Determinaton of protein and amino acid digestibility in foods including implications of gut microbial amino acid synthesis

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Abstract
To meet the protein and amino acid requirements of individuals and of populations requires information not only about their requirements but also about the capacity of available foods to meet those requirements. Most of our current knowledge of the digestibility of food proteins and the methods to estimate it has been derived from work with animals. Because the microbiota of the large intestine alter the amino acid composition of the digesta, and because only trivial quantities of amino acids are absorbed intact from the large intestine, the current method of choice for assessing amino acid digestibility is ileal digestibility corrected for basal endogenous losses, that is, standardized ileal digestibility. For protein as a whole, however, because nitrogen absorbed in forms other than as amino acids can contribute to the nitrogen economy, the absorption of nitrogen over the whole digestive tract is the more appropriate measure. Most of the methods developed for estimating ileal amino acid outflow in animals are not directly applicable to man: the exception is the use of volunteers with an ileostomy. The flow and composition of ileal digesta in human subjects can also be measured by the infusion of a marker and withdrawal of samples through a naso-intestinal tube. However, this method is too demanding for routine use and is likely to be restricted to validating the application to humans of digestibility data obtained either from animals, of which the pig seems most suitable, or in vitro methods. Microbial activity in the gastrointestinal (GI) tract is not confined to the large intestine: the numbers and metabolic activity of the upper GI microbiota lead to substantial amounts of microbial protein leaving the ileum. It appears however that a large proportion of the amino acids used by the upper GI microbiota are preformed - from the diet or from endogenous materials - rather than from de novo synthesis. Although there are still uncertainties about the impact of microbial activity in the upper GI tract, the amino acid composition of ileal digesta provides the best available basis for estimating the proportion of dietary amino acids available for metabolism.

Key words: Protein: Amino acid: Digestion: Fermentation: Microbiota

To meet the protein and amino acid requirements of individuals and of populations, information is needed not only about those requirements but also about the capacity of available foods to meet them. These two kinds of information have to be expressed in the same terms. As it happens, human amino acid requirements have, for the most part, been investigated using diets based on proteins of high bioavailability such as egg or milk, or on free amino acids, which are assumed to be completely absorbed, or on a mixture of the two. So the estimates of requirements are essentially of the protein and amino acids absorbed, i.e., the metabolic requirements. The compatible information that is required to describe foods is two-fold, comprising the concentration and bioavailability of the protein and amino acids that individual foods or mixed diets contain.

The term bioavailability encompasses three properties of foods that can alter the proportion of an amino acid that can be utilized; these properties are digestibility, which describes the net absorption of the amino acid, chemical integrity, which describes the proportion of the amino acid that, if absorbed, is in a utilizable form, and freedom from interference in metabolism resulting from the presence in the food of substances that limit the utilization of the amino acid. Of these, the greatest source of variation in bioavailability is, in most cases, digestibility.

Most of our current knowledge of the digestibility of foods, and of methods to estimate it, has been derived from work with animals. It may therefore be useful to begin by reviewing what has been learned with animals before considering how far those results and those methods may be applied in the context of human nutrition. Although the motivation for obtaining suitable information is different - in commercial animal production efficient use of nutrients is a major determinant of profit - the overall aim of matching food resources most effectively to nutrient needs is the same. In this review the nutrients to be considered are the dietary indispensable amino acids and the total protein.

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Digestibility: amino acids

The commonly used term “amino acid digestibility” does not of course refer to the degradation of amino acids but is shorthand for the proportion of consumed amino acid that is absorbed.

It is worth emphasizing at the outset that digestibility is not a fixed attribute of a food but reflects an interaction between the food and the person or animal eating it. Of course, the composition of a given food is subject to many sources of variation from one sample to the next, but in a given sample chemical composition can – in theory at least – be determined with high precision. Its digestibility however varies according to how and to whom it is fed, so that the digestibility of a food (or rather of a nutrient within that food) cannot be assigned a unique value relevant to all species or all individuals of a species.

For amino acids the current method of choice is ileal digestibility\(^3\). The road to this consensus has been a long one and this is not the place to rehearse all of the evidence that has led there\(^3–5\) but the reasons for this choice stem from two seminal observations. The first observation\(^6\) was that most faecal nitrogen is in the form of microbial protein. Mason \textit{et al.}\(^7\) estimated from the faecal excretion of diaminopimelic acid (DAPA) that as much as 90% of the faecal N of pigs could be of bacterial origin. Subsequent studies using a variety of microbial markers have confirmed this observation. Consequently, the amino acid composition of faeces tends to be closer to that of microbial protein than to that of undigested food residues, so the amino acid composition of faeces varies little with diet. It was concluded that undigested food residues reaching the large intestine are largely degraded by microbial activity during their relatively long residence, their nitrogen being either absorbed or converted into microbial biomass with an amino acid profile more or less independent of their initial composition.

The second observation was that although nitrogen can be absorbed from the large intestine, there is (except perhaps in neonates) little absorption of intact amino acids. This was first demonstrated by Zebrowska\(^8\), who infused hydrolyzed casein into the caecum of pigs. There was very little increase in faecal nitrogen but some 83–90% of the infused N was excreted in the urine with little if any improvement in N retention, in contrast to the substantial response that has led to how and to whom it is fed. It is worth emphasizing at the outset that digestibility is not a fixed attribute of a food but reflects an interaction between the food and the person or animal eating it. Of course, the composition of a given food is subject to many sources of variation from one sample to the next, but in a given sample chemical composition can – in theory at least – be determined with high precision. Its digestibility however varies according to how and to whom it is fed, so that the digestibility of a food (or rather of a nutrient within that food) cannot be assigned a unique value relevant to all species or all individuals of a species.

The required measures are therefore of nitrogen and amino acid intakes, of ileal amino acid outflow and of faecal nitrogen loss, together with appropriate values for the endogenous components, which will be considered in more detail in later papers\(^23–28\).

The measurement of ileal amino acid outflow

Methods for measuring ileal amino acid outflow have been reviewed elsewhere\(^25–28\) and only a brief synopsis will be given here. The methods can be divided into two classes: those in which the whole outflow is collected and those in which only a sample is collected, which is related to the whole flow by use of an indigestible marker. To collect the whole flow requires an ileostomy or ileo-rectal anastomosis and the continuous collection must be sufficiently long for day-to-day variation to be minimized. Whilst obviating the need for a marker, each of these approaches brings its own potential problems. Removal of the ileal digesta means that the potential for recycling nitrogen from the large intestine is lost, with possible consequences for the nitrogen transactions of the small intestine. Although this objection may be
that may alter the nature and extent of digestion there so that they more closely resemble those of the caecum and colon. This may be more relevant with an ileostomy than with an ileo-rectal anastomosis as Hennig et al. (32) reported little change with time in ileal microbial numbers during a 5-month period after ileo-rectal anastomosis in pigs. A further and perhaps decisive concern is the welfare of animals with an ileo-rectal anastomosis.

Digesta collection involving indigestible markers allows shorter and simpler procedures. At its simplest, animals can be killed some time after a test meal and the digesta in the terminal ileum can be removed and the concentrations of amino acids and of marker estimated. Although simple, this approach has several practical problems. First, each animal can be used only once, so that variation between animals cannot be reduced as can by repeated within-animal measurements. Second, especially with small animals, there may be insufficient digesta at the very end of the ileum for the necessary analyses. Collecting digesta from a longer segment of the ileum means that some will be less completely digested.

Commonly used markers include chromic oxide, titanium oxide and acid-insoluble ash. Other heavy metals and rare earth elements have also been used. Dual-phase markers have also been advocated on the premise that amino acids are contained in both liquid and solid phases of the digesta and both need to be represented. Although there have been some comparisons of markers there does not seem to be any consensus as to the relative merits of the various substances.

All methods involving indigestible markers involve the assumption that the marker is distributed in the digesta in the same way as the amino acid (or other nutrient) being measured. This does not necessarily require that the digesta be homogeneous but that the proportion of the intake of marker in the sample be the same as the proportion of the total amino acid flow that is in that sample. This in turn requires the assumption that all the marker fed should reach the point of sampling. Although some investigators have reported virtually complete recovery (30) others have reported recovering significantly less than 100% (37–39), especially of chromic oxide. Marker recovery may well be homogeneous but that the proportion of the intake of marker in the sample be the same as the proportion of the total amino acid flow that is in that sample. This in turn requires the assumption that all the marker fed should reach the point of sampling. Although some investigators have reported virtually complete recovery (30) others have reported recovering significantly less than 100% (37–39), especially of chromic oxide. Marker recovery may well vary amongst marker substances and methods of preparation and administration but complete recovery of the chosen marker is often not tested.

To collect digesta repeatedly and without interrupting normal digesta flow various cannulas have been developed, the simplest being a T-cannula in the terminal ileum. When the T-branch is opened a variable proportion of the flow is diverted to the outside where it can immediately be chilled until analyzed. For most diets this works well but, especially with fibrous materials, there may be a fractionation of the digesta so that the material exiting the cannula is no longer representative of the whole flow. To overcome this problem a post-valvular cannula has been developed (35,34). This is placed just distal to the ileo-caecal junction but so arranged as to divert, while the sample is being collected, practically the whole outflow through the ileo-caecal valve to the exterior. The advantages of this technique now seem to make it the method of choice with pigs. Whatever method is used to obtain digesta it is essential that digesta are collected as soon as they emerge and further bacterial activity prevented: failure to do so results in exaggerated NSP digestibility and presumably other fermentative changes (35).

Apparent, true and standardized ileal digestibility

Subtracting ileal amino acid outflow from amino acid intake gives apparent digestibility. However, because of the contribution of endogenous amino acids to the ileal outflow - a contribution that is neither constant nor proportional to the protein intake - the value obtained depends on the way the food has been given. In particular, the test protein must be included in the diet above a certain threshold concentration to obtain plateau values of apparent digestibility (40–42). However, to obtain digestibility values for individual foods that can be used additively to predict the digestibility of a complete diet, apparent digestibility values must be corrected appropriately for the endogenous losses. For this purpose endogenous losses are considered as consisting of three components, a basal loss that is independent of the diet and which forms part of the animal’s requirement, an additional loss that is induced by giving protein (or peptides, or amino acids) per se (regardless of their source or digestibility) and a further loss which can be accounted as a charge against the particular food, reducing the proportion that is available to meet metabolic requirements. The time-honoured term true digestibility refers to the value corrected for all endogenous losses: the more practical measure is the one corrected only for basal losses, that is, standardized digestibility (53). These issues, as well as endogenous losses and their measurement form the subject of other contributions to this symposium (23,24) and will not be discussed further here.

Measurement of ileal outflow in human subjects

Clearly, most of the methods used to estimate ileal amino acid flows in animals cannot be considered with human subjects. The exception is the use of volunteers who have an ileostomy. This allowed the first direct comparison between measurements in man and those in animals given the same diets (33). Although, as mentioned above, the physiology and microbiology of the ileum are altered following ileostomy, the qualitative effects of these changes on protein digestion and ileal amino acid outflow are not yet known. For example (43), the ileal digestibility of pectin and hemicellulose in ileostomates was 0.87 and 0.45 and the daily output of DAPA in ileostomy fluid was 31 mg, about 40% of that in faeces, which suggests a distribution of fermentation between the small and large
Surrogates for humans

The principal candidates for providing digestibility data applicable to man are other animals and in vitro systems. The primary requirement of course is simply that the proposed system should provide digestibility values that are either the same as those obtained in man, or that can be transformed numerically by a consistent formula. However, given the paucity of data obtained with human subjects a more feasible starting point is to set out certain contributory requirements.

(1) The digestive system should be similar to that of man.
(2) The system should accept diets as consumed, and not require additional processing.
(3) Diets should be completely consumed, with no selection.

Animals that are commonly used in digestibility studies are the rat, the chicken and the pig. The first requisite probably excludes birds, which have a gizzard and do not express lactase activity; they also have a shorter retention time of food in the GIT. It probably also excludes rats unless coprophagy is prevented. In addition, to prevent food selection by rats diets must be finely ground or otherwise processed before feeding; processes that may alter digestibility.

The digestive system of the pig differs from that of man principally in having a large caecum; however, this is not relevant to ileal digestibility. Pigs do not normally practice coprophagy and, true to their reputation, readily eat unprocessed diets without selection.

Possible limitations of ileal digestibility: the role of the upper intestinal microbiota

As discussed earlier, the reason for rejecting faeces in favour of ileal digesta as the basis for assessing amino acid digestibility is the modifying effect of the large intestinal microbiota on the amino acid composition of the digesta. The upper digestive tract, however, has its own microbiota. Only since the development of culture-independent methods have the extent and diversity of this population been apparent.

Many of the species have never been cultured and we do not yet have a clear picture of the impact of this population on the nitrogen transactions that are of interest to us in terms of digestion and absorption of amino acids. However, there is evidence that its impact, in pigs at least, is considerable. It was estimated, from the concentration of DAPA in the ileal effluent of pigs, that the microbial biomass leaving the ileum was a quarter of that in the faeces. Other estimates in pigs show that up to half the nitrogen leaving the ileum may be in microbial biomass. Studies of microbial metabolic activity, measured as adenylate energy charge, are as high in the ileum as in the large intestine in pigs. Related to this microbial activity, substantial digestion of various non-starch polysaccharides has been shown to occur proximal to the ileo-caecal junction, providing an energy source for microbial amino acid synthesis. It is important to bear in mind that these observations were made in cannulated animals in which there may have been some modification of the normal microbial population and its environment.

Although there have been several studies of the microbiology of human ileostomy fluid there is understandably rather little information about the normal human ileal microbiota. Ileal digesta have been reported to contain $10^7-10^8$ bacteria/mL, two to three orders of magnitude less than in faeces. More recent analyses using nucleic acid-based methods have revealed the diversity of organisms in the ileum, the similarities and differences between ileal and colonic flora, and considerable differences between individuals, but each study included samples from only two or three individuals, taken during surgery or at autopsy. Very recent reports based on ileal digesta sampled by naso-intestinal tube provide a clearer picture of the microbiota of the normal ileum.

What are the consequences of this microbial activity on the fate of dietary amino acids (and of amino acids in endogenous proteins that may potentially be reabsorbed)? What is the impact of microbial amino acid synthesis in the upper GI tract on estimates of ileal amino acid digestibility?

The gastrointestinal microbiota can both degrade and synthesize amino acids and, given the heterogeneity of the intra-luminal environment, both may occur simultaneously. When Nesheim & Carpenter pointed out many years ago that microbial degradation of amino acids in the gut would lead to a spurious over-estimate of digestibility, they were referring to faecal nitrogen but the same is true for ileal amino acid digestibility. Equally, a net synthesis of amino acids by the microbiota would lead to an under-estimate of digestibility. Both possibilities need to be considered.

Amino acid catabolism by the upper intestinal microbiota

It is technically difficult to account for the nitrogen trans-actions of the GI microbiota, both because of the very diverse populations that coexist, each with its own activities and because the activities of the microbiota are often confounded with those of the gut tissues. Thus, comparisons of amino acid
disappearance in the upper GIT (i.e. intake minus ileal outflow) with portal uptake\(^1\) include five components: dietary amino acid absorption, absorption of endogenously secreted amino acids, microbial amino acid metabolism (both degradation and synthesis) and amino acid utilization by enterocytes. These have not yet been satisfactorily disentangled. A further complication to assessing microbial amino acid degradation is that, although luminal ammonia is generated from the amino acids of dietary (or endogenous) proteins, it is also produced by the hydrolysis of urea. Typically, in non-ruminants, some 20-30% of whole-body urea synthesis is recycled through the gut\(^9\); the ammonia generated is either reabsorbed and returned to the urea pool or utilized by the enteric flora. As noted previously\(^9\) the observation\(^7\) that there is an almost equal return of the carbon and nitrogen moieties of urea to the urea pool, together with evidence of urea synthesis in enterocytes\(^7\) suggests that urea hydrolysis may occur close to the brush border, the ammonia generated constituting a pool different in both size and turnover from that produced in the degradation of amino acids. The proportion of the ammonia generated by urea hydrolysis in the human GI tract that returns to the urea pool has generally been estimated to be 10-25%\(^5,56\) but one estimate was as high as 70%\(^57\). Interestingly, the proportion was not affected by protein intake\(^56\) or starvation\(^7\). Taken together, the evidence suggests that, in a healthy human, 30-70mgN.kg\(^{-1}.d^{-1}\) from urea hydrolysis contributes to luminal ammonia. Both urea and ammonia pass from the ileum into the large intestine\(^55\) but urea makes only a small contribution to faecal ammonia\(^56\) or faecal microbial protein\(^7\).

Measurements of ammonia concentration in the ileal digesta of pigs show an increase with dietary protein\(^7\), suggestive of greater microbial degradation of amino acids, but without data on luminal ammonia turnover the microbial flux of amino acid N to ammonia cannot be estimated from these data.

Other breakdown products of amino acids include a variety of amines, some of which have been measured in the ileal digesta of pigs\(^8\). Again, though, their quantitative significance is hard to assess without data on rates of production. The extent of microbial amino acid degradation in the upper GI tract therefore remains an area of uncertainty.

**Amino acid synthesis by the upper intestinal microbiota**

The impact of microbial amino acid synthesis in the upper GI tract on ileal digestibility depends on the origin of the amino acids in the microbial protein leaving the ileum. There is an important distinction between microbial amino acids, meaning simply amino acids that form part of microbial protein, and microbially-synthesized amino acids, i.e., those the microorganisms have formed by *de novo* synthesis, whether the carbon and nitrogen for these syntheses are derived from the degradation of amino acids (dietary or endogenous) or from non-protein materials and non-amino nitrogen. If the microbiota simply incorporate into their protein preformed amino acids - whether of dietary or endogenous origin - that would otherwise have been absorbed the passage of these amino acids into the large intestine represents a loss just as much as if they were not part of microbial protein. However, it is clear that the microbiota of the small intestine do synthesize amino acids *de novo*, as evidenced by the incorporation of \(^{15}\)N from ammonium chloride or urea into lysine (which does not transaminate) and of \(^{13}\)C into indispensable amino acids\(^8\). It is also known that these microbially-synthesized amino acids are absorbed, in both animals and man\(^9\), and may thereby contribute to metabolic requirements\(^2\). However, in the assessment of ileal digestibility, microbially-synthesized amino acids in ileal digesta are treated as if they were undigested dietary amino acids and to that extent introduce an error into the estimate.

There is very little quantitative information on the proportion of microbial amino acids in the upper intestinal microbiota that are synthesized *de novo*. As suggested previously\(^3\), a tentative estimate can be inferred from the relative \(^{15}\)N enrichments of microbial amino acids in ileal digesta when subjects were given \(^{15}\)NH\(_4\)Cl\(^2\). In the most widespread pathway of microbial lysine synthesis the \(-\alpha-N\) of lysine is derived from aspartate, the \(\epsilon-N\) from glutamate. One would therefore expect that microbially-synthesized lysine would have a \(^{15}\)N enrichment intermediate between those of aspartate and glutamate. However, it was much lower than either, roughly one tenth of the average. In an experiment directly addressing this question\(^9\) the contributions of urea, endogenous protein and dietary protein to microbial valine leaving the ileum of growing pigs were assessed by isotope dilution following 4-day infusions of L-[\(^{15}\)N]valine (to label endogenous valine) and of [\(^{15}\)N]urea to label valine synthesized *de novo* and by transamination. These results also suggested that more than 90% of microbial valine in the ileum was derived from dietary and endogenous proteins, not synthesized *de novo*. However, these pigs were given a high-protein (19%) diet and it is not clear whether, with a less adequate protein supply, ammonia would have become a more important source of N for microbial amino acid synthesis. From this admittedly very limited evidence it appears that a large proportion of the amino acids in the protein of the upper GI microbiota are incorporated directly from the diet or from endogenous materials rather than being synthesized *de novo*.

Thus there seems to be an important quantitative difference between the upper and lower parts of the GI tract in the impact of microbial activity on amino acid utilization, presumably stemming from differences in the substrates available, in the physiology of the organ, especially its ability to absorb amino acids, in the residence time of the digesta, as well as in the numbers and composition of the resident microbiota. Despite the remaining uncertainties then it seems that the amino acid composition of ileal digesta provides the best available basis for estimating the proportion of dietary amino acids absorbed.

In this regard, the statement in the FAO/WHO/UNU technical report on protein and amino acid requirements\(^3\):

"...Furthermore, since according to circumstances the net effect of the amino acid metabolism associated with bacterial biomass can be to either remove from
or add to amino acids passing through the terminal ileum, ileal digestibility of individual amino acids is unlikely to be a more reliable measure of the systemic availability of dietary protein than faecal digestibility."

seems to overestimate the consequences of ileal microbial activity and undervalue the importance of excluding the far greater impact of the large intestine. It seems more logical to conclude that ileal digestibility, whilst not a perfect measure of net amino acid absorption, nonetheless takes us considerably closer to that ideal than amino acid digestibility over the whole gut.

The strength of these arguments in favour of ileal digestibility has led to the widespread adoption of the system in the feeding of non-ruminant animals and tables of the ileal digestibility of amino acids in a wide range of proteins used in animal feeding have been generated. However, at the present time, comparable data on ileal amino acid digestibility in humans is extremely scarce.

**Integrative bioavailability assays**

As mentioned earlier digestibility is not the only determinant of bioavailability, although it is usually the most important. Other factors, such as the chemical availability of individual amino acids, notably lysine (and other amino acids) in heat-damaged proteins, can be estimated separately. Alternatively, bioavailability may be estimated by one of several bioassays that integrate the several factors that limit the extent to which a dietary amino acid is absorbed and available for metabolism. Using a response that is sensitive to amino acid supply these approaches compare individual foods to pure amino acids, usually in the form of a slope-ratio assay. A suitable response, such as growth rate or body protein accretion or AA oxidation, of animals fed the test ingredient is related to the AA intake, and the slope of the regression line is compared with that obtained with the pure amino acid. All diets used in this assay must be first-limiting in the amino acid in question. However, despite the attractions of a method that integrates all the factors that may limit the utilization of dietary amino acids, the approach has the practical disadvantage that a separate assay is required for each amino acid, using a series of diets in which only that amino acid is limiting. It also tends to confound bioavailability with issues of nutrient imbalance, complicating the interpretation of the results. Furthermore, these assays are laborious and costly and, compared with ileal digestibility measurements, the estimated availability values generally have higher standard errors of determination.

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**References**


