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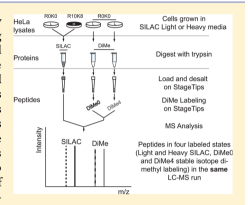
Comparing SILAC- and Stable Isotope Dimethyl-Labeling Approaches for Quantitative Proteomics

Ho-Tak Lau, Hyong Won Suh, Martin Golkowski, and Shao-En Ong*

School of Medicine, Department of Pharmacology, University of Washington, Box 357280, Seattle, Washington 98195, United States

Supporting Information

ABSTRACT: Stable isotope labeling is widely used to encode and quantify proteins in mass-spectrometry-based proteomics. We compared metabolic labeling with stable isotope labeling by amino acids in cell culture (SILAC) and chemical labeling by stable isotope dimethyl labeling and find that they have comparable accuracy and quantitative dynamic range in unfractionated proteome analyses and affinity pull-down experiments. Analyzing SILAC- and dimethyl-labeled samples together in single liquid chromatography—mass spectrometric analyses minimizes differences under analytical conditions, allowing comparisons of quantitative errors introduced during sample processing. We find that SILAC is more reproducible than dimethyl labeling. Because proteins from metabolically labeled populations can be combined before proteolytic digestion, SILAC is particularly suited to studies with extensive sample processing, such as fractionation and enrichment of peptides with post-translational modifications. We compared both methods in pull-down experiments using a kinase inhibitor, dasatinib, and tagged GRB2-SH2



protein as affinity baits. We describe a StageTip dimethyl-labeling protocol that we applied to in-solution and in-gel protein digests. Comparing the impact of post-digest isotopic labeling on quantitative accuracy, we demonstrate how specific experimental designs can benefit most from metabolic labeling approaches like SILAC and situations where chemical labeling by stable isotope-dimethyl labeling can be a practical alternative.

KEYWORDS: SILAC, stable isotope dimethyl labeling, quantitative proteomics, metabolic labeling

■ INTRODUCTION

Over the past decade, mass spectrometry (MS)-based proteomics has become the primary analytical technology to study proteins in complex mixtures. Improvements in nanoscale liquid chromatography coupled to MS (LC-MS) analyses, development of faster and more sensitive MS instrumentation, and identification and quantification of proteins and their posttranslational modifications (PTMs) by MS have contributed to the widespread adoption of MS-based proteomics. Quantifying protein abundance in complex mixtures can be achieved with stable isotope labeling or label-free approaches.^{1,2} Because label-free quantification often requires larger numbers of analytical replicates and depends on highly consistent sample processing,³ stable isotope-labeling approaches are still more widely used in quantitative proteomics studies. The latter allows multiple labeled samples to be combined and analyzed as a single sample, reducing run-to-run variability from sample injection, ionization suppression, or stochastic sampling of complex mixtures by the MS instrument that may differentially impact individual MS runs; such factors, if not addressed by proper experimental design, can negatively affect comparative analyses.

In 1999, Oda et al. used ¹⁵N-enriched medium to metabolically label and quantify changes in protein expression and site-specific phosphorylation in yeast.⁴ Metabolic labeling through the incorporation of stable ¹³C and ¹⁵N enriched

amino acids, 5-7 particularly arginine and lysine, is especially suited to whole-proteome labeling of live cells, proteolytic digestion with trypsin, and downstream quantification by MS. Because SILAC is a simple and robust method to encode cell populations with quantifiable labels, it has been widely adopted in cell culture systems and has been creatively adapted for use in model organisms such as mice, newts, worms and zebrafish.¹¹ Recent developments include neutron encoding SILAC that uses mass defects of different stable isotopes within the same amino acid to encode multiple cell states and allows higher multiplexing. 12 SILAC is known to be a very accurate and precise quantitative method, 13 likely because it allows the mixing of differentially labeled samples early in the experimental workflow, reducing variable sample losses from each experimental step. The primary limitation, however, is that SILAC cannot label nondividing cells and human samples and is expensive to apply in small mammals.

Chemical labeling methods target specific reactive groups, such as primary amines, in proteins or peptides to introduce chemical tags encoded with differential masses. Isobaric tagging strategies, including the commercially available iTRAQ¹⁴ and TMT¹⁵ reagents, are increasingly common because they allow multiplexing of up to 10 samples with high-resolution

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instruments; the reporter ions generated upon fragmentation are quantified at the MS2 level. While multiplexing has been shown to provide increased sampling depth when compared with an MS1-quantification approach, ¹⁶ several groups have described compromised quantitative accuracy with isobaric tags in complex mixtures from coisolation and cofragmentation of interfering precursor ions. ^{17–19} Methods to mitigate this effect have been proposed; these involve an additional MS3 scan to obtain quantitative reporter ions and would therefore affect the instrument duty cycle ²⁰ or require specialized equipment. ²¹ A practical issue is the high cost of iTRAQ and TMT reagents for labeling milligrams of peptides, as is often required in phosphoproteomics ^{16,22} and studies of substoichiometric protein PTMs.

Stable isotope dimethyl labeling is a chemical labeling method for quantitative proteomics introduced in 2003;²³ it labels the N-terminus of proteins/peptides and ε -amino group of lysine through reductive amination. The reductive dimethylation reaction is rapid and specific²³ and has the added advantage that required reagents are commercially available at costs significantly lower than multiplexed chemical labeling reagents.^{24,25} Like other chemical labeling methods, stable isotope dimethyl labeling is applicable to a wide variety of samples. It was recently used in a large-scale phosphoproteomics study from mouse liver tissues.²⁴ Despite these attractive features, dimethyl labeling has not been as widely adopted as other commercial chemical labeling reagents. We sought to compare the performance of SILAC and stable isotope dimethyl labeling in quantitative proteomics applications.

We first compared the labeling efficiencies of three dimethyl labeling workflows using in-solution labeling ^{23,25} and postlabeling desalting on C18 StageTip²⁶ as well as a modified oncolumn labeling protocol²⁵ using the C18 StageTip. With our optimized on-StageTip dimethyl-labeling protocol, we used SILAC-labeled HeLa lysate as both source material and internal standards²⁷ to compare the dynamic range of quantification, precision, and repeatability of SILAC and dimethyl labeling. We demonstrate that SILAC provides improved quantification accuracy because it allows combined sample processing in a much earlier stage in the experimental workflow compared with chemical labeling approaches. We compare SILAC- and dimethyl-labeling approaches in biochemical pull-down experiments using an immobilized version of small molecule kinase inhibitor dasatinib as well as a recombinant protein interaction domain (GRB2-SH2) as affinity baits. Our study demonstrates that SILAC and dimethyl labeling have comparable performance in quantitative proteomics applications and highlights specific reasons for choosing one method over another.

MATERIALS AND METHODS

Cell Culture and SILAC Labeling

HeLa cells were cultured in SILAC Dulbecco's modified Eagle's medium (DMEM) as previously described²⁸ at 37 °C, 95% relative humidity, and 5% CO₂. SILAC custom DMEM lacking lysine and arginine (Caisson Laboratories, Logan, UT) was supplemented with 10% dialyzed FBS (Sigma) and either light (L-lysine and L-arginine (Fisher)) or heavy ([\begin{subarray}{c} \begin{subarray}{c} \begi

Protein Extraction and Trypsin Digestion

With the exception of the affinity capture experiments, HeLa cells were lysed in 6 M guanidine hydrochloride (Gnd-HCl). Proteins were reduced and alkylated with 1 mM tris(2-carboxyethyl)phosphine (TCEP) and 2 mM chloroacetamide (CAM), then diluted seven-fold with 100 mM triethylammonium bicarbonate (TEAB) (Sigma). Proteins were digested at pH 8.5 with 1:50 (w/w) trypsin (Promega, V5113)/substrate and incubated overnight at 37 °C with shaking. Another portion of 1:50 trypsin/substrate was added to the digests on the following day and incubated for an additional 2 h. Digested peptides were desalted using C18 StageTips. 26

Chemical Labeling

The on-column stable isotope dimethyl-labeling protocol²⁵ was adapted to label StageTip-bound peptides with CH₂O (Sigma) or C²H₂O (CIL). Labeling reagent comprising 17.5 μ L of 4% (v/v) CH₂O/C²H₂O formaldehyde, 17.5 μ L of 0.6 M sodium cyanoborohydride (Sigma), 70 μ L of 50 mM Na₂PO₄ (Fisher), and 245 μ L of 50 mM Na₂HPO₄ (Fisher) was freshly prepared for each sample. Immediately after the C18 StageTip desalting step with 50 μ L of StageTip solvent A (5% acetonitrile, 0.1% trifluoroacetic acid in H₂O), peptides were labeled by applying 300 μ L of labeling reagent to the StageTip and spinning through at ~2200g for 10 min. The labeling step was followed by a single wash with 100 μ L of StageTip solvent A. In-solution stable isotope dimethyl labeling was performed as described.²⁵

Ratio Mixing Experiments

Four populations of HeLa cells were SILAC-labeled and lysed in 6 M Gnd-HCl as previously described. For each mixed ratio experiment, 20 µg of light or heavy protein was reduced, alkylated, and digested and used as a common pool. Two light samples were dimethyl-labeled (either CH₂O or C²H₂O) on StageTip. After elution from StageTips, peptides were mixed in seven L/H ratios (10:1, 5:1, 2:1, 1:1, 1:2, 1:5, 1:10). For each MS run, at least 0.1 μ g of each peptide sample was injected, for example, in ratio 1:10 (L/H), 0.1 μ g of light sample was mixed with 1 μ g of heavy sample, and in 1:5, 0.1 μ g of light sample was mixed with 0.5 μ g of heavy sample. Where mentioned in the text, forward and reverse label-swap experiments refer to replicate experiments performed where experimental state and stable isotope labels are swapped to allow systematic errors due to labeling to be identified. SILAC- and dimethyl-labeled samples were then mixed for the combined runs, that is, containing peptides from both labeling approaches or analyzed separately as standalone SILAC or dimethyl-labeling runs, as illustrated in Figure 2A.

NanoLC-MS/MS Analysis and Quantification

All samples were unfractionated unless otherwise indicated. Samples were loaded to self-pulled (P2000 Sutter Laser puller, Sutter Instrument, Novato, CA) 360 μ m OD × 75 μ m ID 10 cm columns with a 10 μ m tip and packed with 3 μ m Reprosil C18 resin (Dr. Maisch, Germany) using a pressure cell (NextAdvance, Averill Park, NY). Peptides were analyzed with 90 min gradients of 3–35% acetonitrile at 200 nL/min nanoLC–MS (Thermo Dionex RSLCnano, Sunnyvale, CA) on an Orbitrap Elite (Thermo, Bremen Germany). Orbitrap FTMS spectra ($R=30\,000$ at $400\,m/z;\,m/z\,350-1600;\,3e6$ target; max 500 ms ion injection time) and Top15 data dependent CID MS/MS spectra (1e4 target; max 250 ms injection time) were collected with dynamic exclusion for 180 s

and an exclusion list size of 500. The normalized collision energy applied for CID was 35% for 10 ms.

Data Analysis

MaxQuant v.1.3.0.5²⁹ and the associated Andromeda search engine³⁰ was used to search a Uniprot human database (July 2012 with 88 849 entries). Search parameters used were: Trypsin/P with two missed cleavages, fixed carbamidomethylated cysteines, and oxidized methionines as a variable modification. Initial FTMS and ITMS MS/MS tolerances were set at 20 ppm and 0.5 Da, respectively. Protein and peptide FDRs were 1%, minimum peptide length was seven amino acids, and a minimum of two peptide ratios were required to quantify a protein. Resulting data were analyzed with the Perseus³¹ and R environments.

GRB2-SH2 Affinity Purification

The SH2 domain (residues 60-152) of Human Growth receptor-bound protein 2 (GRB2) was cloned with a Cterminal FLAG epitope tag in a pET28a vector for bacterial expression. 6xHis-GRB2-SH2-FLAG was expressed by inducing BL21Star cells with 0.1 mM IPTG at 37 °C for 3 h in a shaking incubator at 250 rpm. Expressed protein was purified using standard His-tag purification procedures with NTA-agarose. Cells were sonicated with a microprobe sonicator tip in a buffer containing 500 mM NaCl, 25 mM Tris pH 7.0, 10% glycerol, 5 mM β -mercaptoethanol, and HALT protease inhibitors (Pierce) and centrifuging at 14 000g for 45 min. The supernatant containing the soluble protein was incubated with 200 μL (50% v/v slurry) of NTA-agarose beads (Qiagen) for 1 h in a batch purification. Two washes with 10 bead volumes of wash buffer containing 500 mM NaCl, 25 mM Tris pH 7.0, 5 mM β -mercaptoethanol, and either 10 mM or 25 mM imidazole, pH 7.4 were performed before eluting in buffer containing 200 mM imidazole, pH 7.4. The eluted protein was dialyzed against 50 mM Tris pH 8.0 to remove imidazole. The GRB2-SH2-FLAG bait protein was aliquoted, snap-frozen in liquid nitrogen, and stored at −80 °C.

HeLa cells were cultured in SILAC DMEM medium as previously described. After five cell doublings, cells were washed with PBS and serum-starved for 6 h. After serum starvation, the cells were stimulated with 150 ng/mL of epidermal growth factor (EGF) for 10 min, while the control population was treated with an equal volume of PBS. Cells were lysed with modified RIPA buffer (modRIPA)(50 mM Tris-HCl, pH 7.8, 150 mM NaCl, 1% NP-40, 0.25% sodium deoxycholate, 1× Halt protease inhibitor (Pierce), and 1× Halt phosphatase inhibitor (Pierce)). Equal amounts of light and heavy HeLa lysate were added to 1500 pmol of FLAG-tagged GRB2-SH2 fusion protein immobilized on 70 µL of anti-FLAG M2 Magnetic Beads slurry (50% w/v) (Sigma), respectively. After 90 min on an end-over-end rotator at 4 °C, the beads were washed three times with modRIPA; the first wash was aspirated and discarded; in the second wash, the SILAC samples were combined by resuspending beads in the SILAC heavy tube with 1 mL of modRIPA and transferring the entire volume to the SILAC light tube; this wash was aspirated, and a third and final 1 mL volume of modRIPA was used to wash the combined beads. Bead-bound proteins were reduced and alkylated as previously described, heated to 70 °C in LDS sample buffer (Life Technologies), and resolved on a Bolt 4-12% Bis-Tris Plus Gel (Life Technologies). The gel was stained with Coomassie Blue for 1 h and destained overnight with water. Each sample lane was divided into five slices and cut further into small 1 mm cubes. Gel cubes were destained with 1:1 (v/v) of 50 mM ammonium bicarbonate and ethanol. After destaining, ethanol was used to dehydrate the gel cubes. Proteins were digested with trypsin (13 ng/ μ L) in triethylammonium bicarbonate overnight. In-gel digestion was performed as previously described. Desalting, reductive dimethyl labeling, and LC–MS/MS analysis were performed as previously described.

Dasatinib Affinity Purification

Dasatinib affinity bait was synthesized as described.³³ HeLa lysates were prepared as described in the GRB2-SH2 domain pull-down (above). Lysates were preincubated with DMSO (control) or a dasatinib solution in DMSO (competitor, 50 µM, final) for 20 min at 4 °C on an end-over-end rotator. A 50% (v/ v) bead slurry of the dasatinib affinity matrix (~50 µM immobilized dasatinib final) was then added to both tubes of control and soluble dasatinib-treated lysates. The mixtures were mixed in an end-over-end rotator at 4 °C for 3 h. Pull-downs were washed with buffer containing 50 mM Tris-HCl, pH 7.8, and 150 mM NaCl, as previously described. Light and heavy SILAC samples were first washed separately with 1 mL of modRIPA to remove most of the soluble competitor. Beads from the SILAC light and heavy samples were then combined in the second wash and processed as a single sample thereafter. Note that stable isotope dimethyl-labeling samples were processed separately until after the chemical labeling step. Captured proteins were reduced (1 mM TCEP) and alkylated (2 mM CAM) on-bead, digested, and desalted on C18 StageTips as described.³³ Stable isotope dimethyl labeling was performed on C18 StageTips as previously described. SILAC and stable isotope dimethyl labeling samples were combined and analyzed with LC-MS/MS.

■ RESULTS

Previous comparisons of different chemical labeling methods to SILAC have analyzed these workflows in separate MS runs^{22,34} and found chemical isotopic labeling methods to be very comparable to SILAC. Additionally, comparisons with label-free quantification with spectral counts^{35,36} or extracted MS peak intensities³⁷ have recently been reported. These analytical comparisons may be affected in several stages from the preparation of the peptide sample through to the detection of ions in the mass spectrometer but particularly in sample injection and loading, ionization conditions, and stochastic sampling in data-dependent MS. We decided, therefore, to combine SILAC-labeled and dimethyl-labeled samples for single LC–MS/MS analyses to minimize analytical differences.

Comparing in-Solution and On-StageTip Dimethyl Labeling

We sought to develop a convenient and robust dimethyllabeling workflow compatible with peptide amounts typical in affinity pull-downs and gel-based separations. Hsu et al.'s original dimethyl-labeling protocol labeled peptides insolution.²³ Boersema et al. subsequently developed protocols for on-column and online labeling of peptides.²⁵ Because each workflow may have varying labeling efficiencies and yields, we compared these different dimethyl labeling workflows for protein identification and quantification.

We prepared a single trypsin digestion of HeLa lysate and divided this into aliquots for technical replicates. Digested proteins were labeled by dimethyl-labeling reagent with two different isotopic compositions (CH_2O or C^2H_2O), and the

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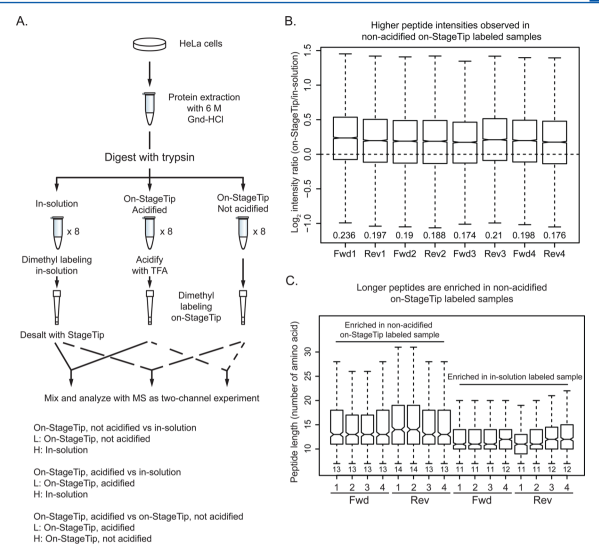


Figure 1. Comparison of dimethyl labeling workflows. (A) Experimental design. HeLa cells were lysed in 6 M Gnd-HCl and proteins digested with trypsin. The digested peptides were aliquoted into eight tubes as technical replicates for each labeling workflow. Comparisons of two workflows in two-state L/H with label swaps were performed for a total of eight replicate experiments (four "forward" experiments and four "reverse" experiments). Forward experiments are listed. After labeling and desalting, samples were mixed 1:1 and analyzed by MS. (B) Higher peptide intensities observed in nonacidified on-StageTip versus in-solution dimethyl labeling. Median ratios for each replicate are at the bottom of each boxplot. The boxplots were normalized, as we observed a consistent pattern between forward and reverse experiments that is associated with the heavy labeling reagent (log 2 C²H₂O/CH₂O = 0.0239) but not the workflow (Supplementary Figure 1A in the Supporting Information). This normalization factor was derived from all 24 runs in this data set. (C) Differences in length of peptides observed between workflows. Median numbers of amino acid residues are listed at the bottom of each boxplot.

signal intensities of peptides from different workflows were compared (Figure 1A).

We observed higher peptide signal intensities in on-StageTip labeled peptides than the in-solution labeled peptides (Supplementary Figure 1A in the Supporting Information). Surprisingly, when comparing the two on-StageTip labeling workflows, we observed slightly higher signal intensity from the nonacidified (pH \sim 7.5) samples (Supplementary Figure 1A in the Supporting Information) over the two other samples. We expect that this difference is most likely due to the selectivity of the C18 Empore resin to peptides at different pH. The greatest difference in peptide signals, therefore, was observed when we compared on-StageTip labeling without acidification to insolution labeling (Figure 1B).

Because we observed differences in peptide intensities between workflows, we also examined the lengths and hydrophobicities of peptides in our data set that had the largest differences in MS signal. For each MS run, the 500 peptides with highest or lowest ratios represent peptides enriched in the nonacidified on-StageTip labeled workflow over the in-solution labeling workflow, respectively. We found that peptides enriched in nonacidified on-StageTip labeled samples were higher in intensity (Figure 1B), longer (Figure 1C), and more hydrophobic (Supplementary Figure 1C in the Supporting Information) than acidified on-StageTip or insolution labeling. In general, we recommend the on-StageTip labeling protocols for their convenience and because we observed equivalent, or even slightly better yields than insolution labeling (Supplemental Figure 1A in the Supporting Information). We used the nonacidified on-StageTip labeling workflow for the experiments described in this paper.

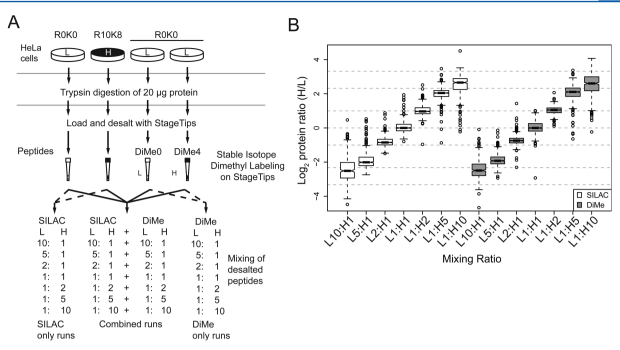


Figure 2. Comparison of protein quantification accuracy and dynamic range of SILAC and dimethyl labeling. (A) Experimental design. Four cell populations were grown separately in SILAC medium, as illustrated. SILAC samples are digested and desalted separately, serving as a control through all stages of sample preparation except for dimethyl labeling step. Samples were analyzed by MS in combined or SILAC/dimethyl labeling (DiMe) separated runs. DiMe0 and DiMe4, dimethyl labeling with CH_2O and C^2H_2O , respectively. (B) Boxplot of log_2 ratio of SILAC and dimethyl labeling combined runs. Results are combined from four replicates. Ratios are normalized so that the median of H1:L1s is centered at log_21 . The quantification ranges of SILAC-labeled peptides and dimethyl-labeled peptides are very similar, and both showed ratio compression. Dotted lines indicate the expected log ratios from log_2 10 to $log_2(1/10)$.

Quantification Accuracy and Dynamic Range of Dimethyl Labeling

We designed the experiment illustrated in Figure 2A to compare quantitative performance of SILAC and stable isotope dimethyl labeling. Cell lysates were prepared from four populations of SILAC-labeled HeLa (3 light: R0K0 and 1 heavy: R10K8). The lysates were digested separately with trypsin and desalted with StageTips. Two of the light StageTip bound peptides (R0K0) were further labeled with dimethyl labeling. After elution from the StageTips, peptides were mixed at seven different ratios (from L/H 10:1–1:10), as shown in Figure 2A. We analyzed SILAC- and dimethyl-labeled samples in single MS runs to minimize the variables introduced by the MS analyses.

We performed four combined SILAC and dimethyl labeling run experiments, yielding 801 \pm 84 and 821 \pm 43 proteins quantified for SILAC and dimethyl labeling, respectively. To ensure that increased complexity of the peptide mixture would not affect our measurement of quantitative reproducibility and accuracy, we also ran the same samples separately (Figure 2 and Supplementary Figure 2 in the Supporting Information). We identified a subset of our identified peptides with very low H/L SILAC ratios in SILAC and dimethyl-labeling combined runs (Supplementary Figure 3 in the Supporting Information) and found that these peptides (~7.5% of all were N-terminal acetylated and contained no lysines) are not labeled in aminedirected labeling strategies and may be wrongly classified as outliers if not appropriately handled in the data set. In agreement with a previous study,³⁴ our results demonstrated a comparable accuracy from the two methods (Figure 2B). The median L/H ratios were very close to the expected ratio for two-fold ratio mixes, but a mild ratio compression was observed

at L/H ratios of 5:1 and 1:5. At L/H ratio 10:1 and 1:10, however, the quantification accuracy was severely affected, limiting the dynamic range of both methods to six-fold differences in our analyses with unfractionated whole proteomes. From our analysis of SILAC- and dimethyl-labeled samples in separated runs, we verified that the ratio compression is not due to the single-run approach (Supplementary Figure 2 in the Supporting Information). We and others have previously reported the ratio compression phenomenon in SILAC. 35,37,38 Our results demonstrate that dimethyl labeling also suffers from ratio compression and has a similar dynamic range. The compression of stable isotopelabeling ratios is a general feature arising from the complexity of peptide mixtures in LC-MS analyses,³⁹ which is exacerbated by the increased complexity from stable isotope-labeled peptide pairs. 16 The application of longer separation gradients or orthogonal peptide fractionation would help to reduce such phenomenon by decreasing interference of quantified peaks from overlapping peptides and hence improving signal-to-noise ratios and quantitative accuracy; for instance, we observed a much wider dynamic range of quantification in our affinity pulldown experiments, described later in this manuscript.

Precision and Repeatability by SILAC and Dimethyl Labeling Quantification

In this study, we define precision as the variability within one experiment and repeatability as replicates performed on different days. It is often thought but has not been definitively shown that SILAC has a better precision and repeatability than chemical labeling methods because its workflow allows the mixing of samples in the stage of intact cells or, more commonly, after protein extraction. In contrast, in chemical labeling methods, samples can only be combined after the label

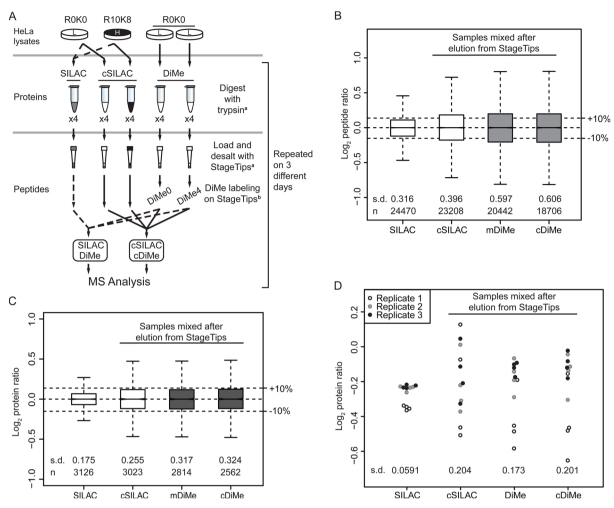


Figure 3. Comparison of precision and repeatability. (A) Experimental design. Four individual digestions were set up for each sample as illustrated. The full experiment was repeated in triplicate, producing 12 data points for each quantification mode considered. cSILAC denotes SILAC-labeled cells processed as in chemical labeling workflow, where samples were mixed after desalting on StageTips. (a,b) Steps where error was introduced in cSILAC and dimethyl labeling and error introduced in dimethyl labeling only, respectively. Precision (B and C) and repeatability (D) of SILAC workflow is higher than the chemical labeling workflow. (B) Peptide and (C) protein ratios from each experiment were normalized so that the medians are centered at Log₂ 1. Normalized ratios from the triplicate experiments were combined and illustrated in the boxplot. cSILAC and SILAC samples mixed after elution from StageTip. DiMe and cDiMe represent the same dimethyl-labeled peptide sample mixed with SILAC and cSILAC sample, respectively. (D) Repeatability. Each point represents the median protein ratio from each MS run. Replicates indicate the experiments performed on different days. s.d., standard deviation; n, number of peptides or proteins in the sample.

incorporation step, which is typically at the peptide level. The combined processing of SILAC-labeled samples at the intact protein level means that losses from sample handling affect all protein populations simultaneously. Because of that, metabolic labeling workflows should therefore be less prone to quantification inaccuracies and would be more reproducible when compared with chemical labeling workflows.

We wanted to directly compare the effect of the stage of sample mixing in SILAC- and dimethyl-labeling workflows on the precision and repeatability in quantitative proteomics applications using the experiment illustrated in Figure 3A. To reduce biological variability, we prepared a single batch of HeLa lysate, which was used for three experimental replicates performed on different days. For each experiment, four separate trypsin digestions were prepared from the lysate. This experimental design results in 12 sets of data (Figure 3 and Supplementary Figure 4 in the Supporting Information). To show the effects of mixing of samples early or late in the sample processing, we also processed a set of SILAC samples using the

dimethyl-labeling workflow (cSILAC in Figure 3). Furthermore, to ensure that the differences between SILAC and cSILAC are not due to variables introduced in separated MS runs, we mixed the same preparation of dimethyl-labeled samples to SILAC and cSILAC runs (DiMe and cDiMe in Figure 3, respectively). As shown in Figure 3B,C, the quantitative precision is improved when samples can be mixed before trypsin digestion (s.d. of log, SILAC ratio is 0.175 in contrast with 0.255, 0.317, and 0.324 of cSILAC, DiMe, and cDiMe, respectively). However, it should be noted that even though the chemical-labeling workflow leads to a lower precision in quantification, the first and third quartiles or 68% of the quantified peptide ratios are within 10% of the mixing ratio, which is still well within the typical quantitative errors (<20-25% CV) observed in targeted MS assays.⁴⁰ Surprisingly, repeatability of the SILAC workflow is nearly four times better than the chemically labeling workflow (Figure 3D, s.d. of SILAC = 0.0591 vs 0.204, 0.173, and 0.201 of cSILAC, DiMe, and cDiMe, respectively). We demonstrate, therefore,

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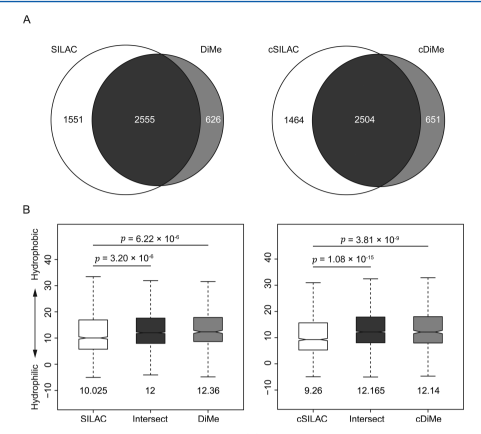


Figure 4. More hydrophilic peptides are identified in SILAC samples. (A) Venn diagrams of unique peptides combined from three replicates (experiment described in Figure 3A). SILAC samples and cSILAC samples have 793 (26%) and 678 (23%) more peptides than DiMe samples and cDiMe samples, respectively. (B) Boxplots of hydrophobicity scale of peptides. For each peptide, hydrophobicity coefficients of amino acids measured in C18 were added up to assign hydrophobicity scale. ⁴⁵ Peptides were grouped according to the Venn diagram in panel A. Medians between groups were compared, and the *p* value of Welch's *t* test was calculated. Medians of each group are listed under the boxplot.

that SILAC improves the precision and repeatability of quantification, which can provide more confidence in interpreting quantitative proteomics experiments (discussed later in this paper).

Effects of Dimethyl Labeling

During our analyses, we consistently observed higher numbers of peptides and proteins identified in SILAC-labeled samples than in dimethyl-labeled samples. We analyzed the experimental data from the "Precision and Repeatability Experiments" to better understand this observation. We identified a total of 4106 unique peptides in SILAC samples but only 3181 unique peptides in dimethyl-labeled samples or a ~23% loss in number of peptide identifications from dimethyl-labeled samples (Figure 4A). A similar result was also observed in the experiment comparing dimethyl-labeled samples with SILAC samples mixed after StageTip desalting (cSILAC, Figure 4A) where we identified 3965 cSILAC peptides and 3155 dimethyl-labeled peptides, representing a ~20% reduction in peptide identifications.

One possible explanation for the fewer peptides identified in dimethyl-labeled samples is that the extra step causes loss of peptide, leading to lower peptide ion abundances and hence fewer precursors detected and sampled by the mass spectrometer. We calculated the log₂ of peptide signal intensities of common peptides identified from SILAC- and dimethyl-labeled samples (peptide intensity dimethyl labeling/peptide intensity SILAC) and compared the median of the log ratios to zero (dimethyl labeling/SILAC 1:1). We found that

the signal intensity from dimethyl-labeled samples is 5% lower ($p = 6.36 \times 10^{-10}$) than that of SILAC samples (Supplementary Figure 5A in the Supporting Information).

We believe that the 5% decrease in signal intensity of the peptides common to SILAC and dimethyl labeling would not explain the 20% fewer peptide identifications. So we compared the properties of peptides that are specific to either SILAC- or dimethyl-labeled samples. We compared peptide properties including hydrophobicity, ^{41–45} partition energy, ⁴⁶ mass, length, and transfer free energy ⁴⁷ of these peptides (Supplementary Figure 5B,C in the Supporting Information). We found differences in peptide mass, length, transfer free energy, partition energy, and most prominently, hydrophobicity.

We compared the hydrophobicity of peptides using the C18 RP-HPLC indices from Wilce (Figure 4B). We found that peptides enriched in SILAC samples are significantly more hydrophilic than dimethyl-labeled peptides ($p \leq 1 \times 10^{-5}$, Welch's two sample t test). We observed the same difference in hydrophobicity in the pair of cSILAC- and dimethyl-labeled samples (Figure 4B), suggesting that the dimethyl-labeling step may cause the loss of hydropholic peptides. We also observed these differences in peptide hydrophobicity using alternate hydrophobicity indices from other research groups (Supplementary Figure 5C in the Supporting Information). $^{41-45}$

Comparing SILAC and Dimethyl Labeling in Affinity Pull-Down Experiments

Affinity bait enrichment combining MS-based quantitative proteomics is a very sensitive and specific approach for

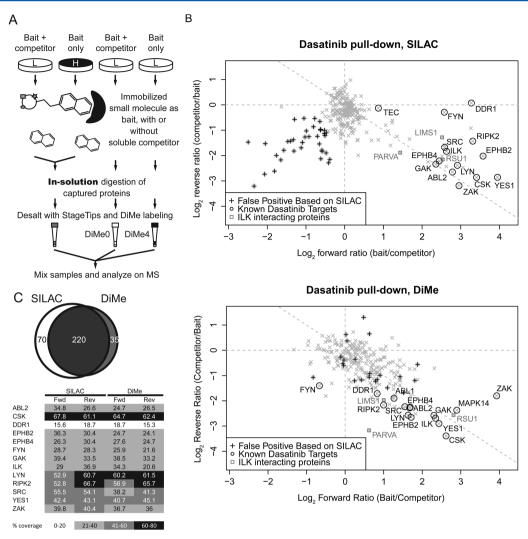


Figure 5. Dasatinib pull-down. (A) Experimental design of "forward" experiment. Four populations of HeLa cells were grown in SILAC medium. Dasatinib was immobilized to carboxy-functionalized sepharose beads. Dasatinib (soluble competitor) or DMSO alone (vehicle control) was added to HeLa cell lysate before affinity enrichment. Captured proteins were digested on-bead. Trypsin-digested peptides were desalted on StageTips. Respective samples were labeled with dimethyl labeling. SILAC- and dimethyl-labeled samples were mixed and analyzed by MS. (B) Scatterplot of protein ratios from the "forward" and "reverse" replicates of a label swap experiment. (C, top) Number of identified peptides from SILAC- and dimethyl-labeled samples. (C, bottom) Percent sequence coverage of quantified dasatinib targets.

identification of bait-interacting proteins. The bait molecule in affinity bait experiments can be small molecule, ³² peptide, protein, ⁴⁸ or nucleic acid. ⁴⁹

We wanted to examine the impact of using SILAC or dimethyl labeling for proteomics applications using affinity bait pull-down experiments with bait molecules that have well-characterized targets: (1) the small molecule kinase inhibitor dasatinib (Figure 5)⁵⁰⁻⁵² and (2) the SH2 domain of GRB2 (Supplementary Figure 6 in the Supporting Information). 48,53,54 We identified known bait-interacting proteins with both SILAC and dimethyl labeling in both pull-down experiments.

In the dasatinib experiment, we were able to quantify 286 and 250 proteins with SILAC and dimethyl labeling, respectively. Out of more than 40 known targets of dasatinib, 50 SILAC and dimethyl labeling led to quantification of 14 and 15 protein targets in HeLa cells, respectively. In addition to the direct target of the small molecule, interacting proteins of ILK $(\alpha$ -parvin/PARVA, PINCH1/LIMS1, and RSU1) $^{55-57}$ were also quantified by both methods. Sequence coverage of targets

was similar in both SILAC-labeled and dimethyl-labeled samples (Figure 5C and Supplementary Figure 6C in the Supporting Information), suggesting the lowered peptide identification by dimethyl labeling has a minimum impact on the quantification of enriched target proteins in samples of lower peptide complexity. We observed, however, that the lower precision of dimethyl labeling leads to a wider distribution of nonspecific proteins when compared with SILAC (Figure 5C). A wider distribution of protein ratios for nonspecific proteins may negatively impact the identification of specific targets of the affinity bait. 58 Furthermore, experimental outliers such as keratin contaminants (with low SILAC H/L ratios) or sample "carryover" from the LC system can be easily detected in label-swap experiments with SILAC because they originate from an non-SILAC-labeled source and their H/L ratios remain low in both label-swap replicates (points marked '+' in scatterplot, Figure 5B). Such contaminants would be labeled in chemical labeling approaches and would not be distinguishable from indigenous proteins in the sample; note that the proteins marked '+' in the SILAC experiment, as

obvious outliers are now detected within the experimental group in the dimethyl-labeling experiment in the lower panel of Figure 5B.

In the GRB2-SH2 pull-down, HeLa cells were treated with EGF or PBS (control). Captured proteins were resolved by SDS-PAGE; SILAC samples were combined and run in a single lane, while samples for dimethyl labeling were run in separate lanes. The samples were fractionated by slicing the gel lanes into slices of similar staining intensity. For dimethyl-labeled samples, we tried our best to cut bands of equal size between lanes for comparison, and proteins were digested in-gel before dimethyl labeling on a C18 StageTip. Our SILAC- and dimethyl-labeling experiments quantified 171 and 165 proteins, respectively, with 118 proteins in common. EGFR and SHC1 were identified as GRB2-SH2 domain interacting proteins in both methods. Although our previous experiments indicate that the dynamic range of peptide quantification in both SILAC and dimethyl labeling is limited in complex peptide mixtures (Figure 2B), we note that in affinity capture experiments with an enriched subproteome, both methods are capable of quantifying abundance changes of up to 16-fold (Figure 5B and Supplementary Figure 6B in the Supporting Information).

To our knowledge, this is the first application of dimethyl labeling applied to in-gel proteolytic digestion in quantitative proteomics; this indicates that dimethyl labeling can be a convenient way to quantify protein abundance differences from gel-separated samples after the fact, even if this was not part of the original experiment design.

DISCUSSION

SILAC has been rapidly adopted as an approach in MS-based proteomics because of its simple and robust performance. As a metabolic labeling approach, however, its use is mostly constrained to cell culture systems. Because of the ease of generating stable isotope labeled proteomes, the proteomics community has used it as a spike-in standard⁵⁹ and for comparing different chemical labeling methods and label-free methods. In our study, we compared SILAC and the chemical labeling method, stable isotope dimethyl labeling, by combining and analyzing these labeled samples in the same nanoLC-MS runs. Our results demonstrate that SILAC and dimethyl labeling achieve comparable quantitative performance. Both methods are affected by ratio compression and have a limited dynamic range; ratios of 1:10 samples were measured ~1:6, although we note that this issue was exacerbated by the use of unfractionated whole proteome analyses, and we demonstrate that the dynamic range can be improved by orthogonal fractionation steps prior to MS analysis.

Under our experimental conditions, dimethyl-labeled peptides had decreased peptide signal intensity than SILAC-labeled peptides. We also observed a reduced number of peptides and proteins identified in dimethyl-labeled samples. By comparing the properties of peptides differentially enriched between SILAC- and dimethyl-labeled samples, we observed that dimethyl labeling resulted in diminished recovery of hydrophilic peptides.

The greatest advantage of SILAC over chemical-labeling methods is that samples can be mixed in an early stage during the sample processing. Theoretically, the SILAC workflow reduces both sample loss and variability from sample processing, leading to a higher precision and repeatability in SILAC than chemical-labeling methods. By comparing SILAC samples mixed in different stages of the proteomic processing

workflow, we show that sample mixing at an early experiment step leads to higher precision and repeatability in quantification. A similar reduction in precision has also been demonstrated in mTRAQ previously.²²

We compared SILAC- and dimethyl-labeling in proteomics affinity pull-down experiments with the small-molecule kinase inhibitor, dasatinib, and GRB2-SH2-FLAG fusion protein. We demonstrated that SILAC can facilitate the identification of potential targets due to the tighter distribution of the nonspecific binders. Furthermore, false-positives whose protein ratios do not "flip" in label-swap experiments are easily detected in SILAC but not in dimethyl labeling. We describe our on-StageTip dimethyl-labeling protocol and applied it to insolution and in-gel protein digests, providing an easy way to incorporate stable isotope label-based quantification in standard gel-based experiments.

Metabolic labeling methods like SILAC allow samples to be combined, even at the level of intact cells, and it is possible to perform complex experimental procedures, like subcellular fractionation, with little detrimental effect on quantitative measurements. The recently developed neutron encoding SILAC approach¹² combines the advantages of metabolic labeling with higher multiplexing and has the potential to provide both high quantitative precision and better sensitivity than the original SILAC approach, although it requires a mass spectrometer that can acquire very high-resolution MS scans (R > 480 000) on a time-scale compatible with nanoscale liquid chromatography. We expect that neutron-encoded quantitative proteomics methods will see increased adoption as MS technology advances and costs of neutron encoded reagents fall. Because SILAC cannot be applied as readily in tissue or clinical samples, chemical labeling is a good and practical substitute. We did not compare dimethyl labeling with isobaric chemical tags like iTRAQ or TMT that use reporter-tag-based quantification from MS2 spectra because we reported on the direct comparison of mTRAQ and iTRAQ elsewhere.16 Because there are pros and cons in quantifying stable isotope labeling methods at both the MS1 or MS2 levels, the choice of the quantification method depends on experimental and analytical parameters as well as practical considerations, such as the cost or the type of available MS instrument and downstream data analysis software.

ASSOCIATED CONTENT

S Supporting Information

Figure S1. Comparison of peptides in different workflows. Figure S2. Boxplot of \log_2 ratio of SILAC- and dimethyllabeling samples analyzed in separated MS runs. Figure S3. Histogram of H/L peptide ratios including N-terminal acetylated peptides. Figure S4. Boxplot of \log_2 protein ratios for 1:1:1:1 experiment. Figure S5. Comparison of peptide intensities and hydrophobicities between SILAC and stable isotope dimethyl labeling workflows. Figure S6. FLAG-tagged GRB2-SH2 domain pull-down. This material is available free of charge via the Internet at http://pubs.acs.org.

AUTHOR INFORMATION

Corresponding Author

*E-mail: shaoen@u.washington.edu. Tel: 206-616-6962. Fax: 206-616-4230.

Author Contributions

[†]H.-T.L. and H.W.S. contributed equally.

Notes

The authors declare no competing financial interest.

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