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Barcoding for Isobaric Tagging of Carboxylic Acids and Amines Metabolites

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Abstract

Conventional isobaric platforms in metabolomic analyses are designed to label a single functional group, typically primary amines. This specificity restricts the range of detectable analytes in a single injection, limiting overall chemical coverage of metabolites, and potentially reducing the depth of biological insight. Using a proof-of-concept 2-plex isobaric barcode tagging scheme, amine-containing metabolites are selectively labeled using acid barcode tags. Acid-containing metabolites (carboxylate) are derivatized using amine barcode tags. This allows for broad coverage of two metabolite classes in a single injection in a 30-minute LC gradient. These barcode tags undergo double fragmentation to generate cyclized products and distinct reporter ions that are specific to each metabolite class, enabling accurate quantitation. This tagging scheme also serves as an untargeted approach for identifying metabolites not present in the user's metabolic library. Tagged acid and tagged amine analytes are mixed at ratios of 1:2:5:10 using various amounts of *Escherichia coli* lysate to produce a 10-fold linear dynamic range, an average linearity (R^2) of 0.99, and an average RSD of 2.64%. Metabolic changes in amine and carboxylic acid metabolism of genetic knockout PanC and wild-type *Escherichia coli* were characterized using this method.

1. INTRODUCTION

Accurate LC-MS based quantitative metabolomics is hindered by low throughput, variable matrix effects, instrument signal drift, and retention time shifts. Approaches to mitigate these difficulties often lead to prolonged analysis times.^{1, 2} Chemical isotope labeling (CIL) is² designed to 1) increase ionization efficiency which enhances signal response³ and 2) improve quantitation through isotope ratio mass spectrometry. These tags render a mass difference for the same analyte from various samples, which enables multiplexing and accounts for matrix effect, as well as signal and chromatographic shifts.^{2, 4}

CIL presents challenges to metabolite coverage.^{5, 6} For mass shift isotope labeling, the number of analyte peaks increases proportionally to the number of isotope tags used. As the level of multiplexing increases, the spectral complexity in the full MS increases. This requires substantial

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3 chromatographic resolution to avoid overlapping MS signal from different analytes.⁴ CIL typically
4 reacts with one class of moieties for analysis, which inherently limits metabolite coverage.

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7 Isobaric tagging is used to alleviate spectral complexity in full MS.⁷ Isobaric tags have the same
8 nominal mass but generate distinct reporter ions during MS/MS, allowing differentiation and
9 quantification of analytes.⁸ Isobaric tags are designed to include 1) a reactive moiety that targets
10 a specific functional group for labeling; 2) a balancer group isotopically labeled to ensure the same
11 mass of overall tag; and 3) a reporter group that incorporates unique isotopes enabling relative
12 quantification across different samples.^{4, 9 10 11 12 7} Although isobaric tagging offers throughput
13 and quantitative advantages, their analytical performance can be confounded by isobaric ratio
14 compression. This phenomenon is typically driven by the co-isolation and co-fragmentation of
15 interfering ions, which distorts reporter ion intensities and therefore true abundance differences.

16
17 Neutral loss-based isobaric tags can be used to remedy chimeric spectra and reporter ion
18 distortion associated with conventional isobaric tags (e.g., TMT,⁹ iTRAQ,¹⁰ DiART,¹¹ DiLeu¹²) that
19 often rely on a1-type fragmentation.⁷ However, interfering ions may exhibit compression and
20 distortion even with neutral-loss isobaric tags. Previous work on neutral-loss isobaric tags shows
21 that the reporter group remains attached to the analyte. This imparts a distinct mass shift in the
22 MS/MS scan that distinguishes co-isolated precursors.^{7, 13} A quaternary amine neutral loss
23 isobaric tag has been reported for the quantitation of acid metabolites.² The neutral loss of the
24 trimethylamine produced a cyclized reporter ion, which remains attached to the analyte. Advances
25 in this tag have generated a set of dual-fragmentation, neutral-loss isobaric reagents for the
26 quantitation of thiols.⁷

27
28 A major limitation of using conventional isobaric platforms in metabolomics is their tagging a single
29 functional group, typically primary amines.^{9 10 11 12 7} This specificity restricts the range of
30 detectable analytes in a single injection which limits overall chemical coverage and potentially
31 reducing the depth of biological insight. In this current work, we expand metabolite coverage to
32 analyze 100 targeted amines and carboxylates using a proof-of-concept 2-plex isobaric barcode
33 tagging scheme in a single injection. The barcoding strategy uses moiety-specific chemical tags
34 that serve as identifiers for different classes of metabolites.

35
36 Barcode tags are designed to selectively label either amine-containing or acid-containing
37 (carboxylate) metabolites, which generate distinct reporter ions upon fragmentation that are
38 exclusive for each class of metabolite, as presented in Figure 1. The labeling of carboxylate
39 analyte using a D₃-neucode pyruvate and a D₀-neucode pyruvate-D₃ amine tags is shown in

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


Figure 1 A1. D₀-neucode alpha-ketoisocaproic-D₃ and D₃-neucode alpha-ketoisocaproic acid tags are used for the tagging of amine analytes, as illustrated in Figure 1 B1. When a higher-energy collisional dissociation (HCD) is applied, it causes the tagged analyte to undergo double fragmentation to produce a cyclized product and reporter ions⁷, as shown in Figure 1 A2 and B2. Specifically, amine barcodes produce reporter ions at m/z 84.0814 and 87.1002 (Figure 1 A3), while acid barcodes yield reporter ions at m/z 126.1276 and 129.1465 (Figure 1 B3). These reporter ions serve as identifiers, enabling the identification and accurate quantification of metabolite classes in complex mixtures. This barcoding framework is inherently scalable and could be expanded to high-level multiplexing, enabling high-throughput quantification of metabolites across large sample sets while maintaining class specificity.

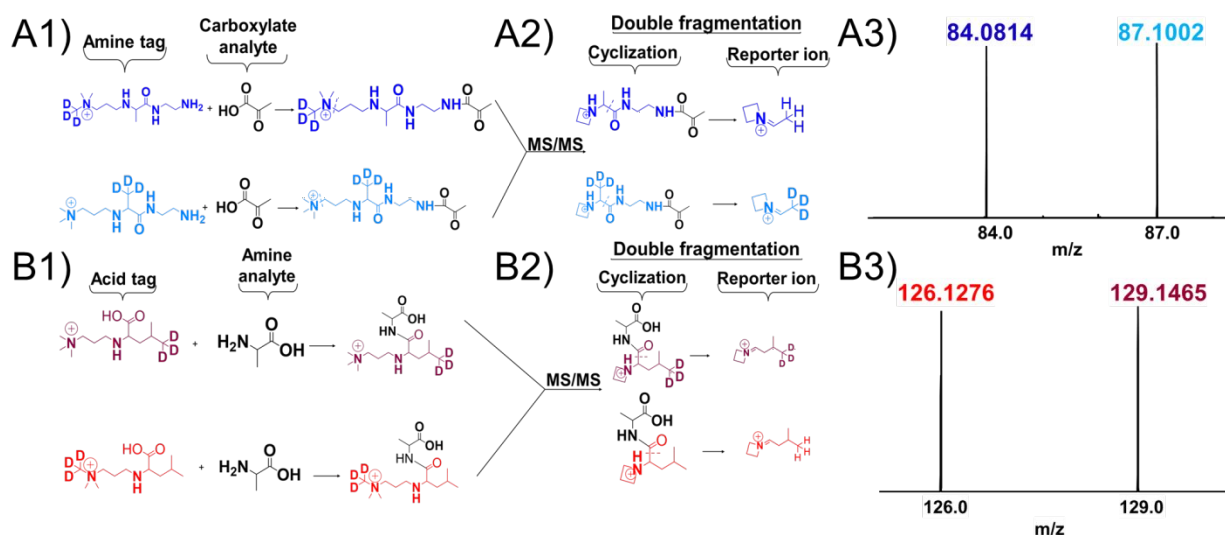


Figure 1. Double fragmentation of (A1) a derivatized carboxylate metabolite using two variants of amine barcode tag and (B1) derivatization of an amine metabolite using two variants of amine barcode tag. The labeled metabolites undergo double fragmentation upon HCD activation (A2 and B2). The first step of fragmentation involves cyclization following trimethylamine neutral loss. Tags undergo a second fragmentation to generate distinct reporter ions at m/z 84.0814 and 87.1002 (A3) for acid metabolites, and at m/z 126.1276 and 129.1465 (B3) for amine metabolites.

2. METHODS

2.1 Reagent and Standard Stock

All acid and amine-containing metabolite standards, 4-(4,6-dimethoxy-1,3,5-triazin-2-yl)-4-methylmorpholinium chloride (DMTMM) and N-methylmorpholine (NMM), Hexafluorophosphate azebentriazole tetramethyl uronium (HATU), hydroxybenzotriazole (HOBT), *N,N*-Diisopropylethylamine (DIPEA), and pyridine were purchased from MilliporeSigma (St. Louis, MO,

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USA). Ammonium acetate, acetic acid, acetonitrile (ACN), and LC-MS grade water were obtained from Fisher Scientific (Pittsburgh, PA USA). Each metabolite stock was prepared at 100 μ M for each standard, and a final concentration of 5 μ M mixed acid and 5 μ M mixed amine-containing metabolites in 80% ACN:20% H_2O was generated. 3-(dimethylamino)-1-propylamine, iodomethane, pyruvate, and alpha-ketoisocaproic acid were purchased from MilliporeSigma (St. Louis, MO USA). Pyruvate- D_3 , alpha-ketoisocaproic- D_3 , and methyl iodide- D_3 were obtained from Cambridge Isotope Laboratories (Boston, MA, USA). Two *Escherichia coli* (*E. coli*) strains: Wild-type (BW 25113, i.e. wt) and PanC gene mutant (JW 0129, i.e. PanC KO) were purchased from Horizon Discovery (Lafayette, CO, USA). The WT and PanC KO (Δ panC750::kan) used in this study were derived from the Keio collection in the *E. coli* BW25113 genetic background and contain a kanamycin-resistance cassette.

M9-media was prepared according to previous reports using the following: potassium phosphate monobasic (KH_2PO_4), sodium phosphate dibasic anhydrous (Na_2HPO_4), ammonium chloride (NH_4Cl), sodium chloride ($NaCl$), magnesium sulfate heptahydrate ($MgSO_4 \cdot 7H_2O$), and calcium chloride dihydrate ($CaCl_2 \cdot 2H_2O$) were purchased from Fisher Scientific (Pittsburgh, PA, USA). D-glucose was obtained from Millipore Sigma (St. Louis, MO, USA). The M9-salts constituted 3.6g Na_2HPO_4 , 3g KH_2PO_4 , 1g NH_4Cl , 0.5g $NaCl$, 491.5mg $MgSO_4 \cdot 7H_2O$, 14.6 mg $CaCl_2 \cdot 2H_2O$, 500mg D-glucose.¹⁴

2.2 Tag Synthesis

2.2.1 D_0 -neucode alpha-ketoisocaproic- D_3 and D_3 -neucode alpha-ketoisocaproic acid tags synthesis

3-(dimethylamino)-1-propylamine (Scheme 1A) was Boc-protected in 10mL ethanol (EtOH) using a microwave for 10 minutes, as shown in Scheme 1B. The tag was preconcentrated using a rotary evaporator, followed by extraction with 50:50 dichloromethane (DCM)/water. The DCM layer was further washed with water (3X) and preconcentrated to obtain tert-butyl (3-(dimethylamino)propyl) carbamate as the intermediate. The Boc-protected intermediate was selectively methylated using excess Iodomethane in 20 mL of tetrahydrofuran (THF) for 30 minutes, yielding the corresponding mono methylated solid product² (Scheme 1B). The tag was washed with THF and filtered under vacuum. The tag was deprotected in 10mL of 4 M Hydrochloric acid (HCl) for 2 hours to form a solid product, which was then filtered and washed with THF. Using 2 equiv of sodium cyanoborohydride and 3 equiv of Sodium bicarbonate ($NaHCO_3$), the tag was functionalized with 1 equiv. of alpha-Ketoisocaproic- d_3 acid by reductive

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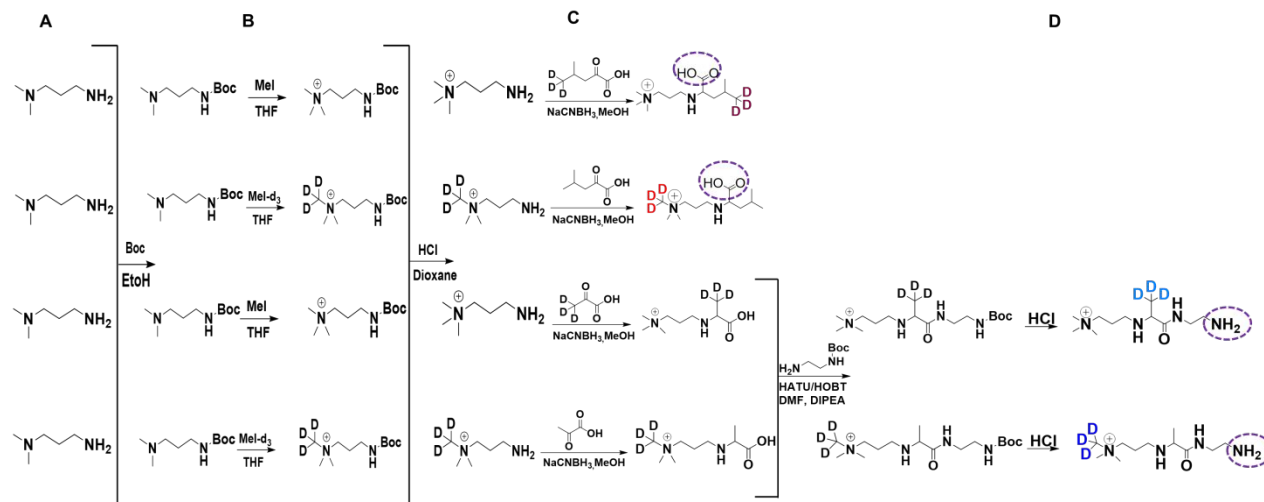
amination in 20mL of methanol (MeOH) for 2 hours (Scheme 1C). The tag was purified by reverse-phase chromatography and characterized using high-resolution mass spectrometry (Figure S1) and nuclear magnetic resonance (Figure S2). Following the same synthesis scheme, the D₃-neucode alpha-ketoisocaproic acid tag was synthesized using iodomethane-D₃ and alpha-ketoisocaproic acid. The exact masses of D₀-neucode alpha-ketoisocaproic-d₃ and D₃-neucode alpha-ketoisocaproic acid tags are 234.2256. The observed m/z values of the reporter ions are 129.1466 and 126.1278, respectively, as shown in Figure 1 B3.

2.2.2 D₀-neucode pyruvate-D₃ and D₃-neucode pyruvate amine tag synthesis

This synthesis used a reaction scheme similar to the one outlined above, incorporating additional steps. Briefly, 3-(dimethylamino)-1-propylamine (Scheme 1A) is Boc-protected (Scheme 1B), methylated, deprotected, filtered and washed with THF. Using 2 equiv of sodium cyanoborohydride and 3 equiv. of NaHCO₃, the tag was functionalized with 1 equiv. of pyruvate-D₃ by reductive amination in MeOH for 2 hours (Scheme 1C). The tag was purified by reverse-phase chromatography. Using 1.5 equiv of DIPEA and 2 equiv of HATU/HOBT, the tag was functionalized with 1.5 equiv of Boc amine by amide coupling in 20mL of dimethyl formamide (DMF), and the reaction was allowed to proceed overnight (Scheme 1D). The tag was purified using an acid-base wash, then dissolved in THF, and deprotected using 5mL of 1 M HCl for 10 min (Scheme 1D). The tag precipitated upon HCl administration and was subsequently rinsed with THF. The tag was then characterized using high-resolution mass spectrometry (Figures S3) and nuclear magnetic resonance techniques (Figures S4). Using a similar synthesis scheme, the D₃-neucode pyruvate tag was synthesized using iodomethane-d₃ and pyruvate. The % yield of all the tags was above 94%. The exact masses of D₀-neucode pyruvate-D₃ and D₃-neucode pyruvate amine tags are 234.2368. The observed m/z values of the reporter ions are 87.0997 and 84.0808 as illustrated in Figure 1 A3.

Scheme 1. Synthetic Routes for Each of the 4 barcode Tags^a





Scheme 1 *a* 3-(Dimethylamino)-1-propylamine, as a starting material, was purchased (A). Protection of starting material using Di-*tert*-butyl decarbonate, then methylated using either Iodomethane or Iodomethane- d_3 (B). Boc deprotection and tag functionalization (C). Boc amine functionalization to tag and Boc deprotection (D).

2.3 Cell Culture: Two strains of *E. coli* were pre-cultured in LB broth at 37°C and 250rpm in a shaker overnight. New cultures of WT and PanC KO using LB broth were carried out using a 1:10 serial dilution from pre-cultured WT and PanC KO *E. coli* with optical densities measured at hourly intervals (Figure S10). Once an optical density of 0.6 was achieved for both *E. coli* strains, the cells were centrifuged at 6300rpm at 4 °C for 15 minutes and lysed using a 2:2:1 ratio of methanol, acetonitrile, and water as the lysing solvent. Briefly, the cells were frozen in liquid nitrogen for 2 minutes and then sonicated for 15 minutes. This was done in three separate cycles and stored at -20°C for one hour to enhance metabolite extraction. Afterward, the cells were centrifuged, and the supernatant was collected and stored at -80°C for further analysis. The *E. coli* strains were cultured and lysed in parallel to mitigate sample preparation bias.

To determine the growth conditions of WT and PanC KO in M9 salts, the two *E. coli* strains were pre-cultured in LB broth for 6 hours in a shaker at 37°C and 250rpm. Cells were centrifuged to remove supernatant. Cell pellets were resuspended and new cultures were carried out using a 1:10 serial dilution of the precultured *E. coli* strains in 750 mL of M9-glucose salts, placed in a shaker at 37°C and 250rpm for 24 hours.

2.4 Tagging Scheme

2.4.1 Amine Derivatization

A 500 μ L stock solution of 5 μ M containing 30 amine metabolite standards was derivatized. For *E. coli* lysate, 30 μ L of 50mM D_0 -neucode alpha-ketoglutaric- d_3 and D_3 -neucode alpha-



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ketoisocaproic acid tags were activated separately using 50 μ L of 75mM HATU and 100mM HOBT in DMF and 24% pyridine for an hour at room temperature. 40 μ L of activated D₀-neucode alpha-ketoisocaproic-d₃ tag was added to 500 μ L of WT *E. coli* lysate, and 40 μ L of activated D₃-neucode alpha-ketoisocaproic tag was added to 500 μ L PanC KO *E. coli* lysate containing 1.5% NMM. The reaction was allowed to proceed for 4 hours at room temperature in a shaker at 25 °C. Following this, the mixture was dried using a vacuum centrifuge, achieving a reaction efficiency of over 95%. Derivatization reaction efficiency was determined using calibration curves generated from untagged metabolites from commercially available standards. Metabolite concentrations were determined in both untagged *E. coli* lysates and tagged lysates. Remaining untagged metabolites from the tagged samples indicated near complete reaction efficiency (>95%). Analysis for the unreacted metabolite is the only way to account for differences in sensitivity of tagged vs untagged metabolites, as there is no commercial standard for the fully tagged counterparts.

2.4.2 Acid Derivatization

70 acid-containing metabolite standards were derivatized at a stock concentration of 5 μ M in 500 μ L. Into a 500 μ L of WT *E. coli* lysate and 500 μ L PanC KO *E. coli* lysate, 35 μ L of 25mM of D₀-neucode pyruvate-d₃ and D₃-neucode pyruvate amine tags were added, respectively. 35 μ L of 75 mM DMTMM, containing 1.2% NMM, was subsequently added to each Eppendorf tube containing the *E. coli* lysate. The mixture was allowed to react in a shaker at 25°C for 4 hours. Thirty minutes into the reaction, 50 μ L of 37 mM DMTMM was added to drive the reaction to > 95% completion. Following this, the mixture was dried using a vacuum centrifuge. All 4 vials of tagged metabolites (acid and amine) were reconstituted in 50 μ L volume and mixed 1:1.

2.5 LC-MS Analysis

2.5.1 Capillary Liquid Chromatography

A frit was fabricated in a 50- μ m inner-diameter capillary column, close to the pulled nano tip.^{15, 16} The tip was created using a laser-pulled method.¹⁷ In brief, a window was created in a 27 cm long fused silica capillary by removing the polyimide coating using an electrical arc. A photopolymerized frit was created at the window under a wavelength of 365 nm for 30 min. Nanospray tips were generated using a Sutter CO₂ laser puller model P-2000. The nano-capillary column was then packed with Acquity UPLC Amide 1.7 μ m particles using a gas pressure cell. The column length was finally cut to 15 cm. The column was attached to a stainless-steel tee to split the flow.⁷ The split was made up of a 50 μ m \times 75 cm open capillary. A 30-minute gradient method was employed at a controlled flow rate of 0.200 mL/min and an injection volume of 5 μ L



(491 nL/min through the capillary column). The mobile phase A comprised 95% acetonitrile (ACN) and 5% water, supplemented with 10 mM ammonium acetate and 0.1% acetic acid. Mobile phase B was 10% ACN and 90% water, which also contained 10 mM ammonium acetate and 0.1% acetic acid.⁷ The gradient used for the separation of tagged amines and carboxylates was: 20% B, 0 min; 20% B, 1 min; 95% B, 5 min; 95% B, 22 min; 20% B, 22.1 min; 20% B, 30 min.

2.5.2 MS Analysis

Samples were analyzed as previously described using a Q-Exactive orbitrap mass spectrometer (Thermo Fisher Scientific, Waltham, MA) coupled to a Thermo Vanquish LC instrument (Thermo Fisher Scientific, Waltham, MA).⁷ All runs were carried out at a spray voltage of 1.75 kV and a capillary temperature of 200 °C.⁷ For parallel reaction monitoring (PRM) runs, a maximum ion injection time of 50ms and an AGC target of $1e^6$, with a resolution of 17.5 K, were used in the MS¹ scan, covering a scan range of 100-600 m/z.⁷ In the PRM scan, a 100-inclusion list of analytes with a maximum ion injection time of 100 ms, AGC target of $1e^6$, resolution of 140 K, and an optimal collision energy of 35 were used. Untargeted runs were carried out using a resolution of 70 K, an AGC target of $1e^6$, a maximum ion injection time of 100ms, and a scan range of 70-800 m/z for the full MS. The data-dependent analysis (DDA)-MS/MS used a 17.5 K resolution, a maximum ion injection time of 50ms, a $2e^5$ AGC target with a DDA setting of 20.0s dynamic exclusion.

2.5.3 Data Extraction and Processing

For targeted metabolomics, the extracted LC-MS data were processed using Xcalibur QualBrowser and Skyline¹⁸. Known metabolites were quantified using the reporter ion observed by PRM. Data analysis was performed in Excel, and graphs were generated in GraphPad Prism 9. Untargeted metabolites were identified using the R package CluMSID.¹⁹ These samples were analyzed using a DDA method

3 RESULTS AND DISCUSSION

The barcoding scheme for multiplexed analysis uses isobaric tagging to quantify 100 carboxylate and amine metabolites in complex biological mixtures from a single injection. Differential reporter ions of amine and acid tags are generated by using different keto-acid precursors (Figure 2), which, upon fragmentation, indicate the presence of an amine or an acid metabolite. The ability of barcode tags to undergo double fragmentation enables the generation of cyclized products and distinct reporter ions for each metabolite class, thereby reducing the potential for spectral overlap and improving resolution in MS/MS analysis.

The general scheme for our barcode tagging uses two strains of *E. coli* (WT and PanC KO), as shown in Figure 2. The barcode acid tags differ from the amine tags in that they use alpha-ketoisocaproic acid (red) and pyruvate (blue) as the central component, respectively (Figure 2A). This yields reporter ions of 126.1276 and 129.1465 m/z for acid tags, and 84.0814 and 87.1002 m/z for amine tags. Each class of tag (acid and amine) uses both isotopic and non-isotopic variants of the keto-acid base, which allows for the two-plex isobaric set to generate distinct reporter ions. The reactive moiety of each tag allows derivatization of amine analytes with acid-tag (red) variants and carboxylate analytes with amine-tag (blue) variants. Because dual fragmentation yields distinct reporter ions, unknown metabolites can be characterized by the presence of appropriate functional groups. These barcode tags are reacted with samples individually and mixed, which increases coverage of the metabolome.

Co-elution of isotopically tagged metabolites is critical to isobaric quantitation. Even small differences in retention times between tags can cause substantial signal variance. Chromatograms show co-elution of tagged metabolites as illustrated in (Figure 2B B1), tagged amine and tagged carboxylate (Figure 2B B1) with near identical peak profiles and retention times. The position of the deuterated methyl group either on the quaternary amine (balancer) or on the keto acid (reporter) of the tags allows for a 2-plex reporter ion in the MS/MS, as presented in Figure 2C.

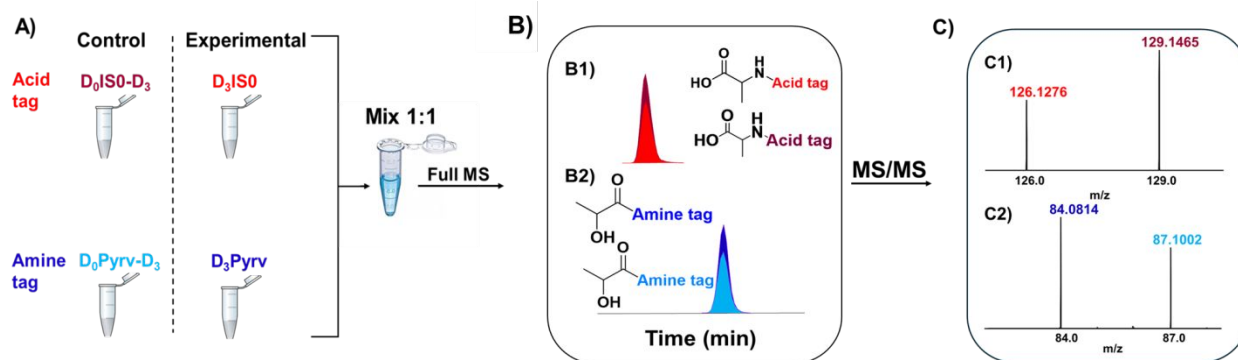


Figure 2: Analytical tagging scheme of amine and acid quantitation. WT and PanC KO *E. coli* were cultured in LB Broth and lysed under the same conditions. (A) 500 μ L of each WT and PanC KO *E. coli* lysate was derivatized. Two vials of each WT and PanC KO *E. coli* lysate were derivatized with either type of acid or amine tag variant, then mixed at a 1:1 ratio. (B) Full MS peaks of tagged (B1) amine and (B2) acid metabolites. (C) Each tag undergoes dual fragmentation, first cyclization, following the loss of trimethylamine, then the reporter ion. Triggers of MS/MS fragmentation to generate reporter ions (C1) of m/z 126.1278 and 129.1466 for amine metabolites (C2) 84.0808 and 87.0997 for acid metabolites.

Quantitative parameters for the tags were characterized to ensure appropriate use in biological matrices. The optimal collision energy for the acid and amine tags was evaluated. Tags themselves were analyzed under different collision energies from the HCD cell in triplicate. A nHCD value of 35 was found to produce the most intense reporter ion, as shown in Figure S5. No retention time shifts due to deuterium incorporation around the quaternary amine tags were observed. This resulted in consistent reporter intensity across the top 50% of each peak, as illustrated in Figures S6 and consistent with previous work on reverse-phase columns.^{2, 7}

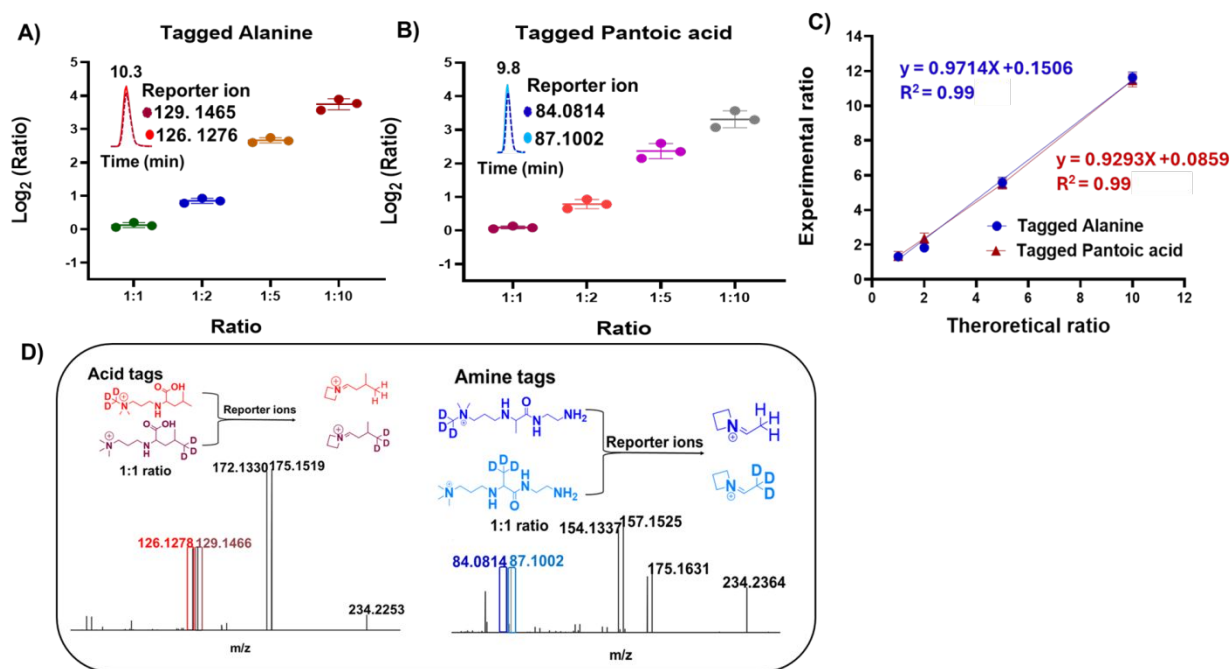


Figure 3. MS/MS reporter ion-based quantification of barcode isobaric tags. **(A & B)** Measured ratios of tagged amine and acid-containing metabolites at mixing ratios of 1:1, 1:2, 1:5, and 1:10. Co-elution of the extracted ion chromatogram of reporter ions of tagged alanine (126.1278 and 129.1466 and pantoic acid (84.0814 and 87.1002) metabolites. **(C)** Linearity and dynamic range of quantification for the experimental to theoretical intensity ratios of amine and acid barcode tags. Each data point represents the mean value of a 1:2:5:10 labeling experiment, with error bars indicating the corresponding standard deviation. **(D)** 1:1 ratio of reporter ions generated from each amine and acid tag variants. Tags undergo dual fragmentation in MS/MS, generating reporter ions of equal intensity for each tag variant.

The analytical performance of this method was evaluated by mixing tagged acid and tagged amine analytes at ratios of 1:2:5:10 using various amounts of *E. coli* lysate. This analytical step used varying volumes of cell lysate and, therefore, cell mass to assess linearity, quantitative accuracy, and to evaluate potential matrix effects in the sample. These data show that the analytical performance is independent of isotopic tag composition. The method had a 10-fold



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3 linear dynamic range for the 100 tagged amine and the tagged acid (Figures 3A and 3B), with an
4 average linearity (R^2) of 0.99 ± 0.027 (Supplemental Table 1). Linearity for tagged alanine was
5 0.99, while tagged acid pantoic acid showed a linearity of 0.99 (Figure 3). To determine the
6 reproducibility of each tag variant, identical samples were mixed in a 1:1 ratio, yielding an average
7 percentage reaction RSD of 2.64 ± 0.789 (Supplemental Table 1). Overall, these results are
8 consistent with previously reported values in the literature.^{2, 7 20, 21} While both tags contain a
9 tertiary amine to enhance signal intensity, there is a slight difference in sensitivity between them
10 as seen by the slope of their calibration curves with an average tagged amine metabolites 0.9647
11 ± 0.0116 , and the average tagged carboxylate 0.9349 ± 0.0144 . This may be of interest
12 when exploring larger levels of multiplexing.

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14 The barcode tags were then applied to WT and PanC KO *E. coli* strains to determine if metabolic
15 differences could be uncovered from a genetic knockout. The WT strain was designated as the
16 control sample, and a mutant strain, designated PanC KO, was used as the experimental sample.
17 The PanC KO strain carries a genetic knockout in the pantothenate biosynthesis pathway,
18 specifically affecting the gene encoding pantothenate synthetase (*PanC*)²² as illustrated in Figure
19 4A. This enzyme catalyzes the final step in the biosynthesis of pantothenate (vitamin B5), an
20 essential precursor for coenzyme A.

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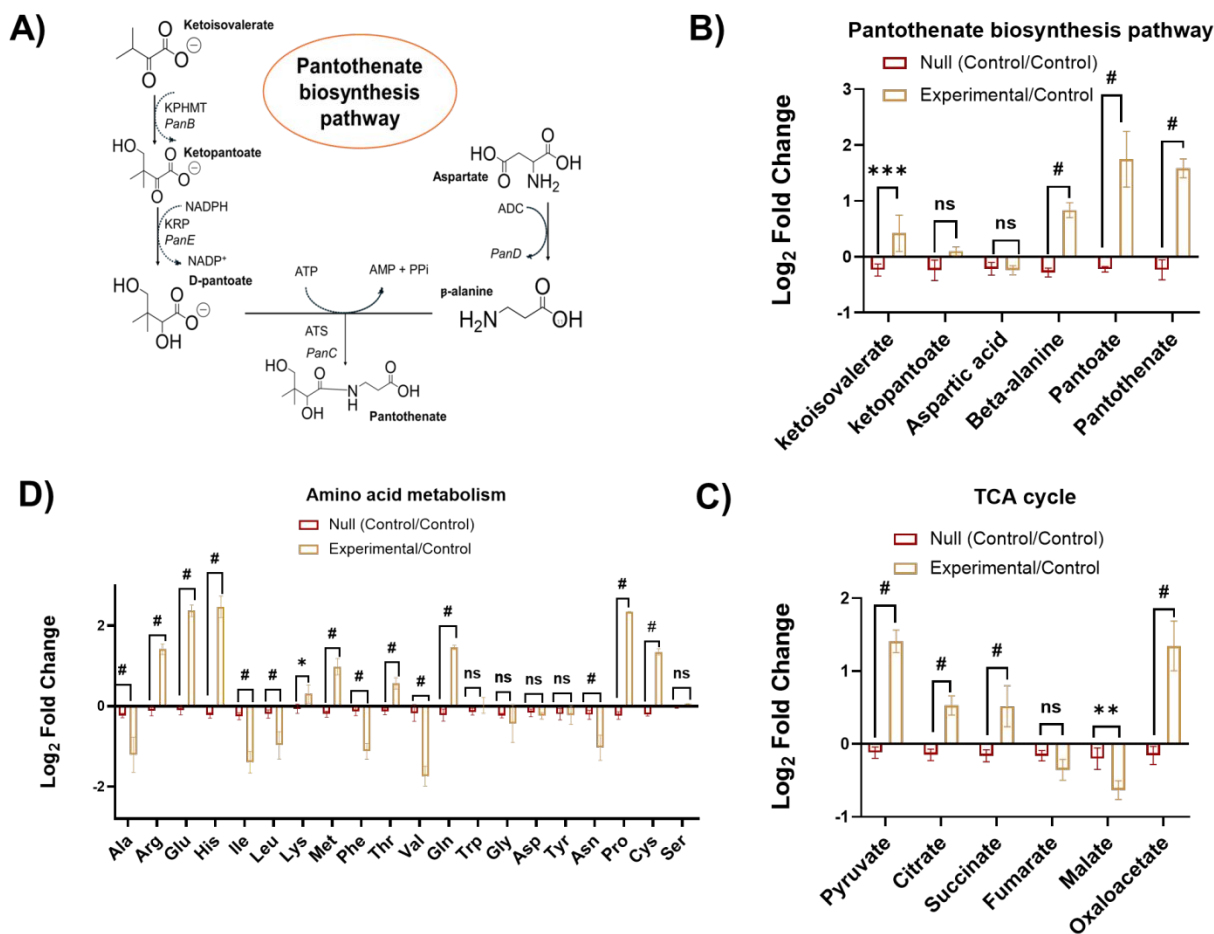


Figure 4. Log₂ fold change of metabolites relative to control compared with a control/control null distribution. Log₂ fold change (Log₂ FC) values for the null distribution were calculated by comparing control samples against other control replicates to estimate baseline variability under no biological change. Experimental Log₂ FC values were calculated relative to the control condition. Bars represent mean values across replicates, and error bars indicate \pm SD. The horizontal line at Log₂ FC = 0 denotes 1:1 and no change relative to control. ns represent no significant difference, * ($p \leq 0.05$), ** ($p \leq 0.01$), *** ($p \leq 0.001$), # ($p \leq 0.0001$).

A total of one hundred known amine- and acid-containing metabolite standards were used to broadly characterize the strains (Table S2). Targeted metabolomic analysis revealed pathway-specific metabolic reprogramming in the PanC KO relative to control (yellow), while the null (control/control, red) comparison remained centered near zero across metabolites, indicating minimal technical or baseline variation. All amino acids, including beta-alanine, were quantified using the acid tag variant. The use of the amine tag for derivatizing the carboxylic group on the amino acids with the DMTMM coupling reagent yields poor reaction efficiency, possibly due to dimerization. The null (control/control, red) represents the derivatized amine and acid-containing



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3 metabolites in WT *E. coli* using D₀ of the tag variants, ratio to the derivatized amine and acid-
4 containing metabolites in WT *E. coli* using D₃ of the tag variants.

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7 Significant increases of metabolites in the pantothenate biosynthesis pathway were observed,
8 particularly pantothenate (Figure 4B). This observation is consistent with prior work showing that
9 PanC KO grown in excess pantothenate elevates intracellular pantothenate and CoA levels
10 through transport/uptake in spite of altered expression.²³ Despite the loss of pantothenate
11 synthetase activity in PanC KO *E. coli*, the cell can import pantothenate from LB Broth via
12 transporters like PanF.^{23, 24} LB Broth is composed of yeast extract, and this extract contains
13 pantothenate, enough to aid in the growth of PanC KO *E. coli*.²⁵⁻²⁷ When M9 salt was used for
14 PanC KO culture, no growth was observed for 24 hours (data not shown). The seeming
15 discrepancy between PanC KO and the elevation in pantothenate is likely due to importing
16 pantothenate from the media. While not a conclusive part of this study, the lack of growth of PanC
17 KO in M9-glucose minimal media and previous reports support this hypothesis.

18
19 To explore the potential impact of elevated pantothenate levels in the PanC KO on CoA enzyme
20 activities, we measured the signal intensities of acetyl-CoA, succinyl-CoA, and malonyl-CoA in
21 both WT and PanC KO strains. Reduced availability of free CoA reduces acetyl-CoA formation,
22 leading to decreased acetyl-CoA levels in the mutant (Figure S7). The findings showed that the
23 PanC KO exhibited increased levels of succinyl-CoA and malonyl-CoA (Figure S7). Given that
24 pantothenate is a precursor in the CoA biosynthetic pathway, the uptake and conversion of
25 extracellular pantothenate from LB Broth suggest an increase in CoA levels^{27, 28}. Feedback
26 regulation at pantothenate kinase plays a crucial role in maintaining CoA homeostasis²⁷.
27 Consequently, in PanC KO, cells may rely more on external uptake, leading to a distinct
28 rebalancing of intermediate metabolite pools compared to the WT²⁷.

29
30 These regulatory changes have the potential to enhance the flow of carbon into the TCA and fatty
31 acid pathways, leading to accumulation of CoA derivatives such as succinyl-CoA and malonyl-
32 CoA, and increases in various metabolite levels due to downstream reaction and altered CoA
33 utilization.^{29, 30} The increased pantothenate levels likely enhanced coenzyme A (CoA) availability,
34 which is consistent with elevated levels of TCA cycle intermediates, including pyruvate, citrate,
35 succinate, and oxaloacetate,³⁰ while fumarate and malate may be depleted due to their use in
36 biosynthetic processes,³¹ as illustrated in Figure 4C.

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38 The CoA availability and altered metabolite abundances lead to enhanced catabolism of
39 branched-chain amino acids such as Val, Leu, and Ile,³² while the elevated levels of amino acids,
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3 including Glu, Gln, and Arg, support nitrogen balance and anabolic demands^{33 34}(Figure 4D).
4 These results are consistent with those reported in the literature, showing that *E. coli* disruptions
5 in pantothenate synthesis can be bypassed through uptake mechanisms or alternative regulatory
6 adaptations that elevate pantothenate and other metabolite levels in the pathway^{29, 35, 36}.

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10 In the fatty acid and dicarboxylate metabolites (Figure S8A), an increase in fatty acids level such
11 as acetate, caprylate, decanoate, lactate, and hydroxypropionate is consistent with altered fatty
12 acid synthesis associated with perturbed CoA homeostasis.³⁷ The PanC KO strains are known to
13 exhibit changes in fatty acid-associated metabolites due to CoA's central role in acyl-group
14 activation and fatty acid metabolism, leading to the accumulation of specific organic acids when
15 canonical pathways are constrained.^{29, 37-39} Concurrent decreases in dicarboxylates such as
16 propionate, suberate, and glutarate suggest selective depletion of metabolite pools that depend
17 on CoA-linked processing steps.

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The results of amino acid derivatives and polyamine-related metabolites profiles (Figure S8B)
suggest a compensatory response to PanC KO disruption. Increased levels of cysteine and
cystine suggest an enhanced sulfur amino acid availability, which has been linked to stress
adaptation and maintaining redox balance, under conditions of perturbed CoA metabolism^{40, 41}.
Increased ornithine and 1,3-diaminopropane indicate remodeling of polyamine metabolism.
Polyamine synthesis is coupled to nitrogen balance and is known to respond to perturbations in
central metabolism, such as CoA availability.^{42, 43}

The significant increase in hydroxyphenylacetate, hydroxyphenyllactate, pyroglutamate, methyl-
oxoalate, hydroxy, and keto-acid derivatives (Figure S8C) derived from aromatic and
branched-chain amino acids is consistent with altered amino acid processing in the PanC KO
strain. These metabolites accumulate when amino acid pathways are rebalanced or when
downstream reactions requiring CoA are limited, leading to diversion into alternative derivative
pools.^{44, 45} This pattern has been observed in *E. coli* strains with disruptions in pantothenate or
CoA biosynthesis, where amino acid-derived organic acids increase as part of a broader
metabolic adjustment.^{27, 39}

The results show a significant increase in N-acetylated amino acids in PanC KO (Figure S8D),
indicating enhanced amino-acid acetylation. The depletion of acetyl-CoA in the PanC KO strain
suggests its usage to enhance amino-acid acetylation.⁴⁶ This is consistent with the increased
availability of CoA-derived acetyl groups, which are known to support amino acid modification and
intracellular homeostasis in *E. coli*.^{30, 47}

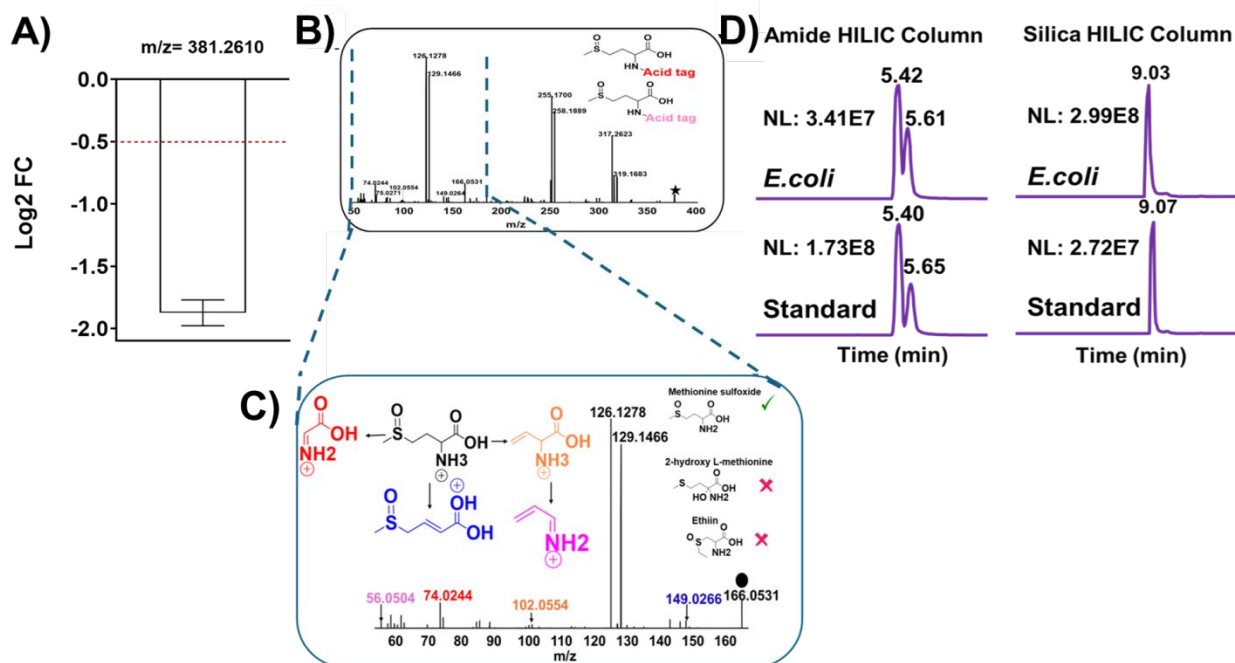


Figure 5. Untargeted barcode isobaric tagging in a separate injection for metabolite quantitation. An untargeted m/z quantified using the R package CluMSID (A). MS/MS spectrum of labeled untargeted mass containing a unique amine reporter ion, indicating that these metabolites have been labeled by an acid barcode tag (B). The black circle over 166.0531 is the m/z of the untagged metabolite. The mass of the tag is subtracted from the untargeted mass, and searching the database yielded three metabolites with the same mass (black circle), and the fragmentation pattern is consistent with methionine sulfoxide (C). Validation of methionine sulfoxide using level one confirmation (D).

Our proof-of-concept barcode tagging scheme not only targets one hundred specific metabolites but also serves as an untargeted approach. Using the DDA method, over 150 masses were quantified, expanding analytical capabilities beyond the initial target list. The identification of these additional untargeted masses is based on the detection of unique reporter ions of the barcode isobaric tags, highlighting the versatility and potential of this tagging scheme. An untargeted m/z of 381.261 (Figure 5A) containing reporter ions of 126.1278 and 129.1466 (Figure 5B) indicates the presence of an amine-containing metabolite. To identify this metabolite, the mass of the acid tag was subtracted from the untargeted mass, resulting in an m/z of 166.0531 (Figure 5C). This value was then cross-referenced with a metabolic database, leading to potential hits as methionine sulfoxide, 2-hydroxy-L-methionine, or ethiin metabolites, all of which share the same mass (Figure 5C).

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3 The fragmentation pattern indicates that the unknown mass is methionine sulfoxide. To explore
4 this further, the barcode acid tag was used to label both the methionine sulfoxide standard and
5 the methionine sulfoxide present in the *E. coli* lysate. Preliminary confirmation at level one shows
6 a consistent retention time and peak shape between the methionine sulfoxide standard and the
7 methionine sulfoxide in the *E. coli* lysate across different amide and silica columns (Figure 5D).
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11 Similarly, gamma-aminobutyric acid, which was not present in the initial 100 metabolite standards,
12 was quantified using an untargeted approach and validated at the one-level confirmation using a
13 chemical standard, as shown in Figure S9B. These examples show the potential for uncovering
14 novel metabolic changes from genetic knockouts as well as identifying new metabolites that are
15 not present in the user's metabolic library.
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19 4. CONCLUSION

20 A proof-of-concept 2-plex barcode isobaric tags has been developed, enabling wide coverage of
21 metabolites. These tags undergo double fragmentation to produce a cyclized product, followed
22 by a distinct reporter ion that serves as an identifier for acid- and amine-containing metabolites.
23 The acid tags are used for the derivatization of amine-containing metabolites, while the amine
24 tags are used for the tagging of acid-containing metabolites. In the PanC KO experimental strain,
25 significant changes in intracellular metabolite abundances were observed relative to the control,
26 confirming that the changes are specifically associated with PanC KO uptake. This 2-plex barcode
27 isobaric tagging scheme also doubles as an untargeted barcode isobaric tagging platform,
28 enabling metabolite identification and quantitation when standards are unavailable.
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33 This method could be expanded up to 128-plex multiplexing with broad metabolite coverage.
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Data for this article, including processed data, raw data files, sample collection, sample preparation, study design, protocol, study metadata, and method, have been uploaded to the Metabolomics Workbench under datatrack ID 7373. Study ID and DOI will be provided once the Metabolomics Workbench review process is complete and the submission is processed.

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