Stoichiometry of Site-specific Lysine Acetylation in an Entire Proteome*

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Background: Lysine acetylation sites have been mapped, but information on stoichiometry is lagging.

Results: We developed and utilized the first direct, unbiased method for quantifying site-specific acetylation stoichiometry of a proteome without antibody enrichment.

Conclusion: High stoichiometry is associated with central metabolism, transcription, and translation. Loss of deacetylase CobB affects site-specific and global acetylation stoichiometry, altering acetyl-CoA metabolism.

Significance: Stoichiometry provides functional insight into protein acetylation.

Acetylation of lysine ε-amino groups influences many cellular processes and has been mapped to thousands of sites across many organisms. Stoichiometric information of acetylation is essential to accurately interpret biological significance. Here, we developed and employed a novel method for directly quantifying stoichiometry of site-specific acetylation in the entire proteome of Escherichia coli. By coupling isotopic labeling and a novel pairing algorithm, our approach performs an in silico enrichment of acetyl peptides, circumventing the need for immunoenrichment. We investigated the function of the sole NAD⁺-dependent protein deacetylase, CobB, on both site-specific and global acetylation. We quantified 2206 peptides from 899 proteins and observed a wide distribution of acetyl stoichiometry, ranging from less than 1% up to 98%. Bioinformatic analysis revealed that metabolic enzymes, which either utilize or generate acetyl-CoA, and proteins involved in transcriptional and translational processes displayed the highest degree of acetylation. Loss of CobB led to increased global acetylation at low stoichiometry sites and induced site-specific changes at high stoichiometry sites, and biochemical analysis revealed altered acetyl-CoA metabolism. Thus, this study demonstrates that sirtuin deacetylase deficiency leads to both site-specific and global changes in protein acetylation stoichiometry, affecting central metabolism.

Acetylation of the ε-amino group of lysine residues is now considered a major regulatory protein modification that influences many cellular processes, including protein-protein interactions, protein-DNA interactions, stability, cellular localization, and enzymatic activity (1, 2). Acetylation occurs by enzymatic and nonenzymatic mechanisms (3). Nonenzymatic acetylation occurs through nucleophilic attack of the lysine side chain on esters of the metabolic intermediates, acetyl-CoA and acetyl phosphate (4, 5). In prokaryotes, acetyl phosphate was reported to serve as the major acetyl donor (4). Removal of acetyl modifications requires protein deacetylases (6).

Mass spectrometry-based proteomic studies have mapped thousands of acetylated sites in a wide range of organisms (7–11). Recently, acetyl proteome measurements have utilized labeling strategies, such as tandem mass tags or stable isotope labeling by amino acids in cell culture, followed by immunoenrichment of acetylated peptides and mass spectrometry (8, 12–15). Although these approaches are valuable for comparing relative changes across conditions, they do not provide direct information on stoichiometry at individual sites (1, 14). Stoichiometric information is essential to accurately interpret the biological significance of these acetylation sites. For example, if only relative acetylation can be determined, as is the case with current acetylation methods, then a 1–5% and a 20–100% change would appear identical, i.e. a 5-fold change. The biological significance of these changes is entirely different, especially in cases in which acetylation is inhibitory.

Mass spectrometry has been used to examine phosphorylation stoichiometry in large scale studies (14, 16, 17). One central issue that is addressed by these approaches is that post-translationally modified peptides and their corresponding unmodified counterparts have varying ionization efficiencies, and thus, they cannot be directly compared. The issue of differing ionization efficiency can be addressed by making indirect measurements that incorporate correlated changes in phosphorylated and nonphosphorylated versions of peptides that are corrected for overall protein level changes across experiments with very different amounts of phosphorylation (18). Very recently, this indirect strategy was applied to estimate acetylation stoichiometry in log phase yeast compared with growth-arrested yeast (19). Yet, this strategy requires quantifying modified and unmodified versions of the same peptide, which are difficult to detect even with enrichment. It is biased in that it can only
measure stoichiometry for peptides where PTM² levels are dramatically different across conditions. Furthermore, it is unclear how the estimates behave when the changes in PTM status across conditions are small. In the context of phosphorylation, these challenges can be addressed by treating an identical stable isotope-labeled sample with a phosphatase and measuring and comparing the increased abundance of the unphosphorylated peptide (17). The percent increase provides a measurement of the stoichiometry of a phosphosite that circumvents the need for enrichment and the requirement for a change in the level of a PTM site. Yet no such approach exists for lysine acetylation.

Here, we developed and utilized the first unbiased and direct method for quantifying the stoichiometry of site-specific acetylation at the proteome-wide scale, without enrichment. Our approach uses a stable isotope chemical labeling step that acetylates all unmodified lysines resulting in a “heavy” and “light” acetyl-lysine pair across the entire proteome. This pair is then analyzed using high resolution, high accuracy mass spectrometry to yield proteome-wide, site-specific stoichiometry. Using this approach, we interrogated the acetyl proteome of Escherichia coli and investigated the functional consequences of genetically removing the sole NAD⁺-dependent protein deacetylase, CobB, on both site-specific and global acetylation. We quantified 2206 peptides from 899 proteins in the range of less than 1% up to 98% acetylation. Loss of CobB resulted in a slight increase of global acetylation at low stoichiometry sites, as well as site-specific changes at high stoichiometry sites. Proteins with the highest stoichiometry included those involved in central metabolism, transcription, and translation.

**EXPERIMENTAL PROCEDURES**

**Cell Culture and Sample Preparation—**E. coli BL21 (DE3) wild type and ΔCobB (with empty pQE80) were grown in 2× YT media under ampicillin selection overnight. Cells were harvested, flash-frozen, and stored in −80 °C. Frozen cells were resuspended in ~10 volumes of 8 mM urea solution (8 mM urea, 5 mM DTT, 100 mM ammonium bicarbonate, pH 8) and lysed by sonication. After centrifugation, the protein concentration was determined using Bradford reagent (Bio-Rad).

**AceCS2 Expression—**AceCS2 pQE80 was transformed into E. coli BL21 (DE3), and protein expression was performed as described previously (20).

**Protein Chemical Acetylation and Digestion—**An equal amount of protein (100 µg) was diluted using 8 mM urea solution (2 µg/µl) and incubated at 60 °C, 1000 rpm, for 30 min on the Thermomixer (Eppendorf). Cysteine alkylation with iodoacetamide was performed for 30 min in the dark. Chemical acetylation was performed as described previously, with slight modification (21). An equal volume of ammonium acetate solution (1 mM NH₄OAc, 8 mM urea, 100 mM ammonium bicarbonate, pH 8) was added to each sample. ~20 µM of acetic anhydride (acetic anhydride, acetic anhydride-d₆, or acetic anhydride-¹³C₆-d₆ (Sigma)) was added to each sample and incubated on a Thermomixer at 4 °C, 1000 rpm, for 20 min. After incubation, the pH was raised to ~7 using ammonium hydroxide and checked using litmus paper. Chemical acetylation was repeated two more times, and buffer exchange occurred with 50 mM ammonium bicarbonate (pH 8), 10% methanol using 10K MWCO spin filters (Millipore). Trypsin was added at a 1:100 ratio and incubated overnight, 37 °C, 300 rpm on a Thermomixer.

**LC-MS/MS—**Peptides were separated with a Dionex Ultimate 3000 RSLCnano HPLC using a Waters Atlantis dC18 (100 µm × 150 mm) reverse phase column. The mobile phase consisted of (A) water with 0.1% formic acid and (B) acetonitrile with 0.1% formic acid. Peptides were eluted with a linear gradient of 2–40% B at a flow rate of 0.7 µl/min over 120 min and introduced into a hybrid quadrupole-Orbitrap mass spectrometer (Thermo Q Exactive) by nanoelectrospray ionization (Thermo Nanospray Flex). The MS survey scan was performed in positive ion mode with a resolution of 70,000, AGC of 1E6, maximum fill time of 100 ms, and scan range of 400 to 2000 m/z. Data-dependent MS/MS was performed with a resolution of 17,500, AGC of 1E5, maximum fill time of 64 ms, isolation window of 1.5 Da, underfill ratio of 0.1%, normalized collision energy of 28, dynamic exclusion of 20 s, and a loop count of 20. The source voltage was set at 2000 V and capillary temperature at 250 °C.

**MS Data Analysis—**Data were analyzed using a modified version of the December 25th, 2013, release of open-source, quantitative mass spectrometry analysis tool PVIEW (22, 23). We generated a peptide database with cleavage only at arginine, as trypsin did not cleave at acetylated lysines. Up to two missed cleavages were allowed. MS1 precursor tolerance was set to ±10 ppm, and MS2 fragment mass tolerance was set to ±15 ppm. Peptide identifications were obtained at a false discovery rate of 1%. Estimation of false discovery rate was carried out using only peptide spectrum matches from paired extracted ion chromatograms (XICs).

**Acetylation Stoichiometry Calculations—**Equation 1 was used to calculate stoichiometry of single lysine-containing peptides,

\[
\frac{\text{XIC}_L}{\text{XIC}_L + \text{XIC}_H} \quad \text{(Eq. 1)}
\]

and Equation 2 for double lysine-containing peptides,

\[
\frac{\text{XIC}_L + \frac{1}{2}\text{XIC}_M}{\text{XIC}_L + \text{XIC}_M + \text{XIC}_H} \quad \text{(Eq. 2)}
\]

where XICₗ is the extracted ion chromatogram of the light acetyl peptide; XICₕ is the extracted ion chromatogram of the heavy acetyl peptide (acetic anhydride-d₆ or acetic anhydride-¹³C₆-d₆), and XICₘ, when present, is the extracted ion chromatogram for the double lysine-containing peptide with a light and heavy acetyl-lysine.

Because of the isotopic purity of the commercially available acetic anhydride-¹³C₆-d₆ labeling reagent (97 atom % D, 99 atom % ᵃ¹³C), a global correction factor was applied to the stoichiometry of the ΔS-acetic anhydride-labeled sample.
Protein Acetylation Stoichiometry

\[
\frac{XIC_L + \frac{1}{2}XIC_M}{XIC_L + XIC_M + XIC_H} \times 0.96 - \frac{1}{2}XIC_M + XIC_H
\]

Determination of Chemical Labeling Efficiency—An equal amount of protein (BSA or E. coli lysate with expressed mouse AceCS2) was prepared as described above. An unlabeled sample was used as a control. Protein digestion, LC-MS/MS, and database search were performed as described above.

Unmodified tryptic peptides of the labeled and control sample were monitored and compared to determine labeling efficiency. Briefly, the peak area for a lysine-containing peptide in the control sample was measured. In the labeled sample, the same lysine site would be acetylated to some degree, generating a larger acetyl peptide and also the unmodified peptide proportionate to the degree of chemical labeling. As a result, the unmodified peptide in both the labeled and control samples can be compared to determine labeling efficiency. For normalization across samples, a peptide lacking lysine residues was used. Peak areas for each peptide were determined using Xcalibur 2.2 SP1 (Thermo Scientific).

Stoichiometry Curve Determination—An equal amount of BSA was chemically acetylated with acetic anhydride or acetic anhydride-\(^{13}\)C\(_4\)H\(_4\), and digested as described above. Light and heavy BSA peptides were resuspended to equal concentrations and then mixed at varying ratios corresponding to 1, 5, 10, 20, 25, 40, 50, 60, 75, 80, 90, 95, and 99% acetylation. The BSA samples were then added to a trypsin-digested E. coli proteome at equal concentrations, analyzed by LC-MS/MS, and data processed as described above.

Bioinformatics—Gene ontology (GO) and pathway analysis was performed using DAVID version 6.7 (24, 25). For the enrichment analysis, an unlabeled, trypsin-digested, E. coli proteome was used as the background. Network analysis was performed using STRING version 9.1 with proteins having enrichment of acetylated peptides.

Method Development and Validation—To measure acetylation stoichiometry on the proteome scale, we developed a robust workflow using chemical acetylation with isotopic acetic anhydride followed by trypsin digestion and high resolution mass spectrometry (Fig. 1). The general strategy involves preparing denatured protein extracts from cells and chemically acetylating free lysine residues with isotopic acetic anhydride, followed by trypsin cleavage and MS analysis. Using this process, every lysine bears an acetyl group, either the light version derived from endogenous acetylation or the heavy version originating from in vitro chemical acetylation. The trypsin cleavage step produces chemically identical peptides, which have matching retention times. The corresponding pairs of peptides that differ in mass are resolved using mass spectrometry, and direct stoichiometry is determined by dividing the light peak area over the sum of the light and heavy peak areas.

To ensure accurate determination of stoichiometry, we first evaluated acetic anhydride labeling efficiency within a simple and complex protein sample. An E. coli whole-cell lysate was chemically modified with acetic anhydride and qualitatively assessed for the degree of acetylation. Western blot analysis revealed that acetic anhydride chemically acetylated E. coli proteins across the full range of protein sizes (Fig. 2A). To quantitatively assess the degree of chemical acetylation in a simple protein sample, equal amounts of labeled and unlabeled bovine serum albumin (BSA) were digested and analyzed by mass spectrometry. Unmodified BSA tryptic peptides in both samples were monitored and compared to determine the degree of acetylation. The labeling efficiency across eight out of nine BSA peptides was greater than 98% (Fig. 2B). To determine the labeling efficiency in a complex sample, the mouse AceCS2 protein was recombinantly expressed in BL21 (DE3) bacteria. The cells were lysed; protein was labeled with increasing amounts of acetic anhydride, and the labeling efficiency was calculated for peptides of AceCS2 protein. Across the eight sites in AceCS2, the labeling efficiency was >98% (Table 1). These results indicate that proteins in complex mixtures can be chemically acetylated with high efficiency.

We developed a computational method for stoichiometric determination of acetylation that avoids the need for immunoenrichment. Existing proteome-wide acetylation studies require an enrichment step to increase sensitivity in the PTM analysis (14). Sensitivity is achieved by the increased abundance and subsequent identification of acetylated peptides. In this study, we achieved high sensitivity by coupling a novel computational algorithm, implemented in a peptide quantification software tool called PVIEW, to the chemical acetylation strategy. This algorithm identifies acetyl peptides (heavy or light) using one of two databases (Fig. 3A) and alternates between iterations of heavy-to-light and light-to-heavy XIC pairing (Fig. 3, B–D), using the high abundant species as a proxy to identify the corresponding, lower abundant, peptide pair. In essence, the algorithm identifies the best acetyl peptide and based on charge and isotopic shift, the corresponding peptide pair is identified, matched, and quantified. In this manner, our algorithm performs an in silico enrichment of acetylated peptides.

We demonstrated the capability of measuring acetylation stoichiometry across a broad range. BSA protein was chemically acetylated with either acetic anhydride (light) or isotopic acetic anhydride (heavy) followed by trypsin digestion. Peptides
were then mixed at distinct ratios corresponding to 1–99% acetylation and added to a trypsin-digested *E. coli* proteome. This experiment served to mimic the full range of endogenous acetylation that might be encountered from diverse biological conditions. Samples were analyzed by LC-MS/MS, and the stoichiometries of five BSA acetyl peptides were quantified using our novel quantification algorithm. As a representative, the MS/MS spectrum for the acetyl peptide, ALK\(^{(Ac)}\)AWSVAR, is shown in Fig. 4A. MS1 spectra corresponding to 10, 50, and 90% stoichiometry are shown in Fig. 4, B–D, respectively. The stoichiometry curve encompassing 1–99% stoichiometry is shown in Fig. 4E. Linear regression analysis of the stoichiometry curves from all five acetyl-BSA peptides show a high correlation with \( R^2 \) values ranging between 0.970 and 0.995 (Fig. 4F). These results establish the feasibility of accurately determining stoichiometry across a wide range of values in a complex mixture.

Acetylation Stoichiometry Values from Proteomes of WT and \( \Delta \text{CobB} \) Bacteria—Many bacteria, like *E. coli*, contain a single NAD\(^+\)-dependent protein deacetylase that is the ortholog to CobB from *Salmonella enterica*. In *S. enterica*, CobB regulates metabolic pathways that permit growth on alternative carbon sources such as short chain fatty acids (SCFA) (30, 31) and citrate (32). Here, we applied our new method to investigate the
role of CobB on both site-specific and global acetylation stoichiometries.

We determined protein acetylation of BL21 (DE3) wild type and ΔCobB strains, analyzing three biological replicates for each strain. Cells were grown in rich media, harvested during stationary phase and quantifies the magnitude change between the two conditions, revealing peptides that have higher stoichiometry in either ΔCobB (red lines) or WT (black lines) strains (Fig. 6 and supplemental Tables S1 and S3). This analysis also revealed three groups of peptide stoichiometry profiles as follows: 1) peptides with low acetylation stoichiometry and low magnitude change; 2) peptides with higher acetylation stoichiometry and low magnitude change; and 3) peptides with different stoichiometries and a high magnitude change between conditions. Group 1 represents the majority of peptides, which is reflected by the low average global acetylation stoichiometry (Fig. 6).

Interestingly, group 2 peptides had a higher stoichiometry regardless of CobB status, which suggests these acetyl-lysine sites are not targets of CobB. Finally, group 3 peptides had the highest magnitude change between WT and ΔCobB and included acetyl peptides that increase in the CobB deletion. These include tryptophan-tRNA ligase (TrpS), 30 S ribosomal protein S16 (RpsP), peptide deformylase (Def), xanthine phosphoribosyltransferase (Gpt), and acetyl-CoA synthetase (Acs).

We next determined site-specific acetyl stoichiometry changes upon CobB deletion by generating a stoichiometry line plot that compares the acetylation of WT and ΔCobB peptides and quantifies the magnitude change between the two conditions, revealing peptides that have higher stoichiometry in either ΔCobB (red lines) or WT (black lines) strains (Fig. 6 and supplemental Tables S1 and S3).
Bioinformatic and Network Analysis of Acetylation Stoichiometry—

To examine whether proteins that display substantial stoichiometry are enriched in specific cellular processes, we performed an enrichment analysis of our unique stoichiometry results using the database for annotation, visualization, and integrated discovery (DAVID) (24, 25). Certain metabolic pathways and ribosomal proteins were significantly enriched above background (Fig. 7A). Gene Ontology (GO) analysis also revealed that acetyl-CoA metabolic processes, transcription, translation, and tRNA aminoacylation were significantly enriched (Fig. 7B). A network analysis of the identified pathways was performed using the STRING Database (26, 27). Acetylation stoichiometries along with the number sites were superimposed across the protein interaction network and visualized using Cytoscape (28). Acetylation sites per protein varied across the proteome (Fig. 7C); therefore, we used the highest stoichiometry quantified for each protein in the network. The pentose phosphate pathway, glycolysis, pyruvate metabolism, and the TCA cycle revealed proteins with sites of higher stoichiometry (Fig. 7D). The proteins in these pathways were also enriched in total acetylation, as many proteins in these pathways had more than seven acetyl-lysine sites (Fig. 7, D and F). Proteins involved in transcription and translation displayed many sites of acetylation and high stoichiometries (Fig. 7, E and F). This unique network analysis revealed the range of stoichiometries in major cellular pathways and processes.

Total acetylation site number and high stoichiometry enrichment in central metabolic pathways suggest that CobB deletion affects these pathways. To visualize these results, we plotted the TCA cycle and pyruvate metabolism pathways with acetyl-CoA at the crossroads of these two pathways (Fig. 8). As evident from this analysis, enzymes that

![Diagram of LC-MS/MS data analysis algorithm used for identifying isotopic pairs for stoichiometry determination.](image-url)
TABLE 2
Acetylation stoichiometry values

The number of acetyl peptides was quantified across the experimental conditions and strains. This includes the total number of acetylation sites quantified, 1K and 2K peptides, and total proteins identified.

| Acetic anhydride | Δ3 | Δ5 | ΔCobB | WT | Total |
|------------------|----|----|-------|----|-------|
| Lysine sites detected | 4025 | 2659 | 2573 | 1945 | 2206 |
| Single lysine peptides with stoichiometry | 2221 | 1539 | 1451 | 1103 | 1602 |
| Double lysine peptides with stoichiometry | 902 | 560 | 561 | 421 | 603 |
| Proteins identified | 852 | 576 | 827 | 680 | 899 |

FIGURE 4. Acetylation stoichiometry is detected and quantified across a full range of values. Light and Δ5-acetic anhydride-labeled BSA protein digests were mixed at different ratios corresponding to 1–99% acetylation followed by MS analysis and quantitation. A, MS/MS spectrum for the acetylated BSA peptide, ALK(Ac)AWSVAR, is shown. (Product ions shown are charge 1⁺.) B–D, MS1 spectrum showing light and heavy forms of the peptide in A (charge 2⁺) corresponding to 10, 50, 90% acetylation, respectively, with a mass shift of ~2.5 m/z. E, scatterplot comparing percent volume input (light/total) to the corrected measured stoichiometry of the BSA peptide, ALK(Ac)AWSVAR. Stoichiometry values were corrected due to the isotopic purity of ~96%. F, linear regression analysis of all the acetyl peptides quantified shows the best fit line and high R² value.
either utilize or generate acetyl-CoA exhibited peptides with higher stoichiometry, such as acetyl-CoA carboxylase subunit \( \alpha \) (AccA), biotin carboxylase (AccC), formate acetyltransferase (PflB), phosphate acetyltransferase (Pta), acetyl-CoA synthetase (Acs), citrate synthase (GltA), and aldehyde-alcohol dehydrogenase (AdhE) (Fig. 8). Also, proteins in these pathways displayed a higher number of quantifiable peptides with above average stoichiometry. For example, pyruvate kinase II (PykA), phosphoenolpyruvate synthase (PpsA), Pta, aconitate hydratase 1 and 2 (AcnA and AcnB), \( \alpha \)-ketoglutarate dehydrogenase E1 component (SucA) and AdhE had at least six peptides with a stoichiometry higher than 7%.

Acetyl-CoA is at the junction of central metabolism. It is striking that most of the metabolic enzymes one reaction away from generating or utilizing acetyl-CoA have a higher acetylation stoichiometry in the \( \Delta \)CobB strain during stationary phase, suggesting that CobB is involved in many facets of acetyl-CoA utilization.
**Protein Acetylation Stoichiometry**

*Metabolite Analysis Reveals Increased Levels of Acetyl Phosphate in the CobB Mutant—NAD⁺/H⁺-dependent protein deacetylases (sirtuins) regulate many metabolic enzymes in prokaryotes and eukaryotes (35). In prokaryotes, CobB regulates acetyl-CoA synthetase, which is required for cell growth on acetate (30, 31). Activation of Acs by reversible acetylation is thought to be essential for "the acetate switch" (36). This transition occurs when bacterial cells have depleted carbon sources such as glucose and begin to utilize environmental acetate for growth. If flux through Acs is inhibited, for example by acetylation, then acetate can be diverted through the acetate kinase, Pta pathway to generate acetyl-CoA (Fig. 8). The higher stoichiometry as well as multiple acetylation sites on Pta suggest an inhibitory effect on protein function. Because acetate is rerouted through the acetate kinase-Pta pathway due to Acs inhibition, we speculated that there would be a buildup of acetyl phosphate if Pta function were inhibited by acetylation. To test this, we measured acetyl phosphate in the WT and ΔCobB strains. Indeed, there was a significant increase of acetyl phosphate in the CobB mutant (Fig. 8). This suggests that strains lacking CobB have an inability to efficiently utilize acetyl phosphate in the ΔCobB strain, consistent with inhibition of Pta activity.

**DISCUSSION**

In this work, we developed a novel method to quantify site-specific acetylation stoichiometry across an entire proteome. This method utilizes isotopic acetic anhydride to chemically acetylate all unmodified lysines in a protein sample. Coupling this protein chemistry to high resolution mass spectrometry and novel bioinformatic data processing, acetylation stoichiometry was quantified. Applying this method to bacterial cells, we describe the effects of the bacterial sirtuin member, CobB, on site-specific and global acetylation.

**Site-specific Acetylation Stoichiometry**—One role of the NAD⁺/H⁺-dependent sirtuins is the regulation of metabolism by deacetylation of key transcription factors and metabolic enzymes themselves (6). CobB deacetylates acetyl-CoA synthetase at a conserved lysine residue, thereby activating Acs (30, 31). Acs forms acetyl-CoA from acetate and ATP, thus allowing bacterial cells to utilize alternative carbon sources for energy when glucose is depleted. The active site lysine was not identified in this study, likely due to the small size of the peptide (<350 m/z) upon trypsin cleavage. We did, however, identify and quantify another acetyl site on Acs, Lys-348, which displayed 32.5% acetylation in ΔCobB and 8.3% in WT. Further investigation is needed to determine whether this site is critical for Acs activity as it lies at the dimeric interface.

**Metabolic enzymes of the TCA cycle, pyruvate metabolism, and glycolysis were also highly enriched in acetylation, most notably in the number of acetyl-lysine sites. Of these enriched metabolic pathways, enzymes one reaction away from acetyl-CoA displayed higher stoichiometry, including Pta, AccA, AccC, GltA, AdhE, and Msa (Fig. 8). It is noteworthy that the enzymes, which utilize or generate acetyl-CoA, could also be...**

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**FIGURE 6. Dynamics of acetylation stoichiometry in WT and ΔCobB strains.** Stoichiometry line plot demonstrates peptide level stoichiometry in WT and ΔCobB and the magnitude change between strains. Peptides are shown that have a higher stoichiometry in WT (black lines) and ΔCobB (red lines).
regulated by reversible acetylation. This observation coupled with the known inhibition of Acs, led us to examine and demonstrate increased levels of acetyl phosphate in the CobB-deficient strain, suggesting an inhibitory effect of acetylation on Pta activity.

Of the group 3 peptides (those showing substantially increased acetylation in the /H9004 CobB), RpsP, TrpS, Def, and Gpt had acetylation sites with the largest magnitude change in the CobB mutant. RpsP had a magnitude difference of 47% acetylation between the WT and /H9004 CobB strains. As a component of the 30 S ribosome, RpsP interacts with the RNA component of the ribosome. The /H9280-amino group of Lys-46 appears to make a critical contact with the 2/H11032-OH of a guanine nucleotide of the RNA (37). Upon acetylation, this contact could destabilize the interaction between RpsP and RNA. TrpS generates tryptophanyl-tRNA that is used for protein synthesis, and Lys-312 has a magnitude difference of 64% acetylation between WT and /H9004. There is not a solved crystal structure for the E. coli form; however, modeling with the Yersinia pestis variant shows that the analogous Lys site, Lys-324, is 2.7 Å away from the backbone oxygen of Pro-86, which likely forms H-bonding contact between the two residues (38, 39). Deformylase removes the formyl group from the N-terminal methionine, and the peptide containing Lys-158/161 has a magnitude change of 39%. These sites are on the surface of the protein, and it is not known whether they are critical for protein-protein interaction.

Xanthine phosphoribosyltransferase is involved in purine metabolism. The acetyl site, Lys-30, has a magnitude change of 27% between WT and /H9004 strains. The crystal structure reveals this site is also surface-exposed, and it is unknown whether it plays a role in interactions (40).

**Global Acetylation Stoichiometry**—The ability to measure acetyl-lysine stoichiometry allowed us to estimate the amount of total acetate attached to protein. The average global acetyl stoichiometry was ~7%, and we determined stoichiometry for ~30% of the quantifiable lysines in the E. coli proteome. Using these values and the known calculated values of E. coli for cell volume (BNID 100004), protein concentration (BNID 108263), and the frequency of lysines present in E. coli (BNID 100636) (13), we estimated the amount of acetate on protein to be between 1.7 and 2.8 mM. This is striking because the concentration of acetyl-CoA is estimated to be less than 500 μM in...
E. coli (15). This analysis reveals the high amount of acetate stored on proteins, although future experiments will be needed to assess the turnover rate with respect to acetyl-CoA and acetyl phosphate consumption.

Stoichiometry Versus Relative Quantitation—Knowledge of stoichiometry is a key piece of information to accurately interpret the functional roles of acetylation. Current acetyl proteomic methods rely on relative changes of peptides across conditions. In most cases, researchers have used an arbitrary 2-fold change cutoff as meaningful (4, 11), which can greatly skew interpretation. For example, in our study, the acetyl peptide from glutathione reductase (Gor) containing Lys-105/Lys-112 has a \( /H9004 \) CobB/WT fold change of 9.3. When the stoichiometry is considered, this peptide is 0.5% acetylated in WT and 4.3% in \( /H9004 \) CobB. This results in a magnitude change of 3.8% acetylation between the two conditions, which might only reflect biological variation. For the converse example, the acetyl peptide from 3-phosphoshikimate 1-carboxyvinyltransferase (AroA) containing Lys-354 has a \( /H9004 \) CobB/WT fold change of 0.6. However, the stoichiometry of this peptide is 38.8 and 69.5% for \( /H9004 \) CobB and WT, respectively (Fig. 6). This is a magnitude difference of 30.7% acetylation and is likely to have more of an impact on overall protein function than the former example.

Unique Features of Methodology—Major advantages/improvements of the presented method include the following. 1) Quantitation is independent of protein level because stoichiometry depends on the total protein in a given sample. This
elimates the requirement to normalize to total protein amount. 2) Stoichiometry as well as relative changes can be determined using this method. 3) There is no need for immunoenrichment of acetyl peptides. This avoids any issues toward antibody enrichment bias introduced in the analysis. This issue is circumvented with the pairing algorithm of PVIEW by performing an in silico enrichment of acetyl peptides. 4) Low amount of starting material is used. Because of the minimal sample processing, less protein can be used, facilitating the analysis of limited and difficult to obtain samples. 5) This method has broad applicability, from an isolated purified protein to cell culture or tissue sample.

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REFERENCES

1. Xiong, Y., and Guan, K. L. (2012) Mechanistic insights into the regulation of metabolic enzymes by acetylation. J. Cell Biol. 198, 155–164
2. Yang, X. I., and Seto, E. (2008) Lysine acetylation: codified crosstalk with other posttranslational modifications. Mol. cell 31, 449–461
3. Tanner, K. G., Trievel, R. C., Kuo, M. H., Howard, R. M., Berger, S. L., Allis, C. D., Marmorstein, R., and Denu, J. M. (1999) Catalytic mechanism and function of invariant glutamic acid 173 from the histone acetyltransferase GCN5 transcriptional coactivator. J. Biol. Chem. 274, 18157–18160
4. Weinert, B. T., Isamantavicius, V., Wagner, S. A., Schöll, C., Gummesson, B., Beli, P., Nyström, T., and Choudhary, C. (2013) Acetyl phosphate is a critical determinant of lysine acetylation in E. coli. Mol. Cell. 51, 265–272
5. Wagner, G. R., and Payne, R. M. (2013) Widespread and enzyme-independent Nε-acetylation and Nε-succinylation of proteins in the chemical conditions of the mitochondrial matrix. J. Biol. Chem. 288, 29036–29045
6. Feldman, J. L., Dittenhafer-Reed, K. E., and Denu, J. M. (2012) Sir2 protein function in the mitochondrial matrix. J. Biol. Chem. 287, 42419–42427
7. Kim, S. C., Sprung, R., Chen, Y., Xu, Y., Ball, H., Pei, J., Cheng, T., Kho, Y., Xiao, H., Xiao, L., Grishin, N. V., White, M., Yang, X. J., and Zhao, Y. (2006) Substrate and functional diversity of lysine acetylation revealed by a proteomics survey. Mol. Cell. 23, 607–618
8. Choudhary, C., Kumar, C., Gnadt, F., Nielsen, M. L., Rehman, M., Walther, T. C., Olsen, J. V., and Mann, M. (2009) Lysine acetylation targets protein complexes and co-regulates major cellular functions. Science 325, 834–840
9. Hebert, A. S., Dittenhafer-Reed, K. E., and Denu, J. M. (2012) Substrate and functional diversity of lysine acetylation revealed by a proteomics survey. Mol. Cell. 23, 607–618
10. Beker-Jensen, D. B., Secher, A., Skovgaard, T., kelstrup, C. D., Dmytryiev, A., Choudhary, C., Lundby, C., and Olsen, J. V. (2012) Proteomic analysis of lysine acetylation sites in rat tissues reveals organ specificity and subcellular patterns. Cell Reports 2, 419–431
11. Rardin, M. J., Newman, J. C., Miller, M. L., Jensen, L. J., Gnad, F., Cox, J., Jensen, T. S., Nigg, E. A., Brunak, S., and Mann, M. (2010) Quantitative phosphoproteomics reveals widespread full phosphorylation site occupancy during mitosis. Sci. Signal. 3, ra3
12. Wu, R., Haas, W., Dephoure, N., Huttlin, E. L., Zhai, B., Sowa, M. E., and Gygi, S. P. (2011) A large-scale method to measure absolute protein phosphorylation stoichiometries. Nat. Methods 8, 677–683
13. Steen, H., Jeganathan, J. A., Springer, M., and Kirschner, M. W. (2005) Stable isotope-free relative and absolute quantification of protein phosphorylation stoichiometry by MS. Proc. Natl. Acad. Sci. U.S.A. 102, 3948–3953
14. Olsen, J. V., and Mann, M. (2013) Status of large-scale analysis of post-translational modifications by mass spectrometry. Mol. Cell. Proteomics 12, 3444–3452
15. Danchin, A., Dondon, L., and Daniel, J. (1984) Metabolic alterations mediated by 2-ketobutyrate in Escherichia coli K12. Mol. Gen. Genet. 193, 473–478
16. Olsen, J. V., Vermeulen, M., Santamaria, A., Kumar, C., Miller, M. L., Jensen, L. J., Gnadt, F., Cox, J., Jensen, T. S., Nigg, E. A., Brunak, S., and Mann, M. (2010) Quantitative phosphoproteomics reveals widespread full phosphorylation site occupancy during mitosis. Sci. Signal. 3, ra3
17. Wu, R., Haas, W., Dephoure, N., Huttlin, E. L., Zhai, B., Sowa, M. E., and Gygi, S. P. (2011) A large-scale method to measure absolute protein phosphorylation stoichiometries. Nat. Methods 8, 677–683
18. Steen, H., Jeganathan, J. A., Springer, M., and Kirschner, M. W. (2005) Stable isotope-free relative and absolute quantification of protein phosphorylation stoichiometry by MS. Proc. Natl. Acad. Sci. U.S.A. 102, 3948–3953
19. Weinert, B. T., Isamantavicius, V., Moustafa, T., Schöll, C., Wagner, S. A., Magnes, C., Zechner, R., and Choudhary, C. (2014) Acetylation dynamics and stoichiometry in Saccharomyces cerevisiae. Mol. Syst. Biol. 10, 716
20. Hallows, W. C., Lee, S., and Denu, J. M. (2006) Sirtuins deacetylate and activate mammalian acetyl-CoA synthetases. Proc. Natl. Acad. Sci. U.S.A. 103, 10230–10235
21. Riarson, J. F., and Vallee, B. L. (1972) [41] acetylation. Methods Enzymol. 25, 494–499
22. Khan, Z., Minim, B., Bloom, J. S., Ruse, C., Caudy, A. A., Kruglyak, L., Singh, M., Perlman, D. H., and Tavazoie, S. (2011) Accurate proteome-wide protein quantification from high resolution 13N mass spectra. Genome Biol. 12, R122
23. Khan, Z., Bloom, J. S., Garcia, B. A., Singh, M., and Kruglyak, L. (2009) Protein quantification across hundreds of experimental conditions. Proc. Natl. Acad. Sci. U.S.A. 106, 15544–15548
24. Huang da, W., Sherman, B. T., and Lempicki, R. A. (2009) Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources. Nat. Protoc. 4, 44–57
25. Huang da, W., Sherman, B. T., and Lempicki, R. A. (2009) Bioinformatics enrichment tools: paths toward the comprehensive functional analysis of large gene lists. Nucleic Acids Res. 37, 1–13
26. Franceschini, A., Szklarczyk, D., Frankild, S., Kuhn, M., Simonovic, M., Roth, A., Lin, J., Minguez, P., Bork, P., von Mering, C., and Jensen, L. J. (2013) STRING version 9.1: protein-protein interaction networks, with increased coverage and integration. Nucleic Acids Res. 41, D808–D815
27. Snel, B., Lehmann, G., Bork, P., and Huynen, M. A. (2000) STRING: a web-server to retrieve and display the repeatedly occurring neighbour relationships. Nucleic Acids Res. 28, 3212–3214
28. Saito, R., Smoot, M. E., Ono, K., Ruscheinski, J., Wang, P. L., Livia, S., Pico, A. R., Bader, G. D., and Ideker, T. (2012) A travel guide to Cytoscape plugins. Nat. Methods 9, 1069–1076
29. Lu, W., Claasquin, M. F., Melamud, E., Amador-Noguez, D., Caudy, A. A., and Rabinowitz, J. D. (2010) Metabolomic analysis via reversed-phase ion-pairing liquid chromatography coupled to a stand-alone Orbitrap mass spectrometer. Anal. Chem. 82, 3212–3221
30. Starai, V. J., Celic, I., Cole, R. N., Boeke, J. D., and Esclante-Semerena, J. C. (2002) Sir2-dependent activation of acetyl-CoA synthetase by deacylation of active lysine. Science 298, 2390–2392
31. Starai, V. J., Takahashi, H., Boeke, J. D., and Esclante-Semerena, J. C. (2003) Short-chain fatty acid activation by acyl-coenzyme A synthetases requires Sir2 protein function in Salmonella enterica and Saccharomyces cerevisiae. Genetics 163, 545–555
32. Wang, Q., Zhang, Y., Yang, C., Xiong, H., Lin, Y., Yao, J., Li, H., Xie, L., Zhao, W., Yao, Y., Ning, Z. B., Zeng, R., Xiong, Y., Guan, K. L., Zhao, S., and Zhao, G. P. (2010) Acetylation of metabolic enzymes coordinates carbon source utilization and metabolic flux. Science 327, 1004–1007
33. Olsen, J. V., Ong, S. E., and Mann, M. (2004) Trypsin clefts exclusively C-terminal to arginine and lysine residues. Mol. Cell. Proteomics 3, 608–614

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34. Beck, H. C., Nielsen, E. C., Matthiesen, R., Jensen, L. H., Sehested, M., Finn, P., Grauslund, M., Hansen, A. M., and Jensen, O. N. (2006) Quantitative proteomic analysis of post-translational modifications of human histones. *Mol. Cell. Proteomics* 5, 1314–1325

35. Schwer, B., and Verdin, E. (2008) Conserved metabolic regulatory functions of sirtuins. *Cell Metab.* 7, 104–112

36. Wolfe, A. J. (2005) The acetate switch. *Microbiol. Mol. Biol. Rev.* 69, 12–50

37. Pulk, A., and Cate, J. H. (2013) Control of ribosomal subunit rotation by elongation factor G. *Science* 340, 1235970

38. Kiefer, F., Arnold, K., Künzli, M., Bordoli, L., and Schwede, T. (2009) The SWISS-MODEL repository and associated resources. *Nucleic Acids Res.* 37, D387–D392

39. Kopp, J., and Schwede, T. (2004) The SWISS-MODEL repository of annotated three-dimensional protein structure homology models. *Nucleic Acids Res.* 32, D230–D234

40. Vos, S., Parry, R. J., Burns, M. R., de Jersey, J., and Martin, J. L. (1998) Structures of free and complexed forms of *Escherichia coli* xanthine-guanine phosphoribosyltransferase. *J. Mol. Biol.* 282, 875–889