Available versus digestible dietary amino acids

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Abstract
Available amino acids are those absorbed from the gastrointestinal tract in a form suitable for body protein synthesis. True ileal digestible amino acids are determined based on the difference between dietary amino acid intake and unabsorbed dietary amino acids at the terminal ileum. The accuracy of ileal digestible amino acid estimates for predicting available amino acid content depends on several factors, including the accuracy of the amino acid analysis procedure. In heat processed foods, lysine can react with compounds to form nutritionally unavailable derivatives that are unstable during the hydrochloric acid hydrolysis step of amino acid analysis and can revert back to lysine causing an overestimate of available lysine. Recently, the true ileal digestible reactive (available) lysine assay based on guanidination has provided a means of accurately determining available lysine in processed foods. Methionine can be oxidised during processing to form methionine sulphoxide and methionine sulphone and cysteine oxidised to cysteic acid. Methionine sulphone, but not methionine sulphoxide or cysteic acid, is partially nutritionally available in some species of animal. Currently, methionine and cysteine are determined as methionine sulphone and cysteic acid respectively after quantitative oxidation prior to acid hydrolysis. Consequently, methionine and cysteine are overestimated if methionine sulphone or cysteic acid are present in the original material. Overall, given the problems associated with the analysis of some amino acids in processed foodstuffs, the available amino acid content may not always be accurately predicted by true ileal amino acid digestibility estimates. For such amino acids specific analytical strategies may be required.

Key words: digestibility: availability: amino acids: processing: protein

Background
The term “available” amino acids has widespread use across multiple research fields. Furthermore, within the nutrition field the same term conjures up numerous interpretations. The definition that the authors of this review subscribe to is that of Hurrell and Carpenter(1), discussed in detail by Rutherfurd and Moughan(2) and put succinctly by Batterham(3), whereby available amino acids refer to “the proportion of the dietary amino acids that are digested and absorbed in a form suitable for protein synthesis”, but does not refer to the amount of dietary amino acid actually utilised since utilisation is a function of both the food (dietary protein and other dietary nutrients) and the animal, while availability is a function of the dietary protein alone.

The available amino acid content of foods have been determined using growth based assays, for example the slope ratio assay(4) and more recently by the indicator amino acid oxidation method(5), but these methods can only examine one amino acid at a time and strictly measure utilisation rather than availability. A simpler approach is to determine absorbed amino acids which can be achieved using the ileal digestibility assay where the absorbed amino acids are quantified from the difference between ingested amino acids and those that remain undigested at the terminal ileum after correction for endogenous ileal amino acids (true ileal digestible amino acid content). The main drawback of using true ileal digestible amino acids as a measure of available amino acids is that for some foods, particularly those that have undergone processing or cooking, the technique is not accurate for all of the amino acids. Batterham(5), in a cornerstone study, showed that the ileal digestible amino acid content was an accurate predictor of the available amino acid content (determined using an animal growth assay) for a soyabean meal, but for a processed cottonseed meal, ileal digestible amino acid content overestimated the available amino acid content for lysine, threonine, methionine and tryptophan but not for isoleucine, leucine and valine. Batterham(5) postulated that the observed difference between digestible and available amino acids was a result of amino acids that had been chemically modified during processing being absorbed in a form that could not be utilised. A contrary view was put forward by workers in New Zealand (the authors of this contribution), namely that the difference between available and ileal digestible amino acids instead results from inadequacies of amino acid analysis methodology. It was argued that traditional amino acid analysis methods do not accurately determine all of the amino acids, particularly those that have undergone chemical
modification during processing. For the latter hypothesis to be 
true, the affected amino acids must be susceptible to chemical 
modification during processing, the modified derivatives must 
be either partially or completely nutritionally unavailable and 
the derivatives must interfere with the determination of their 
parent amino acids when using traditional amino acid 
analysis.

True ileal amino acid digestibility

To determine true ileal amino acid digestibility, a human or 
animal is fed a test diet and ileal digesta are collected. The 
amino acid content of both diets and digesta are then 
determined using conventional amino acid analysis and by 
difference digestibility is calculated. Amino acid analysis tech-
niques have remained largely unchanged since first developed 
in the 1950's and involve the hydrolysis of protein in 6 M HCl 
at 110°C for approximately 24 h in an oxygen free environ-
ment. These hydrolysis conditions are severe and some 
amino acids are chemically altered during hydrolysis and 
cannot be quantitatively determined. For such amino acids, 
other hydrolytic strategies must be used. Asparagine and glu-
tamine are determined as their acid derivatives aspartic acid 
and glutamic acid. Tryptophan is determined using base 
hydrolysis which improves tryptophan yields in comparison 
with hydrochloric acid hydrolysis but even then complete 
recovery of tryptophan is seldom achieved. For cysteine and 
methionine, quantitative conversion to cysteic acid and meth-
ionine sulphone using performic acid oxidation must be car-
ried out prior to hydrochloric acid hydrolysis to improve 
yields, however losses of cysteine are still observed. Serine 
and threonine undergo hydrolytic losses such that 10–15 % 
of these amino acids are lost after 24 h acid hydrolysis.

Lysine

Lysine is a dietary essential (indispensable) amino acid for ani-
mals and humans. Lysine also possesses a reactive side chain 
amino group that can react with other compounds present in a 
food to form nutritionally unavailable derivatives (Maillard 
products). This reaction can occur at any temperature but 
generally accelerates at elevated temperatures such as those 
used during heat processing or cooking. Rutherfurd and 
Moughan(6) showed that the decrease in the reactive lysine 
content of heated field peas was more than three times greater 
after heating at 165°C compared to heating for the same length 
of time at 150°C. Other factors can influence the rate of Maill-
lard product formation, including pH, water activity and reac-
tant concentration. Some of these lysine derivatives formed 
during processing or long-term storage are acid labile and can 
revert back to lysine during hydrochloric acid hydrolysis lead-
ing to an overestimate of lysine content. When conventional 
amino acid analysis techniques are employed, the reverted 
lysine and undamaged lysine cannot be distinguished and 
are determined as “total lysine” and consequently total lysine 
estimates may be misleading. Furthermore, digestibility 
coefficients for total lysine, which are derived from the 
total lysine in the diet and digesta, in samples that contain 
significant amounts of acid labile lysine derivatives are con-
founded and should not be used.

A number of assays have been developed to determine 
reactive lysine (the lysine that possesses an unreacted side 
chain amino group), the most common of which include the 
fluorodinitrobenzene (FDNB) method, guanidination 
method and furosine method. However, reactive lysine is not 
completely absorbed from the digestive tract and none of the 
latter methods account for the incomplete digestion and 
absorption from the gastrointestinal tract that is common 
among heat-treated foods. Moughan and Rutherfurd(13) 
developed a method for determining the true ileal digestibility 
of reactive lysine whereby the reactive lysine content of both 
diets and digesta were determined using the guanidination 
reaction (reaction of O-methylisourea with the ε-amine 
group of lysine to form acid stable homogarginine) and sub-
sequent analysis of homogarginine. The guanidination reaction 
with O-methylisourea (OMIU) is specific for the ε-amine 
group of lysine and OMIU does not react with the α-amine 
group of amino acids with the exception of glycine. Con-
sequently, guanidination is suitable for the analysis of 
reactive lysine in digesta where free lysine may be present. 
True ileal digestible reactive lysine can then be estimated by 
multiplying the true ileal reactive lysine digestibility by the 
reactive lysine content of the food or feedstuff. From a the-
etical standpoint, digestible reactive lysine is equivalent to 
available lysine. Moreover, Rutherfurd et al. conducted 
a study to investigate the accuracy of the assay to predict the 
available lysine content of a heated skim milk powder. The 
study, based on body lysine retention in the growing pig, 
employed a test diet and two control diets containing either 
heated skim milk powder or enzymatically hydrolysed 
casein (EHC) and free amino acids respectively as the sole 
nitrogen source. All diets were identical except for the nitro-
gen source and lysine in the EHC-free amino acid-based diet 
was assumed to be completely digested and absorbed and 
lysine was shown experimentally to be the first limiting 
amino acid in all diets. The control diets were formulated to 
contain lysine at the same level as the heated skim milk 
powder diet based on either true ileal total lysine digestibility 
(traditional amino acid analysis), or true ileal reactive lysine 
digestibility (guanidination). Rutherfurd et al. reported 
that the whole body lysine deposition of the pigs fed the 
heated skim milk powder-based diet was similar to that in 
the pigs fed the control diet formulated based on reactive 
lysine digestibility but was significantly and markedly different 
to that for the pigs fed the control diet formulated based on 
total lysine digestibility (Table 1). This study demonstrated 
the inaccuracy of total lysine digestibility and the accuracy of 
reactive lysine digestibility as a measure of lysine avail-
ability and showed that the inaccuracy of the true ileal diges-
tible total lysine measure in processed feedstuffs was due 
to inadequacies in the traditional amino acid analytical 
approach rather than being due to an inherent flaw in the 
ileal digestibility approach.

The difference between digestible total lysine and digestible 
reactive (available) lysine for processed foods and feedstuffs
tested in our laboratory ranges from nil for minimally processed foods such as UHT milk or a protein supplement to 143% for severely heat-treated feedstuffs such as moist cat food (Table 2). Consequently, ileal digestible lysine predicts available lysine accurately for some processed foods and feedstuffs but for many others the overestimation is considerable. As a result, where the accurate prediction of available lysine is required it is advisable that the digestible reactive lysine rather than digestible total lysine be determined particularly since to do so requires only an additional chemical analysis step rather than a separate bioassay.

Methionine and cysteine

The sulphur amino acids methionine and cysteine can be oxidised in foods that undergo processing or prolonged storage to methionine sulphone and methionine sulphoxide (for methionine) and cysteic acid (for cysteine) (24). In addition cysteine can also react with glucose to produce Maillard products (25) and with lysine to form lysinoalanine (26). Methionine sulphone and cysteic acid have been shown to be nutritionally unavailable in rats (24, 27–29), chickens (28) and turkeys (30), although cysteic acid can be used by some species to synthesise taurine (24, 31). Methionine sulphone on the other hand can be partially utilised in rats (24, 29, 32–35), chickens (30), mice (27, 32) and turkeys (30) by the action of methionine sulphone reductase which converts methionine sulphone to methionine in vivo (57–58). The extent of dietary methionine sulphone utilisation is also dependent on the dietary concentrations of methionine, cysteine and methionine sulphone (24, 34, 36, 39) as dietary methionine sulphone will have a sparing effect on methionine so long as the amount of conversion of methionine sulphone to methionine required does not exceed the capacity of the methionine sulphone reductase enzyme.

True ileal sulphur amino acid digestibility takes into account incomplete digestion and absorption but may not give accurate values for availability for protein sources that have undergone oxidation, since the sulphur amino acids (methionine and cysteine) are traditionally determined following performic acid oxidation to quantitatively convert methionine to methionine sulphone and cysteine to cysteic acid which are then determined using amino acid analysis. Consequently, if significant amounts of methionine sulphone or methionine sulphone are naturally present in the food the methionine will be overestimated. The same is true for cysteic acid and cysteine. Recently, Rutherfurd and Moughan (40) proposed that available methionine in oxidised foods could be predicted as shown below:

\[
\text{Available methionine} = \frac{\text{Met}_\text{food} \times \text{Met digestibility} + \text{MetO}_\text{food} \times \text{MetO digestibility} \times \text{MetO utilisation}}{\text{MetO utilisation}}
\]

Where \(\text{Met}_\text{food}\) is the actual methionine content of the food, Met digestibility is the true ileal digestibility of methionine, \(\text{MetO}_\text{food}\) is the methionine sulphone content of the food, \(\text{MetO digestibility}\) is the true ileal methionine sulphone digestibility and MetO utilisation is the utilisation of dietary methionine sulphone based on the methionine, methionine sulphone and cysteine contents of the food. An accurate estimation of available methionine therefore requires the accurate determination of methionine and methionine sulphone in both diets and digesta of a test animal or human as well as an understanding of the quantitative relationship between...

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Table 1. Least-squares means (n = 8) of whole body lysine deposition (g d⁻¹) in pigs fed a heated skim milk powder based diet and one of two EHC control diets

<table>
<thead>
<tr>
<th>Lysine deposition</th>
<th>Heated skim milk powder</th>
<th>EHC Diet A</th>
<th>EHC Diet B</th>
<th>Overall SE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>5.4²</td>
<td>9.1²</td>
<td>0.61</td>
</tr>
</tbody>
</table>

Table 2. Digestible total and reactive (available) lysine contents (g kg⁻¹) for a range of protein sources

<table>
<thead>
<tr>
<th>Protein Source</th>
<th>Total¹</th>
<th>Reactive²</th>
<th>Difference (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moist cat food</td>
<td>27.0</td>
<td>11.1</td>
<td>143-2</td>
</tr>
<tr>
<td>Dry cat food</td>
<td>17.5</td>
<td>9.2</td>
<td>90-2</td>
</tr>
<tr>
<td>Extruded corn cereal</td>
<td>0.15</td>
<td>0.08</td>
<td>88-9</td>
</tr>
<tr>
<td>Rolled oat breakfast cereal</td>
<td>3.7</td>
<td>2.8</td>
<td>35-5</td>
</tr>
<tr>
<td>Dried maize</td>
<td>2.8</td>
<td>2.2</td>
<td>27-3</td>
</tr>
<tr>
<td>Cottonseed meal</td>
<td>12.9</td>
<td>10.3</td>
<td>25-2</td>
</tr>
<tr>
<td>Whole grain bread</td>
<td>2.8</td>
<td>2.4</td>
<td>17-5</td>
</tr>
<tr>
<td>Evaporated milk</td>
<td>23.4</td>
<td>20.5</td>
<td>14-1</td>
</tr>
<tr>
<td>Wheat meal</td>
<td>3.2</td>
<td>2.9</td>
<td>10-3</td>
</tr>
<tr>
<td>Whole milk powder</td>
<td>26.2</td>
<td>24.0</td>
<td>9-2</td>
</tr>
<tr>
<td>Split lentils</td>
<td>13.3</td>
<td>12.3</td>
<td>7-7</td>
</tr>
<tr>
<td>Maize</td>
<td>1.9</td>
<td>1.8</td>
<td>5-0</td>
</tr>
<tr>
<td>Evaporated milk</td>
<td>18.8</td>
<td>19.9</td>
<td>5-2</td>
</tr>
<tr>
<td>Split peas</td>
<td>16.1</td>
<td>15.4</td>
<td>4-6</td>
</tr>
<tr>
<td>Infant formula</td>
<td>8.3</td>
<td>8.6</td>
<td>3-5</td>
</tr>
<tr>
<td>Meat and bone meal</td>
<td>32.5</td>
<td>31.6</td>
<td>2-8</td>
</tr>
<tr>
<td>Soya bean meal</td>
<td>30.6</td>
<td>31.2</td>
<td>1-9</td>
</tr>
<tr>
<td>Whole milk powder</td>
<td>18.6</td>
<td>18.4</td>
<td>1-2</td>
</tr>
<tr>
<td>UHT milk</td>
<td>31.7</td>
<td>31.4</td>
<td>1-0</td>
</tr>
<tr>
<td>Blood meal</td>
<td>85.9</td>
<td>85.1</td>
<td>0-9</td>
</tr>
<tr>
<td>Protein supplement</td>
<td>14.3</td>
<td>14.3</td>
<td>0-0</td>
</tr>
</tbody>
</table>

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Data reported by Rutherfurd et al. (84–86) and Rutherfurd and Moughan (87).

1 Based on the total lysine content of the food or feedstuff and the true ileal total lysine digestibility.

2 Based on the reactive lysine content (guanidination method) of the food or feedstuff and the true ileal reactive lysine digestibility (guanidination method).

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1 Enzymatically hydrolysed casein.
2 EHC Diet A was formulated to contain lysine equal to the digestible lysine content of the heated skim milk powder determined using the conventional ileal digestibility assay (reactive lysine in heated skim milk powder x true digestibility of reactive lysine (determined using the new method) for the heated skim milk powder).
3 Enzymatically hydrolysed casein.
4 EHC Diet B was formulated to contain lysine equal to the digestible lysine content of the heated skim milk powder determined using the new ileal reactive lysine digestibility assay (reactive lysine in heated skim milk powder x true digestibility of reactive lysine (determined using the new method) for the heated skim milk powder).

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the extent of methionine sulphoxide utilisation and the dietary concentrations of methionine, methionine sulphoxide and cysteine. To that end the analysis of methionine, cysteine and methionine sulphoxide will be discussed.

While performic acid oxidation followed by hydrochloric acid hydrolysis is the most common method used for determining sulphur amino acids in foods, in samples where all the methionine molecules are unoxidised, hydrochloric acid hydrolysis alone can be used to release methionine if rigorous removal of oxygen is undertaken prior to hydrochloric acid hydrolysis(41) since the loss of methionine during hydrochloric acid hydrolysis under oxygen free conditions is negligible(42). Methionine sulphoxide can also be reduced to methionine during hydrochloric acid hydrolysis(40,43–45) which is in contrast to the commonly held belief that methionine is oxidised during acid hydrolysis. Rutherfurd and Moughan(40) suggested that this may be due to the relative flux of oxidation of methionine and reduction of methionine sulphoxide as a result of the extent to which oxygen is removed from the samples being hydrolysed. Consequently, hydrochloric acid would be an unsuitable hydrolysing agent for foods that contain significant amounts of methionine sulphoxide.

Several hydrolytic methods have been reported for determining methionine sulphoxide and these include hydrolysis with methanesulphonic acid(43–44,46–47) or p-toluenesulphonic acid(48), and alkaline hydrolysis(29,38–39,45,49). A method has also been reported based on barium hydroxide hydrolysis where methionine and methionine sulphoxide can be determined in the one analysis procedure(50). Rutherfurd and Moughan(40) reported that hydrolysis in methanesulphonic acid resulted in a substantial loss (50 %) of methionine sulphoxide in an oxidised fishmeal after 24 h methanesulphonic acid hydrolysis which contrasts with the findings of some workers who have reported near complete recovery of methionine sulphoxide using methanesulphonic acid hydrolysis in purified proteins(46) and rumen samples spiked with pure methionine sulphoxide(45). Given the conflicting reports for the analysis of methionine sulphoxide, it would appear that more work needs to be conducted on methods for the accurate analysis of methionine sulphoxide. Furthermore, and to determine ileal methionine sulphoxide digestibility, any promising method would also need to be optimised for digesta samples.

Finally, while the discussion above describes the inadequacies of current analytical methods for determining methionine in proteins that contain methionine sulphoxide or methionine sulphone, it must also be recognised that a significant amount of methionine is used metabolically for purposes other than for protein synthesis, for example, as a methyl donor. Consequently, estimates of utilisable methionine based on body methionine retention such as the slope ratio assay will inevitably underestimate the available methionine content of food proteins as they do not account for the methionine utilised for purposes other than protein synthesis.

**Threonine**

During heating, threonine can react with sugars to form pyrazines, pyridines and pyroles(51) with hundreds of individual compounds being identified(52–55). These structures, however, are unlikely to interfere with the analysis of threonine and such reactions are unlikely to be quantitatively significant under normal conditions of food processing.

The presence of phosphorylated amino acids may lead to an overestimate of the availability of those phosphorylated amino acids, particularly threonine. Phosphothreonine cannot be used directly for the synthesis of body proteins. O- and N-phospho-amino acids occur naturally in many dietary proteins but in addition, controlled phosphorylation is commonly used as a manufacturing technique to alter the functional properties of food proteins(50). Furthermore, uncontrolled phosphorylation may occur when foods are processed at higher temperatures.

Phosphorylated serine, threonine and tyrosine are base labile and partially acid labile(57–58). Under the hydrolytic conditions employed during amino acid analysis (6 M HCl, 24 h, 110°C) recoveries of synthetic phosphothreonine and phosphoserine have been reported to be 36 % and 2 % respectively with 46 % and 81 % of the original phospho-amino acids being dephosphorylated to threonine and serine respectively(59). Similar results were observed for both phosphothreonine and phosphoserine when hydrolysed from phosphothreonine and phosphoserine containing peptides(59). It is likely that for processed protein sources conventional hydrochloric acid hydrolysis overestimates the amounts of threonine and serine if the phosphorylated forms of these amino acids are present and this may have implication for the determination of available threonine and serine if phosphothreonine and phosphoserine cannot be utilised. In addition, as is the case for unphosphorylated threonine and serine, not all the phosphothreonine and phosphoserine can be recovered during hydrochloric acid hydrolysis.

Threonine is a quantitatively important amino acid in the production of intestinal proteins, particularly mucins. As much as 60-100 % of the absorbed threonine is extracted and utilised by the portal drained viscera (PDV) (the intestine, pancreas, spleen and stomach) of pigs(60–61). Furthermore, systemic amino acids would appear to be poorly utilised by the intestine since Dudley et al.(62) showed that mucosal protein synthesis was substantially lower in piglets fed parenterally compared to those fed enterally. Thus the bulk of dietary threonine is used for PDV protein synthesis and the bulk of the threonine used for PDV protein synthesis is likely to be of dietary origin.

Alkaline phosphatase catalyses the dephosphorylation of a wide range of molecules and is present in the enterocytes of the small intestine, predominantly in the duodenum(63–65). However, during a meal containing large amounts of phosphothreonine the activity of intestinal alkaline phosphatase may be insufficient to quantitatively dephosphorylate the absorbed phosphothreonine before it is transported from the brush border to the portal vein. Consequently, the latter phosphothreonine molecules would be largely lost to PDV protein synthesis. The absorbed phosphothreonine may be dephosphorylated in the body but given that most of the threonine requirement is for intestinal protein synthesis the net result
may be poor utilisation of threonine in processed foods that contain large quantities of phosphothreonine.

The impact of phosphorylation on the in vitro digestibility of phosphorylated casein (57), and soya bean proteins (66–67) using a selection of proteases has been reported to be minor. In contrast, Giec et al. (68) reported a 10% decrease in the in vitro digestibility of a yeast homogenate after phosphorylation. To the authors’ knowledge no in vitro studies have been conducted investigating the impact of phosphorylation on protein digestibility. Matheis et al. (57) used a tetrahymena bioassay to show that a broth containing phosphorylated casein reduced the growth of *T. thermophili* by 20% in comparison to a broth containing protease peptone (animal protein hydrolysate) suggesting that with *T. thermophili*, at least, there was a lower utilisation of phosphorylated amino acids compared to their unphosphorylated counterparts.

To summarise, significant amounts of phosphothreonine may be present in some processed food proteins. The presence of phosphothreonine probably does not significantly affect protein digestibility but may impact on threonine utilisation if the phosphorylated threonine cannot be dephosphorylated in the brush border at a rate sufficient to support PVD protein synthesis.

Tryptophan

Tryptophan can also be modified during processing, usually through oxidation, to a range of products depending on the oxidising agents present (69). Weck et al. (70) have shown that the tryptophan oxidation products generated after hydrogen peroxide treatment are not nutritionally available. Tryptophan can be quantitatively recovered during the hydrochloric acid hydrolysis of pure proteins (71), but in foods where significant amounts of carbohydrates are present tryptophan recoveries tend to be low (41). Consequently, base hydrolysis is commonly used for the determination of tryptophan in foods but even then recoveries are incomplete (41).

When analysing samples for tryptophan content, it is common practice to include an internal standard, usually 5-methyl tryptophan, and to correct the determined tryptophan values based on the recovery of the internal standard. This method relies on a similar loss rate for both tryptophan and the internal standard. If the loss rate of the internal standard during hydrolysis is greater than that of tryptophan, then tryptophan content will be overestimated. A disparity in loss rates between tryptophan and the internal standard may explain why ileal digestible tryptophan overestimates available tryptophan in some proteins. However, it is the experience in our laboratory that the loss rates of tryptophan and 5-methyl tryptophan are similar and if they do differ 5-methyl tryptophan degrades more slowly than tryptophan which would lead to an underestimate of tryptophan rather than an overestimate. Overall, more work is required for the accurate analysis of tryptophan in foods and digesta.

**Least-squares nonlinear regression**

While many amino acids are stable under the strong acidic conditions used to hydrolyse proteins for amino acid analysis, some amino acids, for example cysteine, methionine, serine, threonine and tryptophan, are destroyed either partially or totally during hydrochloric acid hydrolysis. Typically hydrolytic losses of 10–15% for serine and 5–10% for threonine are not uncommon after 24 h hydrolysis (72–73). In contrast, branched chain amino acids are difficult to hydrolyse and require more than 24 h hydrolysis for complete liberation. Essentially, the hydrolysis interval of 24 h is a compromise that permits the quantitative hydrolysis and recovery of most but not all amino acids. Hydrolysis for 72–96 h has been used to determine isoleucine and valine content (74–75), while using multiple hydrolysis intervals and extrapolating (linear regression) back to 0 h hydrolysis has been used to estimate serine and threonine content (79). The latter approach assumes that the release of an amino acid from the protein and its subsequent degradation occurs sequentially but it is much more likely that for any given amino acid, liberated residues begin to degrade while other residues are yet to be hydrolysed. Least-squares nonlinear regression, which takes into account the simultaneous hydrolysis and destruction of amino acids has been used to estimate the amino acid contents of foods and other protein sources (42,72,76–80). The method involves determining the amino acid yield for samples that have been

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**Table 3.** Threonine and serine contents (g kg⁻¹) of selected protein sources predicted using either least-squares nonlinear regression or determined after conventional 24 h hydrochloric acid hydrolysis

<table>
<thead>
<tr>
<th>Threonine</th>
<th>Least-squares nonlinear regression</th>
<th>24 h hydrolysis</th>
<th>Underestimation for 24 h hydrolysis (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lysozyme (residues/mol protein)¹</td>
<td>6.9</td>
<td>6.3</td>
<td>9</td>
</tr>
<tr>
<td>Infant formula</td>
<td>53.2</td>
<td>48.6</td>
<td>9</td>
</tr>
<tr>
<td>Skim milk powder</td>
<td>13.5</td>
<td>13.1</td>
<td>3</td>
</tr>
<tr>
<td>Human milk</td>
<td>44.8</td>
<td>39.1</td>
<td>13</td>
</tr>
<tr>
<td>Maize</td>
<td>2.6</td>
<td>2.5</td>
<td>4</td>
</tr>
<tr>
<td>Serine</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lysozyme (residues/mol protein)¹</td>
<td>9.8</td>
<td>8.4</td>
<td>14</td>
</tr>
<tr>
<td>Infant formula</td>
<td>5.57</td>
<td>5.18</td>
<td>7</td>
</tr>
<tr>
<td>Skim milk powder</td>
<td>16.3</td>
<td>15.5</td>
<td>6</td>
</tr>
<tr>
<td>Human milk</td>
<td>46.7</td>
<td>45.7</td>
<td>2</td>
</tr>
<tr>
<td>Maize</td>
<td>3.4</td>
<td>3.2</td>
<td>6</td>
</tr>
</tbody>
</table>

Data reported by Rutherfurd et al. (79), Rutherfurd (80) and Darragh and Moughan (78) and Darragh et al. (72).

¹ The actual threonine and serine content of lysozyme based on amino acid sequence is 7 and 10 residues/mol protein (88).
hydrolysed over multiple intervals and using least-squares nonlinear regression to estimate the amino acid content based on the following model:

\[ B(t) = \frac{A_e b (e^{-kt} - e^{-kt})}{b - 1} + B_o (e^{-kt}) \]

Where \( A_e \) is the protein bound amino acid content, \( B(t) \) is the amino acid concentration at time \( t \), \( B_o \) is the free amino acid concentration prior to hydrolysis, \( b \) is the hydrolysis rate (the rate at which amino acids are released from the protein), \( t \) is the loss rate (the rate at which amino acids are degraded).

In Table 3 the threonine and serine contents predicted using either least-squares nonlinear regression or determined after conventional 24 h hydrochloric acid hydrolysis are given for several protein sources. The threonine and serine content determined using 24 h hydrochloric acid hydrolysis underestimated that predicted using least-squares nonlinear regression by 3–13% for threonine and 2–14% for serine. The main drawback of the least-squares nonlinear regression is the time and cost to run the analyses, given that multiple (between 10 and 25) hydrolysis intervals are required. Consequently, this method is not suitable for routine amino acid analysis of foods. Where it does have a place, however, is when highly accurate amino acid compositional data are required.

Racemisation

Racemisation is another effect of processing, particularly at high pH, leading to the formation of D-amino acids. Amino acids will racemise at differing rates depending on the electron-withdrawing ability of the amino acids side chain with amino acids like cysteine, serine and threonine being the most affected, and the neutral amino acid valine, isoleucine and leucine being the least affected. The proteins within which the amino acids are bound also impact on amino acid racemisation rates. Negligible racemisation occurs in heat-treated proteins in the absence of alkaline treatment after heating at moderate temperatures (120°C for 1 h), and low levels (<3%) of racemisation occurs for most amino acids after heating at high temperatures (230°C for 20 min). D-amino acids are generally not nutritionally available but conventional amino acid analysis determines the D- and L-amino acids collectively thereby overestimating the available amino acid content in alkaline-treated proteins. For proteins that contain significant amounts of the D-amino acid enantiomers, analytical methods that can distinguish between D- and L-amino acids are required. Such methods are discussed in detail by Rutherfurd and Sarwar-Gilani.

Conclusions

The available amino acid content of a food is the amount of dietary amino acids absorbed from the gut in a chemical form that can be used for protein synthesis. True ileal digestible amino acid content is a good predictor of available amino acid content for unprocessed foods and for most amino acids in processed foods. However, for some amino acids, mainly lysine, methionine, cysteine, threonine and tryptophan ileal digestible amino acid content can be a poor predictor of the available content. The differences, by and large, relate to inadequacies of current analytical methodologies to accurately determine the amino acid content in foods and digesta that contain amino acids modified during processing. For these amino acids, other analytical strategies must be adopted if available amino acids are to be determined accurately.

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