Deep, Quantitative Coverage of the Lysine Acetylome Using Novel Anti-acetyl-lysine Antibodies and an Optimized Proteomic Workflow* [5]

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Introduction of antibodies specific for acetylated lysine has significantly improved the detection of endogenous acetylation sites by mass spectrometry. Here, we describe a new, commercially available mixture of anti-lysine acetylation (Kac) antibodies and show its utility for in-depth profiling of the acetylome. Specifically, seven complementary monoclones with high specificity for Kac were combined into a final anti-Kac reagent which results in at least a twofold increase in identification of Kac peptides over a commonly used Kac antibody. We outline optimal antibody usage conditions, effective offline basic reversed phase separation, and use of state-of-the-art LC-MS technology for achieving unprecedented coverage of the acetylome. The methods were applied to quantify acetylation sites in suberoylanilide hydroxamic acid-treated Jurkat cells. Over 10,000 Kac peptides were quantified from a single stable isotope labeling by amino acids in cell culture labeled sample using 7.5 mg of peptide input per state. This constitutes the deepest coverage of acetylation sites in quantitative experiments obtained to-date. The approach was also applied to breast tumor xenograft samples using isobaric mass tag labeling of peptides (iTRAQ4, TMT6 and TMT10-plex reagents) for quantification. Greater than 6700 Kac peptides from over 3000 Kac proteins were quantified from a single stable isotope labeling by amino acids in cell culture labeled sample using 7.5 mg of peptide input per state. This constitutes the deepest coverage of acetylation sites in quantitative experiments obtained to-date. The approach was also applied to breast tumor xenograft samples using isobaric mass tag labeling of peptides (iTRAQ4, TMT6 and TMT10-plex reagents) for quantification. Greater than 6700 Kac peptides from over 3000 Kac proteins were quantified from a single stable isotope labeling by amino acids in cell culture labeled sample using 7.5 mg of peptide input per state. This constitutes the deepest coverage of acetylation sites in quantitative experiments obtained to-date.

Lysine acetylation (Kac)1 is a well conserved, reversible post-translational modification (PTM) involved in multiple cellular processes (1). Acetylation is regulated by two classes of enzymes: lysine acetyltransferases (KATs) and histone deacetylases (HDACs) (2–4). This modification was originally identified as a nuclear event on histone proteins and has been long appreciated for its role in epigenetic and DNA-dependent processes. With the help of a growing number of large-scale acetylation studies, it has become evident that lysine acetylation is ubiquitous, also occurring on cytoplasmic and mitochondrial proteins and has a role in signaling, metabolism, and immunity (1, 4–6). Therefore, the examination of lysine acetylation on nonhistone proteins has gained a prominent role in PTM analysis.

To date, the identification of large numbers of acetylation sites has been challenging because of the substoichiometric nature of this modification (7, 8). Additionally, global acetyla-

1 The abbreviations used are: Kac, lysine acetylation; PTM, post-translational modification; KATs, lysine acetyltransferases; HDACs, histone deacetylases; SILAC, stable isotope labeling by amino acids in cell culture; ELISA, enzyme-linked immunosorbent assay; KLH, keyhole limpet hemocyanin; HEPES, 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid; DTT, dithiothreitol; TFA, trifluoroacetic acid; DMSO, dimethyl sulfoxide; BSA, bovine serum albumin; MeCN, acetonitrile; EDTA, ethylenediaminetetraacetic acid; PMSF, phenylmethylsulfonyl fluoride; BCA, bicinchoninic acid; RT, room temperature; bRP, basic reversed phase; HPLC, high pressure liquid chromatography; MeCN, acetonitrile; IAP, immunooaffinity purification; FA, formic acid; iTRAQ, isobaric tags for relative and absolute quantitation; TMT, tandem mass tags; IMAC, immobilized metal affinity chromatography; Gly, glycine; Asp, aspartic acid; Glu, glutamic acid; K-ε-GG, di-glycine remnant.
tion is generally less abundant than phosphorylation and ubiquitylation (1). The introduction of antibodies specific for lysine acetylation has significantly improved the ability to enrich and identify thousands of sites (9–14). A landmark study by Choudhary et al. used anti-Kac antibodies to globally map 3600 lysine acetylation sites on 1750 proteins, thereby demonstrating the feasibility of profiling the acetylome (10). A more recent study by Lundby et al. investigated the function and distribution of acetylation sites in 16 different rat tissues, and identified, in aggregate, 15,474 acetylation sites from 4541 proteins (12).

Although anti-acetyl lysine antibodies have been a breakthrough for globally mapping acetylation sites from a single sample, as is now routinely possible for phosphorylation and ubiquitylation (13, 15–18). To improve the depth-of-coverage in acetylation profiling experiments there is a clear need for (1) alternative anti-acetyl lysine antibodies with higher specificity, (2) optimized antibody usage parameters, and (3) robust proteomic workflows that permit low to moderate protein input. In this study, we describe a newly commercialized mixture of anti-Kac antibodies and detail a complete proteomic workflow for achieving unprecedented coverage of the acetylome from a single stable isotope labeling by amino acids in cell culture (SILAC) labeled sample as well as isobaric tags for relative and absolute quantitation (iTRAQ)- and tandem mass tag (TMT)-labeled samples.

EXPERIMENTAL PROCEDURES

Anti-Acetyl Lysine Antibody Development and ELISA Analysis—Two degenerate peptide libraries were used to immunize New Zealand White rabbits (library one: XXXXKacXXXXXX for immunizing eight rabbits; library two: XXXKacXXXXXKacXXXXXX for immunizing four rabbits, where X represents a position in the peptide synthesis where a mixture of all 20 amino acids was used except for tryptophan, cysteine, and tyrosine). The peptide libraries were conjugated to keyhole limpet hemocyanin (KLH) with adjuvant to immunize rabbits. Test bleeds were screened for acetyl lysine specificity by ELISA using a lysine acetylated peptide library and nonacetylated peptide library as control. Rabbits showing good reactivity and unique acetyl lysine motif specificity were identified and monoclonal antibodies were generated using a proprietary method for monoclonal production developed at Cell Signaling Technology, Danvers, MA (CST). The final mixture of acetyl lysine antibodies is a mixture containing seven clones (catalog number CST #13416). ELISA analysis of each clone in the mixture was performed according to the procedure described previously (19). Briefly, 100 μl of 1 μM synthetic lysine acetylated and nonacetylated peptides were used to coat wells in a 96-well plate. Coating was carried out overnight at 4 °C. Each purified clone was used at 1:1000 dilution. The plate was incubated for 2 h at 37 °C. An alkaline phosphatase-conjugated goat anti-robot secondary antibody (CST #70545) was added and incubated for 1 h at 37 °C prior to adding p-nitrophenyl phosphate (Sigma-Aldrich) for color development. Absorbance at 405 nm was measured on an ELISA plate reader (PerkinElmer).

Peptide Preparation from Mouse Tissue—Mouse liver tissue was obtained from mature BALB/c mice. Tissue was homogenized in urea lysis buffer (9 M sequanol grade urea, 20 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES), pH 8.0, 1 mM β-glycerophosphate, 1 mM sodium vanadate, 2.5 mM sodium pyrophosphate). Ly- sate were reduced by 4.5 mM dithiothreitol (DTT) for 30 min at 55 °C. Reduced lysate was alkylated with 10 mM iodoacetamide for 15 min at room temperature (RT) in the dark. Sample was diluted fourfold with 20 mM HEPES, pH 8, and digested overnight with 10 μg/ml trypsin (Worthington, Lakewood, NJ). Digested peptide lysate was acidified with 1% trifluoroacetic acid (TFA), and peptides were desalted over 360-μg SEP PAK Classic C18 columns (Waters, Milford, MA). Peptides were eluted with 40% acetonitrile in 0.1% TFA, dried under vacuum, and stored at −80 °C.

Peptide Preparation from Jurkat E6–1 Cells—For label-free experiments, Jurkat E6–1 cells (ATCC) were grown in Roswell Park Memorial Institute (RPMI) 1540 media (Gibco, Grand Island, NY), supplemented with 10% fetal bovine serum (FBS) (Gibco), penicillin, and streptomycin (Gibco). For SILAC experiments Jurkat cells were grown in RPMI (Caisson Laboratories), deficient in L-arginine, and L-lysine and supplemented with 10% dialyzed FBS (Sigma-Aldrich), penicillin, streptomycin, and glutamine (Gibco). Cells were grown in SILAC media supplemented with either L-arginine (Arg 0) and L-lysine (Lys 0), or L-arginine 13C6-15N4-HCL (Arg 10) and L-lysine 13C6-15N2-HCL (Lys 8). Sixteen hours prior to harvest, labeled cells were treated with either 2 μM suberylanilide hydroxamic acid (SAHA) (Sigma-Aldrich) or dimethyl sulfoxide (DMSO) (Sigma). Three biological replicates were prepared for the SAHA treatment experiment with SILAC label swaps. Prior to lysis, cells were pelleted and washed twice in 1× phosphate buffered saline (PBS).

Jurkat cells pellets were lysed at 4 °C in a lysis buffer containing 8 M urea, 50 mM Tris-HCl pH 8.0, 150 mM NaCl, 1 mM ethylenediaminetetraacetic acid (EDTA) (Sigma), 2 μg/μl Aprotinin (Sigma-Al- drich), 10 μg/μl Leupeptin (Roche, West Grove, PA), 1 mM phenylmethylsulfonyl fluoride (PMSF) (Sigma), and 5 mM sodium butyrate. Following lysis, cells were centrifuged at 20,000 × g for 15 min at 4 °C to remove insoluble material. Protein concentration was determined using a bicinchoninic acid (BCA) protein assay (Thermo Scientific, Grand Island, NY).

SILAC labeled samples were prepared by combining 7.5 mg of protein input per SILAC state. Proteins were reduced with 5 mM DTT for 45 min at RT, and subsequently carbamidomethylated with 10 mM iodoacetamide for 30 min at RT in the dark. Prior to digestion the urea concentration was reduced to 2 M with 50 mM Tris-HCl, pH 8. Samples were predigested for 2 h at 30 °C with Endoproteinase Lys-C (Wako Laboratories, Richmond, VA) at an enzyme-to-substrate ratio of 1:50. Samples were digested overnight at 25 °C with sequencing grade trypsin (Promega, Madison, WI) at an enzyme-to-substrate ratio of 1:50. Following digestion samples were acidified with formic acid (FA) (Sigma-Aldrich), and desalted on a 500 mg IC18 Sep-Pak SPE cartridge (Waters) exactly as described in (15, 20). Desalted samples were dried overnight in a vacuum centrifuge and stored at −80 °C.

Basic Reversed Phase (bRP) Chromatography—The offline bRP fractionation was carried out in a manner similar to that previously described (13, 15, 20). In this study, SILAC labeled samples were fractionated using a Zorbax 300 Extend-C18 column (9.4 × 250 mm, 300 Å, 5 μm, Agilent) on an Agilent 1100 series high pressure liquid chromatography (HPLC) system. Samples were reconstituted in 1.8 ml of 5 mM ammonium formate (pH 10.0)/2% (v/v) acetonitrile (MeCN) (bRP solvent A), divided equally into two HPLC vials, and injected successively with bRP solvent A at a flow rate of 3 ml/min. After loading, injected samples were subsequently separated using a 64 min gradient. The gradient consisted of an initial increase to 8% solvent B (1.1% B/min) (90% MeCN, 5 mM ammonium formate, pH 10) followed by a 38 min linear gradient (0.5% B/min) from 8% solvent B to 27% B and successive ramps to 31% B (1% B/min), 39% B (0.5% B/min), and 60% B (3% B/min) using a flow rate of 3 ml/min.
Eighty – 2 ml fractions were collected in a 96-deep well plate (GE Healthcare). For Kac peptide analysis, bRP fractions were pooled in a noncontiguous manner into eight final fractions using the pooling scheme diagrammed in Fig. 3A. For example, to generate the first pooled fraction, original fractions 1, 9, 17, 25, 33, 41, 49, 57, 65, and 73 were combined. A similar pooling strategy was used to generate the remaining seven fractions. Pooled fractions were dried using a SpeedVac concentrator.

**Acetylated Peptide Enrichment from Mouse Tissue—**Immunofinity enrichment of lysine acetylated peptides from mouse liver was performed using the PTMScan protocol as described previously (21). Briefly, 80 µg of each clone in the lysine acetylation antibody mixture from CST #134166 was conjugated to Protein A beads (Roche) overnight at 4 °C and then washed extensively with PBS. A total of 10 mg of mouse liver trypic peptides were dissolved in 1.4 ml of immunofinity purification (IAP) buffer (50 mM MOPS, pH 7.2, 10 mM sodium phosphate, 50 mM NaCl) and centrifuged for 5 min at 12,000 rpm in a MiniSpin microcentrifuge (Eppendorf, Hauppauge, NY). Supernatant was mixed with lysine acetylation antibody beads and incubated for 2 h at 4 °C. The beads were washed twice with 1 ml of IAP buffer and three times with 1 ml of HPLC grade water (Burdick and Jackson, Muskegon, MI). Peptides were eluted from beads with 0.15% TFA (sequential elutions of 55 µl followed by 50 µl, 10 min each elution at room temperature). Eluted peptides were desalted over tips packed with Empore C18 (Sigma) and eluted with 40% acetonitrile in 0.1% TFA. Eluted peptides were dried under vacuum.

**Acetylated Peptide Enrichment from Jurkat E6–1 Cells—**Peptides were enriched for lysine acetylation using the mixture of anti-Kac antibodies described above which were noncovalently coupled to protein A agarose beads (catalog # ICP0388 version supplied as 4 mg/ml) was used for competitive studies. Prior to enrichment, the antibody beads were washed 3× with 1.5 ml of IAP buffer (50 mM MOPS pH 7.2, 10 mM sodium phosphate, 50 mM NaCl). For all Kac enrichment experiments, Jurkat peptide stocks were resuspended in 1.5 ml of IAP buffer and incubated with antibody beads for 1 h at 4 °C with end-over-end rotation. For titration curve experiments conducted with anti-Kac antibody from CST, 2 mg Jurkat peptide aliquots were incubated with 25, 50, 75, 100, or 125 µg of anti-Kac antibody beads. In a parallel experiment 1, 2, 5, 10, and 15 mg of Jurkat peptides were incubated with a fixed amount (50 µg) of anti-Kac antibody beads. For titration curve experiments completed with IMC anti-Kac antibody, 1 mg of Jurkat peptide aliquots were incubated with 20, 30, 40, 50, 60, 100 µg of anti-Kac antibody beads. For the SILAC experiments, eight bRP fractions were each incubated with 50 µg of anti-Kac antibody beads. Following enrichment, anti-Kac antibody beads were washed 4× in 1 ml of 1x PBS. Peptides were eluted from antibody beads with 50 µl of 0.15% (TFA) (2×) and desalted using C18 StageTips. Briefly, each StageTip column was packed with two C18 Empore™ high performance extraction disks (3 µl). Prior to sample loading, StageTips were conditioned with 100 µl of 100% MeOH, 100 µl of 50% MeCN, and 2× with 100 µl of 0.1% FA. Enriched peptides were loaded onto conditioned StageTips, washed 2× with 100 µl of 0.1% FA, and eluted with 50 µl of 50% MeCN, 0.1% FA. Eluted peptides were dried to completeness in a SpeedVac concentrator.

**Mass Spectrometry Analysis—**For mouse liver tissue samples, peptides were dissolved in 0.1% TFA with 5% MeCN and separated on a 100 µm × 15 cm reverse-phase column packed with Magic C18 AQ (100 Å × 3 µm), Michrom, Auburn, CA) into a pulled tip. Peptides were eluted using a 72 min linear gradient of 5–30% MeCN in 0.125% formic acid delivered at 300 nL/min using an Easy nLC (Thermo Fisher Scientific). Tandem mass spectra were collected in a data-dependent manner with either an Orbitrap Elite or a Q Exactive mass spectrometer (Thermo Fisher Scientific). The Orbitrap Elite utilized a top-20 method, collecting MS spectra in the Orbitrap mass analyzer (60,000 resolution) with an automatic gain control target of 1E6 (maximum ion time: 1000 ms) and collision-induced dissociation MS/MS spectra in the ion trap with an automatic gain control target of 4e3 (maximum ion time: 50 ms). The Q Exactive employed a top-10 method, collecting MS spectra at 70,000 resolution and 1E6 automatic gain control with a maximum ion time of 10 ms and higher-energy collisional dissociation MS/MS spectra at 17,500 resolution and 1E5 automatic gain control with maximum ion time of 120 ms and a normalized collision energy of 25. Both instruments scanned precursor masses from 300–1500 m/z and employed dynamic exclusion with a repeat count of 1 and a repeat duration of 30 s. Ions that had a charge of 1 or were unassigned were excluded from MS/MS analysis. The polydimethylsiloxane lock mass (m/z 371.10123) was used as an internal calibrant for all runs.

For all Jurkat cell experiments, desalted peptides were resuspended in 3% MeCN, 0.1% FA and analyzed by online nanoflow liquid chromatography tandem mass spectrometry (LC-MS/MS) using a Q Exactive mass spectrometer (Thermo Fisher Scientific) coupled on-line to a Proxeon Easy-nLC 1000 (Thermo Fisher Scientific). Samples were loaded onto a microcapillary column (360 µm outer diameter × 75 µm inner diameter) containing an integrated electrospray emitter tip (10 µm), packed to 24 cm with ReproSil-Pur C18-AQ 1.9 µm beads (Dr. Maisch GmbH) and heated to 50 °C. The HPLC solvent A was 3% MeCN, 0.1% FA, and the solvent B was 90% MeCN, 0.1% FA. Peptides were eluted into the mass spectrometer at a flow rate of 200 nL/min. For 150 min inject-to-inject LC-MS methods, a linear gradient of 0.3% B/min, followed by a ramp to 60% B (10% B/min) was used. For single-shot, long runs, a 360 min inject-to-inject LC-MS/MS method was utilized. For the 360 min method, a gradient of 0.05% B/min was applied for 150 min, followed by a gradient of 0.09% B/min for 92 min, followed by 0.2% B/min for 42 min, and followed by a ramp to 60% B (10% B/min). The Q Exactive was operated in the data-dependent mode acquiring HCD MS/MS scans (r = 17,500) after each MS1 scan (r = 70,000) on the top 12 most abundant ions using an MS1 target of 3 × 106 and an MS2 target of 5 × 104. The maximum ion time utilized for MS/MS scans was 120 ms; the HCD-normalized collision energy was set to 25; the dynamic exclusion time was set to 20 s, and the peptide match and isotope exclusion functions were enabled.

**Data Analysis—**For the analysis of data generated from mouse liver tissue, RAW files were converted into mzXML format and processed using a suite of software tools developed in-house at CST for the analysis of large-scale proteomic data sets. All precursors selected for MS/MS fragmentation were confirmed using algorithms to detect and correct errors in monoisotopic peak assignment and refine precursor ion mass measurement. All MS/MS spectra were then exported as individual DTA files and searched using SEQUEST (v. 28 (rev. 12), 1998–2007) against the mouse database (20110802_REV_20110628mouse.fasta, NCBI, June 28, 2011; contains 30,120 forward proteins) and their reversed complements. A precursor mass tolerance of 50 ppm and a product ion tolerance of 1.0 Da (collision-induced dissociation spectra) or 0.02 Da (higher-energy collisional dissociation spectra) were allowed. One trypsin terminus was required, and four missed cleavages were allowed. Static carbamidomethylation of cysteine (+57.02146374) was required, and appropriate lysine acetylation (+42.015646863) and methionine oxidation (+15.9949146221) were dynamically allowed with a maximum of four modifications of one type per peptide. Peptide spectral matches were filtered to a 1% false discovery rate using linear discriminant analysis in combination with the target-decoy strategy.
All Jurkat cell data was analyzed using MaxQuant software version 1.3.0.5 (22, 23). Files were searched against the human Uniprot database containing 81,470 entries. This database also included a list of 248 common laboratory contaminants provided by the MaxQuant program. The search parameters were set as follows. The enzyme specificity was set to trypsin with the maximum number of missed cleavages set to 2. The precursor mass tolerance for the first search (used for nonlinear mass recalibration) was set to 20 ppm. The precursor mass tolerance for the main search was set to 6 ppm. Variable modification searches included oxidized methionine, acetylation of internal lysine, and N-terminal protein acetylation. Carbamidomethylation of cysteine was searched as a fixed modification. The false discovery rate for peptide, protein, and site identification was set to 1%. The minimum peptide length was set to 6, and the filter-labeled amino acid and peptide requantification functions were enabled.

The identification and quantification information of acetylated peptides, sites, and proteins was obtained from the MaxQuant “evidence,” and “Acetyl (K) Sites” tables. Only acetylation sites with a localization probability of $>0.75$ were considered for the final SILAC SAHA data set. Significance of the regulated acetylated lysine sites was determined using a moderated T-statistical analysis as previously described (24, 25). Analysis of uniqueness per bRP fraction for Kac peptides was completed using Spectrum Mill. supplemental Table S3 contains information from the MaxQuant “Acetyl (K) Sites” table (28). In addition, either Biological Database Network (http://biodbnet.abcc.ncifcrf.gov/), g:Profiler (http://biit.cs.ut.ee/gprofiler/), or BioMart (http://www.biostat.org/) were used to retrieve ENSEMBL gene identifiers for those Uniprot accessions that were not matched with Uniprot.ws (29, 30).

**Isobaric Labeling Data**—To compare acetylated lysine peptide enrichment across different isobaric labeling techniques, human luminal- and basal-subtype breast tumor mouse xenograft tissue samples were lyzed and digested according to a previous study method that utilized the same type of pooled tissue (31). Luminal and basal xenograft samples were randomly assigned to different channels within each isobaric label experiment, and equivalent peptide amounts of each channel were labeled to generate 4 mg of total peptide material in each four-plex isobaric tags for relative and absolute quantitation (iTRAQ4) (AB Sciex, P/N #86096), six-plex tandem mass tags (TMT6)- (Thermo Scientific, #90068), and 10-plex TMT (TMT10)-labeled (Thermo Scientific, #90406) experiment (1 mg of material was designated to each channel in the iTRAQ4 experiment, 667 μg of peptide was used for each channel of TMT6 and 400 μg of material was designated to each channel in the TMT10 experiment). Offline bRP fractionation of each 4 mg isobaric label experiment was completed using a Zorbax 300 Extend-C18 column (4.6 × 250 mm, 30 Å, 3.5 μm, Agilent) on Agilent, Santa Clara, CA, 1100 series HPLC system. All samples were first separated into 12 bRP fractions and enriched for phosphopeptides as previously described (31). Flow-throughs of the immobilized metal affinity chromatography (IMAC) phosphopeptide enrichments were collected and further combined into four fractions in a noncontiguous manner for sequential Kac peptide enrichment. We used 50 μg of antibody for each Kac enrichment following the same enrichment strategy as described above. All Kac enrichment fractions were analyzed using 140 min inject-to-inject method on a Q-Exact mass spectrometer as previously described (13). We used a normalized collision energy of 28 for iTRAQ- and 27 for TMT-samples. MS/MS scans were acquired at a resolution of 17,500 for iTRAQ4- and TMT6-labeled samples and a resolution of 35,000 for TMT10-labeled samples. MS/MS raw data were searched using Spectrum Mill software against a RefSeq Mouse and Human concatenated database (July 2013 version; contains 56,660 forward proteins). A precursor mass tolerance of 20 ppm and a product ion tolerance of 20 ppm (higher-energy collision dissociation spectra) were allowed. Tryptic enzyme specificity was required, and up to four missed cleavages were allowed. Carbamidomethylation of cysteine as well as iTRAQ- and TMT-tags on lysine residues and the peptide N terminus were analyzed as fixed modifications. Lysine acetylation and methionine oxidation were set as variable modifications. Peptides were identified at a $<1\%$ false discovery rate using a target-decoy search strategy.

**RESULTS**

A Novel Acetyl Lysine Antibody is a Mixture of Seven Monoclonal Antibodies and Shows High Specificity for Acetylated Lysines — An antibody mixture consisting of seven distinct monoclones was recently developed for the enrichment of acetylated peptides. ELISA assays using synthetic acetylated and nonacetylated lysine peptide libraries were performed on test bleeds to determine the specificity of each clone (Fig. 1A, supplemental Table S1) (25). ELISA results show that on average, the clones are $\sim11.5$-fold more specific for acetylated peptides compared with nonacetylated peptides. Enrichment
of Kac peptides from mouse liver tissue using each of the seven monoclones individually and in a mixed format show that on average 3.4-fold more Kac peptides are detected from samples enriched with the mixed antibody reagent relative to enrichment using any individual clone. (supplemental Fig. S1A, supplemental Table S1). Sequence motif analysis of residues flanking Kac sites indicate that there is a general preference for lysine rich regions (Fig. 1B). For some clones a distinct sequence motif proximate to the acetylated lysine is observed. For example, clone 6 has a significant bias for aspartic acid and glutamic acid in the position. Clone 1 and Clone 4 appear to have a significant enrichment of tyrosine in the +1 and −1 position, respectively. Several clones show a bias for underrepresentation of proximate arginine residues. We observed that the sequence motif for the antibody mix is a composite of motifs present in multiple clones (Fig. 1B). These results coupled with the observation that the mixture of antibodies has the highest specificity supports the notion that the combination of multiple clones can allow for broader-range recognition of acetylated peptides (Fig. 1B, supplemental Fig. S1A). Although the non-acetylated peptide signal from clone 3 was much higher than the background signal from the other clones (1, 2, 4, 5, 6, and 7) by ELISA assay (Fig. 1A, supplemental Table S1), the mass spectrometry results clearly highlight the importance of this clone in the final formulation. Based on the major contribution of distinct Kac peptides (supplemental Fig. S1), clone 3 was included in the final formulation providing complementary sequence coverage for additional Kac peptides that were not accessible using only the other six clones (Fig. 1B).

Titration of Antibody and Peptide Input Amount is Critical to Maximize Yield and Specificity for Detection of Lysine Acetylated Peptides—Our previous work focused on large-scale enrichment of ubiquitination sites from cellular lysates highlighted the importance of optimizing the amount of antibody to maximize the yield of Kac-GG peptides (20). Therefore, in parallel experiments, we titrated the amount of anti-Kac antibody mixture and the amount of input protein to study the interplay of these factors on the number of identified Kac and non-Kac peptides. Titration experiments that tested a range of anti-Kac antibody amount revealed that 50 μg of antibody is sufficient and optimal for maximizing the yield of Kac peptides from 2 mg of input (Fig. 2, supplemental Table S2). Although amounts of antibody as low as 25 μg can yield high numbers of acetylated peptides, pipetting this small volume of beads (4 μl) is less reproducible between replicates (Fig. 2). The number of identified acetylated peptides steadily decreased when the anti-Kac antibody mixture amount was increased beyond 50 μg and the number of identified non-acetylated peptides steadily increased. This observation is consistent with our previous findings for the anti-K-GG antibody (20). The titration experiments also show that 50 μg of the anti-Kac antibody is sufficient to enrich Kac peptides from up to 15 mg of peptide input (supplemental Fig. S2, supple-
Taken together, these results indicate that it is imperative to determine the optimal antibody and protein input combination to maximize the yield of Kac peptides.

Alternate anti-Kac antibody reagents are commercially available and have been previously used in global acetylation experiments (5, 9–12). We compared the performance of the anti-Kac antibody clone mixture from CST to a commonly used anti-Kac antibody from ImmuneChem (IMC). The IMC antibody is a polyclonal rabbit antibody, and is one of the leading reagents used for large scale acetylation profiling experiments (9–12, 32). We performed titration experiments for the IMC antibody to establish the optimal amount of reagent needed for Kac peptide enrichment (supplemental Fig. S3A, S3B, supplemental Table S2). Our results show that 60 μg of antibody is optimal to achieve the maximum yield of acetylated peptides from 1 mg of peptide input. Having defined the optimal amounts of both the CST and IMC anti-Kac antibodies to use for a set amount of digested input protein lysate, we set out to compare the results of Kac peptide enrichment with these reagents. We incubated 2 mg of peptides with either 50 μg or 60 μg of anti-acetyl antibody from CST and IMC, respectively. Results indicate that the yield of distinct Kac peptides is on average 4277 (S.D. 77.5) with CST antibody relative to 1957 (S.D. 479.7) with IMC antibody (supplemental Fig. S4A, supplemental Table S2). The percentage of Kac peptides relative to non-Kac peptides identified in a given sample is 41% for CST and 16.8% for IMC. We next evaluated the overlap of Kac peptides enriched with both reagents to understand if the antibodies exhibit complementary specificity. The overlap between any two IAP process replicates was under 41% for CST and 16.8% for IMC. We next evaluated the overlap of Kac peptides enriched with both reagents to understand if the antibodies exhibit complementary specificity. The overlap between any two IAP process replicates was under 41% for CST and 16.8% for IMC. The specificity for Kac peptides per fraction was ~40% as measured by Kac modified over total distinct peptide counts. Although the specificity for identification of Kac peptides per sample is lower in fractionated versus unfractionated samples, we observed a marked improvement in absolute number of identified Kac peptides observed with bRP fractionation.

Similar to previous results for K-ε-GG peptides, pooling of nonadjacent basic RP fractions produced fractions containing consistent numbers of acetylated and nonacetylated peptides (Fig. 3C). Importantly, we found that >80% of identified Kac peptides were uniquely identified in any one of eight bRP fractions (Fig. 3E), resulting in a near linear increase in the numbers of acetylated peptides sequenced with each additional fraction (Fig. 3D). Advantages of having the majority of peptides exist in only one bRP fraction include reduced redundancy of MS peptide sampling, conservation of signal intensity for a given peptide, and reduced peptide co-elution interference. Altogether, these data show that bRP fractionation coupled to anti-Kac antibody enrichment significantly improves the depth-of-coverage for identification of low-abundance Kac peptides.

Large-scale Acetylation Profiling of SAHA Treated Cells Reveals High Numbers of Regulated Acetylation Sites—We benchmarked our acetylation workflow using Jurkat E6−1 cells treated with the HDAC inhibitor SAHA (Fig. 3A). As described above, samples underwent bRP prefractionation prior to Kac enrichment. Using this sample preparation workflow combined with enrichment using the newly generated anti-Kac antibody mixture, we identified and quantified unprecedented numbers of Kac peptides from a single SILAC-labeled sample with 7.5 mg of peptide input per state. On average, we identified over 3000 Kac proteins and quantified over 10,000 Kac peptides with over 8600 localized Kac sites in each of three biological replicates (Table I). For localized Kac sites, an average of 6553 were quantified in any two replicates, and 5524 were quantified in all three replicates (Fig. 4A, supplemental Table S3). At the Kac protein level, the
FIG. 3. **A. Schematic of the proteomic workflow for acetylation profiling.** B, Average number of distinct Kac and non-Kac peptides identified in bRP fractionated and nonfractionated single SILAC samples, enriched from 7.5 mg of peptide input per state. Error bars represent standard deviation for three replicates. C, Average number of distinct Kac and non-Kac peptides identified in each bRP fraction. Error bars represent standard deviation for three replicates. D, Cumulative distribution showing the number of distinct Kac peptides identified with bRP fractionation. Data points show the average of three replicates and error bars represent standard deviation. E, Percentage of distinct Kac peptides identified in one, two, or more than two bRP fractions.
As expected down-regulated (p response to SAHA treatment whereas only 1.8% of sites were quantified in our data set and the Choudhary single replicate of our data (supplemental Fig. S5 our total SAHA data set, and 875 (61%) overlapped with a 1425 Kac sites quantified from the data of Choudhary neChem) and sample processing strategy we used, but also (20.4%) of acetylation sites were up-regulated (acetylation landscape (10, 33, 34). A significant percentage deacetylase inhibitor SAHA induces significant effects on the lysine modification (Fig. 4 age, the occurrence of multiply acetylated proteins, and evolution-ary conservation of lysine acetylation (12, 35).

We also analyzed our acetylome data set for lysine acetylation sequence motifs. In agreement with previous work by Weinert et al. and Lundby et al., we also found that residues flanking acetylation sites are highly enriched in lysine and frequently contain glycine (Gly), aspartic acid (Asp), and glu-tamic acid (Glu) at the −1 position (12, 35). Interestingly, as shown above, the anti-acetyl antibody clone 6 had a strong preference for Asp and Glu at the −1 position (Fig. 1B). However, although it is always important to consider the potential for sequence bias in global proteomic analyses of post-translational modifications, bias is unlikely in the present case because of the use of mixtures of clonally distinct anti-bodies with complementary motif specificities.

**Acetylome Profiling is Compatible with Isobaric Labeling Techniques**—Isobaric labeling techniques such as iTRAQ and TMT offer the ability to quantify peptides and proteins in a highly multiplexed manner and are amenable for use in nearly all sample types. Hebert et al. previously showed that TMT labeling coupled to immunofinity enrichment of Kac pep-tides from mitochondria purified from mouse liver yields an average of 2400 sites from a sample where 1 mg of input material was used per label (11). To further investigate iTRAQ and TMT quantification techniques for global acetylation profiling, luminal and basal subtype breast cancer xenograft tissue samples (see Methods) were homogenized, lysed, and digested as described above. For comparison purposes, a total of 4 mg of tumor material was used in each isobaric labeling plex, with that amount divided equally into each channel of the plex (iTRAQ4, TMT6, TMT10). Each experiment was separated using bRP fractionation, and enriched for Kac peptides. Enrichment for lysine acetylation was completed with 50 μg of anti-Kac antibody (CST) for each of 4 distinct bRP fractions per isobaric label type. We find that bRP fractionation of iTRAQ and TMT labeled peptides leads to a

## Table I

| Rep1 | Rep2 | Rep3 | Average |
|------|------|------|---------|
| # Identified proteins with Kac sites | 3127 | 3221 | 3320 | 3223 |
| # Kac identified peptides | 10409 | 10717 | 11417 | 10848 |
| # Kac quantified peptides | 10135 | 10128 | 11030 | 10431 |
| # Kac identified and localized sites | 8584 | 8831 | 9379 | 8863 |
| # Kac quantified and localized sites | 8400 | 8406 | 9107 | 8638 |

Overlap between any two replicates was on average 2831, and 2642 proteins were identified in all three replicates (Fig. 4B, supplemental Table S3).

The average Pearson correlation value for SILAC ratios across three replicates of the SAHA versus DMSO treated cells was 0.84, indicating a high degree of experimental reproducibility (Fig. 4C). As previously shown, the lysine deacetylase inhibitor SAHA induces significant effects on the acetylation landscape (10, 33, 34). A significant percentage (20.4%) of acetylation sites were up-regulated (p < 0.01) in response to SAHA treatment whereas only 1.8% of sites were down-regulated (p < 0.01) (Fig. 4C, supplemental Table S3). As expected Kac sites up-regulated by SAHA treatment were identified on proteins involved in nuclear processes such as transcription, cell cycle, chromatin assembly and histone modification (Fig. 5A). Interestingly, proteins with down-regulated Kac sites were also enriched in transcriptional and chromatin regulation processes. Up-regulated Kac proteins were significantly enriched in both nuclear and mitochondrial proteins, whereas nearly all down-regulated Kac proteins showed enrichment for nuclear localization (Fig. 5A and 5B).

We used an identical biological setup to previous work by Choudhary et al. to facilitate comparison and to better understand the extent of improvement in our depth-of-coverage for mapping lysine acetylation sites (10). As part of their broad acetylome study, Choudhary et al. examined the effects of SAHA treatment in Jurkat cells using the anti-Kac antibody from ImmuneChem. Searching their data using our data processing pipeline showed that 1425 Kac sites were quantified and localized after SAHA treatment of Jurkat cells. Comparison of our two data sets showed that we identified ~sixfold more acetylation sites in a single replicate than detected in the earlier study (supplemental Fig. S5B). This large difference is due in part to the improved antibody (CST versus ImmuneChem) and sample processing strategy we used, but also because of improvements in MS technology (Q-Exactive versus LTQ Orbitrap XL). Overlap analysis showed that of the 1425 Kac sites quantified from the data of Choudhary et al., 972 (68%) overlapped with the 11,922 Kac sites quantified in our total SAHA data set, and 875 (61%) overlapped with a single replicate of our data (supplemental Fig. S5A and S5B). The differences observed in the specific acetylation sites quantified in our data set and the Choudhary et al. data set are likely because of differences in the specificities of these re-agents as no more than 68% of the Kac peptides identified after enrichment with the ImmuneChem antibody were also identified with the CST antibody. An extensive study completed by Lundby et al. created an organ-wide map of lysine acetylation from 16 different rat tissues (12). In aggregate they identified >15,000 total Kac sites with an average of 3000 to 5000 Kac sites per tissue using the anti-Kac antibody from ImmuneChem. To facilitate comparison of our Kac data (human) to the data presented by Lundby et al. (rat), we completed our evaluation at the Kac protein level. Overlap analysis revealed that 46% of Kac proteins reported by Lundby et al. were identified in our study (supplemental Fig. S5C). Similar to Lundby et al., we identify an average of ~3 Kac sites per protein (supplemental Fig. S6) (12). Taken together, we find that nearly half of all acetylated proteins we identified overlapped to acetylated proteins previously identified from rat tissue (12). This is a result of our high depth-of-cover-age, the occurrence of multiply acetylated proteins, and evolution-ary conservation of lysine acetylation (12, 35).
uniform distribution of peptides across four fractions (supplemental Fig. S7, supplemental Table S4). For the iTRAQ4 experiments we quantified 6753 Kac peptides, 5579 Kac sites, and 2327 Kac proteins. For TMT6 experiments 4191 Kac peptides, 3678 Kac sites, and 1828 Kac proteins were quantified. For TMT10 experiments we quantified 4180 Kac peptides, 3653 Kac sites, and 1804 Kac proteins. Isobaric labeling combined with our optimized acetylation profiling workflow yields extremely deep coverage of Kac sites from total tissue lysates. Labeling using iTRAQ4 yields ~1.5-fold higher numbers of acetylated peptides when compared with labeling with TMT6 and TMT10 (supplemental Fig. S7, supplemental Table S4). Therefore, we find that labeling using iTRAQ4 offers higher sensitivity over TMT6 and TMT10 in acetylation experiments, but with a lower multiplexing capability. Taken together, we show that isobaric labeling techniques are highly amenable for profiling the acetylome and offer increased throughput in profiling of biological samples.

DISCUSSION
We have described the use of a new, commercially available anti-Kac antibody mixture that in combination with our
global PTM workflow results in the identification of unprece-
dented numbers of lysine acetylation sites. Specifically we
were able to enrich over 10,000 lysine acetylated peptides
from moderate sample input. Our findings show that for the
first time the depth-of-coverage attainable for the acetylome
is comparable to that for other well studied PTMs (e.g. phos-
phorylation, ubiquitylation) (13, 16–18, 20).

The high-degree of coverage we achieve can be attributed
to multiple improvements in the proteomic workflow includ-
ing, antibody specificity and handling, off-line fractionation
strategy, and higher sensitivity of new LC-MS/MS technology.
We showed that the new anti-Kac antibody reagent increases
the number of identified acetylated peptides by twofold rela-
tive to another commonly used anti-Kac antibody from Immu-
neChem. In addition to the improved antibody performance,
we attribute the significant increase in the number of Kac sites
quantified relative to the study by Choudhary
et al. in SAHA-
treated Jurkat cells to the highly effective bRP separation
technique, UPLC conditions for online separation, and mass
spectrometry analysis on a Q Exactive instrument (10). Newer
generation of mass spectrometers such as the Q Exactive
significantly improve the detection of low abundant PTMs
because of improvements in speed and sensitivity.

Deep coverage of the acetylome allows for analysis of
primary sequence enrichment proximate to acetylated lysine.
From our work, we find that acetylated lysines occur in lysine
rich regions and are often preceded by Gly, Asp, and Glu
residues as has been previously shown (12, 35). It should be
noted that alternate sequence preferences will likely emerge
for specific cellular compartments as illustrated by Lundby
et al. (12). Finally, we also observed that nearly half of all acety-
lated proteins we identify for SAHA treated Jurkat cells over-
lap at the acetylated protein level to the rat acetylome data set
reported Lundby et al. Ultimately, this degree of overlap is
supported by the evolutionary conservation of lysine acetyla-
tion and our high depth-of-coverage (12, 35).

In this study, we show that both metabolic (SILAC) and
isobaric (iTRAQ or TMT) labeling methods are highly compat-

Fig. 5. A, Gene ontology enrichment of Biological Processes (BP) and Cellular Compartment (CC) terms for SAHA induced up- and
down-regulated Kac sites ($p < 0.01$). Enrichment analysis was performed using DAVID. C, IceLogo motif analysis of Kac peptide sequence windows ($p < 0.01$). The height of amino acid letters corresponds to fold change.
ible with the acetylation enrichment workflow. Having flexibility in quantification strategy is advantageous for broadening the spectrum of sample types that can be profiled for acetylation. SILAC labeling offers the advantage of using relatively larger sample inputs which helps to increase depth-of-coverage. Workflows using isobaric labeling strategies are limited to smaller sample inputs because of reagent costs, but offer increased multiplexing capability and are amenable for use with nearly all samples types.

To further increase the coverage of acetylation sites, possibilities include, extending the number of bRP fractions, increasing sample input, and combining the use of different anti-Kac antibodies in a mixed or serial enrichment format. We envision that the state-of-the-art methodology presented here for deep coverage of the acetylome could be applied to the study of other less abundant PTMs and facilitate improved understanding of PTM cross-talk.

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