminus—K23ac and K32me3 (Table 1). K23 was shown to be acetylated by the presence of the immonium ion at m/z 126. K32 was shown to be methylate by the presence of the MH+-59 ion. Finally, we attempted MS analysis of Asp-N- and Glu-C-digested H3. No interpretable tandem MS data were acquired, although Qq-TOF analysis of Asp-N-digested H3 confirmed that multiple modification-states exist for N-terminal peptide 1–25 (Fig. 5B). We concluded our study of H3 with the knowledge that PTMs were present at the N-terminus that could not be assigned to specific amino acids.

3.6. H4 modifications

Edman degradation of the first 22 amino acids showed that, unlike H2A and H2B, the H4 N-terminus has numerous PTMs (Table 3). K2, K5, K10, and K14, are acetylated in less than 10% of H4. K4ac is an abundant mark that is found in 73% of H4. Both K17 and K18 can be mono-, di- or tri-methylated and 60% of A1 is monomethylated.
Fig. 4. Histone H2B lysines 12 and 16 are acetylated. H2B was propionylated and digested by trypsin. Peptides 12–18 (KEAKKTR) was sequenced by tandem MS. A single MS spectrum contains two modified species, demonstrating that either K12 (b2′′–b3′′ and y3′′–y6′′) or K16 (b1′–b3′ and y3′–y6′) are acetylated. An arrow points to the ammonium ion of acetylated lysine (m/z 126). b72+ and y72+ represent doubly charged ions.

We examined H4 in more detail by MS, to confirm the PTMs observed by Edman as well as to determine whether less abundant PTMs were present. H4 Glu-C, trypsin- and PA-trypsin-digested peptides were fractionated by HPLC, and individual fractions were analysed by MALDI-TOF. Peptides were then sequenced by tandem MS.

PA-trypsin-digested H4 had extensive sequence coverage by tandem MS (Fig. 2E). The data did not suggest the presence of additional PTMs in the core domain or C-terminus of H4. MALDI-TOF demonstrated multiple modification-states for peptides 1–15 and 16–21 (Table 2 and data not shown). Sequencing of peptide 1–15 by tandem MS revealed a modification, K2me2 that was not observed by Edman degradation. We also confirmed that K2, K4, and K10 can be acetylated. Tandem MS analysis of peptide 16–21 demonstrated that K17 is mono- and tri-methylated and K18 is mono-, di- , and tri-methylated (Table 2). Tandem MS spectra representing peptide 16–21 were initially difficult to interpret because the mass of the peptide was 17 Da less than the theoretical mass. Closer examination of these spectra demonstrate that glutamine 16 (Q16) forms a cyclic compound between its free amine and amide carbon, resulting in the loss of ammonia and consequent molecular weight decrease of 17 Da. The cyclization of glutamine to form pyroglutamic acid has been characterized previously by MS [23].

MS analysis of Glu-C-digested H4 was focused on N-terminal peptides 1–8 and 9–22. We were unable to acquire MALDI-TOF data for peptides 1–8 and found only one modified species by tandem MS, demonstrating that A1 is monomethylated and K4 is acetylated (Table 2). Edman degradation showed that this is the most abundant species of peptides 1–8. MALDI-TOF analysis of the fractionated Glu-C digest showed that peptides 9–22 is present in multiple modification states (Fig. 6). Tandem MS sequencing of peptides 9–22 illustrated the combinations in which K10ac, K17me, and K18me may be present (Table 2). We were unable to sequence all of the modified species of peptides 9–22, so PTMs for the following species were not assigned: M + 14 Da, M + 28 Da, M + 98 Da, and M + 112 Da. The significance of these PTMs and the combinations in which they are present is currently unknown, but functional studies of individual marks and their corresponding histone-modifying enzymes will enable us to interpret this data in a meaningful way in the future.

4. Discussion

In this study, we attempted to identify the major covalent modifications present on the four core histones of BF T. brucei. All of the identified PTMs and the methods used to discover them are summarized in Table 3. The N-terminus of H3 was blocked to Edman degradation, and minimal tandem MS data were acquired for modified N-terminal peptides. However, Qq-TOF data of the H3 PA-trypsin and Asp-N digests suggested that several lysines at the H3 N-terminus can be modified. MALDI-TOF data of the
PF H3 N-terminus also suggested that a number of PTMs were present, although no tandem MS data were acquired to assign PTMs to their specific amino acids. Sequencing of trypsin-digested H3 by tandem MS showed that K23 is acetylated and K32 is trimethylated. T. brucei K23 and K32 are the possible homologues of K27 and K36 in other organisms (Fig. 1D), where their modifications are important for gene regulation. There are also predicted homologues of modified lysines 18, 23, 27, and 36 (T. brucei K16, K19, K23, and K32), indicating several possible marks for future study.

Next we investigated the PTMs on T. brucei histone H4. It has been shown that the acetylation of the S. cerevisiae H4 N-terminus at K5, K8, K12, and K16 is classically associated with transcriptional activation [2]. According to our alignment, S. cerevisiae K5, K12, and K16 could correspond to T. brucei K4, K10, and K14, all of which are acetylated (Fig. 1E). Although the T. brucei H4 sequence diverges considerably from the extremely conserved H4 N-terminus of other organisms, three of the four acetylated lysines appear to be conserved.

Importantly, in addition to the conserved acetylated lysines found at the H4 N-terminus, there is an apparent homologue of the methylated H4 K20, T. brucei H4 K18 (Fig. 1E). Trimethyl H4 K20 is a marker of constitutive and facultative heterochromatin and is in competition with the hyperacetylated H4 N-terminus [24]. Monomethyl H4 K20 is present at active gene promoters in association with H4 hyperacetylation, which demonstrates that the degree of methylation may be associated with distinct transcriptional states [25]. We found that T. brucei H4 K18 and the adjacent K17 may both be mono-, di- or trimethylated. K17 and K18 exist together in several modification states (Table 3). It is unlikely that these different methylation levels are created by different methyltransferases because there are only three identifiable putative SET domain-containing methyltransferases in the T. brucei genome [26]. Instead, it is more likely that a single methyltransferase acts on both K17 and K18, and the degree of methylation is determined by cofactors acting on the methyltransferase. We cannot predict whether the degree of methylation or the combination of modification states...
in which K17 and K18 co-exist will result in distinct biological readouts. If so, the amount of variability would allow for intricately regulated downstream effects.

Two modified lysines in the T. brucei H4 N-terminus, K2 and K5, did not have homologues in other organisms. We have shown that H4 K2 can be either acetylated or dimethylated. Histones from other organisms possess a number of lysines that may be either acetylated or methylated as well. Chicken H3 K9, for example, is subject to both types of covalent modification: H3K9me3 is associated with inactive chromatin, whereas H3K9ac is found at the boundaries of heterochromatin [27]. These marks have antagonistic roles in the establishment of different chromatin states. Whether T. brucei H4 K2 is involved in determining chromatin structure is unknown, but we propose that the dual modification status of this lysine makes it an interesting target for further characterization.

The H2A C-terminus contains six acetylated lysines (K115, K119, K120, K122, K125, and K128). Three of these, K120, K122, and K128, are probable homologues of lysines in other organisms, but the homologues of T. brucei K120 and K128 do not appear to be modified (Fig. 1B). T. brucei H2A K122 aligns with human H2A K119, the site of ubiquitination on this histone [21]. In other organisms, H2A ubiquitination is associated with transcriptional silencing [28,29]. We found evidence that T. brucei H2A C-terminus is ubiquitinated—the molecular weight of an H2A peptides 107–133 corresponds to a modified species that has one ubiquitinated and three acetylated lysines. Interestingly, T. brucei K122 is the only lysine at the H2A C-terminus that is adjacent to a possible phosphorylation site, S123. In other organisms, ubiquitinated lysines at the H2A and H2B C-termini are adjacent to phospho-acceptor sites. It has been suggested that phosphorylation influences ubiquitination and de-ubiquitination.

Table 3
Summary of histone modifications in bloodstream forms

<table>
<thead>
<tr>
<th>Histone</th>
<th>N-terminus</th>
<th>Acetyl lysine</th>
<th>Methyl lysine</th>
</tr>
</thead>
<tbody>
<tr>
<td>H2A</td>
<td>A1me1 (60%)&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>K4 (1%)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>A1ac&lt;sup&gt;a&lt;/sup&gt;</td>
<td>K11&lt;sup&gt;b&lt;/sup&gt;</td>
<td>K17&lt;sup&gt;a&lt;/sup&gt;me1 (4%), K17&lt;sup&gt;a&lt;/sup&gt;me2 (8%), K17&lt;sup&gt;a&lt;/sup&gt;me3 (9%)&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>H2B</td>
<td>A1me1 (60%)&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>K2 (2%)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>K2me2&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>A1me2&lt;sup&gt;b&lt;/sup&gt;</td>
<td>K4 (73%)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>K17me1 (4%), K17me2 (8%), K17me3 (9%)&lt;sup&gt;a,b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>S1ac&lt;sup&gt;b&lt;/sup&gt;</td>
<td>K5 (7%)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>K18me1 (6%), K18me2 (4%), K18me3 (18%)&lt;sup&gt;a,b&lt;/sup&gt;</td>
</tr>
<tr>
<td>H3</td>
<td>A1me1 (60%)&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>K10 (7%)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>K18me1 (6%), K18me2 (4%), K18me3 (18%)&lt;sup&gt;a,b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>A1me2&lt;sup&gt;b&lt;/sup&gt;</td>
<td>K14 (1%)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>K18me1 (6%), K18me2 (4%), K18me3 (18%)&lt;sup&gt;a,b&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup> PTMs observed by Edman degradation. The levels of acetylation and methylation, shown in parenthesis, were quantified by comparison to known standards.

<sup>b</sup> PTMs observed by tandem MS. K<sub>me1</sub>, K<sub>me2</sub>, and K<sub>me3</sub> refer to mono-methyl, di-methyl, and tri-methyl lysine, respectively; A<sub>ac</sub>, A<sub>me1</sub>, and A<sub>me2</sub> refer to acetyl, mono-methyl, and di-methyl alanine, respectively; S<sub>ac</sub> refers to acetyl serine.
of its neighboring lysine and also influences whether effector proteins bind ubiquitinated lysine [30]. Based on our data and the alignment of the H2A C-terminus with the C-terminus of H2A from other organisms, we propose that *T. brucei* H2A K122 can be ubiquitinated.

Two recent studies identifying covalent histone modifications in *T. cruzi* and PF *T. brucei* were published recently [13,14]. Both studies uncovered H4 acetylation at K4 and K10 (and K14 in *T. cruzi*) and monomethylation at K18 (and mono- and dimethylation at K17 and trimethylation at K18 in PF *T. brucei*). Additionally, our study detected H4 acetylation at K2 and K5, dimethylation at K2, trimethylation at K17, and dimethylation at K18. Perhaps these marks reflect life-cycle stage variation in BF *T. brucei*. The same may be said for the acetylated lysines found in BF H2A (K4 and K122) and H2B (K4, K12, and K16), which were not detected in PF. Finally, the *T. cruzi* H4 study detected K57 acetylation and R53 dimethylation. After a thorough analysis of *T. brucei* H4, we conclude that these marks are not present and may reflect species-specific variation of histone PTMs.

In higher eukaryotes, histones are highly conserved proteins. The extreme sequence divergence of *T. brucei* histones makes them interesting targets for phylogenetic comparisons. We demonstrate that, despite the sequence divergence, the locations of several modified residues are similar. The information compiled here provides a number of starting points for functional studies, some of which we are beginning to undertake. Future work will focus on identifying the modifications that play key roles in transcriptional regulation in trypanosomes.

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**References**