Studies on a brittle stem mutant of rice, *Oryza sativa* L.; characterization of lignin fractions, associated phenolic acids and polysaccharides from rice stem

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SUMMARY  
Lignin fractions, associated p-coumaric and ferulic acids and polysaccharides have been characterized in the straw and in three lignin fractions isolated from the rice cultivar "Balilla 28" and one corresponding brittle-stem mutant. Significant differences were found between the contents in Klason-, acetyl-bromide and acid-insoluble lignins, in hemicellulosic and cellulosic polysaccharides, in esters and ethers of phenolic acids, in isolation yields of lignin fractions, and in polysidase hydrolysability. Qualitative differences were also observed in the monomeric composition of polysaccharides and lignin preparations. Thus it is not possible to suggest a direct correlation between variations in chemical composition and the brittle-stem character of the mutant. These results are discussed in relation to correlated variations of chemical composition of plant cell walls and heterogeneity of monomeric composition of lignin fractions.

Additional key words: Cell-wall mutation, p-coumaric acid, ferulic acid, hemicelluloses, celluloses, straw.

RÉSUMÉ  
Etude d’un mutant à tige cassante de riz, *Oryza sativa* L.; préparation de fractions de lignine et caractérisation des acides phénoliques et polysides associés.

On a caractérisé par analyses chimiques des fractions de lignine, les acides p-coumariques, féruliques et les polysides associés, qui ont été isolés de la paille d’un cultivar de riz « Balilla 28 » et d’un de ses mutants à tige cassante. Des différences significatives ont été observées entre les deux types de riz pour les teneurs en lignines (Klason, au bromure d’acétyle et insoluble dans les acides), en esters et ethers d’acides phénoliques, en polysides hemicellulosiques et cellulosiques ainsi que pour les rendements d’extraction des fractions de lignine et d’hydrolyse par des polysidases. Le caractère cassant des tiges du mutant ne peut donc pas être simplement mis en relation avec les variations de composition chimique d’un constituant pariétal particulier. Ces résultats sont discutés en fonction des variations corrélatives de composition chimique des parois cellulaires et d’hétérogénéité de composition monomérique des lignines.

Mots clés additionnels : Mutation de la paroi cellulaire, acide p-coumarique, acide férulique, hemicelluloses, celluloses, paille.

I. INTRODUCTION

The use of mutants is now a classical approach for study of biosynthetic pathways in plants: it is of particular interest when lignification processes are concerned since the influence of lignification pattern on properties of lignocellulosic materials can be analyzed by comparison of different genotypes.

Rice, *Oryza sativa* L., is the third gramineous species after maize, *Zea mays* L., (JORGENSEN, 1931) and sorghum, *Sorghum bicolor* L., (PORTER et al., 1978) from which a mutant has been obtained at the level of the lignification process.

The mutant of rice has been obtained by gamma irradiation of the normal rice cultivar “Balilla 28” at the I.N.R.A. Plant Breeding Station of Montpellier (France) and is characterized by a brittleness of the culm appearing only in mature plants without any effect on the high resistance to lodging. The mutant exhibited higher contents of Klason lignin and of hot-
alkali soluble compounds and a lower proportion of α-cellulose than the normal cultivar (DOAT & MARIE, 1977). In agreement with these results concerning the rough chemical and mechanical properties of these rices, we also found that the mutant rice stem (culm) is richer in cell-wall bound p-coumaric esters while wall-linked ferulic ester content was similar in both mutant and normal rice (MONTIES et al., 1981). In addition, the extent of degradation of normal rice straw by both commercial crude cellulases and rumen juice liquor was wider than that of mutant one.

As these last differences may be related to variations in cell wall phenolics but also in polysaccharide composition, a reexamination of the cell wall constituents of normal and mutant rice straws has been undertaken. We report here results concerning the phenolic and polysaccharide compositions of three different lignin fractions and of the corresponding cell walls for straws. Structural characterization of polysaccharide fractions will be published elsewhere. Only nitrobenzene oxidation was used here for monomeric characterization of the rice lignin fractions; results concerning the non-condensed monomeric units obtained in these conditions are thus discussed in relation with the phenomenon of lignin heterogeneity (MONTIES & LAPIERRE, 1981; MONTIES, 1986).

II. MATERIAL AND METHODS

Rices, “Balilla 28” and its corresponding brittle-stem mutant, were grown at Montpellier in the rice plantations of I.N.R.A. using standard conditions of cultivation with N-P-K fertilization corresponding to plantations of I.N.R.A. using standard conditions of cultivation. They were harvested at grain maturity. Rices, “Balilla 28” and its corresponding brittle-stem mutant, were grown at Montpellier in the rice plantations of I.N.R.A. using standard conditions of cultivation with N-P-K fertilization corresponding to plantations of I.N.R.A. using standard conditions of cultivation. They were harvested at grain maturity.

A. Determination of cell-wall phenolics

Extractives were removed from the rough powder by three successive soxhlet extractions with toluene-ethanol mixture (1/1 vol), ethanol and water. The cell wall residue “parietal residue : PR” was then freeze-dried.

B. Lignin determination

Lignin was estimated by three different methods. After 5% sulfuric acid prehydrolysis of the PR, acid-insoluble lignin (AIL) was obtained by 72% sulfuric acid hydrolysis followed by post-hydrolysis in 5% sulfuric acid as previously discussed (MONTIES, 1984). Klason lignin (KL) was obtained by the same procedure but without prehydrolysis. Acetyl-bromide lignin was estimated according to the procedure of JOHNSON et al. (1961) with slight modifications: 25 mg of PR were treated with 10 ml of reagent according to JOHNSON but only 1 ml of the solution was analyzed, after dilution in 25 ml with glacial acetic acid, following the standard procedure of UV absorbance at 280 nm. A calibration curve, corresponding to 5 to 15 mg of guaiacol (2-methoxy phenol, SIGMA puriss.) was used for calculation and standardisation of the results. Data were expressed with reference to the Klason lignin as follows: a milled straw lignin preparation of known Klason lignin content, was submitted to acetyl-bromide solubilization, compared to the guaiacol standard curve, then a coefficient was calculated for conversion of guaiacol equivalent value (GEV) to Klason lignin equivalent (KLE).

C. Phenolic ester and phenolic ether determination

Cell wall bound esters and ethers of phenolic acids were estimated by a combination of alkaline and acid hydrolyses. Esters were first hydrolysed with 2 N NaOH, 20 ml for 100 mg of parietal residue, during 2 h at 35 ± 1 °C with magnetic agitation under nitrogen. The insoluble residue obtained after saponification was recovered by filtration and submitted to a second alkaline hydrolysis for exhaustive hydrolysis of esters. The insoluble final residue, saponification residue (RS), recovered by filtration was washed with water, until about neutrality (pH 6-7), then with 0.1 N HCl and used, without drying, for estimation of ethers. Ethers were hydrolysed with 0.1 N HCl in dioxane-water mixture, 10 ml for the total RS corresponding to 100 mg of RP, during 4 h in a sealed 20 ml tube at 100 °C. Phenolic acids corresponding to the cell-wall ethers were recovered from the acidolysis mixture by ethylether extraction as described by SCALBERT et al. (1985). Phenolic acids corresponding to the cell wall esters were similarly isolated by diethyl ether extraction after acidification of the soluble extracts and washings recovered during the RS preparation. Phenolic acids were, in each case, analyzed by HPLC. After drying on Na2SO4, diethyl ether extracts were concentrated under vacuum and recovered with methanol containing 1,2,3-trimethoxybenzaldehyde (Aldrich, puriss) at the concentration of 0.5 or 1.0 µg/µl as internal standard according to the concentration of the samples. Phenolic acids were separated by reverse phase HPLC with a Resolve C18-5 µ Waters column with a mixture of methanol-water (30-70 vol) containing 1% of acetic acid either in isocratic conditions at a flow rate of 1 ml.min⁻¹ or by gradient chromatography as described previously (MONTIES et al., 1981; SCALBERT et al., 1985). Purity of the trans- and cis-isomers of p-coumaric and of ferulic acids was controlled by cochromatography, by on-line recording of the UV-spectra of the products during the elution with a diode array detector (Hewlett-Packard 1040 A) and comparison of the position and the ratio of the maximum and minimum of absorbance between 250 to 450 nm. cis-isomers were only observed when exposure of samples to sunlight was not carefully controlled.

D. Fractionation of straw lignin

Finely ground parietal residue was ultraground in a Seibtecknik vibratory mill and lignin was isolated into three fractions LM, LE and LR according to
LAPIERRE et al. (1982) and SCALBERT et al. (1985).
The fractionation procedure is shown in table 1.
Conditions of ultragrinding and extraction chosen for
fractionation of poplar wood were used without
modification for straw. Fractionation was repeated
twice giving very reproducible yields. The monomeric
composition of the non-condensed monomeric units
of these lignin fractions was characterized by
nitrobenzene oxidation and analysis of the resulting
aromatic aldehydes by HPLC as previously reported
(MONTIES et al., 1981a; SCALBERT et al., 1985).

E. Carbohydrate analysis

Cell wall residue polysaccharides were hydrolyzed
according to SAEMAN et al. (1954) while carbohydrate
moieties associated with lignin fractions LM, LE and
LR were hydrolyzed with 2 N trifluoroacetic acid
(TFA) for 75 min at 125 °C (ALBERSHEIM et al.,
1967). Liberated neutral sugars were analyzed as their
alditol-acetate derivatives (SAWARDEKER et al., 1965)
by GC on a glass column (180 × 0.2 cm) packed with
3% SP 2340 coated on Supelcoport (100-120 mesh) at
225 °C (BRILLOUET et al., 1982).

The determination of acid-insoluble lignin after
5% sulfuric acid prehydrolysis allows, also, the
estimation of two rough polysaccharide fractions
which are hydrolyzed with 5% and 72% sulfuric
acid respectively. These two fractions had been
described previously in the case of poplar wood and
of wheat straw (MONTIES, 1984), as “readily acid
hydrolysable and solubilized products” (PAS :
“produits aisément solubilisables” in French ; MON-
ties, 1984) and as “poorly acid hydrolysable and
solubilized products” (PDS : “produits difficilement
solubilisables” in French : MONTIES, 1984). The
content of PAS and PDS has been calculated here, as
previously, by difference between the content in ash
(A), acid insoluble lignin (AIL), and lignocellulose
recovered after prehydrolysis (LC) by the relations :

\[ \text{PAS} = \text{PR} - \text{LC} \quad \text{and} \quad \text{PDS} = \text{LC} - \text{AIL} - A \]

Ash was weighed after mineralization at 550 °C for
3 h.

III. RESULTS AND DISCUSSION

The chemical composition of the parietal residue
from mutant and normal rice culms is given in
table 2.

Whatever the analytical procedure, the lignin
content of the mutant was, in each case, significantly
higher than that from the normal. Relative lignin
contents appeared, however, different depending on
the method used. In accord with the results previously
reported on poplar wood and wheat straw (MONTIES,
1984), the Klasson lignin content was significantly
higher than the acid-insoluble lignin content. As
previously discussed, KL content is probably over-
estimated due to condensation with other cell wall
components (DILL et al., 1984; MATSUMOTO et al.,
1984) while, on the other hand, AIL content is
certainly an underestimation of total lignin because of
the loss of acid-soluble monomeric products and
oligomeric fractions. The acetyl-bromide lignin
(KEL), given in table 1, was obtained after stan-
dardization with a milled straw lignin sample as
previously suggested by JOHNSON et al. (1961) or by
MORRISON (1972). Obviously, the KEL value allows a
direct comparison of the results obtained by these
different methods, but, even after these calculations,
KEL value has to be considered only as a relative one
due to possible variations of specific UV absorbance

| Fractions | Lignins | Acid-hydro-
| --- | --- | lysable products |
| KL | Acetyl-
| Bromide-insoluble | Acid-
| KL* | KEL* | AIL* | PAS* | SO₃H₂ | SO₃H₂ | PDS* | Ash 550° |
| Normal | 12.5 | 9.9 | 9.3 | 38.5 | 46.2 | 2.8 |
| Mutant | 14.9 | 12.6 | 11.4 | 43.5 | 37.3 | 4.3 |

* Abbreviations and definitions in chapter II.
between lignin fractions, to systematic occurrence of non-lignin components such as polysaccharide in lignin preparations and finally to uncertainties in KL determination discussed before. Acetyl-bromide determination of lignin is usually considered as an estimation of total lignin because parietal residue is solubilized without any pretreatment. The fact that, in table 2, the acetyl-bromide lignin value (KEL) is nearer the acid-insoluble lignin (AIL) than the Klason lignin (KL) value is thus unexpected. As, up to now, no detailed UV characterization of rice-straw lignin fractions are available, these results cannot be discussed more precisely.

During the determination of the acid-insoluble lignin, it is worthwhile to estimate the weight losses after treatment by 5 % and by 72 % sulfuric acid. The loss of dry matter after prehydrolysis of the cell wall residue may be considered roughly as the amount of polysaccharides which are readily hydrolyzed by dilute acid, i.e. arabino-glucuronoxylans in the case of grass straw (JOSELEAU, 1980). The hydrolysis step by 72 % sulfuric acid removes essentially the cellulose and xylan moieties which resisted the prehydrolysis treatment. From table 2, the weight loss after prehydrolysis was higher for the mutant than for the normal rice. In contrast, the decrease of dry matter on 72 % sulfuric acid hydrolysis was lower for the mutant rice than for the normal one, in agreement with previous results (MONTIES et al., 1981). Indeed the mutant rice stem was richer in hemicellulose polymers and poorer in cellulose than the normal one, which contained more cellulose and a lower proportion of hemicelluloses. This preliminary conclusion was sustained by the relative composition of monosaccharide constituents of the cell wall polymers given in table 3. The cell wall from mutant rice contained more xylose and arabinose than than from normal rice; these two sugars are constitutive of the hemicellulosic fraction of grass straw. On the other hand, glucose, originating mainly from cellulotic glucans, occurs in greater relative quantity in the cell wall from normal rice stem. Using other rough methods, DOAT & MARIE (1971) reported in agreement with our results that the mutant of rice had a higher hot alkaline extract, hemicellulosic products, and a lower α-cellulose content.

Table 3 also shows that the xylose/arabinose ratio, which is indicative of the degree of branching of xylan chains, was higher in the mutant (4.8) than in normal rice (4.0). Therefore hemicelluloses from the brittle-culm mutant seemed more linear than in the normal one. However, this observation needs to be confirmed by methylation analysis of cell wall polysaccharides.

Results concerning the characterization of phenolic acids bound to the stem cell walls are shown in table 4 and figure 1. In addition to the confirmation of the occurrence of p-coumaric and ferulic esters in rice cell wall, table 4 shows also the occurrence of ethers of the same phenolic acids. Ethers of p-coumaric and of

| TABLE 4 |
| Phenolic acids bound to the stem cell wall of the mutant and normal rice. Contents are given in per cent of dry RP*.

<table>
<thead>
<tr>
<th>Saponification residue (RS)*</th>
<th>p-coumaric acid</th>
<th>ferulic acid</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>esters</td>
<td>ethers</td>
</tr>
<tr>
<td>Normal</td>
<td>65</td>
<td>0.83</td>
</tr>
<tr>
<td>Mutant</td>
<td>55</td>
<td>1.19</td>
</tr>
</tbody>
</table>

* Abbreviations in chapter II.

![Figure 1](image)

**Figure 1**

Isolation and characterization of the ethers of phenolic acids associated with cell-wall preparations of rice stem. Results corresponding to normal rice; identical qualitative results have been obtained with the mutant.

A) High-pressure liquid chromatography separation of phenolic acid solubilized after acidolysis following exhaustive alkaline hydrolysis : PC = p-coumaric acid (trans form), FE = ferulic acid (trans form), A = acidolysis products, E = internal standard ; 3, 4, 5-trimethoxyximacaldéhyde.

B) Absorption spectra of phenolic acids isolated from rice stem.

* Séparation par chromatographie liquide haute pression des acides phénoliques libérés par acidolysé après hydrolyse alcaline exhaustive : PC = acide p-coumarique, FE = acide férruleux, A = produits d'acidolysé, E = éthane interne (3, 4, 5-triméthoxyximacaldéhyde).

B) Spectres d'absorption des produits isolés de tige de riz.

| TABLE 3 |
| Relative composition of polysaccharide sugar constituents in the cell wall residue from culm of normal and mutant rice.

<table>
<thead>
<tr>
<th>Sugar</th>
<th>Glucose</th>
<th>Xylose</th>
<th>Arabinose</th>
<th>Galactose</th>
<th>Mannose</th>
<th>Rhamnose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>70.4</td>
<td>22.5</td>
<td>5.6</td>
<td>1.5</td>
<td>tr*</td>
<td>tr*</td>
</tr>
<tr>
<td>Mutant</td>
<td>59.7</td>
<td>31.4</td>
<td>6.5</td>
<td>2.0</td>
<td>0.2</td>
<td>0.1</td>
</tr>
</tbody>
</table>

* tr : traces about 0.02 to 0.05.
ferulic acid bound to rice cell wall are reported here for the first time; this result is in line with the occurrence of the same type of ethers in wheat straw cell wall, as recently reported by SCALBERT et al. (1985). In the case of the two rice cultivars reported here, the trans-isomers of \textit{p}-coumaric and ferulic acids were characterized only by cochromatography and on-line UV-spectrophotometry during elution.

Significant differences in the distribution of cell wall-linked phenolic acids are also shown in table 4. While the amount of ferulic esters was the same in both rices, the amount of esters of \textit{p}-coumaric acid was increased in the mutant, which is in complete agreement with data reported earlier by MONTIES et al. (1981); the \textit{p}-coumaric to ferulic esters ratio was, however, higher than those previously reported. The opposite figure was observed for the ethers; the amount of \textit{p}-coumaric ethers was the same in both rices while ethers of ferulic acid were reduced in the mutant. As, however, the data reported here concern only one sample of rice, these differences in distribution between ethers and esters require confirmation with other samples of rice grown in different conditions.

In addition, table 4 shows a higher yield of recovery of saponification residue (RS) in the case of the normal rice. Recovery yields of RS could be related to the higher content of mutant rice cell wall in hemicellulosic glucuronarabinoxylans; indeed most of the xylan, and also part of the lignin, underwent solubilization under saponification conditions. These last observations were well correlated with PAS data obtained from acid prehydrolysis reported in table 2 and 3.

Table 4 summarizes the fractionation results of lignin from rice culm following the procedure of table 1; three lignin fractions (milled straw lignin, enzyme lignin and residual lignin) were prepared. As cell wall polymers may be associated with the lignin fractions, recovery yields must be compared with caution. The LM fractions were obtained in the same yield for both rices while yields of LE and LR have been always found higher for the normal than for the mutant, which is surprising since the mutant culm is more lignified than the normal one. The most prominent difference between the two rices was the greater sensitivity of the mutant cell walls to \textit{Trichoderma viride} cellulase hydrolysis compared to the normal one. This differential sensitivity is clearly shown in table 5 by the higher yield in undialyzable products (UP) for the mutant and by the corresponding lower yield in final residue (FR) which was about half that of the normal one. These data were rather unexpected since the more lignified culm of mutant was supposed to be more resistant to enzyme hydrolysis than the normal one. In the previous study of MONTIES et al. (1981), no significant difference was found, after cellulase hydrolysis, between the normal and the mutant rice. As, however, commercial \textit{Oxyporus} cellulase preparation was used in place of commercial \textit{Trichoderma} cellulase used in the present fractionations and as enzymatic treatment of 24 h was chosen in place of repeated and extensive cellulase hydrolysis used in the present procedure (table 4), these differences of yield may be explained by the well established structural and kinetic differences occurring between commercial cellulase preparations (AZUMA et al., 1983).

Due to the relatively low quantities of LR fractions isolated from the mutant, only LM and LE lignin fractions were characterized here.

Significant differences in monomeric composition of LM and LE fractions were observed by nitrobenzene oxidation as shown in table 6. In both cases, the syringaldehyde to vanillin ratio and the \textit{p}-hydroxybenzaldehyde to vanillin ratio were higher for the mutant than for normal rice. Similar differences have previously been reported for these two ratios (MONTIES et al., 1981a). Nevertheless, no significant differences in monomeric composition could be observed in table 6 between LM and LE fractions isolated from both type of rice. Even if analysis of phenolic aldehydes obtained by nitrobenzene oxidation only allows the characterization of the non-condensed monomeric units of lignin, these last results clearly show that monomeric heterogeneity was not observed between lignin fractions isolated from the two rice straws. Recently the same results were reported by SCALBERT et al. (1986) who were

### Table 5

<table>
<thead>
<tr>
<th>Lignin fractions</th>
<th>Total yield (KL)</th>
<th>Final residue FR</th>
<th>Undialysed products UP</th>
</tr>
</thead>
<tbody>
<tr>
<td>LM</td>
<td>1.4</td>
<td>3.5</td>
<td>1.3</td>
</tr>
<tr>
<td>LE</td>
<td>1.3</td>
<td>2.5</td>
<td>0.6</td>
</tr>
<tr>
<td>LR</td>
<td>49.6</td>
<td>30.2</td>
<td>8.0</td>
</tr>
</tbody>
</table>

Abbreviations in chapter II according to table 1, yields in per cent of dried paretial residue (RP).

### Table 6

<table>
<thead>
<tr>
<th>Rice lignin fractions</th>
<th>Vanillin</th>
<th>\textit{p}-OH-benzaldehyde</th>
<th>Syringaldehyde</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>3.5</td>
<td>0.43</td>
<td>1.14</td>
</tr>
<tr>
<td>Mutant</td>
<td>3.0</td>
<td>0.53</td>
<td>1.37</td>
</tr>
<tr>
<td>LE</td>
<td>3.2</td>
<td>0.44</td>
<td>1.16</td>
</tr>
<tr>
<td>Mutant</td>
<td>4.9</td>
<td>0.49</td>
<td>1.27</td>
</tr>
</tbody>
</table>

Yield in per cent of corresponding dry lignin fraction.

Rendement en pourcentage de la masse des fractions de lignine.
unable to show lignin heterogeneity between fractions isolated from wheat straw. This inability to detect heterogeneity of monomeric composition in grass straw lignins is rather surprising when considered with reference to evidence and confirmations of the occurrence of such heterogeneity in the case of angiosperm woods such as normal and reaction wood of poplars (LAPIERRE & MONTIES, 1982; LAPIERRE et al., 1982). Furthermore, monomeric heterogeneity between lignin fractions isolated from oak woods has recently been found by MONTIES & LAPIERRE (in preparation), confirming the occurrence of lignin heterogeneity in angiosperm woods.

The inability to observe this phenomenon in the case of grass straws may result, for example, from the fact that the cytological and cytochemical structures of culm is far more complex than those of wood. The rough fractionation procedure shown in table 1 would thus allow the isolation of heterogeneous fractions of lignin when applied to samples containing a small number of cellular types, as in the case of wood xylem, but not when applied to more complex mixture of tissues, such as in the culm of grass straws. Another, but less likely, hypothesis is that the fractionation procedure chosen for woods is not well fitted for straws and requires specific adaptation.

The chemical composition of the polysaccharides associated with the rice lignin fractions is shown in table 7. Clearly, in each case, lignin preparations were mainly associated with arabinoxylans. No significant differences were observed between preparations isolated from normal and from mutant rice straws. The arabinose to xylose ratio was significantly higher, in each rice, in LE than in LM; however in LR, this ratio was nearer of those in LE than those in LM. These differences probably depend on cellulase treatment and cannot be used directly for discussion on the structure of lignin polysaccharide associations.

IV. CONCLUSION

The results presented in this study clearly show that the brittle-culm mutation of rice not only changes, quantitatively and qualitatively, the phenolic fraction (lignin and bound phenolic acid) of the culm cell wall but also the polysaccharides (hemicellulosic arabinoxylan and cellulose). As variations in many different chemical factors are associated with the brittle-culm character of the mutant, it seems difficult to correlate specific changes in properties of the lignocellulosic material, such as in vitro digestibility or mechanical properties, with only one chemical factor such as lignification. A strong correlation between lignin and hemicellulose concentrations has been reported in ten varieties of temperate grasses by MORRISON (1980); the varieties with higher lignin concentration showed higher linear hemicellulose content. The differences reported here between the mutant and the normal cultivar of rice agree with these observations but perhaps fortuitously; it would thus be interesting to search if the same correlation is also found in other lignin mutants of Gramineae, such as corn and sorghum.

Furthermore, when the results reported here are compared with those of the two previous studies published by DOAT & MARIE (1977) and by MONTIES et al. (1981a), a very good agreement appears which may indicate that strong genetic control occurs during the biosynthesis of the plant cell wall. The same type of conclusion has been suggested previously from comparison of the chemical composition of wheat straw harvested after very different conditions of growth (BAYET & MONTIES, 1977). The occurrence of such a strong genetic control may be of great importance in research programs concerning the biodegradability of the gramineous cell wall applied for example to pathogen resistance or to increase of digestibility by animals.

In this case, very careful studies of the correlations between variations in the fine chemical composition of plant cell wall would be required; correlations between variations in lignin and hemicellulose have been recalled previously. The fact that the rough composition of the cell wall polysaccharide may change without significant variation in the content of associated ferulic esters and p-coumaric ethers is a second example of such fine variation; the occurrence of a possible genetic control remains to be established in this last case however. Furthermore, subtle changes may also occur after modification of the lignification characters at the level of not only the content and the monomeric composition of total lignin but also the extractability and composition of lignin fractions. Much work remains to be done particularly to characterize variations in the lignin fractions easily solubilized during mild alkaline and acid treatment of straws. Wheat-lignin-labelling experiments by incorporation of U-14C-phenylalanine have, indeed, shown quantitative and qualitative differences of labelling between lignin fractions (BAYET, 1980; AGOSIN et al., 1986). Lignin heterogeneity not observed here in the case of rice and previously in the case of wheat may thus be shown using different analytical methods.

In conclusion many comparative studies of the fine chemical composition of the cell wall remains to be done not only with rice but also with other Gramineae such as wheat, and with “mutants” for lignification; such studies may provide more exhaustive expla-
nations on the molecular mechanisms involved in the physico-chemical properties of lignocellulosic materials.

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