Carbon and nitrogen isotope fractionation of amino acids in an avian marine predator, the gentoo penguin (Pygoscelis papua)

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Abstract
Compound-specific stable isotope analysis (CSIA) of amino acids (AA) has rapidly become a powerful tool in studies of food web architecture, resource use, and biogeochemical cycling. However, applications to avian ecology have been limited because no controlled studies have examined the patterns in AA isotope fractionation in birds. We conducted a controlled CSIA feeding experiment on an avian species, the gentoo penguin (Pygoscelis papua), to examine patterns in individual AA carbon and nitrogen stable isotope fractionation between diet (D) and consumer (C) (\(\Delta^{13}C_{C-D}\) and \(\Delta^{15}N_{C-D}\), respectively). We found that essential AA \(\delta^{13}C\) values and source AA \(\delta^{15}N\) values in feathers showed minimal trophic fractionation between diet and consumer, providing independent but complimentary archival proxies for primary producers and nitrogen sources respectively, at the base of food webs supporting penguins. Variations in nonessential AA \(\Delta^{13}C_{C-D}\) values reflected differences in macromolecule sources used for biosynthesis (e.g., protein vs. lipids) and provided a metric to assess resource utilization. The avian-specific nitrogen trophic discrimination factor (TDF\(_{\text{Glu-Phe}}\) = 3.5 ± 0.4\(^{\circ}{\text{o}}\)) that we calculated from the difference in trophic fractionation (\(\Delta^{15}N_{C-D}\)) of glutamic acid and phenylalanine was significantly lower than the conventional literature value of 7.6\(^{\circ}{\text{o}}\). Trophic positions of five species of wild penguins calculated using a multi-TDF\(_{\text{Glu-Phe}}\) equation with the avian-specific TDF\(_{\text{Glu-Phe}}\) value from our experiment provided estimates that were more ecologically realistic than estimates using a single TDF\(_{\text{Glu-Phe}}\) of 7.6\(^{\circ}{\text{o}}\) from the previous literature. Our results provide a quantitative, mechanistic framework for the use of CSIA in nonlethal, archival feathers to study the movement and foraging ecology of avian consumers.

Introduction
Resource acquisition and allocation are fundamental requirements for all animals and significantly influence the behavior of individuals, the ecological and evolutionary trajectories of populations, and the functioning and resilience of entire ecosystems (Paine 1966; Polis and Strong 1996; Tinker et al. 2008). There is, for instance, both theoretical and empirical support for the roles of food chain length and primary producer composition in structuring food web architecture, mediating the relationship between species diversity and ecosystem function, and regulating fisheries productivity and biogeochemical fluxes (Post 2002a; Vander Zanden and Fetzer 2007; Young et al. 2013). The ability to quantify food web architecture is critical to understanding ecosystem structure and function, particularly in light of past and future changes in climate and anthropogenic disturbance.

Stable isotope analysis (SIA) has become a widely used tool for examining food web architecture across diverse
ecosystems (Layman et al. 2012). This approach is based on the principle that a consumer’s bulk carbon (δ13C) and nitrogen (δ15N) isotope composition reflects that of its diet coupled with some degree of trophic fractionation (Δ13C_{C:D} and Δ15N_{C:D}) between diet and consumer (Boecklen et al. 2011; McMahon et al. 2013a). While the bulk tissue SIA approach has provided many advances in the fields of trophic and movement ecology (Boecklen et al. 2011; McMahon et al. 2013b), one of the biggest challenges to interpreting bulk stable isotope data is determining whether changes in a consumer’s stable isotope values are due to changes in diet/trophic position, variability in trophic fractionation factors, shifts in isotope values at the base of the food web (δ13C_{baseline}, δ15N_{baseline}), or some combination of these factors (Post 2002b; McMahon et al. 2013a, b). Teasing apart the relative influences of trophic and baseline variability on stable isotope values is particularly challenging in complex ecosystems with dynamic baselines and when studying highly mobile predators that feed within multiple isotopically distinct food webs (McMahon et al. 2013b).

Recent advances in the compound-specific stable isotope analysis (CSIA) of individual amino acids (AAs) have allowed for more detailed studies of diet and trophic dynamics that avoid many of the challenges of bulk SIA outlined above (McMahon et al. 2013a; Chikaraishi et al. 2014). For example, the δ13C values of essential AAs, which animals must acquire directly from their diet (Reeds 2000), show little to no trophic fractionation between diet and consumer (Hare et al. 1991; Howland et al. 2003; McMahon et al. 2010). Further, the high degree of metabolic diversity in essential AA synthesis pathways among primary producers (Hayes 2001; Scott et al. 2006) leads to unique AA δ13C signatures that get recorded in upper trophic level consumers, virtually unmodified. As a result, essential AA isotope fingerprinting can be used to quantify carbon flow through food webs (Larsen et al. 2009, 2013). In contrast, nonessential AAs can either be de novo biosynthesized from a bulk carbon pool or directly routed from dietary protein, with typically highly variable Δ13C_{C:D} values across taxa and diet types (Hare et al. 1991; Howland et al. 2003; McMahon et al. 2010). Previous studies suggest that nonessential AA Δ13C_{C:D} values are likely related to diet composition and quality, although additional work is needed to understand the underlying mechanisms (McMahon et al. 2010; Newsome et al. 2011).

With respect to δ15N, individual AAs are commonly divided into trophic and source AAs (after Popp et al. 2007), based on their relative fractionation with trophic transfer (Δ15N_{C:D}). Trophic AAs, most commonly represented by glutamic acid (Glu), undergo significant isotopic fractionation during transamination/deamination, which often provide greater sensitivity for defining trophic position than bulk SIA (McClelland and Montoya 2002; Chikaraishi et al. 2007, 2009). Conversely, the canonical source AA, phenylalanine (Phe), shows no trophic fractionation between diet and consumer because its metabolic processing does not form or break C-N bonds (McClelland and Montoya 2002; Chikaraishi et al. 2009). Thus, Phe δ15N provides a proxy for the sources and cycling on nitrogen at the base of food webs (δ15N_{baseline}) (Sherwood et al. 2014; Vokhshoori and McCarthy 2014). Together, the δ15N value of Glu and Phe can be used to estimate consumer trophic position while accounting for differences in δ15N_{baseline} without needing to independently characterize and analyze the baseline structure of a food web (McClelland and Montoya 2002; Popp et al. 2007; Chikaraishi et al. 2009). The accuracy of this approach fundamentally depends on the accuracy of the trophic discrimination factor (TDF_{Glu-Phe} = Δ15N_{Glu} − Δ15N_{Phe}) used to calculate trophic position. Accumulating evidence, however, suggests that TDF_{Glu-Phe} values may in fact vary widely across taxa (Germain et al. 2013; Bradley et al. 2014; Hoen et al. 2014; McMahon et al. in press). For instance, Lorrain et al. (2009) hypothesized that AA fractionation may be different in avian consumers because they found that the literature TDF_{Glu-Phe} value of 7.6%/oo greatly underestimated the CSIA-based trophic position of wild penguins in their study. Yet, no controlled feeding experiments have examined patterns of AA fractionation in birds.

Our main objective in this study was to examine individual AA δ13C and δ15N fractionation patterns for an avian species. We conducted a controlled feeding experiment on captive-reared gentoo penguins (Pygoscelis papua Forster 1781) fed a high protein Atlantic herring (Clupea harengus Linnaeus 1758) diet. We analyzed feathers because they can be sampled nonlethally, are metabolically inert after synthesis (e.g., Mizutani et al. 1990), and are the most common tissue used in avian SIA studies (Boecklen et al. 2011). Using the newly derived avian-specific AA fractionation factors, we then examined wild penguin foraging ecology and trophic dynamics in the Southern Ocean, including identifying the sources of primary production supporting wild penguins using essential AA δ13C values and examining how our newly derived avian-specific TDF_{Glu-Phe} values affected estimates of wild penguin trophic position. The results of our study provide new empirical support for the use of CSIA to examine nutritional and foraging ecology, trophic dynamics, and movement ecology of avian consumers.
Materials and Methods

Feeding experiment and field collections

A controlled feeding experiment was conducted on 10 captive adult gentoo penguins (five male and five female) at Omaha’s Henry Doorly Zoo and Aquarium, Omaha, Nebraska. Penguins were reared on a consistent, high-protein diet of wild-caught Atlantic Herring for 10 months prior to the start of molt. During the study, we hand-fed penguins ad libitum allowing us to record the mass of herring each individual consumed per day during the 30 days prior to molt and during the molt period. Prior to the start of molt, we measured the mass of penguins to the nearest 10 g in order to calculate dietary intake relative to body mass. We calculated the length of the molt period as the number of days between when flippers swell in size and old feathers began to lift and stand out from the body to the end of molt when new body feathers were fully grown. We measured the weight (g) and standard length (mm) of five randomly sampled individual herring per month over the 3 months leading up to and during the molt period (January to March 2008). We collected three newly grown breast feathers from each adult gentoo penguin following their annual molt in late March 2008. During the austral summer of 2010/11, we collected breast feathers of five wild adult gentoo penguins (three males, two females) from a colony on King George Island, Antarctica (62°09′S, 58°24′W).

Sample preparation and analysis

Proximate analysis of crude protein, fat, and carbohydrate content were conducted on herring muscle (n = 3 replicates) at the New Jersey Feed Laboratory, Trenton, New Jersey (AOAC 2005). Amino acid composition (AOAC 2005) of herring and captive penguin feathers was also determined at the New Jersey Feed Laboratory (n = 3 replicates). All feathers were cleaned of surface contaminants using a 2:1 chloroform:methanol rinse. Whole herring samples were homogenized and dried for 48 h in an oven at 60°C. Lipids were extracted from dried herring using a Soxhlet apparatus with a 1:1 petroleum ether/ethyl ether solvent mixture for 8 h (Seminoff et al. 2007). Bulk stable isotope results reported here are from a subset of the samples reported in Polito et al. (2011a).

For CSIA, approximately 3 mg of fish and feather tissue was acid-hydrolyzed in 1 mL of 6 N HCl at 110°C for 20 h to isolate the total free AAs. Samples were evaporated to dryness under a gentle stream of N2. The total free AAs were derivatized by esterification with acidified isopropanol followed by acylation with trifluoroacetic anhydride (Silfer et al. 1991) and brought up in dichloromethane (DCM) for stable isotope analysis. For AA δ13C analyses, the derivatized AAs were injected on column in split mode at 250°C and separated on a DB-5 column (50 m × 0.5 mm inner diameter; 0.25 μm film thickness; Agilent Technologies, Santa Clara, CA) in a Thermo Trace Ultra gas chromatograph (GC) at the University of California, Santa Cruz, CA. The separated AA peaks were analyzed on a Finnegan MAT DeltaPlus XL isotope ratio monitoring mass spectrometer (irm-MS) interfaced to the GC through a GC-C III combustion furnace (960°C) and reduction furnace (630°C). For AA δ15N analyses, the derivatized AAs were injected on column in splitless mode at 250°C and separated on a BPX5 column (60 m × 0.32 mm inner diameter, 1.0 μm film thickness; SGE Analytical Science, Austin, Texas, USA) in the same CG-C-irm-MS interfaced through a combustion furnace (980°C), reduction furnace (650°C), and a liquid nitrogen trap.

We analyzed the δ13C and δ15N values of eleven individual AAs, accounting for approximately 76% and 65% of the total hydrolysable AAs in feathers and herring muscle, respectively. For carbon, we assigned glutamic acid (Glu), aspartic acid (Asp), alanine (Ala), proline (Pro), glycine (Gly), and serine (Ser) as nonessential AAs, and threonine (Thr), leucine (Leu), isoleucine (Ile), valine (Val), and phenylalanine (Phe) as essential AAs. For nitrogen, we assigned Glu, Asp, Ala, Leu, Ile, Pro, Val, Gly, and Ser as trophic AAs, and Phe as the only source AA. Note that threonine (Thr) δ15N values do not behave similarly to either of these main groups (Hare et al. 1991; McClelland and Montoya 2002; Styring et al. 2010; Germain et al. 2013). Therefore, we have listed Thr as a metabolic AA (according to Germain et al. 2013) and do not discuss Thr δ15N extensively in this paper. Acid hydrolysis converts glutamine (Gln) and aspartic acid (Asn) into Glu and Asp, respectively, due to cleavage of the terminal amine group, resulting in the measurement of combined Gln + Glu (referred to hereby as Glu), and Asn + Asp (referred to hereby as Asp). While some researchers refer to these groupings as Glx and Asx, we chose our terminology here to be consistent with other CSIA studies.

We analyzed 10 individual penguins and three replicate herring samples (one composite sample of five individuals at the beginning, middle, and end of the experiment) from the controlled feeding study, and five individual wild penguins from King George Island, Antarctica. All CSIA samples were analyzed in triplicate along with AA standards of known isotopic composition (Sigma-Aldrich Co., St. Louis, MO, USA). Standardization of runs was achieved using intermittent pulses of a CO2 or N2 reference gas of known isotopic value. Mean reproducibility of
a laboratory algal standard across all individual AAs was 
± 0.66‰ for δ13C and ± 0.34‰ for δ15N.

Data analysis

Stable isotope values are expressed in standard delta (δ) notation with respect to Vienna Pee Dee Belemnite and air references for δ13C and δ15N, respectively. We looked for differences in herring length, weight, and bulk tissue δ13C and δ15N values over the three months leading up to and during the molt period with separate one-way analysis of variance (ANOVA) and Tukey’s honestly significant difference (HSD) post hoc tests (z = 0.05). We compared mean molt periods and consumption of diet as a percent of body mass before and during molt between female and male penguins using separate unpaired two sample t-tests (n = 5 individuals per sex). Trophic fractionation factors (Δ13N(C-D) and Δ13C(C-D)) were calculated for feathers as Δ13X(D) = δ13X(D) - δ13X(δ), where δ13X(C) and δ13X(D) represent the δ13N or δ13C values of the consumer and diet, respectively. We used separate one-sample t-tests to determine whether bulk tissue and individual AA Δ13C(C-D) and Δ13N(C-D) values were significantly different from zero (z = 0.05) (n = 10 individuals). We then compared Δ13C(C-D) and Δ13N(C-D) values among individual AAs with separate one-way ANOVAs and Tukey’s HSD post hoc tests (z = 0.05) (n = 10 individuals). We tested the hypothesis that diet quality (i.e., AA composition) affected trophic fractionation using linear regressions between AA imbalance (difference in nonessential or trophic AA mol % in diet versus consumer) and trophic fractionation (Δ13C(C-D) or Δ13N(C-D)).

We used an isotopic fingerprinting approach (Larsen et al. 2013) to identify the primary producers at the base of the food chain supporting wild gentoo penguins. Briefly, we used published essential AA δ13C data from eukaryotic microalgae, cyanobacteria, and marine macroalgae (Larsen et al. 2013) as the most likely primary producers supporting penguin food chains in our study. For both the primary producer end members and the wild penguins, individual essential AA δ13C values were normalized to their means to allow for comparisons of essential AA δ13C patterns across groups. We identified the most likely primary producer carbon sources contributing to wild penguin essential AA δ13C patterns using the prediction function of a linear discriminant function analysis (LDA) of normalized essential AA δ13C values (Thr, Ile, Val, Phe, Leu). For calculating the probability of group membership of the classifier samples, we used a leave-one-out cross-validation approach (Larsen et al. 2009).

We examined how calculation method and TDFGlu-Phe value affected estimated TPCSIA for five species of wild penguins from the Southern Ocean: gentoo penguins (P. papua) from this study and northern rockhopper (Eudyptes chrysocome moseleyi Mathews & Iredale 1921), southern rockhopper (E. c. chrysocome Forster 1781), king (Aptenodytes patagonicus Miller 1778), and Adélie (P. adeliae Hombron & Jacquinot 1841) penguins from Lorrain et al. (2009). We calculated TPCSIA using the single TDFGlu-Phe approach of Chikaraishi et al. (2009):

\[ \text{TP}_{\text{CSIA}} - \text{single TDF} = 1 + \left( \frac{\delta^{15}N_{\text{Glu}} - \delta^{15}N_{\text{Phe}} - \beta}{\text{TDF}_{\text{Glu-Phe}}} \right), \]

where \( \delta^{15}N_{\text{Glu}} \) and \( \delta^{15}N_{\text{Phe}} \) represent the stable nitrogen isotope values of penguin Glu and Phe, respectively, and \( \beta \) represents the difference in δ15N between Glu and Phe of primary producers (3.4‰ for aquatic cyanobacteria and algae [McClelland and Montoya 2002; Chikaraishi et al. 2010]), and TDFGlu-Phe is the literature value of 7.6‰. We also calculated TPCSIA using a multi-TDFGlu-Phe approach that included our newly derived avian-specific TDFGlu-Phe value:

\[ \text{TP}_{\text{CSIA}} - \text{multi TDF} = 2 + \left[ \frac{\delta^{15}N_{\text{Glu}} - \delta^{15}N_{\text{Phe}} - \text{TDF}_{\text{Glu-Phe/plankton}} \cdot \beta}{\text{TDF}_{\text{Glu-Phe/penguin}}} \right], \]

where TDFGlu-Phe/plankton = 7.6‰, typical of plankton and other lower trophic level marine organisms (e.g., Chikaraishi et al. 2007), and TDFGlu-Phe/penguin represents the avian-specific TDFGlu-Phe value derived from this study. We used a two-way ANOVA (species and calculation method as the independent variables and trophic position as the dependent variable) with a Tukey’s HSD post hoc test (z = 0.05) to compare TPCSIA values of the five species of wild penguins calculated using the single and multi-TDFGlu-Phe equations. All statistics were performed in R version 3.0.2 using RStudio interface version 0.98.501 (R Core Team 2013).

Results

Feeding experiment characterization

Penguins in our controlled feeding experiment were fed a herring diet high in protein (72%) and fat (18%) and low in carbohydrates (0.2%) (Table 1). Herring length (187.4 ± 4.3 mm; one-way ANOVA, \( F_{2,12} = 0.9, P = 0.43 \)), weight (77.9 ± 6.7 g; one-way ANOVA, \( F_{2,12} = 1.3, P = 0.30 \)), bulk tissue carbon isotope values (−16.9 ± 0.1‰; one-way ANOVA, \( F_{2,12} = 1.1, P = 0.35 \)), and bulk tissue nitrogen isotope values (13.7 ± 0.4‰; one-way ANOVA, \( F_{2,12} = 1.7, P = 0.23 \)) did not change significantly during the course of the
feeding study (Table S1). We found notable differences in AA composition between herring muscle (diet) and penguin feather (consumer), particularly for the nonessential AAs proline, glycine, and serine (Table 1). Most essential AAs, with the exception of valine (>5% imbalance), had relatively similar molar abundances in the diet and consumer tissues (<2% imbalance) (Table 1).

The molt period for penguins in our study ranged from 12 to 18 days (mean 15.0 ± 1.8 days, Table S4) but did not differ between sexes (unpaired two sample t-test, $t_{8} = 0.70, P = 0.51$). Penguins consumed 5.2–8.8% (mean 6.9 ± 1.4%) of their body mass in herring per day during the 30 days prior to molt and 0.5–4.0% (mean 1.5 ± 1.0%) of their body mass in herring per day during molt (Table S4). Again, there were no differences between sexes in dietary intake prior to (unpaired two sample t-test, $t_{8} = 0.48, P = 0.64$) or during molt (unpaired two sample t-test, $t_{8} = 1.13, P = 0.29$).

Table 1. Mean amino acid molar composition (mol % ± SD) for herring muscle and penguin feather, as well as mean proximate analysis of crude protein, fat, and carbohydrate content (% by weight ± SD) of herring muscle (n = 3). Essential amino acids (carbon) designated with $^\text{E}$ and source amino acids (nitrogen) designated with $^\text{S}$.

| Amino Acid | Herring Muscle | Penguin Feather |
|------------|----------------|-----------------|
| Alanine    | 4.2 ± 0.3      | 5.2 ± 0.5       |
| Aspartic acid | 7.5 ± 0.6   | 6.4 ± 0.7       |
| Glutamic acid | 10.9 ± 0.8  | 7.1 ± 0.9       |
| Glycine    | 3.4 ± 0.1      | 11.5 ± 0.6      |
| IsoleucineE | 3.1 ± 0.3     | 4.5 ± 0.4       |
| Leucine    | 5.9 ± 0.4      | 7.7 ± 0.6       |
| PhenylalanineES | 2.8 ± 0.2 | 2.1 ± 0.5       |
| Proline    | 2.7 ± 0.1      | 10.8 ± 0.4      |
| Serine$^\text{S}$ | 2.6 ± 0.2    | 7.5 ± 0.3       |
| ThreonineE | 3.0 ± 0.2      | 4.7 ± 0.4       |
| Valine$^\text{E}$ | 3.2 ± 0.3    | 8.7 ± 0.5       |
| Protein    | 72.4 ± 0.4     | –               |
| Fat        | 18.1 ± 2.1     | –               |
| Carbohydrate | 0.2 ± 0.1    | –               |

Table 2. Mean $\delta^{13}$C values and $\delta^{15}$N values (‰ ± SD) in bulk tissue and individual amino acids of herring muscle and penguin feather (n = 3 composite samples for herring and n = 10 individuals for penguins) along with mean $\Delta^{13}$C and $\Delta^{15}$N values (‰ ± SD) between herring (diet) and penguin (consumer) in the controlled feeding experiment. The results of one sample t-tests ($\alpha = 0.05$) to determine whether individual amino acid $\Delta^{13}$C$_{C,D}$ and $\Delta^{15}$N$_{C,D}$ values were significantly different from 0‰ ($t$ statistic and significance [$^*P < 0.05$, $^**P < 0.01$, $^***P < 0.001$]) are in parentheses after the trophic fractionation factors (n = 10 individuals). Essential amino acids (carbon) designated with $^\text{E}$ and source amino acids (nitrogen) designated with $^\text{S}$.

| Amino Acid       | Herring $\delta^{13}$C | Penguin $\delta^{13}$C | $\Delta^{13}$C$_{C,D}$ (t statistic) | Herring $\delta^{15}$N | Penguin $\delta^{15}$N | $\Delta^{15}$N$_{C,D}$ (t statistic) |
|------------------|------------------------|------------------------|--------------------------------------|------------------------|------------------------|--------------------------------------|
| Bulk             | −16.9 ± 0.1            | −15.9 ± 0.3            | 1.0 ± 0.3 (10.6$^{***}$)              | 13.7 ± 0.2             | 17.2 ± 0.4             | 3.5 ± 0.4 (27.2$^{**}$)              |
| Alanine          | −11.8 ± 0.6            | −10.4 ± 1.1            | 1.5 ± 1.1 (4.4$^{**}$)                | 21.5 ± 0.5             | 24.8 ± 0.5             | 3.4 ± 0.5 (21.3$^{**}$)              |
| Aspartic acid    | −12.4 ± 0.8            | −12.9 ± 1.0            | −0.5 ± 1.0 (−1.6$^{**}$)              | 17.5 ± 0.2             | 21.9 ± 0.6             | 4.4 ± 0.6 (21.7$^{**}$)              |
| Glutamic acid    | −11.9 ± 0.7            | −12.0 ± 0.9            | −0.1 ± 0.9 (−0.3$^{**}$)              | 20.6 ± 0.5             | 24.4 ± 0.6             | 3.8 ± 0.6 (20.9$^{**}$)              |
| Glycine          | −1.4 ± 0.4             | 0.4 ± 0.4              | 1.8 ± 0.4 (13.7$^{***}$)              | 3.4 ± 0.5              | 5.3 ± 0.9              | 1.9 ± 0.9 (6.6$^{**}$)               |
| IsoleucineE      | −11.6 ± 0.6            | −11.7 ± 0.3            | −0.1 ± 0.3 (−0.5$^{**}$)              | 20.3 ± 0.7             | 25.7 ± 0.5             | 5.4 ± 0.5 (33.0$^{***}$)             |
| LeucineE         | −27.0 ± 0.3            | −27.2 ± 0.3            | 0.2 ± 0.3 (−1.4$^{**}$)               | 20.7 ± 0.4             | 25.6 ± 0.2             | 4.9 ± 0.2 (64.6$^{**}$)              |
| PhenylalanineES  | −24.4 ± 0.4            | −24.3 ± 0.3            | 0.1 ± 0.3 (0.6$^{**}$)                | 1.7 ± 0.4              | 2.0 ± 0.5              | 0.3 ± 0.5 (19.9$^{**}$)              |
| Proline          | −13.8 ± 0.6            | −13.0 ± 0.9            | 0.1 ± 0.9 (0.4$^{**}$)                | 20.7 ± 0.7             | 25.9 ± 0.4             | 5.2 ± 0.4 (41.2$^{**}$)              |
| Serine           | −3.4 ± 0.4             | 5.9 ± 0.8              | 2.4 ± 0.8 (9.2$^{***}$)               | 3.7 ± 0.2              | 5.6 ± 0.4              | 1.9 ± 0.4 (13.7$^{**}$)              |
| ThreonineE       | −7.2 ± 0.7             | −7.3 ± 0.2             | 0.1 ± 0.2 (−1.4$^{**}$)               | −10.3 ± 1.4            | −21.7 ± 1.1            | −11.4 ± 1.1 (−31.7$^{***}$)          |
| ValineE          | −20.9 ± 0.3            | −20.6 ± 0.3            | 0.3 ± 0.3 (3.6$^{**}$)                | 22.5 ± 0.3             | 26.5 ± 0.6             | 4.0 ± 0.6 (22.1$^{***}$)             |

Carbon isotopes in captive penguins and diet

Nearly all essential AAs had trophic fractionation values that were not significantly different from 0‰ and were lower than bulk tissue carbon trophic fractionation (Table 2, Fig. 1). Valine was the only essential AA with a $\Delta^{13}$C$_{C,D}$ value significantly greater than 0 (0.3‰, 95% CI = 0.13–0.55‰), although this value is probably not ecologically significant given analytical uncertainty. All nonessential AAs, on the other hand, had highly variable trophic fractionation values (Table 2, Fig. 1) with significant differences among individual nonessential AAs (one-way ANOVA, $F_{5,54} = 18.1, P = 1.7e^{-10}$). The nonessential AAs Gly and Ser, which can be synthesized through glycolytic pathways, and Ala, which can be synthesized via glycolytic and Kreb cycle pathways, had $\Delta^{13}$C$_{C,D}$ values that were significantly higher than 0‰ for all individuals (Table 2). Conversely, there was far more individual variability in trophic fractionation for the nonessential AAs Glu, Asp, and Pro synthesized via Kreb cycle intermediates (s.d. ~1.0‰, Table S5), with some individuals having $\Delta^{13}$C$_{C,D}$ values greater than 0‰ and others having $\Delta^{13}$C$_{C,D}$ values less than 0‰. Despite notable differences
in AA abundance between herring muscle and penguin feather (Table 1), the slope of the relationship between AA imbalance and nonessential AA \( D_{13CC-D} \) (\( f(x) = -0.1x + 0.5, r^2 = 0.24, t_8 = 0.98, P = 0.35 \)) was not significantly different from 0\(^{\%}\) (Fig. 2).

Carbon isotope signatures of wild penguins

Wild-caught penguins around King George Island, Antarctica had essential AA \( \delta^{13}C \) patterns (Table 3) suggesting that eukaryotic microalgae were the carbon source at the base of the food web supporting penguin production (probability = 99\(^{\%}\)) according to our isotopic fingerprinting LDA model (Fig. 3). The model predicted little to no protein contribution from prokaryotic cyanobacteria or benthic macroalgae.

Nitrogen isotopes in captive penguins and diet

Individual AAs showed a much wider range in AA \( \delta^{15}N_{C-D} \) values (−11.4\(^{\%}\) to 5.4\(^{\%}\)) than was found in bulk tissue \( \delta^{15}N_{C-D} \) value (Table 2; Fig. 4). The source AA Phe was the only AA with a \( \delta^{15}N_{C-D} \) value not significantly different from 0\(^{\%}\) (Table 2; Fig. 4). All AAs that were a priori identified as trophic AAs as well as two AAs often considered source AAs (Gly, Ser) had \( \delta^{15}N_{C-D} \) values significantly higher than 0\(^{\%}\) (Table 2; Fig. 4). There were significant differences in \( \delta^{15}N_{C-D} \) among individual trophic AAs (one-way ANOVA, \( F_{6,63} = 23.2, P = 2.9e^{-14} \)). However, similar to carbon, the slope of the relationships between AA imbalance and trophic AA \( \delta^{15}N_{C-D} \) (\( f(x) = -0.1x + 4.3, r^2 = 0.15, t_8 = 1.1, P = 0.31 \)) was not significantly different from 0\(^{\%}\) despite differences in AA abundance between herring muscle and penguin feather.

CSIA trophic position calculations of wild penguins

Estimated TP\(_{CSIA}\) values of five wild penguin species varied significantly depending on the equation and value of TDF\(_{Glu-Phe}\) used (two-way ANOVA, \( F_{1,20} = 86.0, P = 1.1e^{-8} \)) (Fig. 5). The TP\(_{CSIA}\) values of the five species estimated from the single literature TDF\(_{Glu-Phe}\) value of 7.6\(^{\%}\) (eq.1) were significantly lower than estimates from the multi-TDF\(_{Glu-Phe}\) equation (eq. 2) (Tukey’s HSD post hoc test, \( P < 0.05 \)). There were also significant differences in TP\(_{CSIA}\) among the five penguin species (two-way ANOVA, \( F_{4,20} = 18.6, P = 1.6e^{-8} \)) (Fig. 5). There was no significant interaction term between species and method of TP\(_{CSIA}\) calculation (two-way ANOVA, \( F_{4,20} = 2.6, P = 0.07 \)).
Table 3. Mean $\delta^{13}$C values and $\delta^{15}$N values ($\mu$ ± SD) in bulk tissue and individual amino acids of wild gentoo penguin feathers ($n = 5$ individuals) collected from King George Island, Antarctica. Essential amino acids (carbon) designated with $\text{E}$ and source amino acids (nitrogen) designated with $\text{S}$.

|            | Gentoo $\delta^{13}$C | Gentoo $\delta^{15}$N |
|------------|------------------------|------------------------|
| Bulk       | $-23.6 \pm 0.2$        | $9.8 \pm 0.2$          |
| Alanine    | $-20.1 \pm 0.7$        | $18.3 \pm 0.6$         |
| Aspartic acid | $-16.2 \pm 0.6$    | $14.0 \pm 0.7$         |
| Glutamic acid | $-14.9 \pm 1.4$       | $18.9 \pm 0.5$         |
| Glycine$^\text{E}$ | $-10.6 \pm 0.5$    | $9.5 \pm 0.8$          |
| Isoleucine$^\text{E}$ | $-14.9 \pm 0.9$    | $20.0 \pm 0.5$         |
| Leucine$^\text{E}$ | $-26.3 \pm 0.7$       | $19.3 \pm 0.6$         |
| Phenylalanine$^\text{E}$ | $-23.9 \pm 0.9$   | $5.4 \pm 0.5$          |
| Proline    | $-17.0 \pm 1.2$        | $20.5 \pm 0.4$         |
| Serine$^\text{S}$ | $-7.1 \pm 1.2$       | $11.1 \pm 0.8$         |
| Threonine$^\text{E}$ | $-11.7 \pm 0.7$      | $-15.5 \pm 1.2$        |
| Valine$^\text{E}$ | $-24.1 \pm 0.6$       | $21.1 \pm 0.6$         |

Discussion

The stable isotope analysis of individual compounds is a powerful and rapidly expanding approach to study food web architecture, including food chain length, resource utilization, and biogeochemical cycling (McMahon et al. 2013a). We found that essential AA $\delta^{13}$C values and source AA (Phe) $\delta^{15}$N values in feathers showed little fractionation between diet and consumer, and thus provide excellent proxies of $\delta^{13}$C$_{\text{baseline}}$ and $\delta^{15}$N$_{\text{baseline}}$ in a non lethal, archival tissue that is widely used in avian ecology. Conversely, nonessential AA $\Delta^{13}$C$_{\text{D}}$ values varied significantly according to their biosynthesis pathway and reflected the macromolecule sources being utilized for biosynthesis. When calculating trophic position of wild penguins, we found that using a multi-TDF$_{\text{Glu-Phe}}$ equation with our new avian-derived TDF$_{\text{Glu-Phe}}$ value (3.5‰) produced more ecologically realistic trophic position estimates than using a single literature value of 7.6‰.

Carbon trophic fractionation in amino acids

Our results support the general patterns of heterotrophic consumers across a wide range of taxa in marine (McMahon et al. 2010; Newsome et al. 2011) and terrestrial systems (Hare et al. 1991; Howland et al. 2003). We found that modest bulk carbon isotope trophic fractionation was underlain by virtually no trophic fractionation in essential AAs and larger, more variable trophic fractionation in nonessential AAs. The fidelity with which essential AAs reflect dietary sources is one of the major reasons why CSIA is such a powerful tool for dietary reconstruction (McMahon et al. 2013a,b). Here, we demonstrate that the essential AA isotope fingerprinting approach (Larsen et al. 2009, 2013) can be done with nonlethally sampled feathers of wild penguins. Our LDA model predicted that eukaryotic marine microalgae were the dominant primary producers supporting the gentoo penguin food chain around King George Island, Antarctica. This result, while not surprising given that penguins in this region inhabit a diatom and krill-centric pelagic food web (Ducklow et al. 2007), provides a clear demonstration of the potential uses of essential AA $\delta^{13}$C values.
in feathers as accurate tracers of baseline primary production sources.

Unlike essential AAs, nonessential AAs showed highly variable trophic fractionation, both among individual nonessential AAs as well as among individual penguins. Nearly all nonessential AAs had nonzero $\Delta^{13}C_{C,D}$ values when examined at the level of individual penguins, which indicates significant de novo biosynthesis of these AAs rather than direct routing from the diet. This was somewhat surprising given that direct isotopic routing of dietary nonessential AAs is energetically favorable when consuming a high-protein diet (Ambrose and Norr 1993; Tieszen and Fagre 1993; Jim et al. 2006). The prevalence of de novo biosynthesis may be a function of captive penguins dramatically restricting their food intake during the 2–3 weeks of fasting associated with feather synthesis (Polito et al. 2011a).

During de novo biosynthesis, the “scrambled egg” hypothesis assumes that ingested macromolecules (protein, lipids, carbohydrates) are dissembled into a bulk carbon pool and then reassembled into AAs (Martinez del Rio et al. 2009; Newsome et al. 2011). Therefore, we would expect that all individuals feeding on the same diet would have similar $\Delta^{13}C_{C,D}$ values. However, recent evidence suggests that animals feeding on heterogeneous diets with significant contributions from multiple macromolecules can differentially use the carbon skeletons of these dietary constituents, depending on where in the metabolic process individual AAs are synthesized (O’Brien et al. 2002; McMahon et al. 2010; Newsome et al. 2011). The patterns of nonessential AA trophic fractionation in our study appeared to be related to AA biosynthetic family and the macromolecules used to synthesize them.

The penguins in our study were fed a diet of fish with significant contributions of protein and lipids that differ in their $\delta^{13}C$ values. Lipids are depleted in $^{13}C$ compared to protein, with $\delta^{13}C$ values $4–8\%_\text{oo}$ lower than whole tissues (DeNiro and Epstein 1977). The glycolytic AAs (Gly, Ser) had consistently positive trophic fractionation values that were significantly greater than the bulk trophic fractionation value. Protein likely provided the $^{13}C$-enriched source of carbon for the de novo biosynthesis of Gly and Ser via 3-phosphoglycerate (Howland et al. 2003; Jim et al. 2006; McMahon et al. 2010). The Kreb cycle AAs (Glu, Asp, Pro), on the other hand, had more variable $\Delta^{13}C_{C,D}$ values among individual penguins, suggesting greater plasticity in the utilization of different macromolecule carbon sources. For instance, some individual penguins in our study showed positive Kreb cycle AA $\Delta^{13}C_{C,D}$ values, suggesting de novo biosynthesis from a protein carbon source similar to the glycolytic AAs. Conversely, other individuals showed negative Kreb cycle AA $\Delta^{13}C_{C,D}$ values, suggesting reliance on a different macromolecule pool. The negative $\Delta^{13}C_{C,D}$ values for Kreb cycle AAs in some individuals suggest that these individuals were using more $^{13}C$-depleted lipids as the carbon source for biosynthesis. Oxidation of $^{13}C$-depleted dietary lipids results in the production of $^{13}C$-depleted acetyl coenzyme A, which is further oxidized in the Kreb cycle to produce $^{13}C$-depleted keto acids used to synthesize Glu, Asp, and Pro. The enhanced use of lipids in some individuals may indicate nutritional stress during molting. Marine birds store most of their body fuel as fat, primarily triglycerides, which are liberated to free fatty acids to deal with nutritional stress associated with molting and migration (Cherel et al. 1992).

We would expect to see a significant relationship between AA composition and trophic fractionation, where a larger AA imbalance necessitates enhanced biosynthesis and thus larger nonessential AA $\Delta^{13}C_{C,D}$ values (McMahon et al. 2010). However, utilization of a variety of dietary macromolecules with different $\delta^{13}C$ values provides an explanation for the lack of relationship between AA.
imbalance and nonessential AA trophic fractionation. Variability in nonessential AA carbon isotope values reflects the complexity in resource utilization among species and individuals, and demonstrates the value of CSIA for examining the underlying drivers of nutritional ecology.

**Nitrogen trophic fractionation in amino acids**

A major focus of CSIA in recent years has been the influence of biochemical and physiological processes on consumer stable nitrogen isotope fractionation (McMahon et al. 2013a). Our results generally support the conclusions of previous controlled feeding experiments indicating that bulk trophic fractionation factors ($\Delta^{15}N_{C,D}$ values = $3.5 \pm 0.4\%_{oo}$ in this study) reflect the relatively large fractionation of trophic AAs linked to glutamate metabolism (mean $\Delta^{15}N_{C,D}$ across all trophic AAs = $4.4 \pm 0.8\%_{oo}$ in this study) and no fractionation of the source AA Phe (McClelland and Montoya 2002; Chikaraishi et al. 2009). The source AA Phe showed minimal fractionation between diet and consumer, preserving a record of $\delta^{15}N$baseline in feathers. Unlike carbon, where the lack of fractionation of Phe is due to the inability of animals to complete the complex enzymatic pathway necessary to generate the phenol side chain (Gibson and Pittard 1968), minimal nitrogen isotope fractionation is related to C-N bond integrity during the metabolic processing of Phe in animals (e.g., conversion of Phe to tyrosine via phenylalanine 4-monooxidase [Bender 2012; Chikaraishi et al. 2007]). As such, Phe $\delta^{15}N$ values provide a robust tracer of the sources and cycling of N at the base of the food web, independent of the unique synthesis pathways of essential AAs characteristic of distinct primary producer groups (Sherwood et al. 2014; Vokshoori and McCarthy 2014). Note, Gly and Ser, which have previously been identified as source AAs, exhibited trophic fractionation values significantly larger than $0\%_{oo}$. Our results support the growing body of literature indicating that these AAs should not be considered “source AAs” given their variable $\Delta^{15}N_{C,D}$ values, from $<1\%_{oo}$ to $>8\%_{oo}$ across a wide range of species (Chikaraishi et al. 2009; Germain et al. 2013).

While trophic AAs in our study had $\Delta^{15}N_{C,D}$ values significantly greater than $0\%_{oo}$, as reported in the previous literature (McClelland and Montoya 2002; Chikaraishi et al. 2007, 2009), the magnitude of trophic fractionation for penguins was much lower than reported for many previously reported consumers. Diet quality and consumer’s mode of nitrogen excretion have both been hypothesized to affect trophic AA $\Delta^{15}N$ values (Germain et al. 2013; Bradley et al. 2014; McMahon et al. in press). In a controlled feeding experiment on fishes, McMahon and colleagues (in press) found that Glu $\Delta^{15}N_{C,D}$ values were significantly related to two important metrics of diet quality, diet protein content and AA imbalance between diet and consumer. Glutamic acid $\Delta^{15}N_{C,D}$ values were significantly lower for consumers fed higher-quality diets with AA compositions that more closely matched the consumer tissue. In addition, several recent studies have shown low Glu $\Delta^{15}N_{C,D}$ values for ureotelic/uricotelic/urea-retaining consumers (Germain et al. 2013; Hoen et al. 2014). Excess amino acids are catabolized to ammonia, which is either excreted in the case of most aquatic fishes and invertebrates or converted to urea or uric acid by marine mammals and birds (Balinsky 1972). Glutamate, whose keto acid $\alpha$-ketoglutarate is the primary amino acceptor for a number of AAs, plays a lesser role in the excretion of uric acid (Moe 2006). Therefore, the reduction in glutamate transamination among uric acid-producing organisms, such as penguins, may result in a reduction in isotopic fractionation, producing depleted glutamate $\delta^{15}N$ values (Styng et al. 2010). The roles of diet quality and nitrogen excretion mode in determining TDFGlu-Phe values are not mutually exclusive, as both lead to reductions in glutamic acid fractionation during trophic transfers.

**CSIA-based trophic position estimations**

The CSIA approach to estimating trophic position has the distinct advantage of providing independent assessment of the number of trophic transfers and $\delta^{15}N_{baseline}$ values for a consumer from a single sample without needing to a priori characterize and analyze the baseline of a food web. However, this approach fundamentally relies on accurate TDFGlu-Phe values to produce accurate TPCSIA estimates. Given the minimal fractionation of the source AA Phe coupled with the relatively low trophic fractionation of the trophic AA Glu, we found that the TDFGlu-Phe for penguins ($3.5 \pm 0.4\%_{oo}$) was significantly lower than the previously reported literature TDFGlu-Phe Value of $7.6\%_{oo}$ (Chikaraishi et al. 2007). Our results add to a growing body of literature indicating that TDFGlu-Phe values are not consistently $7.6\%_{oo}$ across all taxa and diet types (Bloomfield et al. 2011; Germain et al. 2013; Bradley et al. 2014; Hoen et al. 2014; McMahon et al. in press). These results likely help explain discrepancies in the previous literature between TPCSIA estimates using a single TDFGlu-Phe value of $7.6\%_{oo}$ and expected trophic positions of upper level marine predators (Lorrain et al. 2009; Dale et al. 2011; Choy et al. 2012).

It is imperative to use appropriate TDFGlu-Phe values to accurately estimate consumer trophic position with
CSIA. To illustrate this point, we examined the \( T_{\text{CSIA}} \) of five species of wild penguins (king Adélie, and northern and southern rockhopper penguins from Lorrain et al. (2009) and gentoo penguins from this study) that were strictly carnivorous and foraged on both primary and secondary consumers, that is, minimum \( T > 3 \) (Cherel et al. 1993, 2007, 2008; Polito et al. 2011b). \( T_{\text{CSIA}} \) estimates of all five species of wild penguins calculated using a single \( T_{\text{DFGlu-Phe}} \) value of 7.6\(^{15}N\), were less than \( T \sim 3.0 \), which was difficult to reconcile with expected trophic positions from bulk SIA and extensive gut content analyses (Cherel et al. 1993, 2007, 2008; Polito et al. 2011b). Furthermore, there were no significant differences in \( T_{\text{CSIA}} \) among the five species when calculated with the single \( T_{\text{DFGlu-Phe}} \) of 7.6\(^{15}N\). Again, this is not ecologically realistic, as king penguins, which typically feed on pelagic fishes (Cherel et al. 1993), should not ecologically realistic, as king penguins, which typically feed on crustaceans (Tremblay and Cherel 2003; Polito et al. 2011b). Conversely, estimates of \( T_{\text{CSIA}} \) using a multi-\( T_{\text{DFGlu-Phe}} \) value that included our avian-specific \( T_{\text{DFGlu-Phe}} \) provided significantly better agreement with the expected ecological role of penguins in Southern Ocean food webs (Cherel et al. 1993, 2007, 2008; Polito et al. 2011b). However, the multi-\( T_{\text{DFGlu-Phe}} \) equation still appeared to underestimate penguin trophic positions. This may be because \( T_{\text{DFGlu-Phe}} \) values are also lower in other trophic levels (e.g., fish) within the penguin food chain (Bradley et al. 2014; Hoen et al. 2014; McMahon et al. in press). Additionally, biases in trophic position estimates from bulk SIA and gut content analyses may account for some of these discrepancies (Layman et al. 2012). None the less, the \( T_{\text{DFGlu-Phe}} \) values derived from feathers in this study resulted in more realistic \( T_{\text{CSIA}} \) estimates for five species wild penguins, four of which were based on previously published AA \( \delta^{15}N \) data from blood (Lorrain et al. 2009). This suggests that the avian-specific \( T_{\text{DFGlu-Phe}} \) values derived from feathers may hold true for other avian tissues as well, although further testing is warranted to confirm such patterns.

In summary, our study provides empirical support for the development and application of CSIA-based studies of avian ecology and illustrates the importance of using appropriate CSIA parameters when characterizing ecosystem properties, such as trophic position and food chain length (DeAngelis et al. 1989; Cabana and Rasmussen 1994; Young et al. 2013).

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**Data Accessibility**

All data used in this manuscript are present in the manuscript and its supporting material.

**Conflict of Interest**

The authors have no conflict of interests to declare.

**References**

Ambrose, S. H., and L. Norr. 1993. Experimental evidence for the relationship of the carbon isotope ratios of whole diet and dietary protein to those of bone collagen and carbonate. Pp. 1–37 in J. B. Lambert and G. Groupe, eds. Prehistoric human bone. Springer, Berlin.

AOAC. 2005. Official methods of analysis. 17th ed. Association of Official Analytical Chemists International, Arlington, VA.

Balinsky, J. B. 1972. Phylogenetic aspects of purine metabolism. Amino Acids 50:993–997.

Bender, D. A. 2012. Amino acid metabolism. 3rd ed. John Wiley & Sons, Hoboken, NJ.

Bloomfield, A. L., T. S. Elsdon, B. D. Walther, E. J. Gier, and B. M. Gillanders. 2011. Temperature and diet affect carbon and nitrogen isotopes of fish muscle: can amino acid nitrogen isotopes explain effects? J. Exp. Mar. Biol. Ecol. 399:48–59.

Boecklen, W. J., C. T. Yarnes, B. A. Cook, and A. C. James. 2011. On the use of stable isotopes in trophic ecology. Annu. Rev. Ecol. Evol. Syst. 42:411–440.

Bradley, C. J., D. J. Madigan, B. A. Block, and B. N. Popp. 2014. Amino acid isotope incorporation and enrichment factors in pacific bluefin tuna, *Thunnus orientalis*. PLoS ONE 9: e85818.

Cabana, G., and J. B. Rasmussen. 1994. Modeling food chain structure and contaminant bioaccumulation using stable nitrogen isotopes. Nature 372:255–257.

Cherel, Y., J. P. Robin, A. Heitz, C. Calgari, and Y. Le Maho. 1992. Relationships between lipid availability and protein utilization during prolonged fasting. J. Comp. Physiol. B. 162:305–313.
Chikaraishi, Y., C. Verdon, and V. RidouxF. 1993. Seasonal importance of oceanic myctophids in king penguin diet at Crozet Islands. Polar Biol. 13:353–357.
Chikaraishi, Y., K. A. Hobson, C. Guinet, and C. Vanpé. 2007. Stable isotopes document seasonal changes in trophic niches and winter foraging individual specialization in diving predators from the Southern Ocean. J. Anim. Ecol. 76:826–836.
Cherel, Y., S. Ducatez, C. Fontaine, P. Richard, and C. Guinet. 2008. Stable isotopes reveal the trophic position and mesopelagic fish diet of female southern elephant seals breeding on the Kerguelen Islands. Mar. Ecol. Prog. Ser. 370:239–247.
Chikaraishi, Y., Y. Kashiyama, N. O. Ogawa, H. Kitazato, and N. Ohkouchi. 2007. Metabolic control of nitrogen isotope composition of amino acids in macroalgae and gastropods: implications for aquatic food web studies. Mar. Ecol. Prog. Ser. 342:85–90.
Chikaraishi, Y., N. O. Ogawa, Y. Kashiyama, Y. Takano, H. Suga, A. Tomitani, et al. 2009. Determination of aquatic food-web structure based on compound-specific nitrogen isotopic composition of amino acids. Limnol. Oceanogr. Methods 7:740–750.
Chikaraishi, Y., N. O. Ogawa, and N. Ohkouchi. 2010. Further evaluation of the trophic level estimation based on nitrogen isotopic composition of amino acids. Pp. 37–51 in N. Ohkouchi, I. Tayasu and K. Koba, eds. Earth, life, and ecosystems: the West Antarctic Peninsula. Philos. Trans. R. Soc. B Biol. Sci. 362:67–94.
DeNaurois, M. J., and S. Epstein. 1977. Mechanism of carbon isotope fractionation associated with lipid synthesis. Science 197:261–263.
Ducklow, H. W., K. Baker, D. G. Martinson, L. B. Quetin, R. M. Ross, R. C. Smith, et al. 2007. Marine pelagic ecosystems: the West Antarctic Peninsula. Philos. Trans. R. Soc. B Biol. Sci. 362:67–94.
Germain, L. R., P. L. Koch, J. Harvey, and M. D. McCarthy. 2013. Nitrogen isotope fractionation in amino acids from harbor seals: implications for compound-specific trophic position calculations. Mar. Ecol. Prog. Ser. 482:265–277.
Gibson, F., and J. Pittard. 1968. Pathways of biosynthesis of aromatic amino acids and vitamins and their control in microorganisms. Bacteriol. Rev. 32:465–492.
Hare, E. P., M. L. Fogel, T. W. Jr Stafford, A. D. Mitchell, and T. C. Hoering. 1991. The isotopic composition of carbon and nitrogen in individual amino acids isolated from modern and fossil proteins. J. Archaeol. Sci. 18:277–292.
Hayes, J. M. 2001. Fractionation of carbon and hydrogen isotopes in biosynthetic processes. Rev. Mineral. Geochem. 43:225–277.
Hoen, D. K., S. L. Kim, N. E. Hussey, N. J. Wallsgrove, J. C. Drazen, and B. N. Popp. 2014. Amino acid δ15N trophic enrichment factors of four large carnivorous fishes. J. Exp. Mar. Biol. Ecol. 453:76–83.
Howland, M. R., L. T. Corr, S. M. Young, V. Jones, S. Jim, N. J. Van Der Merwe, et al. 2003. Expression of the dietary isotope signal in the compound-specific δ13C values of pig bone lipids and amino acids. Int. J. Osteoarchaeol. 13:54–65.
Jim, S., V. Jones, S. H. Ambrose, and R. P. Evershed. 2006. Quantifying dietary macronutrient sources of carbon for bone collagen biosynthesis using natural abundance stable carbon isotope analysis. Br. J. Nutr. 95:1055–1062.
Larsen, T., D. L. Taylor, M. B. Leigh, and D. M. O’Brien. 2009. Stable isotope fingerprinting: a novel method for identifying plant, fungal, or bacterial origins of amino acids. Ecology 15:3526–3535.
Larsen, T., M. Ventura, N. Andersen, D. M. O’Brien, U. Piatkowski, and M. D. McCarthy. 2013. Tracing carbon sources through aquatic and terrestrial food webs using amino acid stable isotope fingerprinting. PLoS ONE 8:e73441.
Layman, C. A., M. S. Araujo, R. Boucek, C. M. Hammerslag-Peyer, E. Harrison, Z. R. Jud, et al. 2012. Applying stable isotopes to examine food-web structure: an overview of analytical tools. Biol. Rev. 87:545–562.
Lorrain, A., B. S. Graham, F. Ménard, B. Popp, S. Bouillon, P. Van Breugel, et al. 2009. Nitrogen and carbon isotope values of individual amino acids: a tool to study foraging ecology of penguins in the Southern Ocean. Mar. Ecol. Prog. Ser. 391:293–306.
Martinez del Rio, C., N. Wolf, S. Carleton, and L. Z. Gannes. 2009. Isotopic ecology ten years after a call for more laboratory experiments. Biol. Rev. 84:91–111.
McClelland, J. W., and J. P. Montoya. 2002. Trophic relationships and the nitrogen isotopic composition of amino acids in plankton. Ecology 83:2173–2180.
McMahon, K. W., S. R. Thorrold, T. S. Elsdon, and M. D. McCarthy. In Press. Trophic discrimination of nitrogen stable isotopes in amino acids varies with diet quality in a marine fish. Limnol. Oceanogr. 60.
McMahon, K. W., M. L. Fogel, T. S. Elsdon, and S. R. Thorrold. 2010. Carbon isotope fractionation of amino acids
in fish muscle reflects biosynthesis and isotopic routing from dietary protein. J. Anim. Ecol. 79:1132–1141.

McMahon, K. W., L. L. Hamady, and S. R. Thorrold. 2013a. Ocean ecosgeochemistry: a review. Oceanogr. Mar. Biol. Annu. Rev. 51:327–374.

McMahon, K. W., L. L. Hamady, and S. R. Thorrold. 2013b. A review of ecosgeochemistry approaches to estimating movements of marine animals. Limnol. Oceanogr. 58:697–714.

Mizutani, H., M. Fukuda, Y. Kabaya, and E. Wada. 1990. Carbon isotope ratio of feathers reveals feeding behavior of cormorants. Auk 1990:400–403.

Moe, O. W. 2006. Uric acid nephrolithiasis: proton titration of an essential molecule? Curr. Opin. Nephrol. Hypertens. 15:366–373.

Newsome, S. D., M. L. Fogel, L. Kelly, and C. Martinez del Rio. 2011. Contributions of direct incorporation from diet and microbial amino acids to protein synthesis in Nile tilapia. Funct. Ecol., 25:1051–1062.

O’Brien, D. M., M. L. Fogel, and C. L. Boggs. 2002. Renewable and nonrenewable resources: amino acid turnover and allocation to reproduction in Lepidoptera. Proc. Natl Acad. Sci. USA 99:4413–4418.

Paine, R. T. 1966. Food web complexity and species diversity. Annu. Rev. 51:327–374.

McMahon, K. W., L. L. Hamady, and S. R. Thorrold. 2013a. Ocean ecosgeochemistry: a review. Oceanogr. Mar. Biol. Annu. Rev. 51:327–374.

McMahon, K. W., L. L. Hamady, and S. R. Thorrold. 2013b. A review of ecosgeochemistry approaches to estimating movements of marine animals. Limnol. Oceanogr. 58:697–714.

Mizutani, H., M. Fukuda, Y. Kabaya, and E. Wada. 1990. Carbon isotope ratio of feathers reveals feeding behavior of cormorants. Auk 1990:400–403.

Moe, O. W. 2006. Uric acid nephrolithiasis: proton titration of an essential molecule? Curr. Opin. Nephrol. Hypertens. 15:366–373.

Newsome, S. D., M. L. Fogel, L. Kelly, and C. Martinez del Rio. 2011. Contributions of direct incorporation from diet and microbial amino acids to protein synthesis in Nile tilapia. Funct. Ecol., 25:1051–1062.

O’Brien, D. M., M. L. Fogel, and C. L. Boggs. 2002. Renewable and nonrenewable resources: amino acid turnover and allocation to reproduction in Lepidoptera. Proc. Natl Acad. Sci. USA 99:4413–4418.

Paine, R. T. 1966. Food web complexity and species diversity. Annu. Rev. 51:327–374.

McMahon, K. W., L. L. Hamady, and S. R. Thorrold. 2013a. Ocean ecosgeochemistry: a review. Oceanogr. Mar. Biol. Annu. Rev. 51:327–374.

McMahon, K. W., L. L. Hamady, and S. R. Thorrold. 2013b. A review of ecosgeochemistry approaches to estimating movements of marine animals. Limnol. Oceanogr. 58:697–714.

Mizutani, H., M. Fukuda, Y. Kabaya, and E. Wada. 1990. Carbon isotope ratio of feathers reveals feeding behavior of cormorants. Auk 1990:400–403.

Moe, O. W. 2006. Uric acid nephrolithiasis: proton titration of an essential molecule? Curr. Opin. Nephrol. Hypertens. 15:366–373.

Newsome, S. D., M. L. Fogel, L. Kelly, and C. Martinez del Rio. 2011. Contributions of direct incorporation from diet and microbial amino acids to protein synthesis in Nile tilapia. Funct. Ecol., 25:1051–1062.

O’Brien, D. M., M. L. Fogel, and C. L. Boggs. 2002. Renewable and nonrenewable resources: amino acid turnover and allocation to reproduction in Lepidoptera. Proc. Natl Acad. Sci. USA 99:4413–4418.

Paine, R. T. 1966. Food web complexity and species diversity. Annu. Rev. 51:327–374.

McMahon, K. W., L. L. Hamady, and S. R. Thorrold. 2013a. Ocean ecosgeochemistry: a review. Oceanogr. Mar. Biol. Annu. Rev. 51:327–374.

McMahon, K. W., L. L. Hamady, and S. R. Thorrold. 2013b. A review of ecosgeochemistry approaches to estimating movements of marine animals. Limnol. Oceanogr. 58:697–714.

Mizutani, H., M. Fukuda, Y. Kabaya, and E. Wada. 1990. Carbon isotope ratio of feathers reveals feeding behavior of cormorants. Auk 1990:400–403.

Moe, O. W. 2006. Uric acid nephrolithiasis: proton titration of an essential molecule? Curr. Opin. Nephrol. Hypertens. 15:366–373.

Newsome, S. D., M. L. Fogel, L. Kelly, and C. Martinez del Rio. 2011. Contributions of direct incorporation from diet and microbial amino acids to protein synthesis in Nile tilapia. Funct. Ecol., 25:1051–1062.

O’Brien, D. M., M. L. Fogel, and C. L. Boggs. 2002. Renewable and nonrenewable resources: amino acid turnover and allocation to reproduction in Lepidoptera. Proc. Natl Acad. Sci. USA 99:4413–4418.

Paine, R. T. 1966. Food web complexity and species diversity. Annu. Rev. 51:327–374.

McMahon, K. W., L. L. Hamady, and S. R. Thorrold. 2013a. Ocean ecosgeochemistry: a review. Oceanogr. Mar. Biol. Annu. Rev. 51:327–374.

McMahon, K. W., L. L. Hamady, and S. R. Thorrold. 2013b. A review of ecosgeochemistry approaches to estimating movements of marine animals. Limnol. Oceanogr. 58:697–714.

Mizutani, H., M. Fukuda, Y. Kabaya, and E. Wada. 1990. Carbon isotope ratio of feathers reveals feeding behavior of cormorants. Auk 1990:400–403.

Moe, O. W. 2006. Uric acid nephrolithiasis: proton titration of an essential molecule? Curr. Opin. Nephrol. Hypertens. 15:366–373.

Newsome, S. D., M. L. Fogel, L. Kelly, and C. Martinez del Rio. 2011. Contributions of direct incorporation from diet and microbial amino acids to protein synthesis in Nile tilapia. Funct. Ecol., 25:1051–1062.

O’Brien, D. M., M. L. Fogel, and C. L. Boggs. 2002. Renewable and nonrenewable resources: amino acid turnover and allocation to reproduction in Lepidoptera. Proc. Natl Acad. Sci. USA 99:4413–4418.

Paine, R. T. 1966. Food web complexity and species diversity. Annu. Rev. 51:327–374.

McMahon, K. W., L. L. Hamady, and S. R. Thorrold. 2013a. Ocean ecosgeochemistry: a review. Oceanogr. Mar. Biol. Annu. Rev. 51:327–374.

McMahon, K. W., L. L. Hamady, and S. R. Thorrold. 2013b. A review of ecosgeochemistry approaches to estimating movements of marine animals. Limnol. Oceanogr. 58:697–714.

Mizutani, H., M. Fukuda, Y. Kabaya, and E. Wada. 1990. Carbon isotope ratio of feathers reveals feeding behavior of cormorants. Auk 1990:400–403.

Moe, O. W. 2006. Uric acid nephrolithiasis: proton titration of an essential molecule? Curr. Opin. Nephrol. Hypertens. 15:366–373.

Newsome, S. D., M. L. Fogel, L. Kelly, and C. Martinez del Rio. 2011. Contributions of direct incorporation from diet and microbial amino acids to protein synthesis in Nile tilapia. Funct. Ecol., 25:1051–1062.

O’Brien, D. M., M. L. Fogel, and C. L. Boggs. 2002. Renewable and nonrenewable resources: amino acid turnover and allocation to reproduction in Lepidoptera. Proc. Natl Acad. Sci. USA 99:4413–4418.

Paine, R. T. 1966. Food web complexity and species diversity. Annu. Rev. 51:327–374.
Stable Isotope Fractionation of Penguin Amino Acids

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Table S2. Mean δ¹³C values (‰ ± SD, n = 5 individuals) for bulk tissue and individual amino acids of wild-caught Atlantic herring (Clupea harengus) fed to gentoo Penguins (Pygoscelis papua) prior to and during molt (January to March 2008) at the Omaha’s Henry Doorly Zoo and Aquarium, Omaha, Nebraska. Essential amino acids designated with E.

Table S3. Mean δ¹⁵N values (‰ ± SD, n = 5 individuals) for bulk tissue and individual amino acids of wild-caught Atlantic herring (Clupea harengus) fed to gentoo Penguins (Pygoscelis papua) prior to and during molt (January to March 2008) at the Omaha’s Henry Doorly Zoo and Aquarium, Omaha, Nebraska. Source amino acids designated with S.

Table S4. Individual gentoo penguins (Pygoscelis papua) reared at the Omaha’s Henry Doorly Zoo and Aquarium, Omaha, Nebraska in 2007/2008. Mass (kg) of penguins was measured just prior to molt.

Table S5. δ¹³C values (‰) for bulk tissue and individual amino acids from feathers of gentoo penguins fed Atlantic herring at the Omaha’s Henry Doorly Zoo and Aquarium, Omaha, Nebraska in 2007/2008. Essential amino acids designated with E.

Table S6. δ¹⁵N values (‰) for bulk tissue and individual amino acids from feathers of gentoo penguins fed Atlantic herring at the Omaha’s Henry Doorly Zoo and Aquarium, Omaha, Nebraska in 2007/2008. Source amino acids designated with S.