Large-scale identification of protein histidine methylation in human cells

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ABSTRACT

Methylation can occur on histidine, lysine and arginine residues in proteins and often serves a regulatory function. Histidine methylation has recently attracted attention through the discovery of the human histidine methyltransferase enzymes SETD3 and METTL9. There are currently no methods to enrich histidine methylated peptides for mass spectrometry analysis and large-scale studies of the modification are hitherto absent. Here, we query ultra-comprehensive human proteome datasets to generate a resource of histidine methylation sites. In HeLa cells alone, we report 299 histidine methylation sites as well as 895 lysine methylation events. We use this resource to explore the frequency, localization, targeted domains, protein types and sequence requirements of histidine methylation and benchmark all analyses to methylation events on lysine and arginine. Our results demonstrate that histidine methylation is widespread in human cells and tissues and that the modification is over-represented in regions of mono-spaced histidine repeats. We also report colocalization of the modification with functionally important phosphorylation sites and disease-associated mutations to identify regions of likely regulatory and functional importance. Taken together, we here report a system level analysis of human histidine methylation and our results represent a comprehensive resource enabling targeted studies of individual histidine methylation events.

INTRODUCTION

Methylation of histidine is a post-translational modification (PTM) that was first described to occur on actin (1) and myosin (2) proteins around five decades ago. It can occur at two distinct positions denoted as 1-methyl histidine (1MeH) and 3-methyl histidine (3MeH) (here collectively referred to as Hme) (3). Despite being known to the scientific community for long it has gained far less attention than the well-studied protein methylation events on lysine and arginine (3), which are considered as key epigenetic modifications linked to chromatin compaction state and gene activity (4).

Until recently, little was known about the enzymology and significance of Hme. In 2018, SETD3 was uncovered as the first human methyltransferase enzyme targeting histidine and being responsible for the well-established methylation of Actin-H73 (5). This finding was shortly thereafter independently validated and functionally shown to modulate smooth muscle contractility (6). In addition, the human METTL18 (also known as C1orf156) enzyme represents a clear homolog to the established yeast methyltransferase Hpm1 (systematic name YIL110W) targeting a RPL3-H243 in Saccharomyces cerevisiae (7,8), but its enzymatic activity remains to be validated. Finally, human METTL9 has been shown to act as an enzyme with broad specificity generating 1MeH in motifs composed of consecutive histidine residues spaced by small amino acids (9).

Aside from the few recent protein-centric studies focusing on individual Hme events, little is known about the abundance and function of the PTM. PTMs are most frequently studied at a large scale by affinity enrichment of modified peptides, or proteins, followed by mass spectrometry analysis.

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try (MS)-based identification of targeted sites (10). Such approaches have been described for lysine methylation (Kme) (11,12) and are well-established for arginine methylation (Rme) (13). For many PTMs, including Hme, there are no established affinity reagents, creating a need for innovative approaches for characterization. In such cases, querying ultra-comprehensive proteomic datasets for mass shifts corresponding to distinct modification events has recently emerged as a promising alternative (14).

Studies dedicated to global characterization of Hme are until this date absent. To explore the PTM we here mined a panel of ultra-deep human proteome datasets (15) to generate an extensive resource of Hme sites. The analysis revealed that Hme is widespread in human cells, and uncovers its abundance, context, and function in relation to Kme and Rme. To the best of our knowledge, the present study represents the first system level analysis of Hme and is to date the most comprehensive draft of the human histidine methylome.

**MATERIALS AND METHODS**

**Querying proteomic data for methylation events**

Publicly available comprehensive proteomic datasets (ProteomeXchange id: PXD004452) previously published by Bekker-Jensen et al (15) were downloaded from ProteomeXchange (16). The datasets were chosen based on exhaustive proteome depth, obtained through extensive offline peptide fractionation using reverse phase chromatography at alkaline pH and analysis of individual fractions using fast scanning MS methods with a Q-Exactive HF mass spectrometer (15). The analyzed data correspond to LC-MS/MS analysis of tryptic peptides from human tissue biopsies from colon, liver and prostate as well as the human cell lines A549, HCT116, HEK293, HeLa, MCF7 and SY5Y. In addition, an in-depth analysis of data derived from HeLa cell sample digested with a panel of complementary proteases including trypsin, chymotrypsin, Glu-C and Lys-C was included to achieve comprehensive proteomic coverage.

All raw MS files were searched using MaxQuant (17) (version 1.6.0.171) against a database containing the canonical isoforms of human proteins (Uniprot Complete proteome: UP_2017_04/Human/UP000005640_9606.fasta) using the default software settings with few exceptions. To reduce the search space, the number of allowed missed cleavages was restricted to one. In addition to the default variable modifications, corresponding to acetylation of protein N-termini and oxidation of methionine, mono-methylation of lysine and arginine, di-methylation of lysine and arginine, tri-methylation of lysine as well as the custom generated modification mono-methylation of histidine were included as variable modifications.

**Bioinformatic analyses**

Bioinformatic analysis was performed using the Python programming language: Python Language Reference (version 3.8), available at http://www.python.org. The MaxQuant output files were processed, removing annotated contaminants. Modifications identified in one or more biological replicates containing a mono-, di- or trimethylated modification at an arginine, lysine or histidine were defined as a unique methylation site. If a methylated peptide matched to multiple protein entries, all proteins were categorized as methylated in the downstream analysis. To benchmark our identified sites, the publicly available resource PhosphoSitePlus (version 6.5.9.3) (18) was used as a reference. Protein localization data was derived from the SubCellularBarcode project (19) and the subcellular localization of proteins in the cell line MCF7 was chosen as surrogate dataset in order to infer localization of our identified methylated proteins. Complete predicted subcellular localization of all methylated proteins was also done using the computational algorithm BUSCA (20), allowing for protein assignment into 9 distinct subcellular compartments. Identification of methylations colocalizing with phosphorylation sites were achieved by searching curated phosphoproteomic dataset (21) for known phosphorylations at a distance of 10 amino acids, and the functional score of sites was defined as described in original publication. Annotated protein domains were accessed using the resource portal InterPro (version 82.0) (22) and individual methylation sites were mapped to the interior regions of protein domains annotated in the Pfam database (version 33.1) (23). Functional enrichment analysis was conducted using the String database (version 11.0) (24) and multiple testing was corrected for with the Bonferroni method for false discovery rate (FDR). Logo and enrichment analysis of amino acids flanking methylation sites were performed using the iceLogo server and the human precompiled Swiss-Prot peptide sequence composition as reference (25).

**Proteomic characterization of METTL9 knockout cells**

A HAP-1 METTL9 knockout (product number HZGHGC004343c010, Horizon Genomics) and a wild type control cell line (product number C631, Horizon Genomics) were propagated and maintained in IMDM Glutamax media (Thermo Fisher Scientific) supplemented with 10% fetal bovine serum (Thermo Fisher Scientific), as well as 100 U/ml penicillin and 100 U/ml streptomycin. The cells were lysed in a guanidine hydrochloride-based buffer and peptides were prepared for analysis using a Q Exactive HF mass spectrometer (26) as previously described (27).

The LC-MS analysis was performed using an EASY-nLC 1200 HPLC system (Thermo Fisher Scientific) coupled to a Q Exactive HF orbitrap instrument. For each single shot proteome analysis, 500 ng peptide was separated using a 3 h chromatography gradient linearly ramped from 10% to 30% buffer B (80% acetonitrile in 0.1% formic acid) in buffer A (0.1% formic acid) during 170 min, whereafter the gradient was steeply ramped to 45% buffer B during 10 minutes, before column washing and equilibration. The mass spectrometer was set to continuously sample peptides through a Top12-based data dependent acquisition method. Target values for the full scan mass spectra were set to 3e6 charges in the m/z range 375–1500 and a maximum injection time of 25 ms and a resolution of 60 000 at a m/z of 200. Fragmentation of peptides was performed using higher energy
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Figure 1. Biochemistry of protein histidine methylation. Structure of the different methylated forms of histidine. The methylations are enzymatically introduced by histidine (H) methyltransferases (HMT) and may potentially be removed by histidine (H) demethylases (HDM).

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C-trap dissociation (HCD) at a normalized collision energy of 28 eV. Fragment scans were performed at a resolution of 15 000 at a m/z 200 with a AGC target value of 1e5 and a maximum injection time of 22 ms. To avoid repeated sequencing of peptides, a dynamic exclusion window was set to 30 s.

The raw MS files were analyzed using MaxQuant (version 1.6.0.17i) with identical settings to the exploratory searches of published proteomic datasets, statistical analyses was performed using Perseus (28). First, LFQ intensities were imported from the MaxQuant output file denoted ‘protein groups’. Common contaminants, proteins only identified as modified and proteins hitting the reverse decoy database were thereafter removed by filtering. The resulting data matrix was filtered for proteins detected in at least 70% of the replicates in one experimental condition. The data was then log-transformed and missing values were imputed from the lower tail of the abundance distribution using the default setting in Perseus (28). Proteins displaying significance differences between the conditions were identified through a Student’s t-test (P < 0.05) with P-values corrected for multiple hypothesis testing using the Benjamini–Hochberg method. For cluster analysis, LFQ intensities for proteins displaying a significant difference between the conditions were z-scored and row and columns trees were generated using Euclidean distance and Pearson correlation, respectively. Gene ontology analysis of proteins over- and under-represented in METTL9 knockout cells, was performed using the embedded function in Perseus and P-values were corrected using the Benjamini–Hochberg method.

RESULTS

Histidine methylation (Figure 1) is a poorly characterized PTM which has recently attracted notable attention through the discovery of the human histidine methyltransferase enzymes SETD3 (5,6) and METTL9 (9). Large-scale Hme analysis is challenging since there are no available affinity agents to enrich peptides bearing the PTM for MS analysis. Here, we devise an alternative strategy based on mining ultra-deep human proteomic datasets (15) for the modification. This approach enables global identification of cellular Hme events and a subsequent system level analysis of the PTM.

Histidine methylation is widespread in human cells

To explore the abundance of Hme we searched ultra-comprehensive proteomic datasets derived from the commonly used human cell lines A549, HCT116, HEK293, HeLa, MCF7 and SY5Y as well as tissue biopsies from human colon, liver and prostate (15) (Figure 2A). The datasets were selected based on the expression of the recently established human HMT enzymes SETD3 and METTL9 (Supplementary Figure S1). The searches were performed using MaxQuant (17) with Hme defined as a custom PTM. To avoid misidentification of Hme sites, the PTMs mono-, di- and trimethylation of lysine (Kme1, Kme2 and Kme3; referred to as Kme) (Supplementary Figure S2A) as well as mono- and di-methylation of arginine (Rme1 and Rme2; referred to as Rme) (Supplementary Figure S2B) were defined as additional variable modifications (Figure 2A). This approach enables cellular Kme and Rme events to serve as a benchmark for the downstream Hme-centric analyses.

We anticipated that using a broad range of human cell proteome datasets would enable the identification of both general and cell specific Hme events. The exploratory searches revealed 267 and 80 Hme sites across the cell lines and tissue biopsies, respectively (Figure 2B, Supplementary Table S1). Moreover, we found several distinct Hme events in multiple cell lines and tissue biopsies (Figure 2B). This led us to define a core histidine methylome based on sites identified in 50%, or more, of the cell lines and tissue biopsies (Figure 2C). The core methylome includes two sites present in several actin variants, and actin related proteins, corresponding to ACTA1-H103 and established SETD3-target site ACTB-H73 (1,5,6) (Figure 2C and Supplementary Figure S3A). Moreover, the core methylome encompasses APEX1-H151 (Uniprot id P27695), CPT2-H369 (P23786), EXOS7-H275 (Q15024), NP1L4-H4 (Q99733) and RBM22-H183 (Q9NW64) (Figure 2C) and the Hme sites in these non-actin related proteins do not share apparent sequence homology (Supplementary Figure S3B).

A detailed inspection of tandem mass spectra for actin methylation revealed interesting features. First, fragment spectra were of high quality and unambiguously supported methylation of ACTB-H73, with high coverage of up- and down-stream b- and y-ions (Figure 2D). Tandem mass spectra from PTM bearing peptides can contain so-called immonium ions with a mass corresponding to the modified
Exploring histidine methylation

Exploring histidine methylation in human cells and tissues. (A) Methylome profiling workflow. Publicly available ultra-comprehensive proteomics datasets from human cell lines and tissue biopsies were searched for histidine, lysine and arginine methylation using MaxQuant. Identified histidine methylation events were analyzed to explore the abundance and cellular context of the PTM. (B) Clam plot representation of histidine methylations sites identified in cells and tissues. The total number of sites and the number of shared sites between cell lines (left) and tissue biopsies (right) are indicated. (C) Core histidine methylation sites. Sites were categorized as part of the core methylome if identified in more than 50% of the samples in each category. (D) Mass spectras supporting methylation of ACTB-H73. Tandem mass for a tryptic peptide covering Y69-K84 of ACTB unambiguously supporting methylation of H73. The presence of a specific immonium ion corresponding to methyl-histidine is indicated.

residue (29) and their identification may corroborate PTM events (30). Peptides containing unmodified histidine often yield a strong immonium ion with a mass of 110.0718 atomic mass units (amu) when fragmented (31). Strikingly, for the tryptic peptide covering ACTB-H73 we instead observed a clear peak at 124.0875 amu, corresponding to the mass of an Hme immonium ion (Figure 2D). Similarly, we detected an internal Hme site on the likely histidine methyltransferase METTL18 (METTL18-H154) and the corresponding fragment spectra also contained a clear Hme fingerprint immonium ion (Supplementary Figure S4). The observation of METTL18-H154 methylation is interesting as auto-methylation of methyltransferase enzymes is a well-established phenomenon, which has also previously been reported to occur with an amino acid specificity reflecting the enzymes physiological substrate (7).

In summary, the above analysis demonstrates that cellular Hme sites can be identified by querying comprehensive proteomic datasets and that the PTM is widespread in human cells and tissues.

Exploring the HeLa methylome

Intrigued by our observation that Hme is prevalent in human cells we embarked on an in-depth exploratory analysis of the PTM. For this analysis, we focused on human HeLa cells, which is arguably the most used human cell line model in experimental research. We devised the now established approach of querying publicly available comprehensive datasets for Hme. This time, the chosen datasets (15) were specific for HeLa cells and generated using a panel of complementary proteases including, chymotrypsin, Glu-C and Lys-C, aside from trypsin, allowing for unprecedented proteome depth and coverage (32).

Across the different proteases, the analysis revealed support for 2526 distinct cellular methylation events at 2241 sites (Figure 3B). The number of methylation events exceeds the number of sites since several individual arginine and lysine residues were detected with varying degrees of methylation (Supplementary Table S1). Roughly 12% of methylation events correspond to Hme (n = 299) and the modification was less prevalent (n = 299) than Kme (n = 895) and Rme (n = 1332) (Figure 3B). In line with these observations, a previous study has suggested that roughly 14% of protein methylation events occur on histidine (33).

A comparative analysis of our HeLa methylome data to publicly available PTM resources (18) suggests that the bulk of (>80%) of Hme and Kme sites are not previously characterized (Figure 3C). The fraction of novel sites was notably lower for Rme (Figure 3C), which can be expected as established workflows for affinity enrichment and MS characterization of the PTM exist (13). The large number of identified new Hme sites highlights that the generated dataset is suitable for an exploratory systems level analysis of the PTM.
To investigate whether Hme is scattered across the proteome, or frequently occurring on individual proteins, we first analyzed the number of Hme events per Hme protein. This analysis revealed that a single methylation event was identified for most (>80%) Hme proteins and that no proteins were identified with more than two Hme sites, and similar trends were observed for both Kme and Rme (Figure 3D). Next, we explored the abundance of identified methyl proteins. Interestingly, we found that Hme, Kme and Rme were all overrepresented on abundant proteins (Figure 3E) and envision two alternative explanations for this finding. First, the proteomics datasets in this study were generated using data dependent acquisition MS, an approach intrinsically biased to identify abundant peptides and PTMs (34). Alternatively, it has been suggested that certain methyltransferases have evolved specificity towards key abundant cellular proteins to modulate key cellular functions (35). Prominent examples include Kme and Rme in the core histone H3 (36), the key translational elongation factor eEF1A (37,38), and the molecular chaperone Hsp70 (39-41) as well as Hme in actin (1,5,6), targets that were all identified and validated in this study (Supplementary Table S1).

Next, we investigated the subcellular localization of methyl proteins. To this end, we used both experimental data based on the SubCellBarCode resource (19) as well as the predicted localization based on the BUSCA approach (20). Both methods place Hme, Kme and Rme proteins in the cytoplasm, nucleus, mitochondria and in secretory compartments, at comparable frequencies (Figure 3F and Supplementary Figure S5), suggesting that Hme, Kme and Rme are all widespread across cellular structures.
In summary, the above indicates that Hme is prevalent and present in all major cellular compartments.

Domains and motifs targeted by Hme

Intrigued by the finding of Hme events across different cellular compartments, we next used the Pfam database to explore targeted proteins families and domains. Although Hme, Kme and Rme proteins displayed similar subcellular localization profiles (Figure 3F) they are clearly enriched in different Pfam entries (Figure 4A). Reassuringly, the most strongly enriched Pfam entries include to domains where the methylations are well established to exert important function including Actin for Hme (5,6), RNA recognition for Rme (13) and core histone proteins for Kme (4) (Figure 4A). Notably, Hme was found overrepresented in Pfam entries associated with zinc binding properties (E3 Ligase, CCCH-zinc finger; Zinc finger C2H2 type; Zinc finger CCCH type; Zinc finger C-x-C-x-C type (and similar); ZIP zinc transporter) (Figure 4A and Supplementary Figure S6).

Having established that Hme is over-represented in certain zinc binding proteins and domains, we next set out to analyze the sequence context flanking Hme sites using the iceLogo approach (25). We queried the five flanking amino acids for all detected Hme, Kme and Rme sites using the human precompiled Swiss-Prot peptide sequence composition as reference. This analysis revealed distinct sequence preferences for the different methylations (Figure 4B–D). Hme was overrepresented in mono-spaced repetitive histidine (H) sequences (Figure 4B), Rme in glycine (G) rich regions (Figure 4C), and Kme in the context of the acidic residues aspartate (D) and glutamate (E) (Figure 4D).

In summary, the above results demonstrate that Hme is over-represented in specific classes of zinc binding proteins and in the primary sequence context of consecutive mono-spaced histidine residues.

Co-occurrence of methylation with phosphorylation and disease associated mutations

It has been reported that methylation events can cross-talk with other PTM types such as phosphorylation, constructing a regulatory circuit known as methyl-phospho switch (43). For example, the lysine methyltransferase SET7 has been shown to regulate the stability of DNA methyltransferase-1 (DNMT1) (44), a key enzyme in maintaining methylation patterns after DNA replication. DNMT1 is methylated at Lys142 and the adjacent Ser143 can be phosphorylated by AKT1 kinase. These two PTMs
are mutually exclusive and in absence of phosphorylation at Ser143, DNMT1 is methylated at Lys142. The consequence of the methylation is an overall decrease in abundance of the key epigenetic regulator DNMT1. In order to pinpoint specific Hme events of high functional importance we integrated a highly curated phosphoproteomic dataset into our analysis (21). We searched for phosphorylations on serine, threonine or tyrosine within a distance of 10 amino acids from a methylation event and this analysis revealed 1999 co-localizing phosphorylation sites (Supplemental Table S2). A distinct advantage of the curated phosphoproteomic dataset was that it had been evaluated using a novel machine learning model that integrated multiple features relating to conservation and structural properties of the phosphorylation site, that are indicative of functional relevance, thus providing a functional score. This allowed us to evaluate our identified methylation events based on the functional score of the phosphorylation site in close proximity (Figure 5A). We found that Hme, in addition to Kme and Rme, co-localized with phosphorylation sites with an above median functional score (Figure 5A).

In addition, we queried a publicly available database (45) for mutations in amino acid positions undergoing a methylation event in order to identify sites linked to pathological conditions (46). This analysis uncovered mutations of 212 methylation sites that are co-localizing with phosphorylations (Supplemental Table S3). Notably, the functional score of phosphorylation sites colocalizing with methylation site mutations also had an above the median functional score (Figure 5A).

Our analysis suggest that protein methylation co-localizes with functionally important phosphorylation sites, suggesting crosstalk between the PTMs. However, whether crosstalk has a relevance in relation to regulation of cellular function needs to be further experimentally validated for each individual protein.
Identification of a conserved PTM hotspot in actin

Intrigued by the observations that the ACTA1-H75 methylation site colocalizes with the functional high-scoring ACTA1-Y71 phosphorylation site and that it is mutated in severe nemaline myopathy (47), we decided to do an in depth analysis of the region. Structurally, ACTA1-H75 is located in a loop which has been reported to sense nucleotide binding (48), as shown in Figure 5B. The loop is perfectly conserved between humans, fly, plant, worm and yeast, emphasizing its functional importance (Figure 5C). To explore whether the ATP-sensing loop is targeted by additional PTMs, we again queried the PhosphositePlus database. This analysis revealed five annotated PTMs in the H75-containing nucleotide sensing loop and its flanking residues (Figure 5D), corresponding to mono-methylation, ubiquitination, and acetylation of K70 as well as phosphorylation of Y71 and T79. Notably, the bulk of these PTMs have also been observed in mouse actin (Figure 5D), indicating that modification of this loop is evolutionarily conserved.

In summary, we observed disease associated mutations and PTMs at, and in proximity to, ACTA1-H75, a Hme site previously categorized as belonging to the core Hmeome. The findings emphasize the functional importance of the ATP-sensing loop harboring H75 and support a model where multiple PTMs may play a role in actin regulation.

Cellular effects of METTL9 KO

Having uncovered hundreds of cellular Hme events and explored the subcellular distribution and context of the PTM, we sought to also investigate its direct biochemical functions. Human METTL9 was very recently described as a histidine methyltransferase introducing the bulk of 1MeH in mammalian proteomes through methylation of histidine in the context of a HxH motif, where ‘x’ is a preferentially small amino acid such as alanine, glycine or serine (9). The reported specificity of METTL9 corresponds perfectly to our observed over-represented motif for Hme (Figure 4B), highlighting METTL9 gene targeted cells as a suitable model for studies of cellular Hme loss.

To uncover cellular processes regulated by Hme, we therefore obtained a CRISPR-mediated METTL9 knockout (KO) HAP-1 cell line (49) and characterized its steady-state proteome, compared to a wild type (WT) control (Figure 6A). Cellular proteins were extracted and quantified using a label free MS approach (Figure 6A). Principal component analysis of protein intensities revealed a clear separa-
ation of WT and KO cells in the first component (Figure 6B), indicating that the difference between the cellular conditions exceeds the technical experimental variation. Accordingly, hierarchical cluster analysis of proteins with a significant difference (Student’s t-test, P-adj < 0.05) in abundance between the conditions revealed two distinct clusters of over- and under-represented proteins in the KO cells (Figure 6C). To obtain insights into the processes and functions affected in the METTL9 KO cells we performed gene ontology enrichment analyses of the clusters. This analysis revealed vesicle and vesicle-related processes as under-represented and nuclear nucleic acid-associated processes as over-represented in the KO cells (Figure 6D). This suggests a complex and pleiotropic phenotype caused by METTL9 deletion and loss of pervasive Hme1.

**DISCUSSION**

We here report the first large-scale analysis of Hme and demonstrate that the PTM is widespread in human cells and tissues. Our analyses indicate that the PTM is present in all major cellular compartments and that it is overrepresented in specific protein families, in particular in actin and in zinc-binding proteins. Taken together, we present the hitherto most extensive resource on cellular Hme events and perform the first system level analysis of the PTM.

Global PTM analysis almost invariably involves PTM affinity enrichment followed by MS analysis. Such approaches have contributed greatly to the knowledge on phosphorylation (50,51), acetylation (52), ubiquitination (53), SUMOylation (54) and Rme (13) but it has been more challenging to generate robust affinity agents for proteomics characterization of Kme, and to the best of our knowledge, has not yet been tried for Hme. To study Hme, we here deployed an alternative brute-force approach, taking advantage of the high throughput of modern mass spectrometers, and queried ultra-comprehensive proteomic datasets for the PTM. To provide a benchmark for identified Hme events, we also searched the datasets for Rme and Kme. The strength of our approach can be highlighted by comparing our identified Kme-ome to the current state-of-the-art Kme proteomics studies. We identified 895 Kme events in HeLa cells alone (Figure 2B). For comparison, a study Cao et al (12) using lab-specific IgGs for all Kme states identified 552 Kme events in HeLa cells by another acknowledged study by Moore et al (11) using the bispecific Kme1 and Kme2 binding 3xMBT domain revealed 31 Kme events in 293T cells.

A drawback with our approach is the extensive laboratory work associated with off-line fractionation of peptides before MS analysis required to generate ultra-comprehensive proteomic datasets. Moreover, searching comprehensive datasets for several variable PTMs is both computationally challenging and time consuming. The establishment of Hme affinity enrichment workflows for MS would reduce the requirements for MS analysis time and data processing, and drastically reduce the efforts and costs associated with our Hme-omic approach. Thus, we foresee efforts will be made towards the generation of Hme-specific affinity agents and their optimization for Hme-proteomic applications. The affinity agents can be generated through classical animal immunizations but this approach is inherently associated with low reproducibility (55). A more robust and reproducible approach would be to generate recombinant Hme-specific antibodies using phage-displayed recombinant antibody libraries, an approach proven feasible for the PTM sulpho-tyrosine (56).

Affinity enrichment followed by MS is arguably the most widely applied approach to study PTMs. A prominent example of this is for phosphorylations, where affinity agents that specifically bind to the modified functional phosphate group exist (51). These rely on immobilized metal cations with affinity for the negatively charged phosphate group and can consequently be used for enrichment of phosphorylated serine, threonine and tyrosine (10). For protein methylations, which are small and chemically subtle PTMs, there are no such affinity agents available. Antibodies and specific methyl-state binding protein domains have instead been used to enrich Kme and Rme modified peptides for MS (11–13). However, antibodies and domains often display a preference for PTMs in specific contexts. For example, several Rme antibodies have a preference for flanking residues (57) and the affinity of Kme-binding domains can be affected by neighboring PTMs, such as phosphorylation (58). Although the depth of the histidine methylene may conceivably be increased beyond this study through future affinity enrichment-based approaches, the herein identified Hme events, and the subsequent bioinformatic analysis, is not skewed or biased through context-specific affinity agents.

Our proteomic characterization of METTL9 KO cells suggests a complex phenotype with perturbations of both vesicle-associated and nuclear-linked cellular process. A recent study suggested the presence of as many as 2807 candidate METTL9 target sites (HxH, x = A, N, G, S or T) in the human proteome (9). Our observed pleiotropic phenotype for METTL9 KO cells may be linked to the large number of plausible substrates for the enzyme, which will likely be subjects for future studies. The large number of substrates for METTL9 also render our METTL9 targeted cells a poor tool to study biochemical functions of individual Hme events. Instead, we suggest ectopic expression of WT and HxH-mutated METTL9 substrates in cells as a preferred approach.

Actin proteins are subject to a wide range of PTMs, many of which have determined regulatory function (59). One prominent example is SETD3 mediated methylation of ACTB-H73, which corresponds to ACTA1-H75, which modulates actin dynamics by accelerating the assembly of actin filaments, a process preventing primary dystocia (6). Another example, is a unique multi-step N-terminal processing and modification machinery involving N-terminal acetylation of the initiating methionine (iMet), followed by excision of iMet and subsequent acetylation of the residue in position 2 (60). Our integrated analysis of Hme colocalization with machine learning predicted functional importance scored phosphorylation events (Figure 5A) uncovered ACTA1-H75 as colocalizing with ACTA1-Y71 phosphorylation, a site with a high functional score (0.62) (59). Interestingly, we observed five additional PTMs in the six up- and downstream residues, of these four had also been reported to occur in mouse (Figure 5D). To the best of our knowledge, so far only H75 methylation has been the sub-
ject of detailed biochemical studies. The multiple modifications within the loop reassembles the numerous PTMs in the flexible tail of histone H3, a key component of the histone code determining chromatin compaction state and gene activity. Given the extent of PTMs in the functionally important ATP-sensing loop, a similar ‘actin code’ may exist, where multiple dynamic PTMs collectively, or individually, determine the molecular functions of actin.

The protein methylation dataset we have generated may support further studies on Hme and we envision three direct applications. First, synthetic peptides corresponding to the relatively small Hm.a Hme-ome (n = 299) can be generated and evaluated as substrates for new candidate histidine methyltransferase enzymes. The human genome encodes over 200 predicted methyltransferases (61) and given the abundance and sequence diversity of identified Hme events (Supplementary Table S1), a considerable fraction of these may catalyze Hme. Candidate histidine methyltransferases may be cloned, expressed, and isolated from bacterial systems, and evaluated for activity on peptide arrays comprising the Hme-ome. Second, a synthetic peptide library corresponding to the Hme-ome may be used to uncover Hme-driven protein interactions for yet not discovered Hme reader proteins and Hme demethylases through affinity-enrichment MS approaches (27,62). Third, our resource provides the necessary information to design large-scale targeted MS methods (63) for Hme that can be used to further explore the regulation and variation of Hme-ome in human cells, tissues, and biological fluids.

In summary, our data extends the current knowledge of Hme and the study represents the first system level analysis of the PTM. Finally, we encourage the research community to use this resource for large-scale targeted MS and detailed biochemical studies of individual sites to shed further light on the emerging field of protein methylation biology.

Note added in proof: While this manuscript was revised it was confirmed that human METTL18 is a histidine methyltransferase targeting RPL3-His245 (64).

**DATA AVAILABILITY**

The comprehensive proteomic datasets for human cells and tissues queried in this study are available at ProteomeXchange with the dataset identifier PXD004452. The MaxQuant output data files from our analysis as well as the code used to generate Figure 2B-C, Figure 3B-F, Figure 4A and Figure 5A as well as Supplementary Figure S5 and Supplementary Figure S6 are available at Zenodo.com (https://zenodo.org/record/4462883#.YBAItKehGgu; DOI 10.5281/zenodo.4462883).

The MS proteomics data for HAP-1 wild type and METTL9 knockout cells have been deposited to the ProteomeXchange Consortium via the PRIDE (16,65) partner repository with the dataset identifier PXD023075. The entry also contains Perseus session files used for generation of Figure 6B-D as well as Supplementary Figure S1.

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