Analyses of Histone Proteoforms Using Front-end Electron Transfer Dissociation-enabled Orbitrap Instruments*§

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Histones represent a class of proteins ideally suited to analyses by top-down mass spectrometry due to their relatively small size, the high electron transfer dissociation-compatible charge states they exhibit, and the potential to gain valuable information concerning combinatorial post-translational modifications and variants. We recently described new methods in mass spectrometry for the acquisition of high-quality MS/MS spectra of intact proteins (Anderson, L. C., English, A. M., Wang, W., Bai, D. L., Shabanowitz, J., and Hunt, D. F. (2015) Int. J. Mass Spectrom. 377, 617–624). Here, we report an extension of these techniques. Sequential ion/ion reactions carried out in a modified Orbitrap Velos Pro/Elite™ capable of multiple fragment ion fills of the C-trap, in combination with data-dependent and targeted HPLC-MS experiments, were used to obtain high resolution MS/MS spectra of histones from butyrate-treated HeLa cells. These spectra were used to identify several unique intact histone proteoforms with up to 81% sequence coverage. We also demonstrate that parallel ion parking during ion/ion proton transfer reactions can be used to separate species of overlapping m/z that are not separated chromatographically, revealing previously indiscernible signals. Finally, we characterized several truncated forms of H2A and H2B found within the histone fractions analyzed, achieving up to 93% sequence coverage by electron transfer dissociation MS/MS. Results of follow-up in vitro experiments suggest that some of the truncated histone H2A proteoforms we observed can be generated by cathepsin L, an enzyme known to also catalyze clipping of histone H3. Molecular & Cellular Proteomics 15: 10.1074/mcp.O115.053843, 975–988, 2016.

Chromatin is the structural framework that packages DNA into chromosomes within the nucleus of a cell (2). Histones comprise the principal protein component of chromatin and are involved in the regulation of gene expression (3, 4). This epigenetic regulation is achieved through complex patterns of post-translational modifications (PTMs), the incorporation of histone variants, and through controlled histone proteolysis (5–10). Comprehensive characterization of histones by mass spectrometry (MS) has proven technically difficult for a number of reasons. Traditional methods (bottom-up MS) of sequence determination and PTM site localization are not practical. Histone N-terminal regions are rich in lysine and arginine residues, and thus proteolysis using trypsin generates peptides that are too small or that are poorly retained on reverse-phase HPLC C18 resins for subsequent MS detection (11). With the advent of electron transfer dissociation (ETD) and more efficient electron capture dissociation fragmentation methods, which are better suited for larger, more highly charged peptides (12, 13), several studies utilizing other endoproteases to generate longer peptides have emerged (14–16). Although these methodologies do well to preserve the combinatorial PTM profiles of histone tails, in some cases it is still impossible to identify the proteoforms from which these peptides originate. This is why analyzing histones intact, as they exist in the cells from which they are derived, is the best method for identifying unique histone proteoforms.

The results of several recent studies involving top-down analyses of histones highlight the complexity of the histone

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1 The abbreviations used are: PTM, post-translational modification; HCD, higher-energy collisional dissociation; ETD, electron transfer dissociation; ECD, electron capture dissociation; MS/MS, tandem mass spectrometry; SNR, signal-to-noise ratio; FETD, front-end electron transfer dissociation; IIPT, ion/ion proton transfer; m/z, mass to charge ratio; v/v, volume/volume percent; AGC, automatic gain control; RP-HPLC, reverse-phase high-performance liquid chromatography; ppm, parts per million.
An FETD-enabled Orbitrap mass spectrometer was used to analyze the histones derived from butyrate-treated HeLa cells. The instrument was upgraded to perform data-dependent MS/MS acquisitions in FT mode, which allowed for the detection of fragment ions in high resolution. The mass spectrometer was equipped with a front-end ETD/IIPT module, which simplified ETD fragment ion spectra and provided enhanced fragmentation capabilities.

**EXPERIMENTAL PROCEDURES**

Unless otherwise stated, all MS analyses were carried out on an in-house modified (21) hybrid Orbitrap Velos mass spectrometer (Thermo Fisher Scientific, San Jose, CA). In the time that the experiments described below were performed, the instrument was upgraded to include a high-field Orbitrap mass analyzer (26). Experiments involving recombinant H4 and parallel ion parking were performed following the upgrade and so they make use of enhanced FT (27). Otherwise, spectra are acquired in magnitude mode.

**Direct Infusion of Recombinant H4** — Recombinant H4 was purchased from New England Biolabs (Ipswich, MA), desalted using an Amicon® Ultra-0.5 3K centrifugal filter (Millipore, Billerica, MA), and reconstituted in a solution containing 40% acetonitrile and 60% of 0.1% acetic acid in water (all percentages expressed as v/v). The sample was directly infused at a concentration of 5 pmol/μL and ionized using microelectrospray ionization as described previously (28) into the front-end of an in-house modified ETD/IIPT-enabled Orbitrap Elite™. MS/MS spectra were acquired using the following instrument parameters: resolution (r) = 60,000 at 400 m/z; 190–2000 m/z scan range; 1E5 ion counts precursor automatic gain control (AGC) target; 3E5 reagent AGC target; 1–20 fragment ion fills of the C-trap; 1 μscan/spectrum; 2 m/z precursor isolation window centered at 708.4 m/z ([M + H]^+^-1820 charge state of H4); 6–8-ms ETD; 0–20–ms I IPT.

**Histone Extraction from HeLa S3 Cells** — Cultured HeLa S3 cells were treated with 10 mM sodium butyrate overnight and harvested, and the nuclei were isolated as described previously (29). The nuclei pellet was resuspended in 0.4 M NaCl containing 1 mM dithiothreitol (DTT), 0.3 mM 4-(2-aminophenyl) benzenesulfonyl fluoride hydrochloride, 10 mM sodium butyrate, and 5 μM Halt protease inhibitor mixture at a ratio of 10:1 (v/v), salt buffer/pellet. This was incubated with constant rotation for 2 h at 4 °C and then spun at 3000 × g for 5 min. Supernatant was discarded, and the pellet was resuspended in 2.5 M NaCl solution containing DTT, 4-(2-aminophenyl) benzenesulfonyl fluoride hydrochloride, sodium butyrate, and protease inhibitors as above in a ratio of 5:1 (v/v), salt buffer/pellet. The histones were then acid-extracted and TCA-precipitated as described previously (29).

**Off-line Histone Fractionation** — Histones purified from HeLa S3 cells (~100 μg) were dissolved in water and chromatographically separated on a Yldac C18 column (4.6-mm inner diameter, 250-mm length, 5-μm particle size) at a flow rate of 0.8 ml/min using a gradient of 30–60% B in 100 min, 60–100% B in 20 min, and 100–30% B in 10 min (solvent A = 5% (v/v) acetonitrile, 0.2% (v/v) trifluoroacetic acid in HPLC-grade water; solvent B = 95% (v/v) acetonitrile, 0.19% (v/v) trifluoroacetic acid in HPLC-grade water). Fractions were collected using an automatic sample collector in 1-min intervals between 15 and 80 min from the start of the gradient. The UV detector was set to monitor absorbance at 214 nm, and fractions containing histones H1, H2A, H2B, and H4 were identified and appropriately combined based on their known chromatographic retention times (29). These were transferred to microcentrifuge tubes, taken to dryness using a SpeedVac concentrator, and stored at −35 °C.

**On-line RP-HPLC for Mass Spectrometry** — Dried samples were reconstituted in water at concentrations of 5–10 pmol/μL. Aliquots of 1–2.5 pmol of protein along with 100 fmol of peptide standards were pressure-loaded onto a 360-μm outer diameter × 75-μm inner diameter fused-silica micro-capillary analytical column (packed 11-cm of POROSHELL 300SB-C18 resin (Agilent Technologies, Palo Alto, CA) (30) equipped with a laser-pulled electrospray emitter tip (31). The column was washed with −20 column volumes of 0.3% (v/v) formic acid in water. Samples were eluted at 60 nl/min using a gradient of 0–33% B in 10 min, 33–48% B in 60 min, 48–100% B in 10 min and 100–0% B in 10 min. The elution gradient utilized solvent A, 0.3% formic acid in water, and solvent B, 0.3% formic acid, 72% acetonitrile, 18% isopropyl alcohol, and 9.7% water (all v/v). Following reverse-phase HPLC separation, samples were directly ionized using microelectrospray ionization as described (28). For some samples, initial screens revealed that better chromatography might be achieved by enacting simple changes in the elution gradient. These changes included doubling the 0–33% B portion of the gradient to a time of 20 min for samples containing truncated histones and altering the gradient used for H1 to 0–20% B in 10 min, 20–40% B in 60 min, and then 40–100% B in 10 min. All other aspects of the RP-HPLC separation and ionization for mass spectrometry remained the same across all analyses.

**On-line MS Analyses** — Samples were ionized by microelectrospray into the front-end of an in-house modified ETD/IIPT-enabled Orbitrap Velos Pro™. Except for the occasional targeted analyses, the instrument was operated in data-dependent mode in which the MS1 scan was taken from m/z 300–2000 in the Orbitrap mass analyzer (r = 60,000 at m/z 400) followed by a second MS1 scan acquired in the linear ion trap over a limited m/z range, upon which data-dependent
selection was based. In this way, same species/different charge state reselection was reduced. A single ETD/IIPT MS/MS scan was also taken in the Orbitrap. MS/MS scan parameters were set as follows: $r = 30, 0$ or 100% at m/z 400; 200–2000 m/z scan range; 1E5 ion counts AGC target; 3E5 ion counts reagent AGC target; 5–30 fragmention fills of the C-trap; 1–3 μscans; 2–3 m/z precursor isolation window; 4–25 ms ETD (fluoranthene reagent); and 20–40–ms IIFT (sulfur hexafluoride reagent). The instrument was operated using charge state exclusion (+1, +2, and +3) and dynamic exclusion with a repeat count of 1–2, repeat duration of 60 s, and an exclusion duration of 90 s. Analytes of different sizes and charge densities require different ETD/IIPT reaction times, C-trap fill counts, and resolving power; these were varied depending on the fraction being analyzed. For example, when analyzing H1, a 21.9-kDa protein, shorter ETD and IIPT reaction times, more multiple fills, and higher resolution settings were used than those used for the analysis of H4, an 11.4-kDa protein. Duty cycle times also varied accordingly, but in general they were within 6–15 s/cycle.

Some fractions contained larger proteins that required MS/MS scan ranges over 2000 m/z for sequence determination and PTM site mapping. However, the instrument control software did not allow the acquisition of spectra with scan ranges exceeding 2000 m/z for ETD/IIPT MS/MS scan types in data-dependent mode. This prompted us to perform “on-the-fly” targeted analyses based upon full MS data acquired in the initial sample screenings. Instrument method parameters were varied at our discretion based on our evaluation of the MS/MS spectra generated as data were acquired.

High resolution full MS and MS/MS data were manually inspected using Qual Browser software (Thermo Fisher Scientific). Interpretation of all ETD/IIPT MS/MS data was performed manually on the unprocessed raw spectra. Theoretical fragment ion masses were calculated using an in-house-developed fragment mass calculator. Theoretical isotopic distributions were modeled using Isotope Pattern Calculator software (Pacific Northwest National Laboratories, omics.pnl.gov/). Percent sequence coverage was calculated by dividing the number of observed N–Ca bond cleavages by the total number of predicted N–Ca bond cleavages (note that cleavage of the N–Ca bond that is N-terminal to proline does not produce an observable fragment). Masses for all intact proteoforms identified agreed to within 5 ppm and were confirmed by MS/MS. Relative abundance information for each truncated histone species was determined by taking the area under the curve of the extracted ion chromatogram for the most abundant isoform for every charge state present utilizing a 10-ppm mass window (32). Parallel Ion Parking during IIPT—For parallel ion parking experiments, the MS1 scan was taken from m/z 300 to 2000 in the high-field Orbitrap mass analyzer (1E6 ion counts AGC target, $r = 120,000$ at 400 m/z). A 40 m/z window centered at m/z 810 (790–830 m/z) was selected and subject to 140 ms of IIFT. The instrument control software was modified to apply a custom RF waveform to the x-rod of the linear ion trap during IIFT. The waveform included frequencies mapping to m/z 2300–3900, corresponding to the desired final charge state of the precursor. For the sample analyzed (fraction H2B), the [M + 5H]+ charge state was the desired product (−m/z 2770). The amplitude of the waveform was adjusted such that maximum activation of the target product ions was achieved without dissociation or ejection. Product ions were analyzed in the Orbitrap (1E5 ion counts AGC target, $r = 120,000$ at 400 m/z) after five fills of the C-trap.

In Vitro H2A Proteolysis and nano-HPLC-MS/MS Analysis—The in vitro proteolysis of recombinant histone H2A type 2 (Uniprot ID, Q6F1I3) was carried out in 50 mM MES, pH 6.0, and 5 mM DTT. Recombinant cathepsin L (R&D Systems 1515-CY-010) was used in a 5000:1 (histone/enzyme) ratio during a 1-h time course reaction. 20 μg of the sample reaction was quenched with 10% acetic acid after 15, 30, and 60 min. Samples were desalted using in-house packed Stage tips as described previously (29). After Stage tip desalting, samples were resuspended in 0.1% formic acid prior to nano-LCMS/MS analysis. 1 μg of sample was loaded onto a 75-μm inner diameter × 17-cm Reprosil-Pur C18-AQ (3 μm; Dr. Maisch GmbH, Germany) column using an EASY-nLC nano-HPLC (Thermo Fisher Scientific). The HPLC gradient was 0–100% solvent B (A = 0.1% formic acid; B = 95% acetonitrile, 0.1% formic acid) over 60 min at a flow rate of 250 nL/min. nano-LC was coupled to an Orbitrap Fusion mass spectrometer (Thermo Fisher Scientific). MS1 spectra (m/z 300–1200) were acquired in the Orbitrap with a resolution of 120,000 (at 200 m/z) and an AGC target of 5E5. Data-dependent selection was performed using TopSpeed with a cycle time of 3 seconds. MS/MS for charge state +2–5 was performed using higher-energy collisional dissociation (HCD), with a normalized collision energy of 27%, a maximum injection time of 150 ms, and an AGC target of 5E4. Higher charge states were fragmented using ETD, using a reaction time of 20 ms, a maximum injection time of 400 ms, 3 μscans, and an AGC target of 5E5. Both types of MS/MS fragmentation were acquired in the Orbitrap with a resolution of 30,000 (at 200 m/z). Data were analyzed using Proteome Discoverer (version 1.4, Thermo Scientific, Bremen, Germany) with Mascot (Matrix Science, London, UK) as the database search engine (database: human histones, Uniprot, March 2014, 58 entries). Search parameters were as follows: HCD spectra were searched using a precursor ion mass tolerance of 10 ppm and a fragment ion mass tolerance of 0.05 Da; ETD spectra were deconvoluted using Xtract (Thermo Fisher Scientific) and searched using a precursor ion mass tolerance of 10 ppm and a fragment ion mass tolerance of 0.02 Da. Unspecific cleavage (no enzyme) and no modifications were specified. Only identifications with a false discovery rate of <1% (calculated using fixed value PSM validator) were accepted. Chromatograms of the relevant peptides were manually extracted using Xcalibur (Thermo Fisher Scientific) after deconvolution for peptide quantification.

RESULTS

Multiple Fills and IIPT—A Thermo Fisher Scientific Orbitrap Velos ProTM was modified to introduce reagent anions from the front by adding a glow discharge ionization source between the stacked ring ion guide and multipole 00, as described previously (21). The discharge is used to ionize ETD and IIPT reagents (fluoranthene and sulfur hexafluoride, respectively) for ion/ion reactions. By placing the reagent ion source in the front of the instrument, the C-trap is made available to store fragment ions from multiple iterations of ion/ion reactions. Once sufficient amounts of fragment ions are collected, they are sent to the Orbitrap for high resolution mass analysis.

Fig. 1, A and B, shows 8-ms ETD/20-ms IIPT MS/MS spectra of intact recombinant H4 using 1 and 20 fragment ion fills of the C-trap, respectively. SNRs for fragments shared between the two spectra are indicated in parentheses. By increasing the number of fragment ion volumes analyzed from 1 to 20, total fragment ion current is increased by an order of magnitude. This is also demonstrated in supplemental Fig. 1, which shows a trace of total ion current as the number of multiple fills is varied. The spectrum shown in Fig. 1C is an average of 20 spectra like that shown in Fig 1A and represents a cumulative acquisition time of 11.2 s (0.56 s/spectrum).
Fig. 1. **Multiple fills versus spectral averaging.** MS/MS spectra of intact recombinant H4 (precursor charge state +16, m/z 703.40) following ETD (8 ms) and I IPT (20 ms) reactions with 1 and 20 fragment ion fills of the C-trap. SNRs for fragments shared among the spectra are indicated in parentheses. A, spectrum produced from a single iteration of ion/ion reactions (1 fill, 1 μscan, r = 60,000 at 400 m/z, 0.56-s elapsed scan time). B, spectrum produced from 20 iterations of ion/ion reactions (20 fills, 1 μscan, r = 60,000 at 400 m/z, 6.24-s elapsed scan time). C, spectrum produced by averaging 20 (A) spectra (1 fill, 1 μscan, r = 60,000 at 400 m/z, 0.56-s elapsed scan time/spectrum, 11.2-s total acquisition time).
Elapsed scan time for the spectrum shown in Fig. 1B is 6.24 s. Average SNR gains for the fragments indicated are 10.4 in Fig. 1B versus 3.5 in Fig. 1C. Hand-annotated spectra indicating the most abundant fragments observed are shown in supplemental Fig. 2.

The use and benefits of I IPT to charge reduce ETD-produced fragment ions have been described previously (1, 23). Briefly, following fragmentation by ETD, fragment ions are allowed to react with a proton transfer reagent. These reagents act as a base and abstract protons from highly charged fragments. The amount of charge reduction is tuned by varying the amount of time the reaction is allowed to take place. By extending I IPT reaction time such that fragment ions are significantly charge-reduced, the fragment ion signals are increasingly distributed over the full scan range (example shown in supplemental Fig. 3).

Characterization of Intact Histones Derived from Butyrate-treated HeLa Cells—HeLa cells were treated with sodium butyrate prior to histone isolation and separation. Butyrate treatment of cells results in histone hyperacetylation; butanolate acts as an inhibitor of histone deacetylase activity (33). The histones were acid-extracted from the nuclei and purified by offline reverse-phase HPLC fractionation. The histones were identified based on their well known chromatographic retention times (34). Intact protein was directly loaded onto a reverse-phase C18 Poros-shell column and gradient-eluted into the mass spectrometer.

Fig. 2 summarizes the data that can be obtained from a single LC-MS experiment. Fig. 2A shows the base-peak chromatogram of the H2A2 fraction revealing three distinct regions of intact protein elution. Peaks 1 and 2 correspond to H4 in various modified forms containing oxidized and unoxidized methionines, respectively, whereas peak 3 corresponds to various proteoforms of intact H2A. Displayed in Fig. 2, B and C, are the averaged high resolution ESI spectra recorded on peaks 2 and 3 in the base peak chromatogram, respectively. Signals in the observed charge state distribution carry 7–18 and 8–23 positive charges for H4 and H2A, respectively. An ETD/I IPT MS/MS spectra recorded on an [M + 19H]19+ ion of an intact form of H2A is shown in Fig. 2D. This spectrum was obtained using a 4-ms ETD and a 25-ms I IPT and is an average of six spectra, each taking 11.7 s to acquire utilizing 3 μs scans and 15 multiple fills with r = 30,000 at 400 m/z. Note that equivalent spectra taken as the sum of 45 μs scans (transients) of single fragment ion fills would take ~19.8 s/scan. Residues identified by ions of types c and z observed in Fig. 2D are indicated in Fig. 2E (hand annotated version shown in supplemental Fig. 4). These define 80% sequence coverage and identify the intact H2A as H2A.1 (Q93077).

Depicted in supplemental Fig. 5 is a full MS spectrum indicating signals for intact H2A protoforms that were identified and confirmed by ETD/I IPT MS/MS. In addition to H2A.1, also detected were H2A.1-C and H2A.1-H (Uni Parc Q93077 and Q96KK5, respectively). H2A.1-C differs from H2A.1 by the substitution of Thr-16 with a serine residue and by substitution of Lys-99 with an arginine residue. These substitutions give rise to a Δ mass of +13.9904 Da from H2A.1, which is very close to the Δ mass that would be exhibited in the event of a monomethylation (+14.0157 Da). The calculated and observed mass differences for intact H2A.1-C and the monomethylated form of H2A.1 are 3.5 and 5.3 ppm, respectively, making H2A.1-C the better fit by mass accuracy. However, based on this information alone, neither could be conclusively excluded. The MS/MS spectra contained c ions that exhibited a Δ mass of −14 Da following Thr-16 and z ions that exhibited a mass shift of +28 Da following Lys-99. The +28-Da mass shift is also indicative of a dimethylated lysine, but this possibility was ruled out based on the mass accuracy of the fragment ions observed (z313+, 13C m/z 1101.9823: 0.2 ppm for Lys → Arg, 7.8 ppm for Lys → di-methylation Lys). Thus, we were able to conclude that this is H2A.1-C. This demonstrates the importance of obtaining high quality MS/MS spectra for the confirmation of proteoform identity. H2A.1-H differs from H2A.1 by the deletion of the last two C-terminal residues, emphasizing the importance of analyzing the intact forms of these proteins, as digestions would produce identical peptides with the exception of those containing the C termini. This difference was easily discerned upon examination of the z fragment ion series in the MS/MS spectra. H2A.1-C and H2A.1-H are present at levels roughly 20–25 and 15–20%, respectively, relative to H2A.1. Eluting later in the gradient and also confirmed by MS/MS are the respective mono-Lys-acetylated forms of these proteins. The acetylation is localized to Lys-5 in all H2A types.

Fig. 3 displays a spectrum recorded on [M + 16H]16+ (m/z 718.0) from intact H4 using a 4-ms ETD reaction time and 40-ms I IPT. This is an average of eight spectra, each taking ~8.75 s to acquire utilizing 3 μs scans at r = 30,000 (at 400 m/z) with 10 multiple fills of the C-trap per μs scan (~13.9-s acquisition for single fill/transient equivalent spectrum). Fig. 3B offers an expanded view of the 1250–1500 m/z region of the spectrum. With the exception of the extreme C terminus, which is devoid of charge, a near contiguous series of both c’ and z’ can be observed for ~60 residues from either end of the protein, yielding 81% sequence coverage and site localization of five acetylations as well as Arg-20 dimethylation (fully annotated spectrum shown in supplemental Fig. 6).

Compared with other histones, H1 presents a more difficult situation for on-line LC-MS analysis of the intact form. The most common human isoforms of H1 range from 194 to 226 amino acids in length and, due to their larger size, require higher resolution, shorter gas-phase ion/ion reaction times, and multiple fills of the C-trap, and an extended mass range. Initial sample runs revealed the presence of at least two forms of H1, H1.4 and H1.2. MS/MS spectra acquired in data-dependent mode using 2-ms ETD/25-ms I IPT are shown in supplemental Fig. 7 along with the observed sequence cov-
Fig. 2. Example data obtained from a single LC-MS experiment. A, base peak chromatogram obtained from LC-MS analysis of histone fraction H2A2. Peaks corresponding to 100 fmol of co-loaded peptide standards are labeled, and regions of intact protein elution are numbered. B and C, averaged ESI full MS spectra depicting the charge state distributions of intact H4 and H2A, respectively (peaks 2 and 3 from A, respectively). D, ETD/IPT spectrum recorded on $[\text{M + 19H}]^{19+}$ ions from H2A type 1 (P0C0S8) using 4-ms ETD and 25-ms IPT. Precursor m/z 738.0, 3 µscans, 15 multiple-fills, $r = 30,000$, average of 6 spectra, 11.7-s elapsed scan time/spectrum (19.8 s for single fill/transient equivalent spectrum). E, sequence coverage obtained from MS/MS spectrum shown in D.
Fig. 3. ETD/IIPT MS/MS spectrum of intact H4 proteoform. A, ETD/IIPT spectrum recorded on [M + 16H]^{16+}\ ions from H4 (P62805) using 4-ms ETD and 40 ms IIPT. Precursor m/z 718.0, 3 μscans, 10 multiple-fills, r = 30,000, average of 8 spectra, 8.75 sec elapsed scan time/spectrum. B, expanded view of the 1250–1500 m/z region of A (indicated in red). Ions of type c' and z'' are indicated. C, sequence coverage obtained from MS/MS spectrum shown in A. The locations of PTMs are indicated.
verage (H1.4, 72%, and H1.2, 67%, P16403 and P10412, respectively). Additional targeted analyses elucidated several additional unique H1 proteoforms. One of these was a form of H1.2 containing a previously uncharacterized single amino acid substitution, A141T. The spectrum and coverage are shown in Fig. 4 (hand-annotated version shown in supplementary Fig. 8).

**Parallel Ion Parking during IIPT**—Histone fraction H2B presented the most difficult challenge due to the number of unique proteoforms it contained. Fig. 5, A and B, displays the averaged (five spectra, \( r = 120,000 \) at 400 \( m/z \)) high resolution ESI spectra recorded on intact H2B proteoforms eluting over ~45 s. Isolation of the region containing the [M + 23H]\(^{23+}\) ions from H1.2 (P16403) with an undocumented single amino acid substitution (A141T) using 4-ms ETD and 20-ms IIPT. Precursor \( m/z \) 927.5, 1 \( \mu \)scan, 25 multiple-fills, \( r = 100,000 \) at 400 \( m/z \), average of 11 spectra, 9.38-s elapsed scan time/spectrum (27.6 s for single fill/transient equivalent spectrum). B, sequence coverage obtained from MS/MS spectrum shown in A. The location of the Ala → Thr amino acid substitution is indicated with an asterisk.

**Fig. 4.** ETD/IIPT MS/MS spectrum of intact H1 proteoform. A, ETD/IIPT spectrum recorded on [M + 23H]\(^{23+}\) ions from H1.2 (P16403) with an undocumented single amino acid substitution (A141T) using 4-ms ETD and 20-ms IIPT. Precursor \( m/z \) 927.5, 1 \( \mu \)scan, 25 multiple-fills, \( r = 100,000 \) at 400 \( m/z \), average of 11 spectra, 9.38-s elapsed scan time/spectrum (27.6 s for single fill/transient equivalent spectrum). B, sequence coverage obtained from MS/MS spectrum shown in A. The location of the Ala → Thr amino acid substitution is indicated with an asterisk.

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Truncated H2B proteoforms observed in I IPT/parallel ion parking MS/MS spectrum

Observed m/z and theoretical precursor m/z values are shown. The masses reported are those of the most abundant isotopologue in the charge state envelope. UniProt sequences from isoforms of H2B with residues matching those contained in the truncated H2B are given. Note these are not confirmed by MS/MS, and identification is based on accurate mass and MS/MS spectra of other truncated H2Bs seen in LC-MS analyses performed without parallel ion parking. Observed mass accuracies are reported in ppm and were calculated from the most abundant isotopologue.

| Peak | Observed m/z | Precursor m/z | Possible H2B isoform | Residues | Observed mass accuracy (ppm) |
|------|--------------|---------------|----------------------|----------|-----------------------------|
| A    | 2134.4929 (+3) | 801.0661 (+8) | P62807              | 68–125   | 2.16                        |
| B    | 2167.1735 (+3) | 813.3221 (+8) | O60814              | 67–125   | 3.09                        |
| C    | 2172.5071 (+3) | 815.3215 (+8) | P62807              | 67–125   | 2.16                        |
| D    | 2200.1957 (+3) | 825.7057 (+8) | O60814              | 66–125   | 3.32                        |
| E    | 2205.5313 (+3) | 827.7050 (+8) | P62807              | 66–125   | 1.50                        |

Fig. 5D displays the product ion spectrum following 140-ms I IPT/parking. The vast majority of the ion current lies within 2730–2800 m/z range where the “parked” intact H2B [M + 5H] +5 ions lie. An enlarged inset of this region is shown in supplemental Fig. 10. Signals corresponding to precursor ions that were not fully reacted to the +5 charge state or that were not fully parked and underwent further proton transfer to the +4 are indicated. Close inspection of the spectrum shown in Fig. 5D reveals several signals of lesser charged species (labeled A–E). Their observed charge state and m/z as well as their calculated m/z and charge state prior to I IPT are given in Table I. Comparisons of the observed masses to calculated theoretical values suggest that these signals are derived from truncated forms of H2B that co-eluted with intact H2B during the time these spectra were acquired (identity and mass accuracy of the most abundant isotopologue in each envelope are shown in Table I). Peptides containing residues 66–125, 67–125, and 68–125 derived from two isoforms of H2B match within 5 ppm mass error. Expected signals for these species A–C are not observed in Fig. 5C (also see supplemental Fig. 9), presumably due to the density of unique species within the window. Note that the assigned identities for these truncated H2Bs are not confirmed by MS/MS and are based on mass accuracy and on MS/MS-confirmed identities of the intact total. The +17 charge state is indicated. B, expanded view of the region indicated in red (m/z 790–830) from A. C, spectrum produced upon isolation of ions possessing m/z 790–830. Truncated proteoforms of +8 charge are labeled D and E. D, MS/MS spectrum produced upon isolation of ions possessing m/z 790–830 and 140-ms I IPT with charge-reduced precursor “parked” in the 2300–3900 m/z range. The inset shows an expanded view of newly observed truncated histone H2B proteoforms (labeled A–C).
H2Bs observed in this fraction. These truncated H2Bs could be derived from several other proteoforms of intact H2B due to their sequence similarity.

Characterization of Truncated H2B and H2A Proteoforms Derived from Butyrate-treated HeLa Cells—Eluting after the intact H2B proteoforms were several fairly abundant forms of truncated H2B. The base peak chromatogram and the sequence of the most abundant intact H2B (H2B-1C, P62807) observed in the fraction are shown in Fig. 6. The solid black lines indicate “primary” cleavage sites mapped by MS. The gray lines indicate additional less abundant cleavage sites.

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Characterization of Truncated H2B and H2A Proteoforms Derived from Butyrate-treated HeLa Cells—Eluting after the intact H2B proteoforms were several fairly abundant forms of truncated H2B. The base peak chromatogram and the sequence of the most abundant intact H2B (H2B-1C, P62807) observed in the fraction are shown in Fig. 6. The solid black lines indicate “primary” cleavage sites mapped by MS. The gray lines indicate additional less abundant cleavage sites. This is not a comprehensive account of all the cleavages that are observed, but rather only those that are most abundant. Complementary N-terminal peptides, 1–25 and 1–51, are present and elute near the front of the gradient. They are less abundant and are more difficult to characterize because their signals are split among multiple acetylated forms (+1–3 lysine-acetyl confirmed by MS/MS). The acetylations are shared among multiple sites, including Lys-11, Lys-12, Lys-15, Lys-16, and Lys-20, which are all sites that were confirmed by MS/MS in the intact form and that have previously been identified by MS in an earlier study (35). Truncations derived from two additional H2B isoforms (possibly H2B.2-E, Q16778, and H2B.1-K, O60814) were also present. Note that the sequences of the intact forms found in the sample differ by only one residue (V39I for H2B.2-E or S124A for H2B.1-K), and so distinction between truncated peptides can only be made if both unique residues are contained within the peptide. If only one of these residues is present, then only one of the two additional isoforms can be excluded. There is also no way to distinguish the modified (acetylated) form of the intact protein from which the C-terminal peptides are derived. Up to 93% sequence coverage is obtained by ETD/IIPT MS/MS of the three most abundant C-terminal containing peptides (90%, 26–125; 93%, 52–125; and 84%, 50–125). The spectrum and sequence cov-
erage of the 26–125 peptide are shown in supplemental Fig. 11.

Additionally, dozens of truncated H2As were observed in the H2A1 fraction. These ranged in size from 10 to 75 residues in length, and each account for less than 1% of the total ion current relative to intact H2A from the same fraction. The base peak chromatogram is shown in Fig. 7, A–C, and indicates where truncated species elute relative to intact H2A. Fig. 7D shows the sequence of the most abundant intact H2A observed in the fraction (H2A2-A, Q6FI13). Again, the solid black lines in Fig. 7 indicate “primary” cleavage sites, and the less abundant cleavage sites are indicated with gray lines. This is not a comprehensive account of all the truncated histone forms observed, only the most abundant. Supplemental Table 1 lists these truncated proteoforms along with their respective abundances. The most abundant of these was the N termi-
Histones containing proteoform cleaved C-terminal to Gly-44. In- 
tact forms of H2A identified on the fraction were H2A2A and 
H2A2C (Q16777), which differ by their last 5–6 C-terminal 
residues. Because of this, there is no way to determine 
whether the N terminus-containing truncations were derived 
from H2A2A or H2A2C. MS/MS spectra of truncated species 
are shown in supplemental Fig. 12.

Identification of Histone H2A Clipping Enzyme by in Vitro 
proteolytic Assay—Given the large variety of truncated his-
tone proteoforms identified, we further investigated whether 
cathepsin L, a protease responsible for histone H3 and H3.3 
cipping (9, 37, 38), is capable of cleaving histone H2A. To do 
so, we performed an in vitro assay incubating recombinant 
histone H2A with cathepsin L and characterized the cleavage 
specificity using nano-LC coupled to an Orbitrap Fusion with 
FETD. Supplemental Fig. 13 and supplemental Table 2 show 
that the signal of intact histone H2A decreased by ~3 orders 
of magnitude after 1 h of incubation, while the clipped forms 
increased in abundance over time. Specifically, we deter-
dined the clipped proteoform Gly-44 (cleaved after glycine 
44) as one of the most abundant cleaved proteoforms and 
confirmed its sequence by ETD MS/MS (supplemental Fig. 
13C). We also observed several other C-terminal peptides 
(supplemental Table 2). Taken together, our data suggest that 
cathepsin L might have a larger variety of targets than cur-
rently described.

DISCUSSION

In the commercial configuration of the Orbitrap Velos Pro/ 
EliteTM, the reagent ion source is located at the rear of the 
instrument, requiring that reagent anions traverse the C-trap 
to reach the ion trap, where ion/ion reactions are performed. 
By putting the reagent ion source in the front of the instru-
methat, the C-trap is made available for the storage of frag-
ments from multiple iterations of ion/ion reactions prior to 
transient acquisition in the Orbitrap. In this way, we are able to 
transient ion current. The multiple fill 
strategy for improving SNRs has a number of advantages over 
 transient summing or spectral averaging. Chief among these 
is that the time that it takes to acquire spectra of multiple 
C-trap fills is less than the time it takes to acquire transients 
from the same number of single C-trap fills to be averaged or 
summed. Specifically, if a transient takes 380 ms to acquire, 
an additional fragment ion fill can be accomplished in less 
than ½ that time, i.e. ~130 ms for overhead plus time for 
ETD/IPT activation. Additionally, SNRs improve faster using 
multiple fills compared with summing transients or averaging 
spectra and so fewer transients are needed to produce a 
useful spectrum for analysis. This enables the acquisition of 
high quality spectra on a time scale more compatible with 
chromatography.

The power of top-down mass spectrometry lies in its ability 
to provide an unambiguous view of protein expression dy-
namics by distinguishing closely related proteoforms from 
one another while also observing any combinations of PTMs. 
Bottom-up MS strategies are not capable of this, which is why 
there has been a significant push in the field of proteomics to 
develop instruments and methodologies that enable high 
throughput, top-down mass spectrometry of intact proteins. 
Here, we demonstrated the power of FETD-enabled sequen-
tial ion/ion reactions along with multiple fragment ion fills 
through their coupling with on-line HPLC and application to the 
analysis of intact histones. Data-dependent and targeted 
LC-MS experiments were employed to record high resolution 
MS/MS spectra of histones from butyrate-treated HeLa cells. 
Using these spectra, several unique intact histone proteo-
forms were identified with up to 81% sequence coverage. 
Examples shown here contain multiple modifications, includ-
ing single amino acid substitutions, methylations, and acety-
lations. Additionally, many truncated forms of histones H2A 
and H2B were characterized with up to 93% sequence 
coverage.

Histone proteolysis has been implicated in several impor-
tant regulatory processes, yet it remains poorly understood. 
Only eight histone-specific proteases are known, and their 
biological significance has not been extensively defined (39). 
This area of study is clearly poised for significant expansion, 
as evidenced by the identification of two H2A-specific pro-
teases and several novel histone clipping events in just the 
past few years (39–42). Discerning whether any histone-
cleaving event is the result of continuous histone turnover or 
sample degradation versus an alternative biological process 
with epigenetic implications is challenging (36). Our in vitro 
data suggest that cathepsin L might be a potential candidate 
to explain some of the truncated histone H2A proteoforms we 
observed in the butyrate-treated HeLa cell histone samples. 
We found that cleavage C-terminal to Gly-44 produced the 
most abundant truncated H2A proteoforms observed in both 
the in vitro experiments and those derived in vivo from HeLa 
cells. Differences between the in vitro and the HeLa cell data 
in the abundance of other truncated proteoforms might be 
explained by improper folding of the recombinant histone or 
by the absence of a proper nucleosome structure in the in 
vitro experiments. In the future, we hope that the tools we 
describe here will aid in deciphering whether or not these 
truncated histone proteoforms are biologically regulated and 
contribute to epigenetic mechanisms.

That we are able to observe these low level truncated 
histone proteoforms in simple top-down LC-MS experiments 
further demonstrates the utility of our methodologies as it is 
very likely that bottom-up- or middle-down-based proteomics 
approaches would have missed cleavage events like those we 
describe in our analyses. This could happen for a number of 
reasons. First, given the low levels at which these truncations 
were observed (which, for any single truncated form, is less 
than 1% relative to the intact histone), along with the added 
complexity that is generated when samples are digested for 
MS proteomic studies, peptides of this type may have not
been detected or may have simply gone unnoticed by investigators. Several of the generated peptides derived from originally truncated species might have been too small to be retained by the HPLC column. Quantitative analysis of the relative abundance of unique truncated forms would be difficult to perform as evidence for some unique truncated species could be lost or could be below the limit of detection. Also, the ability to correlate the endogenous histone proteoylsis with any specific histone variant or PTM state would not be possible unless those PTMs or sequence variations and the cleavage sites were contained within the same peptide produced in the sample preparative digestion. Only by determining the primary structures of these species, as they existed in the cell, can one draw these potentially important biological parallels.

Furthermore, the use of parallel ion parking during I IPT is demonstrated in an analysis of H2B proteoforms. Chromatographic separation is made difficult by the fact that several H2B proteoforms differ from one another by just a single amino acid residue. At high ETD-compatible charge states, multiple proteoforms are present within a relatively small m/z range. Upon data-dependent selection, often more than one proteoform is isolated and fragmented, resulting in mixed MS/MS spectra, making identification difficult. Additionally, as demonstrated here, overlapping signals make it difficult to tease out the number of unique species present. By using I IPT to charge-reduce the precursors contained within a wide isolation window and parking the products in a desired m/z range, we are able separate species that, prior to I IPT, possessed roughly the same m/z but a significantly different mass. This reveals signals that can be attributed to less abundant H2B proteoforms that were not observed in the MS1 spectrum nor the isolation–only spectrum due to their proximity to more abundant signals derived from other species. This is the first time ion parking has been demonstrated on a commercially available linear ion trap using ion cloud densities that lend themselves to high resolution MS/MS analyses. We believe that parallel ion parking will become a powerful tool in identifying proteins in a mixture when used in conjunction with MS/MS fragmentation methods and multiple fill technology. Most exciting is the potential for this technology to allow differentiation and identification of closely related proteoforms that are difficult to separate chromatographically, like H2B. As demonstrated in supplemental Fig. 10, following I IPT/parking, we are able to spread apart isotopic clusters such that their centers are ~3 m/z apart (compared with <1 m/z prior to I IPT) without losing signal due to further proton transfer. This is a stepping stone to enabling higher resolution isolation on the MS3 level. However, doing so will require further instrument modification to increase the precursor ion capacity of the linear ion trap to be able to observe fragments in MS3 spectra following I IPT.

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