

Three-minute method for amino acid analysis by UHPLC and high-resolution quadrupole orbitrap mass spectrometry

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Abstract Amino acid analysis is a powerful bioanalytical technique for many biomedical research endeavors, including cancer, emergency medicine, nutrition and neuroscience research. In the present study, we present a 3 min analytical method for underivatized amino acid analysis that employs ultra high-performance liquid chromatography and high-resolution quadrupole orbitrap mass spectrometry. This method has demonstrated linearity (mM to nM range), reproducibility (intra-day <5 %, inter-day <20 %), sensitivity (low fmol) and selectivity. Here, we illustrate the rapidity and accuracy of the method through comparison with conventional liquid chromatography–mass spectrometry methods. We further demonstrate the robustness and sensitivity of this method on a diverse range of biological matrices. Using this method we were able to selectively discriminate murine pancreatic cancer cells with and without knocked down expression of hypoxia-inducible factor

1 α ; plasma, lymph and bronchioalveolar lavage fluid samples from control versus hemorrhaged rats; and muscle tissue samples harvested from rats subjected to both low-fat and high-fat diets. Furthermore, we were able to exploit the sensitivity of the method to detect and quantify the release of glutamate from sparsely isolated murine taste buds. Spiked in light or heavy standards (¹³C₆-arginine, ¹³C₆-lysine, ¹³C₅¹⁵N₂-glutamine) or xenometabolites (5-fluorouracil) were used to determine coefficients of variation, confirm linearity of relative quantitation in four different matrices, and overcome matrix effects for absolute quantitation. The presented method enables high-throughput analysis of low-abundance samples requiring only one percent of the material extracted from 100,000 cells, 10 μ l of biological fluid, or 2 mg of muscle tissue.

Keywords QExactive · Biofluid · Cell extract · Tissue extract · Amino acids · Mass spectrometry · PLS-DA

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Abbreviations

3AA	Three-minute method for amino acid analysis
BALF	Bronchioalveolar lavage fluid
CV	Coefficient of variation
GC/MS	Gas chromatography/mass spectrometry
HFD	High-fat diet
HIF1 α	Hypoxia-inducible factor 1 α
HILIC	Hydrophilic interaction liquid chromatography
HPLC	High-performance liquid chromatography
LFD	Low-fat diet
LOD	Limit of detection
LOQ	Limit of quantification
MAP	Mean arterial pressure
MLD	Mesenteric lymph diversion
MRM	Multiple reaction monitoring
MS	Mass spectrometry

<i>m/z</i>	Mass-to-charge ratio
PCA	Principal component analysis
PLS-DA	Partial least-square discriminant analysis
SD	Standard dilution
S/N	Signal to noise ratio
T/HS	Trauma/hemorrhagic shock
T/SS	Trauma/sham shock
TOF	Time of flight
UHPLC	Ultra high-performance liquid chromatography
XIC	Extracted ion chromatogram

Introduction

Amino acid analysis is a critical research endeavor that has attracted a great deal of interest since the late 1950s when Stein and Moore established an analytical workflow to detect and quantify amino acid mixtures from protein hydrolysates (Manning 1993). While historically employed to determine amino acid composition of proteins and peptides, this technique has recently seen expanded utility resulting from the increasingly documented connections between amino acid metabolism and biological responses. In cancer research, for example, glutamine (Cardaci and Ciriolo 2012), serine (D'Alessandro and Zolla 2013), glycine (Amelio et al. 2014a, b) and tryptophan metabolism (Platten et al. 2012) now represent novel targets for drug therapeutics. Additionally, stem cells feed on threonine to promote energy metabolism and mediate epigenetic regulatory post-translational modifications of histones (Chen and Wang 2014), thus providing additional areas for therapeutic intervention. In the fields of transfusion medicine and red blood cell storage, glutamine, glutamate and cysteine are emerging as potential contributors to age-dependent impairments in anti-oxidant potential (Droge 2005) and the accumulation of storage lesions in the blood bank (D'Alessandro et al. 2015a). In the field of neurobiology, glutamate, tryptophan, and tyrosine can directly mediate stress responses such as ischemia/reperfusion injury and mood disorders (D'Alessandro et al. 2014b) since they are precursors to γ -aminobutyric acid, serotonin/melatonin and catecholamines. In the field of food science, glutamate has been associated with the umami taste and improved sensory appreciation of meat and beverages (Yamaguchi and Ninomiya 2000). Examples of the connections between amino acid metabolism and biological responses are therefore both plentiful and diverse.

A long list of methods for amino acid analysis have been published over the last 50 years (Rutherford and Gilani 2009). The technique was pioneered by Moore et al. (1958), with the original iteration achieving amino acid separation on a sulfonated cation-exchange resin. The separated amino acids were then detected by calorimetry

using a post-column reaction with ninhydrin. More recent methods rely upon reversed-phase derivatization through different approaches, such as the use of phenylisothiocyanate (Heinrikson and Meredith 1984), ion chromatography coupled with post-column derivatization that uses ninhydrin (Bidlingmeyer et al. 1984), fluorecamine (Stein et al. 1973), or *o*-phthalaldehyde/2-mercaptoethanol (Benson and Hare 1975; Dorresteijn et al. 1996), or cation-exchange solid-phase extraction followed by propyl chloroformate derivatization and analysis by gas chromatography/mass spectrometry (GC/MS) (Kaspar et al. 2008; Badawy 2012). Alternatively, pre-column derivatization methods have been proposed through the use of *o*-phthalaldehyde, 9-fluorenylmethyl chloroformate, phenyl isothiocyanate, 1-dimethylaminonaphthalene-5-sulphonyl chloride (dansyl-chloride), *N*-alkylnicotinic acid, 3-aminopyridyl-*N*-hydroxysuccinimidyl carbamate or *N*-hydroxysuccinimide (Fürst et al. 1990; Yang et al. 2006; Shimbo et al. 2009).

Recent improvements in capillary electrophoresis, high-performance liquid chromatography (HPLC), and mass spectrometry (MS) technologies have revolutionized the field of metabolomics, consequently providing opportunities for the development of derivatization-free amino acid analysis (Armstrong et al. 2007; Hirayama and Soga 2012). These methods make use of the improved chromatographic selectivity of sub 2- μ m resin particles that offer increased surface area to volume ratios (Armstrong et al. 2007), and reversed-phase ion-pairing chromatography by perfluorinated carboxylic acids (Piraud et al. 2005). In parallel, advances in mass spectrometry technologies now provide enhanced sensitivity down to the low nM range for small molecules, primarily due to the use of such targeted approaches as multiple reaction monitoring (MRM) available on triple quadrupole (Piraud et al. 2005) and ion trap mass analyzers (D'Alessandro et al. 2011), and hybrid quadrupole time of flight (TOF) or Orbitrap mass analyzers.

While these methods are simple, selective, and sensitive, they often rely on sample preparation kits that require amino acid derivatization and subsequent analysis by GC/MS and LC/MS in 6, 10 and 15 min (Kaspar et al. 2008; Shimbo et al. 2009; Badawy 2012; Salazar et al. 2012). Despite increasing the time and effort required for sample preparation, chemical derivatization makes amino acid assignment and quantitation down to the μ M level amenable to routine analysis. Additional LC/MS methods for the detection of underivatized amino acids have been described that rely on 12–15 min analytical times and multiple reaction monitoring (MRM) for amino acid detection (Thiele et al. 2012; Zhou et al. 2013; Yao et al. 2013; Buiarelli et al. 2013; Le et al. 2014).

Although automated sample preparation and derivatization of biofluid extracts improve efficiency,

analytical run time is a critical factor for translational applications of amino acid analysis such as clinical biochemistry (D'Alessandro et al. 2012b) and drug development (D'Alessandro and Zolla 2013). In the present study, we have harnessed the improved resolving power of ultra high-performance liquid chromatography (UHPLC) to optimize a highly sensitive three-minute method for amino acid analysis (3AA). Such method is enhanced by both the chromatographic selectivity of 1.7- μm C18 particles and accurate intact mass determination within a sub 5 ppm range of error using a hybrid quadrupole Orbitrap mass spectrometer. This method has demonstrated linearity (mM to nM range), reproducibility (intra-day <5 %, inter-day <20 %), sensitivity (down to the nM level) and selectivity using commercially available underivatized standards. Although this method is affected by matrix composition, linearity and reproducibility of relative and absolute quantitation are not compromised, as demonstrated in four different matrices of clinical relevance (red blood cells, plasma, bronchioalveolar lavage fluid and mesenteric lymph) by spiked in supplementation of light and heavy standards ($^{13}\text{C}_6$ -arginine, $^{13}\text{C}_6$ -lysine, $^{13}\text{C}_5^{15}\text{N}_2$ -glutamine) or xenometabolites (5-fluorouracil) at different concentrations. Furthermore, this method is quite robust and has been successfully employed to analyze amino acid content in different biological matrices including biofluids, cell extracts, and tissue extracts. Briefly, we provide evidence to support the utility of accurate amino acid quantitation in biological matrices as a high-throughput discovery mode analysis aimed at rapidly discriminating phenotypes through unsupervised statistical analyses such as Principal Component Analysis (PCA).

Materials and methods

Sample preparation

Biofluids

Rat plasma samples were kindly provided by Drs. Hunter Moore and Anthony W. Bacon (University of Colorado Denver). Briefly, trauma was induced in Sprague–Dawley rats through laparotomy and bowel crush, as previously reported (D'Alessandro et al. 2015b). Sham shock (SS—no hemorrhage) or hemorrhagic shock (HS) was induced by controlled hemorrhage until rat achieves mean arterial pressure (MAP) equal to 25 mm Hg (MAP 25). Matched plasma, mesenteric lymph and bronchioalveolar lavage fluid (BALF) samples (100 μl) were collected before (pre) and immediately after 30 min of trauma/hemorrhagic shock (at MAP 25) (post), either in presence or absence of mesenteric lymph diversion (MLD). Ten μl of

each plasma was extracted in 490 μl of ice-cold extraction buffer (methanol:acetonitrile:water 5:3:2 v/v), vortexed for 10 min at 4 °C. Proteins and lipids were subsequently pelleted through centrifugation for 10 min at 10,000g at 4 °C and supernatants collected for subsequent analysis.

Taste bud releasates were kindly provided by Dr. Aurelie Vandenbeuch (University of Colorado Denver). Samples were harvested from isolated mouse tongue epithelium cells stimulated in vitro with KCl in PBS buffer, as previously reported (Vandenbeuch et al. 2010).

Cell extracts

Pancreatic cancer (Panc-1) cells (10^7 cells, biological triplicates) were kindly provided by Dr. Agnieszka Kendrick (University of Colorado, Denver). Cells were grown in DMEM, 10 % fetal bovine serum, 1 % penicillin/streptomycin, 1 % non-essential amino acids and harvested at 90 % confluence in 60-mm dishes.

Murine Pan02 pancreatic adenocarcinoma cancer cells and shHIF1 α knock downs were kindly provided by Dr. Jeniann Yi (University of Colorado Denver—CO, USA). The Pan02 murine pancreas adenocarcinoma cell line (National Cancer Institute, Bethesda, MD) was modified using short-hairpin RNA targeting the HIF1 α gene (Sigma Aldrich) via lentiviral transduction to create Pan02-SH+ cells lacking HIF1 α activity. Cells were cultured in RPMI + 10 % FBS and 1 % penicillin/streptomycin (Cellgro) to 50 % confluence, then exposed to the shRNA/lentiviral particles. Transduced cells were selected using puromycin at a concentration of 3 $\mu\text{g}/\text{ml}$. Successful knockdown of HIF1 α activity using two distinct vectors confirmed using real-time polymerase chain reaction (RT-PCR) at both room temperature and simulated hypoxia. Five biological replicates were obtained for wild type or shHIF1 α (two different clones, hereby referred to as shHIF1 and shHIF2) with cell counts as low as 3.7×10^5 .

All cell pellets were stored at -80 °C and were extracted immediately before analysis in ice-cold lysis/extraction buffer (methanol:acetonitrile:water 5:3:2 v/v) at a ratio of 2×10^6 cells/ml (Maddocks et al. 2013; D'Alessandro et al. 2014a). Samples were then agitated at 4 °C for 30 min and centrifuged at 10,000g for 10 min at 4 °C. Protein and lipid pellets were discarded, while supernatants were stored at -80 °C prior to analyses.

Tissue extracts

Five mg of tissue biopsies were obtained from Drs. Bryan Bergman, Sean Newsom (University of Colorado Denver—CO, USA) and Drs. Gregory D. Cartee and Carlos M. Castorena (University of Michigan). Animals used in this study were part of a larger, recently published investigation

(Castorena et al. 2014). All procedures for animal care were approved by the University of Michigan Committee on Use and Care of Animals. Eight-week-old male Wistar rats (initial body weight ~200–250 g; Harlan, Indianapolis, IN) were individually housed and provided with standard rodent chow (Low-Fat Diet; LFD: 14 % kcal fat, 58 % kcal carbohydrate, 28 % kcal protein; Lab Diet no. 5001, PMI Nutritional International, Brentwood, MO) or high-fat diet [HFD: 60 % kcal fat, 20 % kcal carbohydrate, 20 % kcal protein; Research Diets (D12492), New Brunswick, NJ] and water ad libitum for 2 weeks. Rats were fasted at ~1900 h on the night before the terminal experiment. At ~12:00 h the next day, fasted animals (for ~17 h) from both groups were anesthetized (Castorena et al. 2014), and tissues were harvested and then immediately frozen in liquid N₂. Frozen skeletal muscle samples were weighed (5 mg) and homogenized in 1 ml of ice-cold extraction buffer (methanol:acetonitrile:water 5:3:2 v/v) for 2 min at 30 Hz in TissueLyser (QIAGEN -Valencia, CA, USA). Homogenized samples were vortexed for 30 min at 4 °C. Samples were then centrifuged for 10 min at 10,000g at 4 °C. The supernatants were then collected for the analysis, while Bradford assays were performed on the pellets, as to determine the comparability of the extracted samples and eventually normalize results on protein concentrations.

Amino acid standard characterization

To determine linearity and sensitivity of the approach, we weighed and progressively diluted down to the nM level a subset of amino acid standards, including commercially available L-glutamic acid, and glycine (49621, G8898,—SIGMA Aldrich, St. Louis, MO, USA), MSMLS standard compound library (IROATech, Bolton, MA, USA) and EZ faast amino acid standards (Standard dilution SD1, SD2, SD3—AG0-7184—Phenomenex, Torrance, CA, USA). A total of thirty-five different amino acids were tested using commercially available standards (SD1, SD2, SD3—Phenomenex, Torrance, CA, USA). To determine reproducibility of the approach, we diluted the SD1 mixture to a final concentration of 50 nM for each amino acid with 18 mΩ H₂O and analyzed 10-μl injections five consecutive times. To validate separation of L-leucine and L-isoleucine, 99 % pure amino acids were weighed and used to prepare standard solutions of 1 μM (L0029—TCI America, Portland, OR, USA, 166170050—Acros Organics, New Jersey, USA). Leu and Ile standards were thus analyzed both independently and as a 1:1 mixture. To test retention time reproducibility against standards, samples were either dried under vacuum and resuspended in water or directly analyzed in the lysis/extraction buffer.

1–10 μl of standards at different concentrations, 5 μl of biofluid (either plasma or taste bud releasates), 20 μl of

cell, or 10 μl of tissue extracts were injected into a UHPLC system (Ultimate 3000, Thermo, San Jose, CA, USA) and separated through a 3-min isocratic elution on a Kinetex XB-C18 column (150 × 2.1 mm i.d., 1.7 μm particle size—Phenomenex, Torrance, CA, USA) at 250 μl/min (mobile phase: 5 % acetonitrile, 95 % 18 mΩ H₂O, 0.1 % formic acid; column temperature: 25 °C).

Comparison against two conventional metabolomics methods

Technical consistency (i.e., reproducibility and linearity) of the 3AA method was compared to two conventional chromatographic separations used for small molecule and amino acid analyses. Pancreatic cancer cell extracts (biological triplicates) either undiluted or diluted fivefold, were analyzed using (1) the 3AA method, as well as (2) a 15-min gradient on a Kinetex HILIC column (150 × 2.1 mm i.d., 1.7 μm particle size—Phenomenex, Torrance, CA, USA) and (3) a 23-min gradient on an Acquity UHPLC BEH Amide Column (2.1 × 100 mm, 1.7 μm—Waters, Milford, MA, USA). In (2), samples were analyzed on the Kinetex HILIC at 350 μl/min (mobile phases: A: ACN; B: 18 mΩ H₂O, 20 mM (NH₄)₂CO₃, 0.1 % NH₄OH; gradient: 1.5 min hold at 5 % B; 5–60 % B in 8.5 min; 60–95 % B in 0.5 min at 0.5 ml/min; 95 % hold for 2 min; 95–5 % B in 0.5 min; 5 hold for 2 min; column temperature: 25 °C). In (3), samples were analyzed on the Acquity UHPLC BEH Amide Column (2.1 × 100 mm, 1.7 μm—Waters, Milford, MA, USA) at 500 μl/min (mobile phases: A: 18 mΩ H₂O, 20 mM (NH₄)₂CO₃ pH 4.00; B: acetonitrile; gradient: 3 min hold 85 % B; 85–30 % B in 9 min; 30–2 % B in 3 min; 2–85 % of B in 1 min; 85 % equilibration for 7 min; column temperature: 60 °C).

The UHPLC system was coupled online with a QExactive mass spectrometer (Thermo, San Jose, CA, USA), scanning in Full MS mode (2 μscans) at 70,000 resolution from 60 to 900 *m/z*, with 4 kV spray voltage, 15 sheath gas and 5 auxiliary gas, operated in positive ion mode. Calibration was performed before each analysis using a positive calibration mix (Piercenet—Thermo Fisher, Rockford, IL, USA). Limits of detection (LOD) were characterized by determining the smallest injected amino acid amount required to provide a signal to noise (S/N) ratio greater than three using <5 ppm error on the accurate intact mass. Based on a conservative definition for Limit of Quantitation (LOQ), these values were calculated to be threefold higher than determined LODs.

MS data acquired from the QExactive were converted from .raw file format to .mzXML format using MassMatrix (Cleveland, OH, USA). Amino acid assignments were performed using MAVEN (Princeton, NJ, USA). The MAVEN software platform provides tools for peak picking, feature

detection and metabolite assignment against the KEGG pathway database. Assignments were further confirmed using a process for chemical formula determination using isotopic patterns and accurate intact mass (Clasquin et al. 2012). Analyte retention times were confirmed by comparison with external standard retention times, as indicated above.

Relative quantitation was performed by exporting integrated peak areas values into Excel (Microsoft, Redmond, CA, USA) for statistical analysis including *T* Test and ANOVA (significance threshold for *p* values <0.05) and unsupervised Principal Component Analysis (PCA) (Pan et al. 2007; Fonville et al. 2010), calculated through the MultiBase macro (freely available at <http://www.NumericalDynamics.com>).

Matrix effects: spiked in standards at different concentrations

The effect of matrix-dependent ion suppression was tested by injecting the amino acid standards (SD1) at different amounts (0.5, 2.5 and 5 pmol injected), either alone or as incremental additions to a complex matrix of three freshly withdrawn packed red blood cell extracts. Briefly, whole blood was collected from three healthy donor volunteers, in compliance with the declaration of Helsinki, and red blood cells were sorted from plasma and buffy coat through centrifugation (1500g × 10 min at 4 °C). Additionally, to further test intra-day reproducibility of sample processing, analytical runs, and coefficients of variation (CV—standard deviation/mean) for relative quantitation, a xenometabolite (5-fluorouracil—F6627 SIGMA Aldrich, St. Louis, MO, USA) was added to the lysis buffer used to extract the same samples at a final concentration of 25 µM.

Absolute quantitation using heavy-labeled amino acids

To overcome matrix effects and achieve absolute quantitation, three heavy-labeled amino acid standards (¹³C₆-arginine, ¹³C₆-lysine, ¹³C₅¹⁵N₂-glutamine—99 atom % ¹³C; 95 % compound purity—SIGMA Aldrich, St. Louise, MO, USA) were spiked in the lysis/extraction buffer at a final concentration of 10 µM. This supplemented buffer was used to extract matched plasma, BALF, and mesenteric lymph from control, trauma sham shock (T/SS) or hemorrhagic shock (T/HS) rats prior to or after shock, in presence or absence of MLD, as described above. Each condition contained two biological replicates and a 1:1 ratio of the replicates at each time point. Peak areas for heavy-labeled standards in each sample were used to calculate inter-run CVs.

To demonstrate linearity of this approach, an additional 1:1 biological replicate ratio was prepared with heavy-labeled standards added to a final concentration of 20 µM. Ratios of heavy-labeled standard peak areas from the 1:1

ratio with either 10 or 20 µM standards were used to calculate linearity of quantitation and CVs of fold changes (expected fold change values = 2.00).

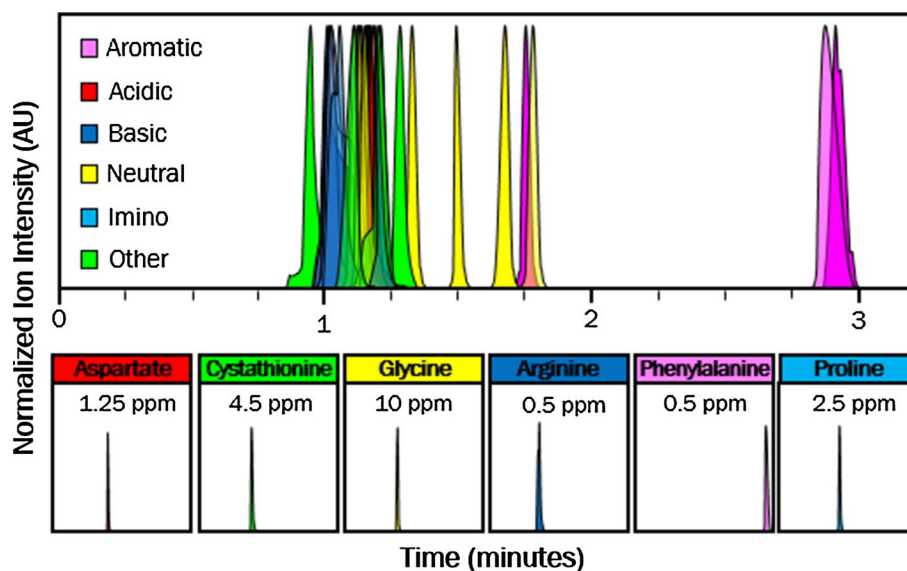
Heavy to light isotopologue ratios for each one of the three amino acid standards were used to determine absolute quantities of arginine, lysine, and glutamine in the two biological replicates, as well as the 1:1 mixture (with a conversion factor of either 10 or 20 µM, depending on the concentration of the heavy-labeled standards).

Results and discussion

Selectivity, linearity, sensitivity and reproducibility of the approach

In Fig. 1 and Table 1, we report an overview of retention times for 32 representative basic (blue), acidic (red), neutral (yellow), aromatic (pink), imino (light blue) and non-essential (green) amino acids along the three min isocratic run. While extracted ion chromatograms (XIC) for a representative class of amino acids are depicted in Fig. 1, XICs for the entire set of profiled amino acids are depicted in Supplementary Fig. 1. Basic and neutral amino acids were weakly retained by the column, with basic amino acids eluting within a 10 s time window around the first min of the elution. Neutral and acidic amino acids eluted from 1.10 to 1.80 min, and aromatic compounds eluted near the end of the run (2.8–2.9 min). Mean peak width was 3 s for most amino acids, while approximately 6–8 s for basic amino acids. Narrow peak width resulting from the isocratic elution increased the signal to baseline ratios, thereby LOD and LOQ in comparison to the 15-min HILIC and 23-min Amide HILIC analysis. Selectivity of the approach is further demonstrated by the ability to independently identify both leucine and isoleucine with baseline separation using both commercially available standard mix and single standards (Figs. 1, 2; Table 1), even though separation of alloisoleucine was not achieved. Other isobaric compounds could be independently resolved including 4-aminobutanoate and L-amino-isobutanoate, sarcosine and alanine (unique retention times indicated in Table 1), providing additional supporting evidence of the specificity of the approach. Accurate intact mass enabled assignment and quantitation of co-eluting amino acids (Table 1). Reproducibility of the approach is illustrated by the CV for each amino acid measured through five consecutive analyses of SD1 (500 fmol injections—Supplementary Table 1). CV was lower than 5 % for every amino acid (3.92 ± 0.01 %) with the exception of L-alanine (6.81 %) (Supplementary Table 1). Inter-day variability, calculated using three injections of SD1 on three consecutive days, resulted in CVs <20 % (13.7 ± 10.8 %—Supplementary Table 2).

Fig. 1 Extracted ion chromatograms of amino acids. Extracted ion chromatograms (XIC) for 32 representative amino acids assayed in this study have been overlaid to demonstrate chromatographic separation (*top*). Amino acid categories are color coded as follows: aromatic (*pink*), acidic (*red*), basic (*blue*), neutral (*yellow*), imino (*light blue*), or other (*green*) amino acids. Six representative XICs are presented to illustrate peak shape (*bottom*). Parts per million (ppm) values used to detect each amino acid are provided for each XIC (color figure online)



In Table 1 we report the compound name, along with the KEGG pathway compound ID, the MS polarity in which all the amino acids were detected (positive), the observed parent ion, retention times, logarithms of linearity for quantitative analysis, quadratic correlation coefficients (R^2 —also highlighted in Supplementary Fig. 2), LOQ and LOD values. The highest LOQ and LOD values were observed for glycine (390 and 130 pmol, respectively), while many metabolites showed linearity over three to five (logarithmic) orders of concentrations, down to the lowest tested injection amount (1 fmol). Importantly, this range is relevant to biological contexts, where metabolite quantities are often in the range of mM to μ M concentrations (Armstrong et al. 2007).

The linearity of quantitation over several orders of magnitude down to even low fmol levels (exemplified by L-glutamate in Fig. 3a, b) expands the capability of this analytical platform to analyze and determine absolute quantitation of extremely dilute or scarce samples. One interesting example of a dilute sample is the neurological releasate from taste buds. A role for glutamate as an efferent neurotransmitter in taste buds has been recently proposed (Vandenbeuch et al. 2010), though only indirect evidence has been provided due to the lack of analytical capability to directly identify this compound in such a dilute sample. While our analysis of neurological releasates (glutamate release from taste buds—Fig. 3a—box) exemplifies the capability of 3AA to assess low-level metabolites, it suggests that the described approach might enable the analysis of other clinically relevant low-volume/abundance matrices. Such matrices could include blastocyst fluid, as previously described (D'Alessandro et al. 2012a), and stem cell populations that currently require pooling from different sources for analysis due to low abundance, which limits the

biological readout of stem cell studies (Panopoulos et al. 2012).

Despite the absence of a gradient, no carryover of any analyte was observed when testing simple or complex biological matrices, including taste bud releasate (Fig. 3a box) and cell extracts (Supplementary Fig. 3). Carry over effects were not observed in any of the tested matrices, whether the samples were resuspended in water or directly injected in the organic sample extraction buffer (methanol:acetonitrile:water). Retention times of amino acids in the samples were consistent with analytical standards only when injecting samples resuspended in water. However, direct injection of samples in the organic extraction/lysis buffer resulted in a minor reduction in retention times (-6% median deviation—3.6 s—Supplementary Table 3), while accuracy was not affected (<5 ppm error on the accurate intact mass). Despite this observation, this method has been developed to provide a high-throughput platform for amino acid analysis and it is worth noting that the implementation of an extra drying/resuspension step would represent a burden in terms of sample processing time.

Notably, results obtained with the 3AA method were consistent with those collected using a longer HILIC gradient, a method that represents one of the current standards of practice in the field of polar compound metabolomics (Cubbon et al. 2010) (Supplementary Fig. 3). Although less stable than RP approaches, the polar compound-friendly chemistry of HILIC makes it an eligible strategy to analyze polar matrices such as biofluids and cell extracts, as extensively reviewed by Cubbon and colleagues (Cubbon et al. 2010). Therefore, the ability of the RP-based 3AA method to replicate results determined using an HILIC-based method establishes 3AA as a more efficient, alternative

Table 1 Amino acid standards

No.	Mix	Compound	KEGG Cmpd ID	Parent ion	Accuracy (ppm)	Median Rt	Log LIN	R ²	LOQ	LOD
1	SD1	L-2-aminoadipate	C00956	162.0761	0.5	1.26	4	0.99	750 fmol	250 fmol
2	SD1	L-alanine	C00041	90.0555	5	1.30	4	0.96	300 fmol	100 fmol
3	SD1	L-arginine	C00062	175.1190	0.5	1.10	4	0.96	75 fmol	25 fmol
4	SD1	L-aspartate	C00049	134.0446	1.25	1.30	5	1.00	7.5 pmol	2.5 pmol
5	SD1	L-3-amino-isobutanoate	C03284	104.0711	3.5	1.18	4	0.97	4.5 pmol	1.5 pmol
6	SD1	L-cysteine	C00097	122.0271	1	1.26	3	0.99	<1 fmol	<1 fmol
7	SD1	L-cystine	C00491	241.0311	0.6	1.16	4	0.98	<1 fmol	<1 fmol
8	SD1	L-citrulline	C00327	174.0874	0.35	1.27	3	0.99	<1 fmol	<1 fmol
9	SD1	4-aminobutanoate	C00334	104.0711	3.5	1.21	4	0.97	4.5 pmol	1.5 pmol
10	SD1	L-glutamate	C00025	148.0604	0.25	1.20	4	1.00	61 fmol	20.3 fmol
11	SD1	Glycine	C00037	76.0400	10	1.14	4	0.99	390 pmol	130 pmol
12	SD1	L-histidine	C00135	156.0765	1	1.22	4	0.99	2 pmol	750 fmol
13	SD1	Trans-L-3-hydroxyproline	C05147	132.0656	3	1.22	3	0.99	2 pmol	750 fmol
14	SD1	L-isoleucine	C00407	132.1020	0.5	1.78	4	0.98	480 fmol	160 fmol
15	SD1	L-leucine	C00123	132.1020	0.5	1.72	4	0.99	420 fmol	140 fmol
16	SD1	L-lysine	C00047	147.1128	0.5	1.05	3	0.98	750 fmol	250 fmol
17	SD1	L-methionine	C00073	150.0584	1	1.64	4	0.99	<1 fmol	<1 fmol
18	SD1	N(pi)-methyl-L-histidine	C01152	170.0924	3.5	1.04	3	0.95	1.5 pmol	500 fmol
19	SD1	L-ornithine	C00077	133.0973	2	1.03	3	0.99	22.5 pmol	7.5 pmol
20	SD1	L-phenylalanine	C00079	166.0863	0.5	2.92	4	0.99	480 fmol	160 fmol
21	SD1	L-proline	C00148	116.0709	2.5	1.25	4	0.99	7.5 pmol	2.5 pmol
22	SD1	Sarcosine	C00213	90.0555	5	1.16	3	0.99	600 fmol	200 fmol
23	SD1	L-serine	C00065	106.0501	4.5	1.24	3	0.99	7.5 pmol	2.5 pmol
24	SD1	L-threonine	C00188	120.0656	3	1.29	4	1.00	1.3 pmol	430 fmol
25	SD1	L-tyrosine	C00082	182.0812	0.25	1.79	4	1.00	<1 fmol	<1 fmol
26	SD1	L-valine	C00183	118.0866	3	1.41	4	1.00	15 pmol	5 pmol
27	SD2	L-asparagine	C00152	133.0609	1.5	1.21	4	0.99	7.5 pmol	2.5 pmol
28	SD2	L-glutamine	C00064	147.0765	0.3	1.19	4	0.98	15 pmol	5 pmol
29	SD2	L-tryptophan	C00078	205.0973	0.3	2.79	3	0.99	<1 fmol	<1 fmol
30	SD3	5-hydroxy-L-lysine	C16741	163.1076	1	1.03	4	0.99	780 fmol	260 fmol
32	SD3	Cystathionine	C00542	223.0745	4.5	1.13	4	0.99	<1 fmol	<1 fmol
33	SD3	Pro-HYP	C10H16N2O4	229.1181	0.5	1.18	4	0.99	<1 fmol	<1 fmol
34	SD3	Gly-Pro	C7H12N2O3	171.0765	1	1.42	4	0.99	750 fmol	250 fmol
35	SD3	Thioproline	C6H12SNO3	173.0920	4.5	1.25	4	0.99	6 pmol	2 pmol

method for the rapid analysis of amino acid content for high-throughput discovery mode purposes.

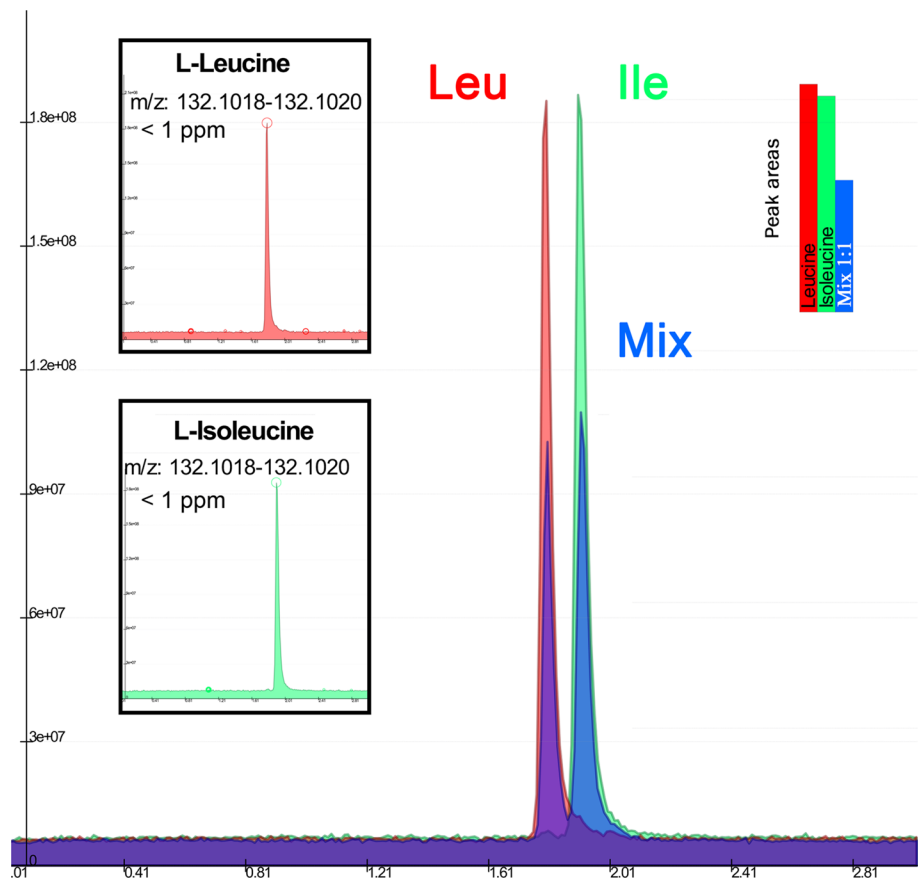
Robustness of the method in different biological matrices

As a proof of principle, the applicability of this method for general discovery through relative quantitation was tested with three unique biological matrices, including biological fluids (plasma), cell extracts (Pan02 cancer cells) and muscle tissue extracts (Fig. 4). All 20 of the classic amino acids were detected in each matrix, showing the general utility of the method for rapid application to diverse biological

samples. Relative quantities of amino acids determined in these matrices were used to run unsupervised statistical analyses such as PCA and PLS-DA, with the goal of discriminating sample groups into clusters (Fig. 4—Sample Plot) based on the relative contribution of each amino acid (Principal Components, PCs, depicted in the Loading Plot).

Plasma is a key biofluid in clinical and biomedical research settings, as it is used as a proxy for monitoring systemic health (Dunn et al. 2014). In the emergency care setting, plasma metabolic profiles have been recently investigated to monitor the severity of trauma in rats (D'Alessandro et al. 2015b) and humans (Peltz et al. 2015). In particular, trauma and hemorrhagic shock have

Fig. 2 Resolution of leucine and isoleucine. Individual extracted ion chromatograms for leucine (*red*), isoleucine (*green*), and a mixture (*blue*) (1 μ M stocks of 99 % pure standards and a 1:1 mixture, respectively) were overlaid to show resolution of the isobaric amino acids over the 3 min isocratic run at sub 1 ppm error on the intact mass



been associated with increased proteolysis, and plasma amino acid accumulation (Peltz et al. 2015). Here, amino acid analysis of plasma extracts in a rat model of trauma/hemorrhagic shock discriminated the pre-shock baseline samples from shock (Fig. 4a—top panel). Many amino acids showed the same increasing trend in response to trauma/hemorrhagic shock, illustrated by the Loading Plot (Fig. 4a—top panel). Alanine levels, which increase in both the shock and resuscitation time points, are depicted in the Extracted Ion Chromatogram (XIC) in the right-hand panel of Fig. 4a. This particular finding has biological relevance to trauma/hemorrhagic shock response; the conversion of pyruvate to alanine by glutamate–pyruvate transaminase could affect the process of converting pyruvate to lactate by lactate dehydrogenase. Importantly, lactate is known to be a key driver of ketoacidosis in response to trauma (Husain et al. 2003) and thus this finding may offer insight into the rate of lactate accumulation during trauma/hemorrhagic shock.

In cell extracts, a key matrix for example in the field of cancer research, quantitation of amino acids such as L-serine allowed for discrimination between control cells (blue bar plots) and two different clones of cells with HIF1 α knock down (red and green bar plots—Fig. 4, middle panels). Serine metabolism is emerging as a key pathway in

cancer cells (Maddocks et al. 2013), in the light of recent evidence linking serine metabolism to p53-family modulated responses (Maddocks et al. 2013; Amelio et al. 2014a, b; D'Alessandro et al. 2014a). Serine anabolism stems from a branch of aerobic glycolysis often referred to as the Warburg effect (D'Alessandro and Zolla 2013), which is the main energy-generating pathway in rapidly proliferating cancer cells.

Providing further evidence for the utility of the 3AA method, relative amino acid quantitation from tissue extracts could independently cluster biological triplicates of muscle tissue samples obtained from rats subjected to low-fat diet (LFD) versus high-fat diet (HFD) (Fig. 4, lower panels). Glutamine (Fig. 4, lower right panel) was found to contribute to the discriminating pattern between LFD (green bars) and HFD muscle samples (blue bars). Interestingly, previous studies on diabetes, insulin resistance, and its effect on rat liver and muscles have suggested a role for L-glutamine supplementation in the induction of insulin resistance in the adipose tissue, while promoting insulin signaling in muscles and liver (Prada et al. 2007).

Of note, all of the samples described herein were prepared from either 10 μ l of biofluid, 5 mg of muscle tissue, or 3.6×10^5 cells and injection volumes represented 0.5, 1 and 11 % of the total extraction volumes, respectively. The

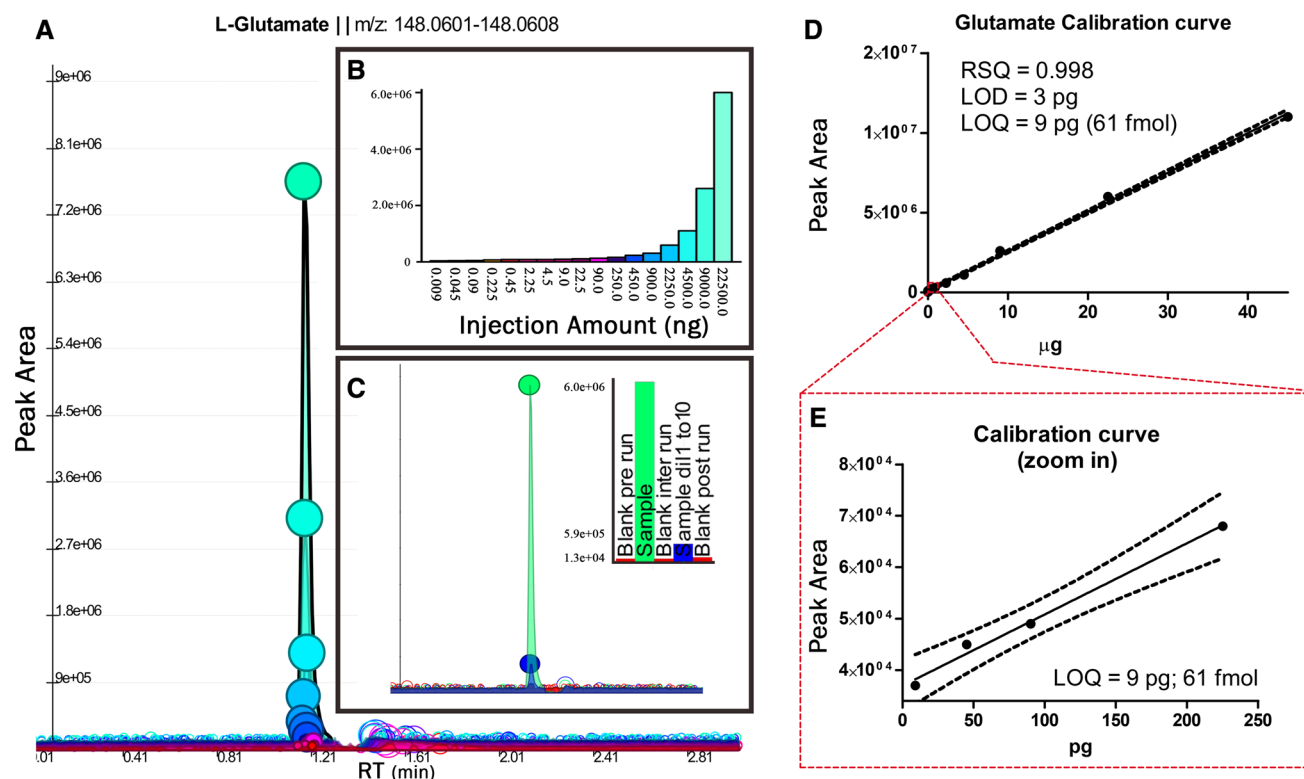


Fig. 3 Glutamate quantitation. In **a** extracted ion chromatograms are shown for 1 μ l injections of L-glutamate ranging from 22.5 μ g/ μ l (153 nmol injected) down to 9 pg/ μ l (61 fmol injected); the *latter injection* represents the limit of quantitation ($3 \times$ Limit of Detection) for this amino acid with the present analytical set up. Peak apexes are indicated by *circles filled with colors* specific for each injection amount. These *colors* are referenced in **b**, which describes the peak area for each injection amount as a *bar graph*. **c** Shows the absence of L-glutamate carry over between injections, by showing the corre-

sponding peak areas for the sample injection, a 1:10 dilution of the sample (in methanol:acetonitrile:water 5:3:2 v/v), and blank runs carried out throughout the analysis (pre-run, inter-run, post-run). The representative *bar graphs* for L-glutamate peak area in the *inset* are ordered chronologically from *left to right*. **d** depicts the calibration curve for L-glutamate. **e** depicts the calibration curve for the 4 lowest injection amounts to show linearity down to Limit of Quantitation using an error of 2.5 ppm in positive ion mode

small fraction of the sample required for analysis demonstrates the sensitivity of this method, which is a requirement to perform amino acid analyses on samples with limited quantities. Through extrapolation, this method could provide comprehensive amino acid analysis from nanoliters of biofluids, micrograms of tissue, and 10^4 cells from culture. Sample preparation and handling methods are currently being optimized for sample sources with such low abundance.

Quantitation and matrix effects

To further validate the 3AA method, we compared results for the undiluted or fivefold diluted pancreatic cancer cell extract samples (biological triplicates) from the 3AA method as well as two UHPLC/MS methods using either normal or Amide HILIC (Yuan et al. 2012) with 15 or 23 min separations, respectively (Supplementary Fig. 3). Relative quantitation results from all platforms were comparable, as depicted using heat map plots in Supplementary

Fig. 3. The C18 and Amide HILIC method showed the most consistent results in terms of relative quantitation (red = fivefold \pm 0.5 increase—Supplementary Fig. 3). Importantly, the length of the analysis using the 3AA analytical method is five- to eightfold shorter than the normal and amide HILIC runs. Therefore, this improvement in workflow efficiency opens up new opportunities in clinical biochemistry (D'Alessandro et al. 2012b) and drug discovery (D'Alessandro and Zolla 2013) whereby rapid readouts will enhance patient care, drug development, testing, and screening.

Because of the rapid separation times of this method and the complexity of the assayed samples, the effects of matrix composition on ionization were tested to confirm linearity of the quantitative approach. These experiments were carried out with the addition of a standard mix at different concentrations (0, 0.5, 2.5 and 5 pmol injected) into the complex matrix of red blood cell extracts (Supplementary Table 4). Red blood cells interact closely with the plasma metabolome and the metabolic complexity of these

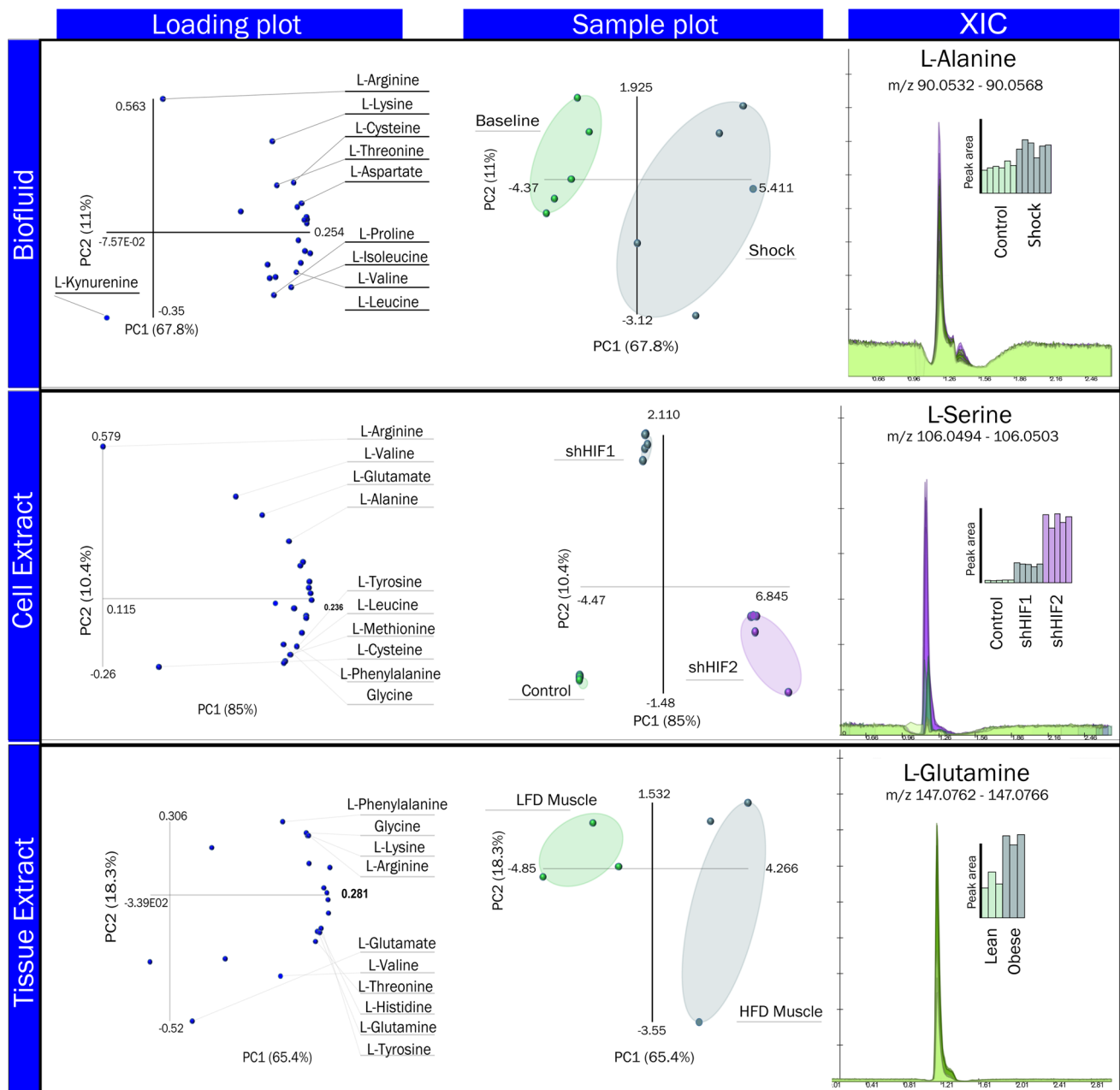


Fig. 4 Three-minute amino acid analysis is predictive of biological phenotypes by PCA in three different biological matrices. From left to right, panels indicate the principal amino acid contributors (Loading Plot, left), the sample clustering pattern resulting from an unsupervised Principal Component Analysis (Sample Plot, center) and the extracted ion chromatogram (XIC) for a representative amino acid in each group (XIC, right). Bar graph colors in the XIC right panel inset correspond to sample cluster colors in the center Sample Plot panel and indicate peak area quantities for the biological replicates in different conditions. Experimental mass to charge ratio (m/z) windows for detection are provided in the XIC right panel inset for

each amino acid to illustrate high mass accuracy. From top to bottom, examples are provided for representative amino acid analyses of biofluid extracts (rat plasma extracts under control–baseline conditions, or in response to trauma/hemorrhagic shock–shock), cell extracts (Pan02 cancer cell lines, five replicates, control versus two different HIF1 α knock down clones), and tissue extracts [skeletal muscle biopsies from three rats subjected to low-fat diet (LFD) or high-fat diet (HFD)], respectively. XICs were obtained from the MAVEN software. PCA plots were prepared using the Excel macro Multibase. The total figure was assembled using Adobe Photoshop CS6

cells has been recently appreciated by our (D'Alessandro et al. 2014c) and other groups (Zhang et al. 2011). Median Pearson's correlation coefficients for quantitative analysis

of all tested amino acids averaged above 0.95 (0.98 and 0.99 in two out of three samples—Supplementary Table 4), showing linearity of quantitation in real samples. However,

signal suppression (2.59 ± 0.66 fold—Supplementary Table 4) was observed in comparison to the same standard mixture injected in the absence of background matrix, suggesting that absolute quantification of amino acids in a real matrix should be performed by establishing calibration curves in the matrix in question.

In the light of the observed matrix effects, a stable isotope dilution approach was employed by adding heavy-labeled internal standards ($^{13}\text{C}_6$ -arginine, $^{13}\text{C}_6$ -lysine, $^{13}\text{C}_5^{15}\text{N}_2$ -glutamine) and xenometabolites (5-fluorouracil) to the lysis/extraction buffer. This enriched buffer was thus used to process matched plasma, BALF and mesenteric lymph samples from trauma rats either undergoing sham or hemorrhagic shock (T/SS or T/HS), in presence and absence of mesenteric lymph diversion (MLD) to prevent distal organ injury after trauma/hemorrhagic shock (Stringham et al. 2014). Peak areas of heavy-labeled standards and xenometabolites were used to determine reproducibility of the analyses in complex matrices (CVs <10 %, Supplementary Fig. 4, Supplementary Table 5). Ratios of the 1:1 mixtures of biological samples extracted in lysis buffer either containing 10 or 20 μM concentrations of heavy-labeled standards were used to confirm linearity of relative quantitation (Supplementary Table 5). Calculated fold changes were highly precise (average CVs <10 %) and mostly accurate (median of all fold changes = 2.07). However, two exceptions were observed in that heavy-labeled arginine in BALF was underestimated and heavy-labeled glutamine in lymph was overestimated (Supplementary Table 5). This finding is consistent with the anticipated matrix dependency of matrix effects. Therefore, even though in 7 out of 9 cases no preliminary optimization was necessary, future applications will require identification of the linear range of quantification in the matrix being studied.

Finally, heavy to light isotopologue ratios for each standard amino acid were used to determine absolute concentrations in each tested sample ($n = 2$, plus two 1:1 mixtures with 10 and 20 μM spiked in samples—Supplementary Table 5). Here we show that, despite the influence of biological variability on experimental measurements, calculated CVs were <15 % for arginine and lysine and <10 % for glutamine in each one of the assayed matrices. While no previous data were available for amino acid concentrations in lymph and BALF to the best of our knowledge, the concentrations of arginine, lysine and glutamine in rat plasma were consistent with the literature (Morrison et al. 1961; Yang et al. 2002; Hiscock and Pedersen 2002).

Conclusion

The rapidly expanding field of metabolomics has shown increasing potential for translational applications in the clinical setting. The feasibility of using metabolomics as a readout

in these applications depends on the capacity to analyze a biologically relevant sub-portion of the metabolome in a robust, quantitative, fast, and reproducible fashion. In particular, rapidity is a key factor to analyze clinical samples from large cohorts of patients, which helps to cope with biological variability-associated issues by increasing the number of observations. In light of the increasingly consolidated focus of amino acids in several research areas including cancer metabolism, neurobiology, fertility research, nutrition, and food chemistry, amino acid analysis represents a key method in the clinical and research setting. Here, we describe a three-minute method for amino acid analysis (3AA) via UHPLC/MS. This method is fast, reproducible, sensitive, linear, and has been validated through the use of underivatized commercially available standard mixtures, injected either alone or in combination with background matrix to test linearity of quantitation in the presence of matrix-dependent ion suppression effects. In cases where biological matrices give rise to isobaric species with similar retention times, product ion transitions can be monitored for further selectivity. We conclude that the method is amenable for high-throughput fast amino acid analysis in routine samples for biological and clinical research (e.g., biofluids, cell and tissue extracts). While in most of these applications relative quantitation might be sufficient for discovery mode applications, here we show two approaches for absolute quantitation through the use of exogenously added light- or heavy-labeled internal standards. This method surpasses the rapidity of chemical derivatization-based approaches while maintaining sensitivity and selectivity as demonstrated by unambiguous assignment of amino acids with sub 5 ppm accuracy, retention time and isotopic pattern distributions to determine elemental composition. Rapidity and accuracy of relative quantitation were superior to normal and amide HILIC runs. Taken together, our method represents a very robust and sensitive tool for amino acid analyses.

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Conflict of interest All the authors disclose no conflict of interests.

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