MASS SPECTRAL FRAGMENTATION OF SOME INDOLE DERIVATIVES

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Abstract - EI mass spectra of oxindoles C, iminoethers D, and dihydro β-carbolines E, arising from chloroindolenines B are analysed.

Chloroindolenines B, obtained from 4-oxo indole (2, 3-α) quinolizidines A, rearrange to oxindoles C, iminoethers D, and dihydro β-carbolines E (TABLE I), in which R=H, C2H5 or CH2CH2CO2R'. Some oxindoles C may also be prepared after HARLEY MASON, through condensation of 2-hydroxy tryptamine with the suitable aldehyde-ester. Iminoethers D (R'=CH3 or C2H5) are available from C, using the appropriate MEERWEIN'S reagent. All these compounds exhibit very simple EI spectrum upon a 70 eV energy electron beam.

Then, it became obvious to compare the behaviour of each class, in order to deduce the fragmentation pattern and mechanism. Only one natural base, i.e. the alkaloid vincatine, is known to possess the same skeleton as ester 1c. Its mass spectrum has already been published. However, oxindoles 1a,b,c, which bear a supplementary lactame carbonyl group at C(3), follow a fragmentation mode drastically differing from that of vincatine.

1. Mass spectral fragmentation of oxindoles C and iminoethers D

![Diagram](image-url)
Oxindoles 1a,b,c and iminoethers 2c and 3a,b,c are concerned in this study. They differ only by
the nature of R on C(20)\(^2,6\). In each case, many stereoisomers exist, due to the chiral carbons
7,20 and 21. Yet, the mass spectra of the stereoisomers are quite similar, and throughout this
paper, the relative configuration of the discussed molecules will not be further taken in account.
Some fragments vary with R, some other do not.

1.1 Fragments non varying with R : ions a,b,b',c,c'

\[ a : \text{m/e 159} \]
\[ b : R' = \text{CH} \text{\_3 : m/e 160} \]
\[ b' : R' = \text{C}_2\text{H}_5 : \text{m/e 174} \]
\[ c : R' = \text{CH} \text{\_3 : m/e 173} \]
\[ c' : R' = \text{C}_2\text{H}_5 : \text{m/e 187} \]

When bearing only one nitrogen (N(1)), they retain carbons from cycles A,B or C.

The abundant ions are:

- from oxindoles 1a-c : ion a, m/e 159
- from iminoether 2c : ions b, m/e 160 and c, m/e 173 (base peak)
- from iminoether 3a-c : ions b', m/e 174 and c', m/e 187 (base peak).

Ion a is commonly encountered in oxindole alkaloids\(^7\). The nature of b,b' and c,c' is deduced from
the gain of 14 m.u. from the methyl iminoether 2 to the ethyl derivate 3.

1.2 Fragments varying with R : ions d,e,f

When bearing only are nitrogen (N(4)), they involve carbons of cycle D. They arise from oxindole 1
and its iminoethers 2 and 3. Their alleged genesis and structures are shown on TABLE 2. Radical
cations d result from the homol~tic cleavage of bonds 6-5 and 7-21 in the molecular ion. These
unstable species rearrange to the more stable ions e and f through the loss of one of the two
radicals attached to C(20).

These assumptions are deduced from the correlative variations of d and R : d\(_1\) (m/e 139) d\(_2\) (m/e 167)
d\(_3\) (m/e 225).

Then d\(_1\)-3 lead to the corresponding ions e through loss of 29 m.u. : e\(_1\) (m/e 110) ; e\(_2\) (m/e
138) ; e\(_3\) (m/e 196).

When the variable radical R is lost from d, ion f arises. Actually, f (m/e 138) is present on the
spectrum of any compound 2 and 3. Moreover, the ubiquitous ion m/e 138 is notably more abundant on
the spectra of compounds 1b and 3b. In these compounds, the two substituents attached to C(20)
are identical; then d₂ leads to identical ions e₂ and f.

2. Mass spectral fragmentation of dihydro-α-carbolines E

Series a consist of methyl esters, series b of ethyl esters.

Compounds 7a and 7b could not be separated: then the spectrum of the mixture was measured.

The fragmentation of compounds E appears to proceed through three competitive mechanisms (TABLE 3):

2.1 Loss of an acrylic ester plus homolytic cleavage: ions g and h

Peaks M -101, i.e. M-(86 + 15) are to be seen on the spectra of 4a,5a,6a,7a, whereas peaks M -115, i.e. M-(100 + 15) are to be seen on the spectra of 4b,5b, and 7b. These ions may result from the simultaneous loss of methyl (a series) vs ethyl (b series) acrylate and a methyl radical.

Ions g give rise to the base-peak in the spectra of 4a,b and 5a,b. Their genesis may result from the cleavage of the 15,20-bond with hydrogen radical transfer on N(4) (loss of an acrylic ester), followed with the loss of a methyl radical from the ethyl chain at C(20), thus leading to the very stable species g.

In the case of diesters 6a and 7a,b, the same process (i.e., loss of an acrylic ester and a methyl radical) can occur: ions g m/e 283 vs 297, but the main process is the loss of an acrylic ester and an acetate ester radical: h m/e 225 (base-peak). The latter process requires less activation energy.

2.2 Loss of an acrylic ester through a MAC LAFFERTY's rearrangement: ions i

The ions i appear at m/e M-86 (4a,5a,6a and 7a) or M-100 (4b,5b,7b), in every case with the same relative intensity (about 30 % of the base peak). These relatively abundant radical cations are suggested to be generated through an electrocyclic process such as path 2 outlined on TABLE 3.

2.3 Homolytic cleavage of the 14-15 bond

The classical scission β to a carbonyl group accounts for the loss of a methyl acetate radical (a series) or an ethyl acetate radical (b series) from the molecular ion and gives rise to ions j. The strikingly high abundance of j from 5a (80 %) and 5b (85 %) could not be interpreted.

The above results are of interest for the structure elucidation and recognition of the intermediates C,D,E in the total synthesis of indole alkaloid performed in this8⁵⁻⁹ and other1⁰ laboratories.
REFERENCES


4. The mass spectra were recorded on C.E.C. n°21-580 low resolution single focusing spectrometer.


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