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INTERPRETING MULTICOMPONENT INFRARED SPECTRA

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INTERPRETING MULTICOMPONENT INFRARED SPECTRA

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ABSTRACT

The complexity of a multi-component spectrum decreases as constituents are removed from it. This decrease is reflected by a reduction in intensity of the first derivative. If the spectrum of a candidate is subtracted stepwise from a multi-component spectrum and a derivative is taken at each stage, then the candidate will be identified in the mixture if the derivative minimizes. An application to the determination of lignin in pulp will be presented together with preliminary work on the spectral identification of recycled fiber.

RESULTS AND DISCUSSION

Consider the spectrum in Fig. 1A which comprises two signals and its derivative. As one of the signals is progressively removed, the absolute value of the overall intensity (positive and negative) of the first derivative decreases until the signal is exactly stripped out in Fig. 1B. Further stripping of the signal increases the intensity of the derivative (Fig. 1C). Thus, the derivative, which relates more closely to spectral complexity than to the intensity, minimizes when a component is exactly removed. Hence, in principle, the intensity of the derivative can be used as a tool for identifying component spectra.

Consider now the spectrum in Fig. 2A which contains components from Figs. 2B and 2C. A computer program was written to subtract various fractions $n$ of the Fig. 2B spectrum from Fig. 2A and to take a derivative after each stage. A plot of the absolute integrated intensity (positive and negative) of the derivative is plotted against $n$ in Fig. 3. The minimum occurs at $n = 0.85$, indicating that 85% of the Fig. 2B spectrum is contained in Fig. 2A. A similar analysis shows that 24% of the Figure 2C spectrum is contained in Fig. 2A.

The technique has been used to determine kappa number from the infrared spectrum of pulp. Several attempts to do so have been reported earlier (3-8). Berben et al. (3) used the absorbance of a characteristic lignin band at 1510 cm$^{-1}$ to estimate kappa number. Since cellulose also absorbs in this region, the band intensity was measured after a spectrum of cotton linters had been subjectively subtracted out. The residual 1510 cm$^{-1}$ absorption was quite small, and since the subtraction step introduces substantial uncertainty, kappa number could be determined to only within ±10 units (3).
Schultz et al. (4) used five absorption bands ratioed to a sixth to express lignin content, and a similar multiple band approach was taken by Grandmaison et al. (5). Backa and Brolin (6) used partial least squares analysis with absorptions at 232 discrete wavenumbers. Sample preparation included milling and drying to constant weight. A precision of 2.75% was obtained after removal of outliers.

Consider the spectra in Fig. 4. The Fig. 4A spectrum is that of an unbleached loblolly pine Kraft pulp. The Fig. 4B trace is a library spectrum of Kraft softwood lignin. Fractions of the Fig. 4B spectrum were incrementally subtracted from Fig. 4A, and a derivative was taken after each step. A plot of the integrated area (positive and negative) of the derivative vs. the scaling factor applied to the lignin spectrum showed a minimum at 0.18, indicating that 18% of the Fig. 4B spectrum is contained in (or overlaps with) Fig. 4A.

The lignin scaling factor needs to be normalized in order to compensate for interferences and for spectral variability. The obvious choice for a reference is cellulose since it is the major component of pulp. A derivative minimization as described above was conducted with a spectrum of cotton linters. The ratio of the lignin:linters minima is designated as \( f'_{\text{lig}} \) and is expected to be a measure of the lignin content relative to the cellulose (cotton linters) content. Values of \( f'_{\text{lig}} \) for several Kraft and one sulfite pulp were regressed against kappa number and led to

\[
\text{kappa no.} = -23.5 + 191 f'_{\text{lig}} \quad (n=14; r=0.99)
\]  

(1)

The relationship is illustrated in Fig. 5. The quality of the fit is unchanged if a spectrum of hardwood lignin is used to determine \( f'_{\text{lig}} \). This is consistent with the general spectral similarity between hardwood and softwood lignin. Consequently, it appears that a single equation may apply to softwoods, hardwoods, and to mixtures thereof.

The magnitude of the intercept may initially appear to be too large, since lignin-free material has a kappa number of zero. However, the spectra of lignin and cotton linters are not orthogonal. We found (through derivative minimization) that 11.2% of the lignin spectrum was contained in a spectrum of cotton linters, i.e. \( f'_{\text{lig}} \) for cotton linters was 0.112. This leads to a value of about 2 for the kappa number of cotton linters, which is of the order of the uncertainty of the estimation.

The \( f'_{\text{lig}} \) values used in Fig. 5 were averaged from two spectra, and duplicates agreed to within 12%. One of the pulps was visibly inhomogenous, and the duplicates differed by 22%. It would seem that the approximately 6 mm² area of the pulp sample exposed to the beam does not fully represent the pulp, and it is likely that greater replication will improve accuracy.

Equation 1 consistently underestimates kappa numbers of \( \text{NO}_2 \) treated \textit{unbleached} pulps by an average of 23 units. Exposure to \( \text{NO}_2 \) probably alters the infrared spectrum of the lignin; certainly, the electronic spectrum is altered as indicated by a yellowing of the exposed pulps. A change in lignin structure would reduce the overlap between the lignin and pulp spectra and lead to an underestimate of lignin content as observed. However, once the \( \text{NO}_2 \) treated pulps are bleached they are accommodated by eq. 1. This indicates that the altered lignin is largely removed during bleaching; the residual native lignin is then again recognized by our algorithm.

An important component of our procedure is that it considers all the spectral information rather than discrete preselected bands. It is unnecessary for a signal in the mixture to be associated exclusively with any one component. Perhaps the most useful feature of our technique is its relative insensitivity to interferences in comparison to absorbance-based techniques. A common variable in pulp is the moisture factor, and in order to determine the effect of water on kappa number, we collected spectra of pulps varying only in moisture content. The results are provided in Table 1; clearly, water does not add significantly to the uncertainty. The "0% moisture" pulp was oven dried at 105°C for 4 hours, a process that removes volatiles and causes structural changes in the pulp. Our technique
Table 1 Variation of kappa number with moisture content

<table>
<thead>
<tr>
<th>Percent moisture</th>
<th>f'lig</th>
<th>f'linters</th>
<th>f'lig</th>
<th>kappa no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.054</td>
<td>0.314</td>
<td>0.16</td>
<td></td>
</tr>
<tr>
<td>4.2</td>
<td>0.108</td>
<td>0.520</td>
<td>0.21</td>
<td>16.6</td>
</tr>
<tr>
<td>33.9</td>
<td>0.099</td>
<td>0.493</td>
<td>0.20</td>
<td>14.7</td>
</tr>
</tbody>
</table>

picks up the change, and the $f'_{lig}$ value in Table 1 for the dried pulp is very different from those of the others.

We are presently exploring the possibility of applying the technique to differentiating recycled from virgin fiber. The basis of our hypothesis is that recycled material should be depleted in hemicellulose and other solubles by virtue of its longer exposure to water. Conversely, one might expect the content of insoluble material such as cellulose to increase with progressive recycling. Hence, the hemicellulose:cellulose ratio should be lower for recycled fiber than for the virgin material, and could, in principle, be used to detect the former. The hemicellulose:cellulose ratio can be obtained by IR spectroscopy, and preliminary data relating it to the content of recycled fiber will be presented.

LITERATURE CITED

Figure 1 Gaussian signals and their derivatives. A Two equivalent Gaussians; B Residual after stripping out a component signal. C Residual after overstripping a component signal.

Figure 2 Spectrum of a mixture A including components from B and C.
Figure 3 Integrated absolute intensities (positive and negative) of derivatives of the Fig. 2A spectrum taken after stepwise subtraction of $n$ times the Fig. 2B spectrum.

Figure 4 Spectra of softwood Kraft pulp (A) and softwood lignin (B).
Figure 5  Dependence of area of the derivative on scaling factor.