Histone sample preparation for bottom-up mass spectrometry: a roadmap to informed decisions

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Abstract

Histone-based chromatin organization enabled eukaryotic genome complexity. This epigenetic control mechanism allowed for the differentiation of stable gene-expression and thus the very existence of multicellular organisms. This existential role in biology makes histones one of the most complexly modified molecules in the biotic world, which makes these key regulators notoriously hard to analyze. We here provide a roadmap to enable fast and informed selection of a bottom-up mass spectrometry sample preparation protocol that matches a specific research question. We therefore propose a two-step assessment procedure: (i) visualization of the coverage that is attained for a given workflow and (ii) direct alignment-between-runs to assess potential pitfalls at the ion level. To illustrate the applicability, we compare four different sample preparation protocols, while adding a new enzyme to the toolbox, i.e., RgpB (GingisREX®, Genovis), an endoproteinase that selectively and efficiently cleaves at the c-terminal end of arginine residues.

Raw data is available via ProteomeXchange with identifier PXD024423.

Introduction

The eukaryotic lineage arose from the symbiotic merger between bacterial and archaeal cells. Herein the bacterial partner contributed genes, metabolic energy, and the building blocks of the endomembrane system, while the archaeal partner provided the potential for complex information processing by adding histones to the eukaryotic experiment[1]. The strongly conserved histone proteins provided the potential for complex information processing by developing a nucleosome-based chromatin structure [2,3]. This, together with the complex language of chemical modifications evolved into the so-called histone code [4]. This not only enabled the organization of vast amounts of DNA, but also the regulation of gene expression through the modulation of the functional state of DNA[1]. In fact, it is hypothesized that the modifications of the histone code arose as a sensing mechanism of the energetic state of the cell; the presence of energy rich donors like acetyl-CoA (acetylation), ATP (phosphorylation), and SAM (methylation) enabled direct chemical modification of the histone backbone[5], thus fully entangling the Prokaryote energy housekeeping with the Archaeal expression system. Importantly, these epigenetic changes can persist over many cell generations, allowing for the differentiation of stable gene-expression patterns in various cell types, essential for multicellular organisms [6]. In other words, histones are at the heart of the eukaryotic information management system and therefore amongst the most complexly modified proteins in the biotic world.

Over the past decades it has become clear that mass spectrometry (MS) is an indispensable technique capable of studying the complex interplay of the histone code [7], i.e. the proverbial grammar of the Eukaryote information management system. In essence, an LC-MS/MS system measures the intensity and certain physicochemical properties of analytes, like mass-to-charge (m/z), retention time (tR), drift
time ($t_0$) and fragmentation pattern. This allows to create a multidimensional data space that captures a picture of the full complexity of the PTM landscape purely built from numbers.

However, as for any proteomics workflow, all parts of a histone workflow are entangled and each combination of sample preparation, data acquisition, and data analysis provides a different image of the histone code. Top-down and middle-down proteomics can omit or simplify many steps in sample preparation workflows, but the combinatorial complexity of coexisting PTMs makes many proteoforms hard to resolve [8–10]. Although middle-down approaches are more feasible and are particularly gaining interest in the field of LC-MS/MS based histone analysis [11,12], bottom-up approaches remain most widely used.

Unfortunately, the golden standard for LC-MS/MS based proteomics, i.e. tryptic digest, is less well suited for the bigger part of the histone code, due to the high abundance of lysine (K) and arginine (R) cleavage sites. Unmodified K and R residues result in peptides too small for efficient LC-separation [8,13], while changes in the modification status of K (and to a lesser extent R) dynamically block digestion, which in turn creates peptidoforms of different length complicating the alignment between runs for label-free quantification. Taken from the general abundance of both cleavage sites and their modification frequencies, a digest with arginine specificity would reduce these problems considerably. To this end, two widespread workarounds have been described. On the one hand, there is enzymatic digestion with clostripain, also known as ArgC—an enzyme that naturally cleaves at the c-terminal end of arginine residues. However, its lack of specificity discredited it for most users [14]. On the other hand, lysine residues can be chemically derivatized to obstruct digestion with trypsin, thus incorporating its superior efficiency and specificity into the workflow, while obtaining an arginine specific cleavage [15]. This most commonly used strategy gives good coverage of the N-tail of histone H3 and H4, but lags behind on coverage of other regions in the histone code and introduces additional chemical noise into the workflow [16–20]. Overall, the histone coverage greatly depends on the sample preparation workflow, regardless of how advanced the LC-MS/MS system and posterior data-analysis might be. Indeed, it is currently impossible to cover the entire histone code with a single workflow.

Here we present a new perspective on histone sample preparation that guides researchers in selecting a protocol that covers a specific stretch of interest, while taking certain limitations such as side reactions into account. As a proof-of-concept we have acquired data from bovine histone standards prepared with four different histone sample preparation workflows, including a workflow with the arginine specific enzyme RgpB (GingisREX, GRX, Genovis), which has not been described for bottom-up LC-MS based shotgun proteomics applications before [21,22]. Additionally, digestion with trypsin, both with and without derivatization of the lysine cleavage sites, as well as digestion with ArgC were included in the experimental design. To allow for a direct comparison of the obtained coverage, all treatments were executed in five replicates and measured in one randomized sample list with pairwise mixtures interspersed as quality control samples (QCs). An in-house tool was built that creates a snapshot of the histone coverage—including post translational modifications (PTMs)—of both in-house and public LC-MS based histone workflows. Additionally, the tool can be used to visualize the coverage of predicted MS/MS spectra, which enabled us to create coverage plots for nine hPTMs using their curated depiction from Uniprot. This comparison between the curated histone code from Uniprot and the experimental histone workflows reveals the discrepancy between what is possible and what is feasible with the current day workflows. Indeed, the experiments show that the coverage of all workflows is incomplete but complementary and that optimization of histone sample preparation is still very much alive and relevant. We therefore provide a stepping stone to an informed decision making strategy regarding sample preparation and experimental design for bottom-up histone mass spectrometry workflows.
Materials and methods

Sample preparation

A bovine histone standard extracted from calf thymus (Sigma-Aldrich, 10223565001) was used to avoid bias towards specific regions of the histone code caused by the extraction of histone proteins from cell lines or tissue. All protocols were executed in five-fold on 20 µg of histone standard.

The ArgC and RgpB protocols—hereafter embedded in and referred to as the ArgC and GRX workflow respectively—were executed according to the instructions of the manufacturer. In brief, 5 µg ArgC (Sigma-Aldrich, 11370529001) was resuspended in 50 µL ultrapure water (18.2 MΩ·cm resistivity) of which 4 µL was added to the samples, followed by 10 µL of activation solution, and addition of digestion buffer (100 mM TrisHCL, 10 mM CaCl2) to a final volume of 100 µL. For the RgpB protocol, cysteine and dithiothreitol were added to the samples to a final concentration of 20 mM and 10 mM respectively. The samples were incubated at room temperature for 30 minutes. Next, 5 µg RgpB (Genovis, GingisREX®) was resuspended in 50 µL ultrapure water of which 4 µL was added. All samples were incubated overnight at 37 °C and vacuum dried.

The tryptic digest samples—hereafter embedded in and referred to as the NoPropTryp workflow—were resuspended in 500 mM Triethyl Ammonium Bicarbonate buffer with an aimed final volume of 50 µL, followed by addition of CaCl2 and acetonitrile to a final concentration of 1 mM and 5% (v/v) respectively. Finally trypsin was added in a 1:20 (w/w) ratio and the samples were incubated at 37 °C overnight and vacuum dried.

The tryptic digest with propionylation of the lysine residues—hereafter embedded in and referred to as the PropTryp workflow—has been described previously by Meert et al.[16] In brief, the samples were resuspended in 20 µL TEAB. First, 20 µL of an isopropyl alcohol : propionic anhydride (7:1) solution was added, followed by incubation at room temperature for 30 minutes. Next, 20 µL of ultrapure water was added, followed by incubation at 37 °C for 30 minutes to quench the reaction. The samples were vacuum dried and digested as described for the tryptic digest. After digestion, the N-termini of the resulting peptides were propionylated accordingly and the samples were vacuum dried. To reverse over-propionylation on serine, threonine, and tyrosine residues, 50 µL 0.5 M hydroxylamine and 15 µL ammonium hydroxide (pH 12) were added, followed by incubation for 20 minutes at room temperature. The samples were acidified (pH 3) with formic acid and vacuum dried.

Sample Acquisition

All samples in the sample list were randomized to avoid systematic variation, blocked to correct for drift in sensitivity of the instrument, and interspersed with quality control samples to monitor instrumental variation. Seven different quality control samples were prepared, four of which were mixtures of all separate digest conditions. In addition, a mixture of GRX and ArgC samples, a mixture of NoPropTryp, ArgC, and GRX samples and a mixture of PropTryp and NoPropTryp samples were prepared. These were used to increase alignment efficiency and quantitative accuracy during analysis with Progenesis QIP.

3 µg of each sample was LC-separated prior to MS-analysis with a Nano Acquity system coupled to an Acquity UPLC CSH-C18 analytical column (1.7 µm, 300 µm x 100 mm, Waters) using a two-step linear gradient of 50 minutes (3-30% B) and 10 minutes (30-40% B). 0.1% formic acid and 3% DMSO in water was used as solvent A and 0.1% formic acid in acetonitrile as solvent B.

All samples were acquired on a Synapt G2-Si (Waters) in HD-DDA mode. Herein fragment ions are mobility separated based on charge state, which allows the instrument to selectively sample single charged fragment ions in the ion beam that enters the TOF-tube. This significantly increases duty cycle.
and thus sensitivity for these fragment ions. This is called wideband enhancement [23]. Precursor scan time was set to 0.2 s and each scan cycle allowed for fragmentation of a maximum of 12 precursor ions with an intensity threshold of 3000, a scan time of 0.1 – 0.3 s and a switchback threshold of 100000. The mass range was set to m/z 50-5000 for both precursor and fragment scans.

Coverage Plot
Spectral libraries were generated according to the workflow of Van Puyvelde et al. [24] Briefly, a protein sequence FASTA file of all histones was in silico digested with trypsin specificity. Both double and triple charged peptide fragment spectra were predicted using MS²PIP [25], the results were written to a spectral library file (*.msp format). There was no need to predict retention times, given the intended purpose of the library.

Using an in-house script (available at https://github.com/swillems/spectral_histones), all known and verified uniprot modifications (exported from uniprot as a .txt file) were parsed into the predicted spectral libraries and exported as an *.MGF file. Coverage plots can be created from both theoretical and empirical MGF files by searching the spectrum files in Mascot Daemon and exporting the results as a *.CSV file, including the start and end point of all identified peptides. The in-house coverage plot script (available at https://github.com/swillems/coverage_plots) also requires the protein database (*.FASTA) used to search the MGF files to map the identified peptides and PTM against the protein backbone. The displayed modifications depend on the search parameters used in Mascot Daemon.

Quantitative analysis with Progenesis QIP
All samples were imported, peak picked and aligned in Progenesis QIP (Nonlinear Dynamics) for quantitative comparison. Each experiment/experimental design had its accompanying QC samples for alignment and normalization purposes (Figure 1). For identification, all MSMS spectra were exported in a single *.mgf file for searching using Mascot (Matrix Science). A dedicated instrument was configured for the search, i.e. Synapt G2-Si, which only allows 1+ fragments, because wideband enhancement was enabled during acquisition [23]. Bovine Histones and cRAP (https://www.thegpm.org/crap/) were downloaded from Uniprot on 09/12/19 (141 sequences; 43,864 residues). Search parameters were selected based on the experimental design under investigation. Progenesis 1 was searched with Acetyl (K), Butyryl (K), Citrullination (R), Crotonyl (K), Dimethyl (KR), GG (K), Methyl (KR), Phospho (ST) and Trimethyl (K) as variable modifications and Propionyl (K) and (N-Term) as fixed. Progenesis 2 and 3 were searched with Acetyl (K), Butyryl (K), Citrullination (R), Crotonyl (K), Dimethyl (KR), GG (K), Methyl (KR), Phospho (ST) and Trimethyl (K) as variable modifications with two missed cleavages. After all identifications were imported back into Progenesis QIP, all LC-MS runs were normalized against histone peptides in order to consolidate the constant protein abundance within the data. Notably, for coefficient of variance calculations, no normalization was used as this would skew the results. Here we relied on the fact that we started from an identical amount of histone sample at the beginning of each workflow.

Plots were generated in Excel using Pivot Table functionalities on the export log-transformed intensities of the features. Principal Component Analysis was done using ClustVis [26].
Results
Comparing the performance of sample preparation workflows requires a single experiment wherein samples are prepared and measured in a randomized way to avoid batch effects. This allows for an objective assessment of the histone coverage, the chemical noise and variation introduced by the workflow, the enzyme specificity and efficiency, the experimental variation and the quantitative accuracy. Here we assess these features for four different workflows (Figure 1), providing a roadmap for future workflow assessment in the process.

Figure 1: Experimental design for direct comparison of different enzymatic treatments. All replicates (n=5) were prepared in parallel and the samples were analyzed in a full factorial design sample list to avoid batch effects. The different QC mixtures allow for direct alignment between runs. The different QC mixtures highlight the dotted lines. Other combinations can be made by using the available QC samples as alignment template. Blue: GRX: Workflow based on digestion with RgpB (Genovis, GingisREX); Red: ArgC: Workflow based on digestion with clostripain (Roche, ArgC); Grey: PropTryp: Workflow based on the blocking of lysine residues with propionyl groups to obtain arginine specific cleavage during tryptic digest; NoPropTryp: Workflow based on regular tryptic digest

The Histone Code Coverage with Trypsin
A Theoretical Tryptic Histone Snapshot
To enable direct comparison of the histone coverage of different sample preparation workflows, we first created a PTM Coverage Tool in Python. To illustrate its applicability, we created an overview of detectable curated modifications of the histone backbone with the search parameters depicted in figure 2. Coordinately, this also assesses the efficiency of the search strategy itself. Therefore, an unmodified spectral library was predicted using MS²PIP [27] and all delta masses of curated histone modifications from Uniprot were parsed into the spectral library at their corresponding location. Note that these noiseless predicted spectra generate a considerable amount of ambiguity above the significance threshold when searched by Mascot using the set of variable PTMs shown. This is due to localization errors and unconsidered PTMs present in the MGF file that are explained through other PTM combinations or point mutations [28]. Therefore, all searches on empirically measured histone peptidoforms below should be considered in light of this first limitation. Despite this limitation, Figure 2 shows that a considerable part of the histone code can be covered theoretically using a workflow with tryptic cleavage.

Taken together, the tool presents the researcher with an end-point report of the full workflow, including the search. The yellow regions highlight where many different peptidoforms are created, which translates into higher spectral counts in a predicted spectral library as shown in Figure 2.
Figure 2. Theoretical coverage of the histone code. A spectral library containing all histone peptidoforms with up to 3 tryptic missed cleavages was predicted using MS²PIP. Next, all delta masses of curated histone modifications from uniprot were parsed into this spectral library at their corresponding location, thus creating a searchable MGF file. This was searched using Mascot and the resulting CSV output, including peptide start and end positions, was used to create a histone coverage plot. This represents what is theoretically possible with a probabilistic scoring algorithm, given the set of variable modifications depicted in the Y-axes. The scale to the right of each plot indicates the spectral count (relative within each histone from red to yellow) of each modification/amino acid residue.

In search of a suitable protocol, the tool can also be used to visualize the detected PTMs in public data, with the current limitation that results need to be parsed in the Mascot *.csv format. As an example, Supplementary Figure 1 depicts the PTMs annotated using Progenesis QIP and Mascot on the AQUA heavy peptide standard as described in Li et al. [15]. In doing so, the list of peptides in the AQUA mixture was converted to one image depicting the detected PTMs with the given search parameters.

Theory Versus Practice

Figure 3 depicts the coverage plot of an empirical workflow using trypsin. Five independent tryptic digests of a commercial bovine histone extract were performed and analyzed using LC-MS/MS, interspersed with QC samples (Figure 1). This provides a very different picture compared to the predicted coverage plot. Indeed, only a fraction of the possible peptidoforms are identified.

This is primarily due to the fact that only a small subset of all possible proteoforms are actually present in the sample, i.e. a calf thymus histone extract. However, equally important is the fact that certain peptidoforms, which are present in the sample, are excluded from annotation because of a number of other reasons: (i) the enzyme could have created unconsidered aspecific cleavages, or these could have been present in the sample to begin with [29], (ii) in vitro induced PTMs like deamidation, oxidation, and formylation were not considered in the search, (iii) some peptidoforms have no retention on the LC column or lie outside the mass range acquired by the instrument, (iv) singly charged precursor peptides were not targeted by the instrument in data-dependent acquisition (DDA) mode, (v) some peptidoforms are too short to generate enough specificity for the probabilistic scoring algorithm to
reach the significance threshold, (vi) gas phase chemistry artefacts (like neutral losses, cation adducts and in source decay) were not considered in the search, ...

Figure 3. Histone coverage of a tryptic digestion. To assess the coverage of an actual experiment, five replicate tryptic digests were merged and searched allowing four missed cleavages and the 9 PTMs depicted in the theoretical search from Figure 1.

Overall it is clear that the epicenters of sequence coverage have shifted considerably, most strikingly so for H3. Still, the core and C-termini, including those of H2A, H2B and H1 are being sampled quite efficiently. This indicates that, although tryptic digest is usually not preferred for histone epigenetic research, it can be of value for certain regions in the histone code, especially when considering workload. Therefore trypsin should always be the first choice if the coverage of the workflow suits the research question. Indeed, numerous histone related discoveries have been described using this workflow. One such example was the identification of H2A-specific protease as being neutrophil elastase, capable of cleaving H2A at valine 114, i.e. 5 amino acids upstream the essential PRC1-mediated K119 ubiquitination [29–31].

Exploring Other Options
This lack of H3 and H4 N-tail coverage in bottom-up histone research is most commonly addressed by blocking tryptic digest at the lysine cleavage site by derivatization with e.g. propionic anhydride which introduces propionyl groups [15]. This results in larger and more hydrophobic peptidoforms, more suitable for LC-MS/MS analysis of those sequence stretches (Figure 4). Since these regions are densely covered with PTMs, this protocol has been the preferred approach for bottom-up histone analysis for nearly 15 years. However, the core and C-termini of most histones are less well/accurately sampled because of the increased peptide length. This not only impairs LC-separation for these peptides, but also hampers accurate identification and localization of PTM combinations with current search algorithms. Due to these erroneous annotations of isobaric combinations, two intensely sampled
regions become overreported with PTMs: histone H2B 1-29 and H1 1-32 [28]. These suspicious cases can only be detected by manually verifying the sequence coverage (Supplementary Figure 2). Overall, caution is required with longer peptide stretches, especially when combinatorial PTMs are considered.

Figure 4. Coverage plot of a tryptic digest following propionylation. (A) When histone lysines and N-termini are derivatized with e.g. propionic anhydride the lysine cleavage sites are blocked with propionyl groups and the coverage of the histone code shifts considerably, with much improved sampling of the H3 and H4 N-tails. (B) Visualization of chemical noise introduced on histone H3 through derivatization. Several non-biological modifications are introduced during sample preparation. The red box highlights different PTMs that are (potentially) changed by the PropTryp workflow. These can no longer be interpreted in a biological context and increase ambiguity and false discovery rate.

Unfortunately, chemical derivatization also introduces chemical noise, which interferes with the quantification and identification of important PTMs [16–18,32]. Figure 4B shows the coverage plot of Histone H3 when searched with a selection of PTMs that includes in vitro introduced PTMs. Indeed, when propionylation is combined with trypsin, propionyl groups of biological origin can no longer be distinguished and methylated lysines are propionylated to butyryl, thus becoming indistinguishable.
The protocol can also introduce overpropionylation (S, T, Y) and formylation as a side reaction, apart from amidations not displayed here [16–20]. This compromises the correct biological interpretation of all these PTMs. While many solutions have been proposed [16–18,32], including more controlled reaction circumstances and the use of heavy labeled or non-biological reagents, entirely excluding side reaction will probably proof to be impossible. This was recently illustrated by the finding that formylation of serine and threonine can even be induced by 0.1% formic acid in the LC buffer [33]. Fortunately, these side reactions do not interfere with relative abundance of single PTMs [18]. Still, they dilute the signal, creating uninformative precursors that take up acquisition time of an instrument that is acquiring in DDA [16] and aggravate the already complex issue of ambiguity in annotation [28].

Comparing three different enzymatic treatments with ArgC specificity

Adjusting the protocol to reduce side reactions and chemical noise

Theoretically, ArgC (Clostripain) generates similar-sized peptides compared to PropTryp without the disadvantages of chemical derivatization. However, the lack of specificity of the enzyme has been known for years [14,34]. Therefore, we also validated another arginine-specific enzyme, i.e. RgpB, also known as Gingipain or GingisREX (Genovis). This enzyme was first described in 1992 as an important virulence factor of Porphyromonas gingivalis and later praised for its cleavage specificity and efficiency at the c-terminal end of arginine [22,35,36]. However, it was only recently commercialized for proteomics applications. As expected, considerably less chemical noise was introduced in either of these workflows (Supplementary Figures 3-6), while methylation, propionylation, and butyrylation remain useful for biological interpretation on all histones. Surprisingly, however, the sequence coverage of both enzymes shifts dramatically compared to PropTryp, as exemplified in Figure 5, which shows histone H3 for all the considered workflows as an example. Coverage plots of biological PTMs detected on other histones in each workflow are shown in Supplementary Figure 7-10.

Figure 5. Direct comparison of the coverage of the histone H3 PTM coverage using the different protocols in this study. From top to bottom: Pred: theoretically predicted tryptic spectral library of H3 searched with the PTMs depicted in the Y-axis; Tryp: empirically digested and measured H3; PropTryp: tryptic digest of H3 following derivatization with propionic acid.
anhydride (arrowheads indicate PTMs impacted by the derivatization); GRX: empirically digested GingesRex enzyme; ArgC: empirically digested with ArgC enzyme. The latter three were all searched with ArgC as enzyme specificity.

Feature detectability
In search of the peptide characteristics that contribute to the shifting detectability, we moved beyond the PTM coverage tool. The experimental design (Figure 1) allows direct comparison of different treatments. Project Progenesis PropTryp contained five PropTryp replicates and five QC injections. ArgC and GRX replicates were aligned directly in project Progenesis GRX_ArgC using a mixture of both treatments as alignment template and quality control (QC). Project Progenesis GRX contained the GRX runs and their QCs only, to study this enzyme in isolation.

A total of 279 histone ions with the expected properties were annotated in the mixed project Progenesis 2, i.e. features derived from ArgC specific peptides without missed cleavages. 47 of these features were annotated through charge state deconvolution, a feature in Progenesis QIP that also allows to annotate singly charged features that were not selected for fragmentation during DDA. To this end, the annotation of the doubly charged precursor is transferred by the software to the co-eluting, singly charged precursor ion. For PropTryp 302 identifications were imported into project Progenesis 1, of which 79 were found through charge state deconvolution.

Despite the presumed identical cleavage specificity, the three workflows show clear discrepancies. This is unsurprising for PropTryp because of the chemical derivatization. The two-dimensional LCMS representation of the three protocols (Supplementary figure 1A-C), mainly surfaces the increased retention of the propionylation reaction, whereby the different peptidoforms of the H3 and H4 N-tails are more efficiently separated and sampled, as described earlier [15].

Figure 6. In-depth analysis of three different enzymatic treatments with Arg-C specificity. (A) The Principle Component Analysis of the feature abundances from the Progenesis GRX_ArgC project containing ArgC (red) and GRX (blue) shows a clear discrepancy between both workflows despite their identical cleavage specificity. The QC (green) is a mixture of both measured interspersed throughout the sample list. (B) This double-sided boxplot shows the inverse pattern of MS1 peaks (orange, right axis) and MS2 spectra (blue, left axis). PropTryp has the least MS2 spectra while it has the most MS1 precursors. Grey line indicates the separate Progenesis PropTryp project. (C) Charge state distribution of the MS1 precursors from (B) for the three workflows, showing a higher proportion of singly charged precursors in PropTryp, which will not be selected by the instrument for fragmentation in DDA mode. (D) Summed MS1 signal of all identified peptidoforms showing that the highest portion of the generated ions is being annotated in GRX, while all treatments were done on the same starting amount of bovine histones. (E) The histone H3K9-R17 peptide stretch XICs show that the singly charged (up) ion species comprises double the amount of...
ions compared to the doubly charged (down) counterpart. Intensities are depicted in the upper left corner, retention time is shown in the X-axis. (f) MS1 signal intensity for each peptide sequence annotated in the different treatments showing the complementarity between PropTryp and GRX.

However, GRX and ArgC also differ considerably. This is equally reflected in the principal component analysis (PCA) of the feature abundances of the direct align-between-runs in project Progenesis GRX_ArgC (Figure 6A). An even more striking difference from the acquisition point of view is that the number of different MS1 precursors of all charge states is in the order of PropTryp > ArgC > GRX, while the number of MS/MS-spectra is in the reversed order GRX > ArgC > PropTryp (Figure 6B). The QC from Progenesis GRX_ArgC illustrates that this is not an instrumental effect, as physically mixing ArgC and GRX results in intermediate values. At least in part this turns out to be a consequence of the charge state distribution of the precursors. Figure 6C shows that nearly 50% of the PropTryp ions is singly charged as opposed to only 20% for the other two enzymes. This is likely caused by the charge blocking effect of the propionyl group and will exclude these precursor ions from fragmentation during regular DDA acquisition.

Indeed, the reduced sampling is also reflected in the summed MS1 signal of all identified peptidoforms (Figure 6D). Supplementary Figure 11D displays the charge state distribution of these annotated ions, confirming the overall pattern of all features depicted in Figure 6C. For GRX, the singly charged ions annotated by deconvolution represent only 10%, as opposed to 40% for PropTryp. At the precursor ion level, the implications are further highlighted. Figure 6E illustrates how e.g. the H3K9-R17 peptide stretch is about 60% singly charged on a SynaptG2Si micro flow source with 3% DMSO in the aqueous buffer. This implies that 2/3 of the signal cannot be annotated and is usually not used for quantification for this peptide.

In conclusion, the differences in retention and charge state distribution together make that GRX and PropTryp cover complementary sequence stretches, as visualized in Figure 6F.

**Enzyme specificity and efficiency**

On top of the increased retention of the histone peptides, the propionylation protocol is also preferred because it uses trypsin as proteolytic enzyme, which assures high efficiency and specificity. To assess the enzyme specificity of ArgC and GRX, the same data was searched with semi-specific cleavage. This results in a total of 1135 different annotated histone features, 142 of which by charge state deconvolution. Supplementary Data 12 shows that this indeed provides a better histone coverage for ArgC.

However, there is an important caveat; these semi-specific peptides cannot readily be used for biological interpretation, as adding them to the dataset would significantly alter normalization, which would in turn impact the relative abundance calculations of individual PTMs. Indeed, when based on all detected features, the calculated log normalization factors between the enzymes are close to zero for all runs. This is an intuitive consequence of the fact that all workflows were executed on an identical amount of commercial bovine histones (Figure 7A). However, Figure 6D already showed that for the identified desired features with specific cleavage, only half the signal was found in ArgC compared to GRX. This implies that, using only the correctly cleaved ions to normalize the data, results in large normalization differences between ArgC and GRX runs (Figure 7A). Normalizing against semi-ArgC specific annotated peptides reduces this gap but does not remove it completely. Thus, ArgC proportionally has more identified aspecific peptides, while the rest of the gap can be explained by further degradation of the sample into peptides that can no longer be annotated by the applied searches.
Figure 7. In-depth Enzyme specificity and efficiency. (A) Calculated logarithmic normalization factors based on all features (left), annotated peptidoforms searched with aspecific cleavage (middle) and annotated peptidoforms searched with specific cleavage (right). (B) Signal intensity (y-axis) of identified peptidoforms with up to two missed cleavages which are indicative of reduced enzyme efficiency. (C) Distribution of peptide length for all workflows.

To assess enzyme efficiency of ArgC and GRX, the data was searched with up to 2 missed cleavages. In total, 83 (7%) additional features were identified, most of which more abundant in ArgC, implying that this enzyme is also less efficient (Figure 7B). However, while only 8% (98/1246) of all annotated ions had a citrullination on arginine, this fraction became 35% (29/83) for the peptides that contained missed cleavages. This also holds for the 3.6% (45/1246) arginine methylations in the total population that is enriched to 23% (19/83) in the missed cleaved population. This surfaces an alternative conclusion, i.e. that in fact GRX is capable of cleaving modified arginines and ArgC is not.

Taken together, because of its comparable efficiency and higher specificity GRX generates larger peptides compared to ArgC (Figure 7C). However, more surprisingly, the detected peptides in GRX are also considerably longer than what is detected with PropTryp. This is mainly due to the increased retention induced by propionylation. More specifically, where this increased retention is a benefit for the hydrophilic short and mid-range peptidoforms of H3 and H4 N-tails, it makes resolving the longer peptides very challenging. The result is that GRX covers overall a larger size range of histone peptides, but at the cost of resolution or even retention of the densely modified H3 and H4 N-tail peptidoforms. This in turn urges for alternative fragmentation and annotation strategies that are akin to middle down approaches. Indeed, longer peptide stretches as depicted in Supplementary Figure 11E tend to only fray on the ends and do not fragment towards the middle when applying collision induced dissociation (CID). We are therefore developing an adjusted GRX workflow, including phenyl iso-cyanate derivatization and electron transfer dissociation (ETD), which should respectively increase retention of smaller peptides and fragmentation of larger peptides.

As a concluding overview, Figure 8 shows the LC-MS feature map of the PropTryp and GRX enzymatic treatments with the peptide charge states color coded. Supplementary Data 13 and 14 highlight the localizations of the most prominent sequence peptidoforms identified in PropTryp and GRX, respectively, providing a true roadmap to these two enzymatic treatments.
**Figure 8.** LCMS 2D representation of eluting features of PropTryp and GRX. The Y-axis depicts the retention time from top to bottom as a function of MS1 scan rate (not linear), the X-axis represents m/z. Features are color coded according to their charge state. The dotted line in GRX depicts the point beyond which peptides with propionyl groups become too hydrophobic to be measured with the LC-setup applied in this study. This illustrates that a set of longer and higher charged peptides are better detectable with GRX.

**Workflow Variability**

Workflow variability—expressed as % covariance (%CV)—is one of the most important metrics to assess during validation of any sample preparation workflow, as it greatly impacts quantification and the required number of replicates in the experimental design [37]. Minimizing this variability becomes even more important when it comes to detecting small biological changes, as is often the case for histone PTMs. It is assumed that longer workflows usually introduce more variation. However, figure 9A shows that the total %CV for each workflow is comparable, despite the additional derivatization steps in the PropTryp workflow. We therefore isolated the instrumental %CV by calculating the %CV on all precursor ions in the QC samples of each treatment (Figure 9B). Strikingly, while instrumental variation is generally assumed to be constant, the instrumental %CV of ArgC and especially GRX was higher compared to the NoPropTryp and PropTryp workflow. To avoid bias, the total %CV (light coloured graph) and instrumental %CV (dark coloured graph) as a function of precursor ion abundance was plotted (Figure 9C), which shows that instrumental variation is higher throughout the dynamic range for ArgC and GRX. This leads to the finding that the different workflows generate a comparable total %CV, but derived from different sources, i.e. sample preparation- and instrumental %CV. We hypothesize that the larger peptidoforms from the GRX workflow are prone to a higher instrumental %CV because of their broad and possibly unstable charge state distribution. In that case, blocking of these higher charges by propionylation of K residues can reduce this effect.
Figure 9. Variability assessment of the workflows. (A) Boxplot of the total % covariance (%CV) as calculated from the separately prepared and measured samples of each workflow. (B) Boxplot of the Instrumental %CV as calculated from the respective quality control (QC) data of each workflow. (C) Total %CV (light coloured graph) and instrumental %CV (dark coloured graph) as a function of precursor ion abundance for each workflow.

Conclusion

Both trypsin and GRX have a high specificity and efficiency, which makes them suited for histone- and proteomics workflows in general. This is in contrast to ArgC, whose aspecificity hampers the accurate quantification of peptides and peptidoforms. Without derivatization, trypsinization can be used to study the core and C-termini of all histones. Still, propionylation remains the preferred protocol to target the N-termini of histone H3 and H4, despite the chemical noise that is introduced by derivatization and the accompanying loss of several biological interpretations. We here also describe a strong shift in the charge-state distribution following propionylation, excluding many precursors from sampling in a DDA MS run. Surprisingly, the latter also reduces instrumental variation of the workflow, which is remarkable since this source of variation is generally assumed to be constant. GRX on the other hand can be considered as a complementary enzyme with the specific benefit of retaining all biological hPTMs and minimizing chemical noise. It is particularly suited for detecting longer peptide stretches, which in turn might require more dedicated fragmentation and annotation strategies. The lack of retention of H3 and H4 N-tail peptidoforms reduces the sampling efficiency of these biologically essential sequence stretch however. But overall, short peptides are currently lost in GRX, while long peptides are lost in PropTryp.
References


