Position-specific $^{13}\text{C}/^{12}\text{C}$ analysis of amino acid carboxyl groups – automated flow-injection analysis based on reaction with ninhydrin

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Rationale: The fundamental level of stable isotopic knowledge lies at specific atomic positions within molecules but existing methods of analysis require lengthy off-line preparation to reveal this information. An automated position-specific isotope analysis (PSIA) method is presented to determine the stable carbon isotopic compositions of the carboxyl groups of amino acids ($\delta^{13}\text{C}_{\text{CARBOXYL}}$ values). This automation makes PSIA measurements easier and routine.

Methods: An existing high-performance liquid chromatography (HPLC) gas handling interface/stable isotope ratio mass spectrometry system was modified by the addition of a post-column derivatisation unit between the HPLC system and the interface. The post-column reaction was optimised to yield CO$_2$ from the carboxyl groups of amino acids by reaction with ninhydrin.

Results: The methodology described produced $\delta^{13}\text{C}_{\text{CARBOXYL}}$ values with typical standard deviations below $\pm0.1$‰ and consistent differences ($\Delta^{13}\text{C}_{\text{CARBOXYL}}$ values) between amino acids over a 1-year period. First estimates are presented for the $\delta^{13}\text{C}_{\text{CARBOXYL}}$ values of a number of internationally available amino acid reference materials.

Conclusions: The PSIA methodology described provides a further dimension to the stable isotopic characterisation of amino acids at a more detailed level than the bulk or averaged whole-molecule level. When combined with on-line chromatographic separation or off-line fraction collection of protein hydrolysates the technique will offer an automated and routine way to study position-specific carboxyl carbon isotope information for amino acids, enabling more refined isotopic studies of carbon uptake and metabolism.

1 | INTRODUCTION

There is a wide ranging interest in the isotopic analysis of amino acids, from medicine to meteorites. For example, there is a long-term interest in the isotopic compositions of amino acids as monitors of metabolism, including indicators of serious medical conditions such as diabetes and disease such as breast cancer. New methods dealing with amino acids are thus timely and of widespread interest.

A number of previous studies have described the stable carbon isotopic analysis of individual amino acids, typically derived by hydrolysis of proteins, from various biological sources. This compound-specific isotope analysis (CSIA) is achieved either by gas chromatography of...
derivatised amino acids coupled with isotope ratio mass spectrometry (IRMS) or by high-pressure liquid chromatography (HPLC) of derivatised amino acids coupled to IRMS using chemical oxidation to convert the carbon present in organic compounds into CO₂ (LC/CO/IRMS). The use of LC/CO/IRMS for CSIA is now a relatively mature technique, especially with regard to the separation of derivatised amino acids using mixed-mode HPLC columns and aqueous mobile phases. Recent research has shown that the CSIA δ₁³C values of derivatised amino acids, derived from hydrolysis of hair samples, can indicate metabolic trends associated with factors such as age and obesity in humans.

While the CSIA approach has yielded important information, it measures the average isotopic information available from amino acids which contain between two (glycine) and 11 (tryptophan) carbon atoms. These atoms are not all equal in ¹³C/¹²C ratios and it has long been recognised that the distribution of stable isotopes within amino acids is neither random nor homogeneous. It is also generally recognised that the fundamental level of stable isotopic information lies at specific positions within molecules. Therefore, it seems that the time is right to take the technology to 'the-next-level' and develop a more detailed position-specific isotope analysis (PSIA) methodology for amino acids.

A number of IRMS techniques have previously been described to measure the stable isotopic composition at specific positions within small molecules. These methods typically require tedious chemical or enzymatic degradations that fragment the molecule and then separate and analyse the fragments for their δ¹³C composition. Methods for PSIA of small molecules based on Stable Isotope Fractionation–Nuclear Magnetic Resonance (SNIF-NMR or rH NMR) have also been developed although these techniques are typically applied to compounds that can be prepared in large amounts and high purity, for example ethanol distilled from European wines. These techniques now provide the basis for the authentication of a number of natural products, and reference materials (RMs) exist for the PSIA of ethanol with certified δH/δD ratios of both methyl and methylene groups (e.g. European Commission, Joint Research Centre BCR-656 and BCR-660). More recently, ¹³C NMR has been developed for PSIA studies of (bio)chemical and physiochemical pathways but, unlike SNIF-NMR, this has the disadvantage of not being linked to international RMs. A detailed discussion of these techniques is beyond the scope of this study but can be found in the citations above. The aims of this study were:

- to interface existing pieces of commercially available equipment to produce a practical flow-injection analysis (FIA)–ninhydrin reaction–IRMS (FIA/NR/IRMS) instrument,
- to develop robust chemistry for the consistent yield of CO₂ from amino acid carboxyl groups,
- to provide a first estimate of the carboxyl carbon isotopic compositions (δ¹³C_carboxyl) values of internationally available amino acid RMs.

The PSIA of the carboxyl group of amino acids was an easy choice because the reaction of ninhydrin (2,2-dihydroxyindane-1,3-dione) reaction required to produce CO₂ from the carboxyl groups of amino acids is already well established. The development of Ruhemann’s purple (RP) diketohydrindamine-diketohydrindylidene enolate) from the reaction of ninhydrin with amino acids was first described over a century ago and the method has been widely used to visualise amino acids, peptides and proteins. Commercial systems are now available for the post-column derivatisation (PCD) of amino acids separated by HPLC. These systems aim for consistent colour development although the reaction may only proceed to 30% completion in 30 s. A red-coloured compound, hydrindantin (a dimer of ninhydrin), also makes a minor contribution to the colour formed during the ninhydrin reaction.

Although some controversy still exists as to the exact mechanism of the reaction of amino acids with ninhydrin it is generally accepted that CO₂ is liberated at a very early stage of the reaction and before any colour development, as shown in Figure 1. Although the formation of CO₂ might be regarded as a side-effect of the ninhydrin reaction, CO₂ from this reaction has been used to quantify amino acids and for both radiocarbon dating and stable isotope analysis of peptide-bonded carbon in proteinoaceous materials. Early studies showed that the reaction of amino acids with ninhydrin at boiling temperatures can liberate carboxyl CO₂ at 100.0 ± 0.2% yields, with higher pH (4.7) conditions being optimum for most of the 20 common amino acids and lower pH (1) being optimum for cystine, glutamate, and lysine. This quantitative aspect has encouraged the use of the ninhydrin reaction for isotopic studies, with no isotope fractionation expected when both ¹³C and ¹²C are completely converted into CO₂.

Generally, the carboxyl group of amino acids show isotopic enrichment patterns specific to their metabolism and provide a means to test labelling expectations such as the metabolic uptake of bicarbonate. For example, carbon isotopic analysis of 15 samples of glutamic acid showed little variation in average composition whereas large and significant differences existed in the δ¹³C_carboxyl values. Other studies showed that δ¹³C_carboxyl patterns were characteristic of different growth conditions and enzyme pathways involved in carbon by autotrophs and heterotrophs. More recently, a study of biopsy tissue concluded that bulk cellular ¹³C composition provided

![Figure 1](image1.png)
a marker for breast cancer, with a characteristic $^{13}\text{C}$ enrichment linked to the urea cycle and arginine build-up.\textsuperscript{3} PSIA of the arginine carboxyl group might well provide a more sensitive cancer indicator than isotope measurements of the whole tissue.

It also seems likely that individual carboxyl groups will be classified as those that do not change through the food web (carboxyl groups of ‘essential’ or ‘indispensable’ amino acids) while carboxyl groups of other amino acids (‘non-essential’ or ‘dispensable’ amino acids) may change much more than average and be more dramatic metabolic indicators. Thus, amino acid PSIA might record the diet and the metabolic response to the diet in highly differentiated ways.

Although the LC/IRMS interface was intended to be used in conjunction with chromatographic separations it is frequently used for FIA, without HPLC separation, for the analysis of pre-conjunction with chromatographic separations it is frequently used for FIA, without HPLC separation, for the analysis of pre-

TABLE 1 Amino acid reference materials used in this study and their compound-specific isotope analysis (CSIA) delta values

| Reference material | Nature | CSIA $\delta^{13}\text{C}_{\text{VPDB}}$ (%) mean | SD |
|--------------------|--------|---------------------|-----|
| USGS40             | L-glutamic acid | -26.39 | ± 0.04 |
| USGS41a            | L-glutamic acid | +36.55 | ± 0.08 |
| USGS64 (G1)        | glycine      | -40.81 | ± 0.04 |
| USGS65 (G2)        | glycine      | -20.29 | ± 0.04 |
| USGS66 (G3)        | glycine      | -0.67  | ± 0.04 |
| D01*               | L-valine     | -10.97 |          |
| USGS73 (V1)        | L-valine     | -20.03 | ± 0.04 |
| USGS74 (V2)        | L-valine     | -9.03  | ± 0.04 |
| USGS75 (V3)        | L-valine     | +0.49  | ± 0.07 |
| TTA*               | L-alanine    | -23.30 | ± 0.12 |
| TTS*               | L-alanine    | -18.65 | ± 0.11 |
| TTW*               | L-alanine    | -19.64 | ± 0.18 |

*see section 2.1.

2 | EXPERIMENTAL

2.1 | Reference materials

Samples of L-glutamic acid, glycine and L-valine RMs were purchased from the US Geological Survey Stable Isotope Laboratory (Reston, VA, USA). These materials (Table 1) are internationally available and have well-characterised CSIA $\delta^{13}\text{C}_{\text{VPDB}}$ values.\textsuperscript{36,37}

A sample of natural abundance L-valine (D01) that was used to prepare L-valine RMs USGS74 and 75 was kindly supplied by colleagues from Indiana University (Bloomington, IN, USA).\textsuperscript{37}

Three samples of L-alanine (TTA, TTS, TTW) were, respectively, purchased from Aldrich Chemical Co., Inc. (St Louis, MO, USA; Lot No. CA03519DO, certified purity >99%), from Sigma Chemical Co. (St Louis, MO, USA; Lot No. 078 K1099, certified purity >98.5%) and from Wako Pure Chemical Industries (Osaka, Japan; Lot No. EWL2621, certified purity >99%). In Tokyo, CSIA $\delta^{13}\text{C}_{\text{VPDB}}$ values were characterised using an off-line method. The method involved sealed tube combustion followed by cryogenic separation of the resulting CO$_2$ using a high vacuum line system, then measurement of the purified CO$_2$ by dual-inlet IRMS. The $\delta^{13}\text{C}_{\text{CARBOXYL}}$ values were characterised using an off-line method consisting of a 1-h boiling reaction with 3.5% ninhydrin in phosphoric acid (pH 2) followed by cryogenic separation of the resulting CO$_2$ and measurement of the purified CO$_2$ by dual-inlet-IRMS. Both CSIA and PSIA dual-inlet measurements were made using a Delta XL isotope ratio mass spectrometer (Thermo Fisher Scientific, Bremen, Germany). Data were normalised to the VPDB-LSVEC scale using CO$_2$ liberated from NBS 19 ($\delta^{13}\text{C} = +1.95$‰) and LSVEC ($\delta^{13}\text{C} = -46.6$‰) (International Atomic Agency, Vienna, Austria) by reaction with phosphoric acid.

Although the four amino acids used in this study (Table 1) are a small subset of the 20 common amino acids, they represented a cross-section of ninhydrin reaction chemistry. For example, glutamic acid is reported to readily over-yield CO$_2$ at higher pH (4.7) whereas alanine and glycine under-yield at low pH (1) and L-valine gives consistent yields across a range of reaction chemistries.\textsuperscript{27,28}

For the automated work in Brisbane, 5 mM solutions of individual amino acids were prepared by dissolving weighed aliquots in 1 mM sulphuric acid (approximately pH 3). Solutions were then degassed by ultrasonication under vacuum to remove any dissolved atmospheric CO$_2$. This ‘sonic vac’ (sonication under vacuum) process proved especially important for isotopically labelled materials for which the dissolved CO$_2$ was very different in carbon isotope composition from the $^{13}$C-labelled carboxyl group.

A number of in-house quality control amino acids (glycine, phenylalanine and aspartic acid) were analysed together with each batch of samples. These amino acids were purchased from Sigma (Castle Hill, NSW, Australia) with certified purities >98%.

The stretch factor for the Brisbane IRMS instrument was assessed using inorganic carbon standards; LSVEC lithium carbonate (~46.6 ‰) and a sodium bicarbonate (LSUB, ~3.28 ± 0.06 ‰) which was calibrated against NBS 19 at the Louisiana State University (Baton Rouge, LA, USA) by the first author. Both LSVEC and LSUB were prepared as 5 mM solutions by dissolving the dry salts in high-purity water.

2.2 | Instrumentation

The FIA/NR/IRMS instrument is shown schematically in Figure 2. Stable carbon isotope ratio measurements were made using a Delta V Advantage isotope ratio mass spectrometer (Thermo Fisher Scientific, Bremen, Germany).

Aliquots of the amino acid solutions were injected into a flow of high-purity water (18.2 MΩ, 0.15 mL min$^{-1}$) using an Accela 600 high-pressure liquid chromatography pump (Figure 2, pump #1) and autosampler (Thermo Fisher Scientific, Waltham, MA, USA). The minimum sample sizes were equivalent to approximately 180 ng of carboxyl carbon per injection.

The HPLC eluent passed through a small-volume (37 µL) non-metallic check valve (Upchurch Scientific/Idex, Rohnert Park, CA, USA) and was combined with a 0.15 mL min$^{-1}$ flow from a modular
PCD system (Rigas Labs, Sindos, Greece) (Figure 2, pump #2). The detailed preparation of the PCD reagent is described below. The PCD unit was modified by replacing the 0.5-ml PEEK reaction coil with a stainless steel coil with a nominal internal volume of 1.5 mL (Figure 2, reactor #1), to allow longer reaction times at elevated temperatures. The total residence time in the PCD coil was approximately 5 min. The mixing tee in the PCD unit was replaced with a smaller mixing tee with 2-μL internal volume (VICI AG International, Schenkon, Switzerland).

The outlet from the PCD unit was connected to an LC IsoLink interface (Thermo Fisher Scientific, Bremen, Germany) which added 25 μL·min⁻¹ of 25% phosphoric acid (Figure 2, pump #3) via another low-volume mixing tee. The overall flow through the CO₂ separation membrane in the LC IsoLink was maintained below the 0.4 mL·min⁻¹ upper limit considered optimum for quantitative transfer of CO₂.³⁸ Acid was added primarily to decrease the solubility of CO₂ but had the added effect of solubilising the RP that formed along with CO₂; RP tended to block and sometimes break the CO₂ separation membrane. Frit filters (5 μm pore size, VICI #1110-5P-5) were used on all liquid input lines (Figure 2); these filter required only infrequent replacement, every 3–6 months. The small-volume reaction coil (0.2-mL oxidation reactor) of the LC IsoLink did not provide sufficient residence time for the ninhydrin reaction and was not used in this application. Gas from the LC IsoLink was transferred to the mass spectrometer via two fused-silica capillaries for sample (1.5 m × 100 μm i.d.) and working gas (1.5 m × 50 μm i.d.).

3 | RESULTS AND DISCUSSION

3.1 | Notes on method development

The first problem to be addressed by this research was how to link the four analytical instruments; HPLC system, PCD unit, LC IsoLink and mass spectrometer. The three liquid-handling units were connected with 0.005″ i.d. PEEK tubing, with the exception of a 30 mm length of larger-bore 0.03″ i.d. PEEK tubing used to connect the PCD reaction coil outlet to the mixing tee where acid was added (shown in bold in Figure 2). This larger-bore tubing was used to prevent clogging from the RP that was most likely to form at this point, as the reaction solution cooled and before downstream acid addition that began to dissolve the RP. Initial experiments resulted in many clogging problems due to RP and a length of tubing was added downstream of the acid addition to allow 1–2 min for RP to dissolve (Figure 2, reactor #2). Later experiments dispensed with this extra reactor when it was found that clogging was minimised if relatively small samples (<8 V peak heights) were analysed, a practice that became routine and also resulted in narrower peaks. Using relatively small samples and adding 25% phosphoric acid immediately following the PCD reaction coil allowed routine analysis over several days, with cleaning typically once a week. Cleaning mostly involved removing hydindantin that precipitated as a side-product of the RP reaction. The cleaning regime involved cooling the PCD reaction coil to room temperature then sequential flushing with 1% potassium hydroxide, 100% water, 100% acetone, and 10% nitric acid. Finally, three separate flushes of high-purity water were used to remove the cleaning agents (especially acetone) that could enter the mass spectrometer via the LC IsoLink CO₂ separation membrane and disturb the background signals of m/z 44, 45 and 46 for many hours.

The second challenge was to optimise the PCD reaction chemistry to produce consistent yields of CO₂ and reproducible δ¹³C_CARBOXYL values. The PCD reaction presented a large number of parameters to be optimised including the concentrations of ninhydrin, organic solvent and acid in the PCD solution, the overall pH and buffer strength of the reagent plus the reaction temperature and time in the coil. Faster (more complete) reactions have been reported at higher pH, higher temperature, longer reaction times and greater concentrations of ninhydrin.²⁶–²⁸

Although commercial reagents are formulated for consistent colour development and fast analysis, not necessarily complete reaction, we began with conditions common to many commercial systems: pH 5.8 and 130°C reaction. An increased concentration of organic solvent (sulfolane) facilitated higher concentrations of ninhydrin in the reagent that had a concomitant effect of faster reaction times. The use of organic solvents such as sulfolane is avoided in LC/CO/IRMS in which any organic carbon present is oxidised to CO₂, but organic solvents such as sulfolane could be tolerated in the PSIA method that specifically targeted carboxyl groups without oxidation. Some organic solvents were found to decompose at high temperatures and sulfolane was
chosen for its high boiling point and good thermal stability, i.e. low CO₂ background in the mass spectrometer. The reagent was buffered using phosphate to stabilise the chemistry, with the amounts of sulfolane and phosphate adjusted such that all components would dissolve overnight to a clear, yellow solution. The final PCD solution was prepared as follows (per litre): sulfolane (200 mL), deionised water (800 mL), ninhydrin (14 g), sodium dihydrogen phosphate dihydrate (28.905 g) and disodium phosphate anhydrous (2.659 g). The solution was sparged overnight with helium to remove residual CO₂ from this slightly acidic (pH 5.8) phosphate-buffered solution.

The normal working temperature for the PCD reactor (130°C) was found to be adequate for routine work with the FIA/NR/IRMS system. To illustrate, Figure 3 shows the relative CO₂ yields and Δ₁³C₉CARBOXYL (Δ = δ₁ – δ₂) values at reaction temperatures of 90, 100 and 130°C, with Δ₁³C₉CARBOXYL representing the difference between the measured δ₁³C₉CARBOXYL values and values obtained at 130°C. At lower temperatures the CO₂ yields dropped dramatically while the Δ₁³C₉CARBOXYL values changed only slightly in the –3 to +1 ‰ range compared with results at 130°C. As the reaction temperature approached 130°C both yields and Δ₁³C₉CARBOXYL values reached asymptotic values that were consistent above this temperature (e.g. Table 2). These patterns were also typical of other amino acids. These relatively minor deviations in the Δ₁³C₉CARBOXYL values (Figure 3) indicate that yields were not too critical in the PSIA reactions, and were consistent with reaction 1 (Figure 1) mostly limiting reaction kinetics, in agreement with a previous study.²² Larger isotope effects and larger Δ₁³C₉CARBOXYL value offsets than those documented for low yields in Figure 3 would be expected if decarboxylation (Figure 1, reaction 2) was rate-limiting.

The PCD reaction coil typically required 2 h to stabilise after which the Δ₁³C₉CARBOXYL values became very precise and repeatable; the average standard deviation (SD) for multiple isotope measurements was typically <0.1 ‰ (n = 3) so the system was precise. There were, however, day-to-day offsets or variations in accuracy, for reasons that were usually unclear but might include incomplete degassing of CO₂ from reaction solutions or incomplete transfer of CO₂ across a contaminated diffusion membrane to the mass spectrometer. Regardless of the detailed explanation, it became apparent that measuring isotopic differences between samples proved to be very reproducible across reaction conditions, even when the absolute daily values varied somewhat. Table 2 illustrates some of the reaction conditions explored over a 1-year period for three glycine standards (USGS 64–66 also known as G1-G3). To correct for varying reaction conditions, a number of in-house quality control materials including ‘Woods Hole glycine’ (WHG) were analysed together with each batch of samples. Using this in-house standard to normalise data, results were reproducible across reaction conditions and at different reaction temperatures.

![Figure 3](image-url)

**Figure 3** The effect of increasing reaction temperature on (A) yield of CO₂ and (B) CO₂ carbon isotopic composition for a number of samples of alanine [Color figure can be viewed at wileyonlinelibrary.com]

| Date      | Mar. 2016   | Feb. 2017   | Mar. 2017   | Mar. 2017   | Mean | Predicted³ |
|-----------|-------------|-------------|-------------|-------------|------|------------|
| WHG       | –27.5 ± 0.16| –28.1 ± 0.2 | –27.8 ± 0.2 | –26.1 ± 0.2 |      |            |
| USGS64 (G1)| –38.0 ± 0.2 | –38.8 ± 0.2 | –38.3 ± 0.3 | –36.3 ± 0.2 |      |            |
| USGS65 (G2)| –24.9 ± 0.1 | –25.3 ± 0.1 | –24.9 ± 0.1 | –23.0 ± 0.2 |      |            |
| USGS66 (G3)| +14.3 ± 0.2 | +13.6 ± 0.2 | +13.5 ± 0.1 | +16.0 ± 0.4 |      |            |
| G1⁴       | –37.5       | –37.7       | –37.6       | –37.3       | –37.5 ± 0.2 |            |
| G2⁵       | –24.2       | –24.3       | –24.5       | –24.0       | –24.2 ± 0.2 |            |
| G3⁶       | +14.2       | +14.7       | +14.7       | +15.1       | +14.6 ± 0.4 |            |
| Δ G2-G1   | 13.4        | 13.5        | 13.0        | 13.3        | 13.3 ± 0.2 | 13.4      |
| Δ G3-G2   | 38.4        | 39.0        | 39.3        | 39.1        | 38.9 ± 0.4 | 39.2      |
| Δ G3-G1   | 51.7        | 52.4        | 52.3        | 52.4        | 52.2 ± 0.3 | 52.6      |

³δ scale has been shifted based on measurements of in-house QC glycine WHG.

⁴From calculations based on the reported formulations of these materials presented in the supporting information, assuming 96% purity of enriched materials used in preparations.
very comparable across more than 1 year of work and under several
time and temperature variations in reaction conditions.

Previous studies have shown that ninhydrin reactions at higher
pH (4.7) can result in low yields of CO₂ for the amino acid cystine
and 5 to 50% over-yields for glutamic acid and lysine. This study
confirmed these results and showed that USGS41a glutamic acid
was especially sensitive to over-yield because the α carboxyl group
was highly 13C-enriched (approximately +280‰) versus natural
carbon in the reaction mix. For this amino acid, over-yield of CO₂
resulted in δ13C(carboxyl) values of approximately +150‰ compared
with an expected value of approximately +280‰. Earlier studies showed that cystine, glutamic acid and lysine give quantitative yields
under controlled low pH conditions, and we found that adding 50 mM
sulphuric acid to the sample and to the HPLC solvent gave 100%
yields, the expected δ13C(carboxyl) values for USGS41a, and consistent
δ13C(carboxyl) values for lysine and cystine.

3.2 Notes on system performance

The water background (m/z 18) of the FIA/NR/IRMS system was
typically 30 to 45 V in the middle Faraday collector. Although greater
than the 12 V specified in the LC IsoLink operating manual this
background appeared to be comparable with the background reported by other LC/IRMS users. The background of CO₂ (m/z 44)
was between 0.4 and 2.2 V and was typically less than the 2.0 V
background reported for LC/CSIA applications. The reduced m/z 44
background allowed observations of δ13C/13C ratio isotope swings that
were typically 3 to 6‰ across peaks.

Injections of 3 to 24 μL of 5 mM amino acid solutions yielded CO₂
with typical peak heights of 3 to 6 V with amplitude/width ratios
between 75 and 150. Blank injections of high-purity water produced
peaks with <1% of the height of sample injections.

The stretch factor of the mass spectrometer was assessed by
injecting solutions of inorganic carbon standards that had known
δ13C values; LSVEC lithium carbonate (−46.6‰) and LSUB sodium
bicarbonate (−3.28 ± 0.06‰). Typically, the measured δ13C values
were in close agreement with the expected values and no stretch
correction was applied.

Peak size correction (linearity) was assessed by injecting varying
amounts of amino acid solutions and was found to be approximately
0.5% across the range of peak sizes and for a number of amino acids.
A linearity correction factor was applied as previously described.

The average yields for the RMs were somewhat variable and
ranged from 96 to 108% (Table 3) although the range always
included 100%. Common amino acids, other than those listed in
Table 3, had similar, near-100% yields – aspartic acid, proline,

| Amino acid  | Yield (%) relative to LSVEC carbonate solution (± 1 sd) |
|-------------|------------------------------------------------------|
| L-alanine   | 102 ± 9                                              |
| L-glutamic acid | 106 ± 20                                          |
| glycine    | 103 ± 3                                              |
| L-valine   | 96 ± 6                                               |

The CSIA uncertainty was included to represent possible physical
inhomogeneity in the RMs and resulting variability in isotope ratios.

The δ13C(carboxyl) value assigned to USGS41a will probably be subject
to the uncertainty in the certified CSIA values of the materials,
the standard deviation derived from replicate off-line PSIA
measurements of L-alanine,
the standard deviation derived from replicate FIA PSIA
measurements.
It was possible to estimate $\delta^{13}_{\text{CARBOXYL}}$ values for the $^{13}$C-enriched RMs (see supporting information) These estimates allowed the comparison shown in Figure 4 These results show an overall close agreement between the calculated and measured $\delta^{13}_{\text{CARBOXYL}}$ values for a number of chemically different compounds (the solid line shows a 1:1 relationship).

Although it is widely recommended that stable isotope measurements are traceable to international reporting scales via two-point calibration, PSIA is an emerging technique and the initial values reported here are based on a single-point calibration, the average of the L-alanine samples in Table 4. It may well become good practice for two-point calibrations to use the approximately 50 % differences that exist between RMs such as the two glycine samples (USGS64 and 66) or the two L-valine sample (USGS73 and 75). We have analysed all the RMs successfully, only varying the reaction conditions somewhat, in particular using a more acidic (50 mM sulphuric acid) carrier and sample diluent for the glutamic acid RMs and avoiding similar acid conditions for the alanine and glycine RMs in Table 4.

The methodology described provides a means to measure the carbon isotopic composition of the carboxyl group ($\delta^{13}_{\text{CARBOXYL}}$ values) of amino acids and has proved to be reproducible over many months of operation. The methodology was found to be very forgiving of most changes in reaction temperature, chemistry and CO$_2$ yield. Exceptions were found for glutamic acid and lysine that over-yielded CO$_2$ with biased and low $\delta^{13}_{\text{CARBOXYL}}$ values. More acidic conditions in the sample and carrier solvents were needed for good isotope results for glutamic acid and lysine. Cystine is also better analysed under acid conditions.

This study presents the first estimates for the $\delta^{13}_{\text{CARBOXYL}}$ values for the carboxyl groups of a number of internationally available amino acid RMs. The results from the automated method were found to be in close agreement with values determined both by off-line measurements and by calculations based on descriptions of how enriched and natural abundance compounds were mixed to produce these RMs. Readers are invited to review and improve on the values presented in Table 4.
Using the method described, the analysis of an individual amino acid can be performed in approximately 7.5 min, such that the isotopic profile of the 17 amino acids, typically present in hair, can be obtained in a little over 2 h, provided that these amino acids have been separated and purified prior to PSIA.

Typical metabolic studies of amino acids analyse mixtures of compounds liberated from biological tissues by hydrolysis with hydrochloric acid whereas the methodology presented here is applicable only to individual amino acids. A next step will be to modify the instrument design to a system compatible with on-line HPLC separation. Using this approach it may be possible to combine carbon and nitrogen CSIA (amino acid) and PSIA (carboxyl group) data to enhance the specificity and power of stable isotopes to elucidate sources and metabolic cycling of amino acids.

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