Critical review of chemical and enzymatic methods for the estimation of nutritive value in roughages

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Summary - The methods used classically for estimating the nutritive value of temperate forages (grasses and legumes) have to be corrected to take into account the specificity of roughages (crop residues, numerous tropical species, trees and shrubs), which may affect the results of assays. Tropical forages have their own physical and chemical features linked to genetic factors. For example, they may contain tannins liable to form protein-tannin complexes and to inhibit certain enzymes. In general, the nutritive values of roughages cannot be predicted as accurately as those of temperate forages. For tannin-free temperate and tropical forages, in vivo organic matter digestibility can be predicted by chemical methods based on the determination of cell wall contents or lignin, but these methods are less accurate than enzymatic methods (cellulase). Digestible protein content (DCP) is closely linked to crude protein content (CP) in temperate grassland legumes and in tropical grasses. To determine DCP of tropical herbaceous or woody dicotyledons, cell wall nitrogen (ADIN: acid detergent insoluble nitrogen), which closely corresponds to indigestible nitrogen content, also has to be determined. Tannins, which are present in many dicotyledons, also modify the overall digestibility of nitrogen, but their effects depend on the types of tannins present and on other factors, and are difficult to predict. Prediction of PDI (Protéines réellement digestibles dans l'intestin grêle, i.e. digestible protein in the small intestine) requires knowing the theoretical degradability of protein in the rumen (Deg) and the true digestibility of dietary proteins (dr) in the small intestine. The methods of Deg prediction using commercially available proteases permit a useful classification of temperate and tropical forages, but must be used carefully especially in tannin-containing forages. Chemical and enzymatic methods of dr prediction (pepsin-pancreatin) have not been widely used for temperate or tropical forages. Tannin determination provides a qualitative interpretation of differences in the nutritive value between tannin-containing forages. However, none of the numerous methods tested to date can be considered as predictive of the nutritive value. This is partly due to the diversity of the phenol groups present and their methods of determination and to interactions with other digestible nutrients, which makes it difficult to predict the effects of tannins on digestion. For these forages, Van Soest's fractionation corrected for tannin content is a reliable predictive model for energy content, while NDIN (neutral detergent insoluble nitrogen) and ADIN determinations provide an initial approach to degradation mechanisms occurring in the rumen and intestine.

Introduction

Roughages are poor in digestible nutrients, either because nutrients such as nitrogen and non structural carbohydrates are present at low concentrations, or because they are poorly digestible, due to various physical or chemical factors such as lignin and polyphenols.

In warm climates, the diet of many ruminants is largely composed (between 90 and 100 %) of forages such as the aerial parts of herbaceous mature plants, crop residues after harvest, and trees and shrubs.

As in temperate zones, grasses form the basis of grazing land in humid and subhumid savannah and arid or semi-arid tropical and Mediterranean steppe. Grass straws and cereal straws (maize, sorghum, rice) are also consumed by ruminants in tropical zones.

Legumes are important in pasture ecosystems. They are generally cultivated for maintaining soil fertility and supplementing ruminants diets. Legumes are found among both cultivated and wild herbaceous forage plants, crop plants such as groundnuts and beans, and tree foliage.

Besides these two plant families, browse plants include many other dicotyledon species.
They contain more nitrogen than grasses, but usually less than legumes. Like legumes, they have variable levels of phenol contents.

Because of their botanical diversity and the pronounced differentiation of their organs and tissues, roughages show considerable morphological, anatomic and physicochemical heterogeneity (table I). This heterogeneity occurs seasonally among plant species, and among organs and tissues. Hence the «rough» characteristics of these forages is neither permanent nor uniform for any particular plant species.

Most of the methods used to predict the nutritive value of forages have been developed in temperate zones, usually for grasses and legumes, in which the range of variation of this value is less wide than in roughages. The composition of these forages usually enables microbial or enzymatic digestion to take place under optimal physiological conditions. In contrast, the specific features of roughages make their digestibility more difficult to measure, and require an adaptation of the prediction methods. In addition, the quality of the estimations is further reduced by differences in the abilities of animal species to use roughages, and the influence of ingestion conditions and other components of diet on digestibility.

After briefly reminding the main features of roughages under poor agronomic conditions, we shall review the adaptations and limits of the chemical and enzymatic methods used to predict their nutritive value.

**Table I. Rough forages compared to temperate grasses and legumes.** (DM: dry matter, tN: total nitrogen, OMD: organic matter digestibility, DCP: digestible crude protein, indCP: indigestible crude protein, S: solubility, Deg: theoretical degradability).

<table>
<thead>
<tr>
<th>Examples</th>
<th>OMD %DM</th>
<th>DCP %DM</th>
<th>indCP %N</th>
<th>S %N</th>
<th>Deg %IN</th>
<th>Condensed Tannin %DM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperate (fresh) grasses</td>
<td>50-70</td>
<td>5-19</td>
<td>4</td>
<td>20-45</td>
<td>65-90</td>
<td></td>
</tr>
<tr>
<td>Legumes</td>
<td>55-65</td>
<td>8-20</td>
<td>4-5</td>
<td>7-60</td>
<td>30-90</td>
<td>0-7</td>
</tr>
<tr>
<td>Cereal staws</td>
<td>35-45</td>
<td>0</td>
<td>3-5</td>
<td>35</td>
<td>60</td>
<td></td>
</tr>
<tr>
<td>Tropical grasses</td>
<td>45-65</td>
<td>0-12</td>
<td>4</td>
<td>20-35</td>
<td>60-65</td>
<td></td>
</tr>
<tr>
<td>(fresh+standing hay)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Browse species</td>
<td>35-65</td>
<td>0-25</td>
<td>4-25</td>
<td>7-67</td>
<td>30-90</td>
<td>0-11</td>
</tr>
<tr>
<td>Oilseed hulls</td>
<td>15-60</td>
<td>0-12</td>
<td>4-16</td>
<td>20</td>
<td>40-55</td>
<td></td>
</tr>
<tr>
<td>Grape marc</td>
<td>25</td>
<td>0-1</td>
<td>0-1</td>
<td>15</td>
<td>&gt; 6</td>
<td></td>
</tr>
</tbody>
</table>

**Differences in chemical composition between temperate forages and roughages in warm climate zones**

This comparison is justified by the fact that our approach involves adapting methods of prediction developed for temperate zone forages.

The first comparison is for grasses for which the genetic and metabolic differences are well known (C₃ versus C₄ plants).

For the other plant families, the approach is more difficult, since the chemical components that act on digestion are both more numerous and more complex. Their spatial distribution in the tissues and cells and their interactions are also important. However, these data have only been studied for a few species that make up ruminant diets. Comparisons between botanical groups are useful since they can help to extend the scope of the prediction methods.

Only the most marked differences in chemical composition between temperate and warm climate forages will be stated here.

**Differences between temperate and tropical grasses**

**Cell wall constituents**

Most of the tropical grasses are C₄ plants, which fix and concentrate CO₂ in 4-carbon organic acids in the mesophyll before photosynthesis, which takes place in the cells of the perivascular sheath. This enables these
plants to save water through more efficient use and to grow more rapidly in high ambient temperatures than temperate C₃ grasses. Conversely, at low temperatures, including in altitude or during the cool period in certain tropical regions, the growth rates of C₄ plants are lower than those of C₃ plants.

The C₄ grasses are richer in cell wall material, but also have more tissues that are poorly digestible or less readily digestible than in C₃ plants (epidermis, sclerenchyma, perivascular parenchyma and vascular bundle), as shown by Wilson et al (1983) in the genus *Panicum* (table II).

Under unrestricted water supply conditions, high temperatures accelerate ageing of tropical grasses. This can be measured in terms of stem growth compared to leaf growth, and chemical differentiation as expressed in their cell wall constituents (lignocellulose).

The poorer digestibility of tropical grasses results therefore from the fact that they are mainly C₄ type plants and that they grow at higher temperatures than temperate C₃ grasses. Other cell wall constituents contribute to the poor digestibility of the cell walls of tropical grasses, for example the silica that coats the epidermis of rice stems and leaves, and the cutin (Van Soest, 1994).

**Digestibility**

On average, wild or cultivated tropical grasses have lower dry matter digestibilities (DMD) than temperate grasses. During active plant growth the difference can reach 10 to 15 points (55 to 65 % versus 60 to 80 %) (INRA, 1989). Over 6 to 9 months of the year in dry tropical zones when tropical forage grasses are not growing, steppe annual grasses and savannah perennial grasses have digestibilities comparable to those of rice and sorghum straws (DMD close to 50 %) and those of wheat and barley straws (DMD close to 40 %),

**Table II.** Leaf attributes of C₄, intermediate (C₃/C₄) and C₃ photosynthetic types of *Panicum* species (According to Wilson et al, 1983).

<table>
<thead>
<tr>
<th>Type of Plant</th>
<th>Dry matter digestibility (%)</th>
<th>Cell wall content (%)</th>
<th>Tissue proportion in leaf cross-section</th>
<th>MES</th>
<th>BS</th>
<th>VAS</th>
<th>EPI</th>
<th>SCL</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Panicum</em> a</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C₄</td>
<td>69</td>
<td>50</td>
<td>43 20 8 27 1.7</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C₃/C₄</td>
<td>70</td>
<td>42</td>
<td>48 18 6 26 1.7</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C₃</td>
<td>76</td>
<td>33</td>
<td>66 10 3 22 0.5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

aMeans of 18(C₄), 3(C₃/C₄) and 6(C₃) Species; bMES, mesophyll; BS, bundle sheath; VAS, vascular bundle; EPI, epidermis; SCL, sclerenchyma.

**Table III.** Typical concentrations of carbohydrates in temperate legumes, and cool and warm-season grasses (adapted from Van Soest, 1982, according to Moore and Hatfield, 1994).

<table>
<thead>
<tr>
<th>Category</th>
<th>Temperate Legumes</th>
<th>Cool-Season Grasses</th>
<th>Warm-Season Grasses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonstructural Carbohydrates (g.kg⁻¹ DM)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Soluble sugars</td>
<td>20-50</td>
<td>30-60</td>
<td>10-50</td>
</tr>
<tr>
<td>Starch</td>
<td>10-110</td>
<td>0-20</td>
<td>10-50</td>
</tr>
<tr>
<td>Fructans</td>
<td>-</td>
<td>30-100</td>
<td>-</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Structural carbohydrates (g.kg⁻¹ DM)</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Cellulose</td>
<td>200-300</td>
<td>150-450</td>
<td>220-400</td>
</tr>
<tr>
<td>Hemicellulose</td>
<td>40-170</td>
<td>120-170</td>
<td>250-400</td>
</tr>
<tr>
<td>Pectin</td>
<td>40-120</td>
<td>10-20</td>
<td>10-20</td>
</tr>
</tbody>
</table>
respectively (Richard et al, 1989).

Cell contents

-Cyttoplasmic carbohydrates

The levels of water soluble carbohydrates such as glucose, fructose and sucrose in tropical grasses are generally low compared to those in temperate zone grasses (3 to 5 % and 6 to 10 % DM respectively). Their polysaccharide reserve is starch, which is mainly stored in the leaves (2 to 5 % DM), whereas temperate meadow grasses mainly store fructosanes in stems and sheaths (2 to 10 % DM), (table III).

-Nitrogen

During early growth, tropical grasses contain slightly less nitrogen than temperate grasses (2.5 to 4.0 % DM), but in tropical grasses, nitrogen content decreases more rapidly with age of plant. At the end of the growth cycle, the nitrogen content may be very low (0.5 % DM). This difference relative to temperate grasses partly derives from the faster growth of tropical zone grasses, with an increase in the proportion of stems and the senescence of leaves, together with generally low levels of soil nitrogen.

The nitrogen of cell walls represents a small part of total nitrogen. Plant proteins can be mainly split into two groups according to their location in the plant tissue: the water-soluble proteins of the cytoplasm and the water-insoluble proteins of the inner membranes of the chloroplasts. Wherever they are located, these proteins are not accessible until the cell membrane or cell wall polysaccharides have been hydrolysed. The proportion of proteins released therefore depends on the accessibility of the chlorophyll tissue to the rumen micro-organisms.

The solubility of nitrogen in a phosphate buffer at pH 6.9 (Véritable and Demarquilly, 1978) is on average 0.30 for temperate grasses versus 0.24 for tropical grasses (Richard, 1987). These means hide very large differences between organs in tropical grasses through organ differentiation and selection by animals. Aii and Stobbs (1980) have found values ranging between 0.15 and 0.43 according to the species, the organ and its growth stage, i.e. an average of 0.28, but they used a slightly different buffer (Burroughs mineral solution).

Soluble proteins are distributed differently in tropical C_3 and temperate C_4 grasses. Ribulose 1,5-diphosphate carboxylase makes up 30 to 80 % of the soluble proteins in C_3 grasses versus 10-20 % in tropical C_4 grasses (Huffaker, 1982). The different distribution of these proteins in temperate grasses (in the mesophyll) and tropical grasses (in the perivascular sheath) (Wilson, 1994) may partly explain the differences in the nitrogen

![LEAVES C₃](image1)

![LEAVES C₄](image2)

Figure 1. Anatomical comparison between C₃ leaves and C₄ leaves (according to Alberts et al, 1990).
degradability of these forages (Redfearn et al., 1995) (figure 1). Moreover the chloroplast membrane proteins of temperate grasses are poorly soluble as they are complexed with chlorophyll, but those of tropical grasses are even less soluble.

Comparison of tropical dicotyledons and temperate grasses and legumes

Cell wall constituents
Compared to grasses, dicotyledons are characterised by average total cell wall contents (Neutral Detergent Fibre, abbreviated NDF) that are lower but which vary widely among species. In contrast, lignin contents are often higher, but at equal lignin levels, the dicotyledons are more digestible than the grasses. Legumes for instance are less rich in cell wall material, and the composition and structure of their lignin are different resulting in a lower effect on cell wall digestibility (figure 2). In grasses the lignin contains more ester bonds and more readily hydrolysable methoxyl groups than that of dicotyledons, which have more ether bonds (Lapierre et al., 1989).

Lignin levels in tropical herbaceous dicotyledons are slightly higher (7 to 14 % DM) than in the usually cultivated legumes including lucerne and clover (7 to 10 %). Lignin levels in shrub forages vary widely (on average 13 ± 6.5 % DM on 700 samples of tropical tree foliage). Given the complexity of lignins, the poor accuracy of ADL (Acid Detergent Lignin) determinations and possible contamination (Maillard reactions, tannins, etc...), the highest values (up to 40 % DM) are to be considered with caution.

Nitrogen
In dry periods, when no young grass is growing, herbivores find their nitrogen supply in herbaceous or shrubby dicotyledons. Whether green or dry, these offer organs, i.e. chlorophyll stems, folioles, inflorescences, fruit and seeds, which are richer in nitrogen (1 to 5 % DM) than grass straws. To an even larger extent than in grasses, this nitrogen is variably distributed between cell walls and cytoplasm and for the cytoplasmic fraction between soluble and insoluble proteins.

In certain dicotyledons, especially in tropical woody species, part of the proteins binds to tannins during ingestive mastication to form complexes of varying stability, making this nitrogen partly unavailable to the animal. Though relatively abundant, the nitrogen in these forages thus exhibits varying digestibility due to physical and chemical factors. The methods for predicting the nitrogen value must therefore directly take into account nitrogen unavailability in the rumen or intestine, or estimate it by characterising the physical and chemical factors that are responsible.

Antinutritional factors and toxic constituents
Many substances synthesised by plants and released during mastication serve to protect them against herbivores by making the plant material less appetising (e.g., astringent effects of tannins), or even toxic. Polyphenols (including tannins) are the most common of these substances. Legumes in tropical zones contain other toxic substances such as lectins, protease inhibitors, nonprotein amines, gums, alkaloids and some amino acids (i.e. mimosine in Leucaena leucocephala). This toxicity depends on the amounts ingested and their metabolism and restricts the use of certain forage species, even though they can have high energy and/or nitrogen values (Manidool, 1983; D'Mello, 1992).
Tannins are complex phenolic polymers capable of precipitating proteins. Tannins are especially abundant in certain legumes (e.g., sweet clover, sainfoin, desmodium) and in many tree and shrub species. They are classified into condensed tannins and hydrolysable tannins.

Condensed tannins are polymers with a flavone nucleus and high molecular weights (1000 to 30,000). They are present in the vacuoles of a network of specialised cells situated under the epidermis of the leaves and stems of some temperate herbaceous, tropical herbaceous and shrub legumes, and in the leaves of forage shrubs in semi-arid zones. As soon as the cell structures are destroyed by mastication, the condensed tannins can bind to soluble proteins such as ribulose 1,5-diphosphate carboxylase, by hydrogen linking in particular (Jones and Mangan, 1977). These complexes are stable at the pH of the rumen, but dissociate to some extent at the acid pH in the abomasum and at the alkaline pH in parts of the small intestine, releasing some proteins, as shown in figure 3. They may therefore be beneficial to the animal by reducing protein degradation in the rumen without adversely affecting true protein digestibility in the intestine. This beneficial action will persist as long as the condensed tannin content does not exceed 4% of the dry matter (Barry et al., 1986; Mangan, 1988). However, at higher levels, the quantity of tannins present may exceed the amounts of protein available for binding and may then inhibit some microbial or digestive enzymes, or bind to cellulose, protecting it partially from degradation in the rumen.

The hydrolysable tannins have lower molecular weights (500 to 3000) and precipitate proteins less readily than condensed tannins. They are partly hydrolysed in the digestive tract, and can be toxic if ingested in large amounts. A feed content of 20% will cause acute poisoning in steers and sheep (Reed, 1995). Hydrolysable tannins can reduce cell wall degradation in the rumen and be hydrolysed in the intestine, releasing substances that are toxic for the liver and kidneys (McLeod, 1974).

Total phenolic substances make up only 1.5 to 2.5% DM in some meadow grasses in temperate regions, but much more in some temperate dicotyledons (3 to 6%; Sechovic, 1990) and up to 10% or more in some tropical forage shrubs (Ahn et al., 1989).

Tannins modify nitrogen and organic matter digestibility at all stages of digestion. It is therefore important to determine their content. However, the results of the

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**Figure 3.** Condensed tannins and protein protection in the rumen (according to D'Mello, 1992). The protein-condensed tanin complex escapes fermentation in the rumen where the pH ranges from 5 to 7 but dissociates on exposure to gastric (pH 2.5 - 3) and pancreatic (pH 8 - 9) secretion.
determinations are difficult to interpret due to the numerous factors involved in the action of tannins.

**Prediction of energy value**

The prediction of the energy value, expressed in net energy (NE) or metabolizable energy (ME) usually consists in estimating the organic matter digestibility (OMD), which is the main factor of variation in the energy value.

The methods used to predict OMD basically consists in defining mathematical relationships to link in vivo digestibility with the results of chemical or enzymatic laboratory tests.

The in vivo digestibility of roughages depends on their specific characteristics, but also on the conditions in which they are used. Forages poor in fermentable nitrogen and in certain minerals such as P, Mg, S and Cu have to be supplemented so that the rumen can function correctly and these forages reach their full «potential» digestibility. Roughages are usually associated with other feeds in animal diets, and numerous interactions can therefore take place. Forages containing tannins have in vivo digestibilities that depend on their proportion in the diet. Moreover, samples taken for analysis usually correspond to the forage «offered», whereas in vivo digestibility correspond to the forage «ingested», whose chemical composition is different due to the choice made by the animal. This difference is more marked for roughages, and depends on the amount of feed on offer (refusal rate) by the experimenter or farmer. This partly accounts for the lower accuracy of prediction of the digestibility of these forages compared to those of temperate zones using laboratory methods.

In contrast, for some forages, especially those rich in tannins and other often badly known secondary compounds, the true digestibility of the cytoplasmic constituents is not total, but highly variable according to the amounts and nature of these secondary compounds. In addition, the digestibility of the cell walls of these forages is also affected by tannins.

The OMD of these forages does not therefore only depend on their indigestible cell wall content, which is highly variable, but also on their concentrations in truly indigestible cytoplasmic constituents.

**Prediction of OMD by chemical methods**

**Nitrogen**

Crude protein content (CP) is often a good predictor of OMD in both temperate and tropical grasses, since it diminishes as the plants growth and age, and also varies inversely with the indigestible cell wall fraction. However, OMD predictions are only accurate if they are carried out on individual plant species, production cycles or seasons, and soil types. The nitrogen content of grasses depends directly on the nitrogen supply from the soil they are growing on, which can be a limiting factor in tropical zones. Wide variations in CP for the same digestibility can be observed within the same species (Guérin, 1987).

Conversely, for most dicotyledons in tropical zones, total nitrogen is a poor predictor of OMD (Guérin, 1987).
**Crude fibre**

Cell wall indigestibility is mainly due to the lignin content. However, the Weende crude fibre values in temperate forages increase according to the cell wall content, at least in the course of any given growth cycle. Therefore crude fibre is generally a good criterion of cell wall indigestibility for a particular plant. However, for the same crude fibre content, plants can exhibit different digestibilities according to the plant species and, even, to the regrowth cycle number. Crude fibre content (CF) will therefore permit digestibility to be accurately predicted only if equations specific to each species are used. Prediction is improved if nitrogen content is included in the prediction equation (Andrieu and Weiss, 1981).

However, determination of CP and CF does not solve the problem of predicting the digestibility of mixed forage or forage of permanent pasture, since specific equations would be required for each case, which is impracticable.

For hay harvested at the end of its first cycle, straws, and old tropical forages in which crude fibre varies relatively little, it is preferable to use lignin content to estimate OMD.

**Cell wall constituents**

Van Soest's fractionation (Van Soest and Wine, 1967) makes it possible to measure the total cell wall content (neutral detergent fiber, NDF), lignocellulose content (acid detergent fiber, ADF) and lignin content (acid detergent lignin, ADL) of forages. In particular, ADF determination allows the evaluation of practically all the cellulose and lignin. For a given species and organ, the relationship between ADF and total crude fibre is usually high, which allows forages analysed by different methods to be compared.

This fractionation also permits an estimation of the degree of cell wall lignification from the ADL/NDF and ADL/ADF ratios. This is one advantage of Van Soest's method over that of Wende, since these ratios vary widely from one browse species to another.

To predict the digestibility of forages, total cell wall content or NDF is a less accurate predictor of OMD than ADF or crude fibre. Lignin content is the variable most closely linked to in vivo digestibility, at least when correctly determined. This is even more obvious for predicting OMD of forages from different species or cycles but belonging to the same botanical families. It is necessary to establish separate relations, at least for grasses and legumes. For the same lignin content, legumes have less indigestible cell wall material and higher digestibility than grasses (Van Soest, 1964; Demarquilly and Andrieu, 1987). The parameters NDF, ADF, ADL have been associated in models which take into account cell wall content and lignification. The summative equations of Goering and Van Soest (1970), for example, offer the advantage of being both accurate and applicable to wide ranges of forages:

\[
DDM = 0.98 \times S + NDF \times [147.3 - 78.9 \log_{10} ([L/ADF] \times 100)] - M
\]

where: DDM = digestibility of dry matter in percent; S = cell content of average digestibility 98 %; L = acid detergent Klason lignin, M = estimation of faecal metabolic losses (12.9 units on average for sheep).

However, this analytical procedure is still imperfect for several reasons:

(i) In legumes, the neutral detergent solution used to determine NDF solubilises pectic substances present in stems at levels between 11 and 22 % (Hatfield, 1992). The cell wall content is thus underestimated. This is of little consequence in grasses, in which the levels of these pectic substances are very low (1 to 2.5 %). For the purposes of predicting digestibility, this underestimation is of limited consequences since the pectic substances are highly digestible.

(ii) Cell wall content (NDF and ADF) can be overestimated if a large proportion of nitrogen remains bound to the cell walls, which is the case in temperate zone forages for legumes, young grass leaves, (Van Soest and Robertson, 1980), and numerous tropical dicotyledons (Guérin et al, 1989). In addition, tropical dicotyledons can contain large amounts of cutins and tannins that form insoluble complexes with neutral and acid detergent solutions (figure 4). The presence of varying quantities of tannins can lead to an overestimation of lignocellulose content (Makkar et al, 1995). Van Soest's fractionation is not suitable for forages containing tannins, though more recently a summative equation corrected for tannin content has been proposed (Conklin et al, 1987). Although prediction is improved, it is still often poor. A specific equation would be needed for each plant species and even each organ, given the diversity of the nature of the tannins and their
levels according to the plant species, organ and growth stage.

(iii) The results of ADL determinations can be adversely modified by errors caused by products of Maillard reactions (Van Soest and Mason, 1991) or by tannins (Van Soest, 1994). These errors can be important, and ADL is certainly overestimated, like in shrub forages (ADL 40 % of DM).

Acid detergent insoluble nitrogen (ADIN) is essentially made up of nitrogen of extensin and enzymes in the primary cell wall (peroxidases). It represents 7 to 9 % of total nitrogen. We can add the nitrogen implicated in the Maillard reactions, and that of the proteins complexed by the tannins. ADIN levels above 9 % of total nitrogen thus indicate indigestibility factors.

Regardless of its origin, ADIN indicates the general level of indigestibility factors that are often especially important in roughages. This explains why ADIN is a more efficient predictor of OMD than total nitrogen or ADF for tropical zones forages that are usually of varying composition (Guérin et al, 1988 ; Silva-Colomer et al, 1989).

Van Soest's fractionation does not lead to the isolation of perfectly pure chemical entities, but it offers the advantage of being rapid, reproducible and can be automated for samples poor in proteins, tannins, lipids and/or starch. Other more accurate methods (method of Jarrige, 1961, determination of phenolic acids) may be used, but these are often time and money consuming. However, whatever the fractionation method used, the digestible and indigestible fractions are not separated as they are by in vitro, in situ and enzymatic methods.

**Prediction of OMD by enzymatic methods**

The specificity and advantage of ruminal digestion are mainly linked to the cellulytic activity of the microbial flora. In the 1960's, Donefer et al (1963), Jarrige and Thivend (1969) used commercially available cellulytic enzyme preparations, often extracted from fungi, to reproduce this activity. Since then, many enzymatic methods have been proposed to predict feed digestibility. They differ in the nature of the enzyme preparations and

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**Figure 4.** Digestibility of neutral detergent fiber (NDF) fed to mule deer and white-tailed deer as a function of the fiber's lignin and cutin content (according to Robbins et al, 1987).

High-phenolic forages are (A) red alder leaves (B) red-osier dogwood leaves, and (C) fireweed flowers.
whether a pre-treatment is necessary or not (chemical or enzyme) (Aufrère and Michalet-Doreau, 1990). These methods are widely used for forages, and have also been applied to by-products, concentrate and mixed feeds produced by agro-food industry. For various types of forages, in temperate or tropical zones, prediction is higher than with chemical methods (table IV) and comparable to that obtained in vitro (Terry et al, 1978; Aufrère and Demarquilly, 1989; Navaratne et al, 1990). In addition, cellulase methods can be used for mixtures and permanent pastures. They also offer the advantage of measuring improvement in the digestibility of hays and straws treated with ammonia, unlike the usual chemical methods. For these forages, which have low cell contents, a simplified method can be applied, with no pre-treatment (Rexen, 1977).

For straws, however, different equations must be used according to whether they are treated or not (Table IV).

For forages containing tannins, OMD prediction is poor when cellulolytic enzymes are used. Two possible reasons can be suggested for this:

(i) The enzymes are used at pH values different from that prevailing in the rumen, enabling possible release of tannins bound to proteins. The diagram of Mangan (fig 3) shows the varying degrees of reversibility of the protein complexation according to the pH of the digestive compartment. Tannin-protein bonds favoured at the rumen pH (5 to 7) dissociate at pH values outside the range 4 to 7. In the pepsin-cellulase method, for example, pepsin pre-treatment (pH approx. 3) could cleave protein-tannin complexes and lead to an overestimation of OMD for these forages.

(ii) Some kinds of tannins might have inhibiting effects on the enzyme activity, especially that of cellulases (Smart et al, 1961; Mandels and Reese, 1963).

However, for tree foliage species with or without tannins, pepsin-cellulase digestibility is closer to in vitro digestibility than chemical criteria in general, though its accuracy is very poor (about 6 points) (table V). Both enzymatic and chemical methods lead to satisfactory accuracy only if the different species are taken separately.

Besides their ease of use, and their favou-

Table IV. Prediction of organic matter digestibility (percentage) by using chemical and enzymatic methods.

<table>
<thead>
<tr>
<th>Nature of samples</th>
<th>n</th>
<th>Method used*</th>
<th>RSD</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperate forages</td>
<td>10</td>
<td>crude fibre</td>
<td>2 to 6</td>
<td>review of Giger-Reverdin, 1995</td>
</tr>
<tr>
<td>Temperate forages</td>
<td>20</td>
<td>crude fibre, MAT</td>
<td>2 (species)</td>
<td>Andrieu and Weiss, 1981</td>
</tr>
<tr>
<td>Temperate forages</td>
<td>20</td>
<td>ADF</td>
<td>1.6 to 5.3</td>
<td>review of Giger-Reverdin, 1995</td>
</tr>
<tr>
<td>Temperate forages</td>
<td>20</td>
<td>pepsin-cellulase</td>
<td>1.4 to 3.5</td>
<td>review of Aufrère and Michalet-Doreau, 1990</td>
</tr>
<tr>
<td>Tropical forages</td>
<td>45</td>
<td>pepsin-cellulase</td>
<td>2.7</td>
<td>Goto and Minson, 1977</td>
</tr>
<tr>
<td>Tropical forages</td>
<td>30</td>
<td>ADL</td>
<td>6.9</td>
<td>Kronauer and Bickel, 1981</td>
</tr>
<tr>
<td>Tropical forages</td>
<td>20</td>
<td>N, ADF</td>
<td>2.7 to 3.8</td>
<td>Guerin et al, 1989</td>
</tr>
<tr>
<td>Tropical hays</td>
<td>20</td>
<td>CB, MAT</td>
<td>7.2</td>
<td>Zoungrana, 1995</td>
</tr>
<tr>
<td>Tropical hays</td>
<td>20</td>
<td>pepsin-cellulase</td>
<td>4.5</td>
<td>idem</td>
</tr>
<tr>
<td>Untreated straw</td>
<td>36</td>
<td>ADL</td>
<td>2.3</td>
<td>Demarquilly and Andrieu, 1987</td>
</tr>
<tr>
<td>Untreated straw</td>
<td>36</td>
<td>pepsin-cellulase</td>
<td>2.7</td>
<td>idem</td>
</tr>
<tr>
<td>Ammonia treated straw</td>
<td>55</td>
<td>NDF, MAT, Phenols</td>
<td>2.8</td>
<td>Guillermin et al, 1988</td>
</tr>
<tr>
<td>Ammonia treated straw</td>
<td>55</td>
<td>pepsin-cellulase</td>
<td>4.1</td>
<td>idem</td>
</tr>
</tbody>
</table>

* MAT: Matière Azotée Totale; ADF: Acid Detergent Fiber; ADL: Acid Detregent Lignin; N: Nitrogen; CB: Cellulose Brute; NDF: Neutral Detergent Fiber.
rable repeatability and reproducibility, these enzymatic methods allow a satisfactory classification of forages and accurate prediction of in vivo digestibility except for forages containing tannins for which adaptations are necessary.

**Nitrogen value prediction**

As shown in Table I, tropical grasses and cereal straws are less rich in total nitrogen than temperate ones. Their digestible nitrogen and rumen nitrogen degradability are also lower. Some shrub and herbaceous legumes behave like lucernes or clovers, while others are poorly digestible. These latter can contain up to 10% condensed tannins, and often a large proportion of the nitrogen is found in the faeces.

The apparent digestibility of nitrogen was long used to measure the nitrogen value of feeds. However, most of the current methods of evaluation of the nitrogen value recommend taking into account the quantity of proteins absorbed in the small intestine, which corresponds to the quantity of amino acids available to the animal. This implies knowing:
(i) The degradability of feed proteins in the rumen, and
(ii) The digestibility of feed proteins in the small intestine.

**Table V.** Use of chemical and pepsin-cellulase methods to predict in vitro organic matter digestibility (in vitro CMD) of browse leaves (Steingass and Arbelot, 1994).

<table>
<thead>
<tr>
<th>Species</th>
<th>n</th>
<th>In vitro digestibility (gaz-test method) mean value</th>
<th>laboratory methods</th>
<th>R²</th>
<th>RSD</th>
</tr>
</thead>
<tbody>
<tr>
<td>All species (90)</td>
<td>438</td>
<td>48 ± 12</td>
<td>ADF</td>
<td>0.43</td>
<td>8.9</td>
</tr>
<tr>
<td></td>
<td>438</td>
<td></td>
<td>pepsin-cellulase</td>
<td>0.66</td>
<td>6.4</td>
</tr>
<tr>
<td><em>Pterocarpus erinaceus</em></td>
<td>15</td>
<td>46 ± 8</td>
<td>ADIN</td>
<td>0.81</td>
<td>3.8</td>
</tr>
<tr>
<td></td>
<td>15</td>
<td></td>
<td>pepsin-cellulase</td>
<td>0.92</td>
<td>2.6</td>
</tr>
<tr>
<td><em>Spondias mombin</em></td>
<td>10</td>
<td>47 ± 4</td>
<td>ADF</td>
<td>0.70</td>
<td>2.5</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td></td>
<td>pepsin-cellulase</td>
<td>0.75</td>
<td>2.3</td>
</tr>
<tr>
<td><em>Daniellia oliveri</em></td>
<td>15</td>
<td>42 ± 8</td>
<td>N</td>
<td>0.96</td>
<td>1.7</td>
</tr>
<tr>
<td></td>
<td>15</td>
<td></td>
<td>pepsin-cellulase</td>
<td>0.80</td>
<td>3.6 (n.s)</td>
</tr>
<tr>
<td><em>Acacia nilotica</em></td>
<td>13</td>
<td>52 ± 6</td>
<td>ADIN</td>
<td>0.65</td>
<td>3.9</td>
</tr>
<tr>
<td></td>
<td>13</td>
<td></td>
<td>pepsin-cellulase</td>
<td>0.60</td>
<td>4.2</td>
</tr>
</tbody>
</table>

* ADF: Acid Detergent Fiber; ADIN: Acid Detergent Insoluble Nitrogen; N: Nitrogen.

Figure 5. Relationship between apparent digestibility of nitrogen and nitrogen bound in acid detergent insoluble fiber (ADIN) (according to Van Soest, 1994). Values, obtained from 80 digestion trials with sheep or cattle, are expressed as a percentage of total feed nitrogen. The hays and silages without heat damage (n=28) give a different regression line from forages (n=52) that were heated.
The true digestibility of forage crude protein is generally very high (about 92-93 % for green forages). The apparent digestibility is lower, because the faeces contain nitrogen-containing substances of endogenous or microbial origin, whose quantities are proportional to the dry matter intake. For this reason, digestible crude protein content (DCP) is closely linked to the total nitrogen content (CP = N x 6.25) of the forage as expressed by the relation:

\[ \text{DCP} = \text{CP} - \text{inDCP}, \]

where inDCP is indigestible crude protein, mainly of endogenous origin.

The inDCP varies slightly, representing 40 to 50g per kg of DM (Demarquilly et al., 1981). The CP in feed that is truly indigestible corresponds essentially to nitrogen linked with ADF (i.e. under 9 %). The equations are the same for temperate and tropical grasses (Chenost, 1975; Minson, 1982; Richard, 1987).

Conversely, many dicotyledon species, especially woody species, contain proteins that are not readily available since they are enclosed in highly lignified tissues (Koné, 1987; Koné et al., 1989), or complexed with tannins or by Maillard reactions. In all these cases, ADIN is a reliable approach to true indigestible nitrogen and provides a correction to the standard prediction from CP for heated forages (figure 5).

The presence of tannins is also a cause of nitrogen indigestibility, and various scientists have endeavoured to develop methods to predict nitrogen digestibility from the determinations of different phenol fractions (Zimmer, 1993), in particular those that are separated through Van Soest’s fractionation: soluble condensed tannins, those dissolved in neutral detergent solution (NDS) and those insoluble either because they precipitate out in the neutral detergent solution or because they remain bound to cell walls (figure 6). The NDF/ADF - ADF/NDF sequence provides a first approach to tannin content (Van Soest et al., 1987). The tannins obtained by this method are those linked to the cell walls and proteins, and have marked effects on digestibility.

Although condensed tannins have been shown to play a major role in digestibility, it is still difficult to deduce methods for predicting...
the nitrogen value, or even to classify species in terms of condensed tannin effects. However, condensed tannin content is useful for comparing agroforestry cultivars, since the effects of increasing amounts of condensed tannins has been clearly shown in vivo.

Tannins can also be studied indirectly by their precipitating action (Grillet et Villeneuve, 1994). The methods most often applied use bovine serum albumin. This method will not separate the beneficial effect of protection against rumen microbes from the negative effects on overall digestibility. It is a simple method for detecting tannins which also act on ingestion by their astringent properties. However, this method, like the chemical determination of tannins, will not predict nitrogen digestibility. Even so, with the help of this method, Robbins et al (1987) set up a model to correct the standard DCP prediction.

Finally, ammonia-treated hays and straws have a noticeably higher indCP content than the corresponding untreated forages. The faeces of the animals fed with these forages contain, in addition to endogenous and microbial nitrogen, (i) nitrogen from ammonia bound to cell walls and not released in either the rumen or the intestine, (ii) soluble dietary nitrogen derived from NH₃ treatment which has not been used by microbes, and (iii) nitrogen from Maillard reactions between nitrogenous constituents and carbohydrates, especially in wet treated hays (Demarquilly and Andrieu, 1987).

### Prediction of rumen degradability (Deg)

In the new nitrogenous systems, it is necessary to know the degradability of nitrogen in the rumen (Deg), which depends on the rate of degradation of the forage in the rumen and the outflow rate of the particles.

### By chemical methods (solubility)

The solubility of proteins in solvents has been abundantly used to estimate protein degradability (Henderickx and Martin, 1963; Johnson, 1976; Wohlt et al, 1976; Crooker et al, 1978; Waldo and Goering, 1979; Vérété et Demarquilly, 1978; Stern and Satter, 1982). However, this solubility varies with the nature of the solvent, its pH and incubation temperature. The first PDI system (protein digested in the small intestine) (INRA, 1978) relied on measuring nitrogen solubility by the method of Durand (Vérété and Demarquilly, 1978). This measurement characterises the immediately soluble nitrogen fraction. There is a close relationship between in situ degradability and solubility within any given group of feeds (Aufrère et al, 1991), whereas the relations are less clear among different

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**Table VI. Theoretical fractionation of nitrogen feedstuffs (according to Krishnamoorthy et al, 1995).**

<table>
<thead>
<tr>
<th>Nitrogen Fraction</th>
<th>Nutritional Property</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buffer Soluble N</td>
<td>Rapidly degraded in the rumen</td>
</tr>
<tr>
<td>(borate-phosphate buffer, pH 8)</td>
<td></td>
</tr>
<tr>
<td>BIN</td>
<td>Slowly degraded in the rumen</td>
</tr>
<tr>
<td>in borate phosphate buffer, pH 8</td>
<td></td>
</tr>
<tr>
<td>PSN</td>
<td>Total N Degraded in the rumen</td>
</tr>
<tr>
<td>(extracted from Streptomyces griseus)</td>
<td></td>
</tr>
<tr>
<td>in borate phosphate buffer, pH 8</td>
<td></td>
</tr>
<tr>
<td>PIN</td>
<td>Rumen undegraded N</td>
</tr>
<tr>
<td>PIN - ADIN=</td>
<td>Rumen undegraded but digestible in intestine N</td>
</tr>
<tr>
<td>ADIN =</td>
<td>Indigestible N</td>
</tr>
</tbody>
</table>

PSN: Protease Soluble N; BIN: Buffer Insoluble N; PIN: Protease Insoluble N; ADIN: Acid Detergent Insoluble N.
groups and in mixed feeds.

These solubility measurements have been gradually abandoned in favour of enzymatic methods, since they fail to take into account the slowly degradable protein fraction in the rumen.

By enzymatic methods
These methods use poorly specific commercial proteases, of bacterial, plant, animal or fungal origin (Aufrère and Michalet-Doreau, 1990). They have mainly been tested on concentrates (Pichard and Van Soest, 1977; Chamberlain and Thomas, 1979; Poos-Floyd et al, 1985; Aufrère et al, 1991; Susmel et al, 1989) and temperate forages (Aufrère et al, 1989), but rarely on roughages. They are inexpensive compared to the reference methods (measurements in nylon bags), and offer the following advantages: there is neither microbial contamination of bag residues, nor loss of particles through the bag mesh.

Although set values of Deg were proposed in 1988 for each major type of forage, the nitrogen degradability of green forages in temperate countries varies according to the growth stage, species and season (LeGoffe et al, 1993; Amrane and Michalet-Doreau 1993). The Deg can be predicted accurately from the nitrogen content and by measuring the quantity of protease solubilised after one hour (Aufrère and Cartailler, 1988).

The application of these methods to forages containing tannins raises certain problems.
(i) In the method selected by Aufrère et al (1991), the commercial protease, of bacterial origin, is used in a borate-phosphate buffer at pH 8. At this pH the enzyme activity is optimum, but the tannin-protein complexes are unstable, which may cause an overestimation of the true enzyme degradability. Other methods use lower pH values between 5.5 and 7 (Assoumani et al, 1992), but have seldom been tested on forages.
(ii) In in vitro studies, some authors (Tagari et al, 1965; Oh and Hoff, 1988) have shown an inhibiting effect of tannins on the activities of proteolytic enzymes, which might distort the results obtained with enzyme methods using proteases.

However, for tree foliage rich in tannins, the enzymatic method enables the accurate ranking of species and organs and can be used for screening (Zoungrana, 1995).

Prediction of intestinal digestibility (dr)
Determination of true digestibility in the intestine (dr) requires knowing the quantity of dietary nitrogen that has escaped degradation in the rumen and is truly indigestible in the small intestine (PIAnd).

In the French system, these PIAnd values are estimated from indCP values minus faecal nitrogen of microbial and endogenous origin. In other systems, PIAnd values are estimated from nitrogen remaining in the bags after a long period in the rumen or from ADIN considered as indigestible. Enzymatic methods (pepsin-pancreatin) applied to samples after rumen digestion provide a good estimation of intestinal digestibility for concentrate feeds (Antoniewicz et al, 1992; Calsamiglia and Stern, 1995). In contrast, for forages, Van Straalen et al (1993) found a lower indigestible protein content with the mobile bag method than with the pepsin-pancreatin method.

If little work has been done on dr measurement for temperate forages, even less has been done on tropical forages (except for Mgheni et al, 1994), even though the dr is certainly highly variable for forages rich in tannins since the tannin-protein bonds can be partly hydrolysed in the intestine.

The methods that have become most commonly used for nitrogen degradability measurements in roughages are those that take into account cell wall constituents as indicated by the very accurate prediction (RSD = 3.6, $R^2 = 0.98$) obtained between Deg and ADIN on tropical shrubs by Fall-Tourd and Michalet-Doreau, (1995). Recently, Krisnamoorthy et al (1995) have presented a more elaborate model to predict nitrogen degradability of tropical by-products, taking into account both degradation in a protease at pH 8 and nitrogen bound to NDF (NDIN) and ADF residues (table VI). This model, which is still being developed, takes into account the degradability of nitrogen, limited by whether cell wall tannins or heating.

Conclusion
The nutritive value of roughages (tropical forages, crop residues and especially tree foliage) is often predicted with less accuracy than that of temperate forages, even if its variability is as wide. This is partly due to the
botanical heterogeneity of the samples. Moreover, the methods for predicting the nutritive value have almost all been developed using samples of temperate zone forages. They are often applied to tropical forages without making the necessary adaptations to allow for the specific physical and chemical features of plants growing in tropical climates.

Van Soest's fractionation, despite its imperfections, is still a useful and global method for measuring the energy value, while NDIN and ADIN offer a first approach to the mechanisms of nitrogen degradation in the rumen and intestine.

Enzymatic methods can find important applications in laboratory analysis to predict the nutritive value of roughages. They are inexpensive relative to the reference in vivo methods, and are also faster and easier to implement routinely. They are generally accurate, repeatable and reproducible, but need to be calibrated against samples measured in vivo. When applied to tropical zone forages containing neither tannins nor anti-nutritional substances, the accuracy of the prediction of energy value is close to that obtained for temperate zone forages. The prediction of the nitrogen value with proteolytic enzymes has been less studied for temperate and tropical forages. The use of commercially available enzymes at their optimal pH rather than at rumen pH raises problems. The values obtained will be distorted if tannin-protein complexes are hydrolysed, and the commercial enzymes may be inhibited to some extent by tannins. At present it is difficult to predict the nutritive value of forages, especially woody forages, on the basis of their tannin content. In the future, reliable methods for predicting the nutritive value of forages taking into account the tannin content have thus to be developed.

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