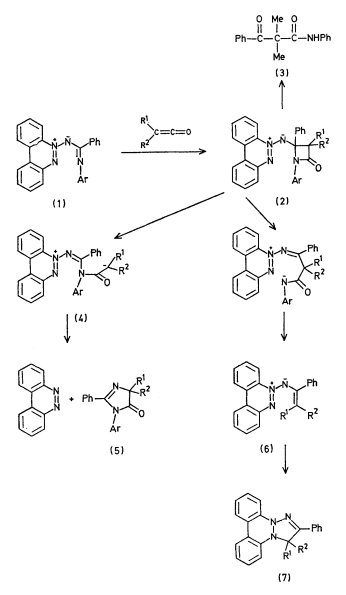
Cycloadditions of Extended Dipoles: Reaction of Imidoylazimines with Ketens

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Summary Ketens undergo 2+2 cycloaddition to the C=N bond of the imidoylazimines (1) to give the β -lactams (2) which are cleaved on heating to give either benzocinnoline and 1,2,4,4-tetrasubstituted imidazolin-5-ones (5) or the triazolines (7), depending on the keten substituents.

CYCLOADDITIONS involving extended dipolar systems offer the possibility of a simple route to medium ring heterocycles; the periselectivity observed in such processes is therefore of practical, as well as of current theoretical interest.¹



† Generated in situ by the action of triethylamine on the appropriate acid chloride.

The imidoylazimines $(1)^2$ can in principle undergo 5+2, 3+2, or 2+2 cycloaddition with ketens. However, addition of diphenyl-, phenylmethyl-,[†] and dimethyl-ketens[†] to the imidoylazimines (1; Ar = Ph, o-tolyl, and p-NO₂·C₆H₄) in benzene at room temperature gave the yellow 2+2 cycloadducts (2) quantitatively. Evidence for the β -lactam structure is the i.r. carbonyl absorption at 1750—1730 cm⁻¹, retention of the characteristic benzo-cinnoline N-imide ¹H n.m.r. absorption pattern,³ and the ready hydrolysis of the adduct (2; Ar = Ph, R¹ = R² = Me) to give benzocinnoline N-imide and the ketoamide (3).

The adducts (2) are unstable and decompose on prolonged standing or heating to give products which depend on the keten substituents R^1 and R^2 . The diphenylketen adducts (2; $Ar = R^1 = R^2 = Ph$ and Ar = o-tolyl, $R^1 = R^2 = Ph$) in refluxing benzene gave benzocinnoline quantitatively and the imidazolinones (5; $Ar=R^1=R^2=Ph$ and Ar=o-tolyl, $R^1 = R^2 = Ph$), m.p. 158° (70%), and m.p. 138° (75%), respectively. These imidazolinones were also produced unambiguously but in lower yields from diphenylglycine and the N-arylbenzimidoyl chloride. Formation of the imidazolinones is readily explained in terms of the highly stabilised zwitterionic intermediate (4) resulting from C-C bond cleavage of the β -lactam. Significantly, in the overall sequence $(1) \rightarrow (2) \rightarrow (5)$ the imidoylazimine has functioned effectively as an imidoylazide. Such reactivity is precluded for the latter which exist as the very stable tetrazoles.

The dimethylketen adduct (2; Ar = Ph, $R^1 = R^2 = Me$), although hydrolytically less stable than the diphenyl analogue, is somewhat more stable thermally and undergoes a different mode of breakdown. Heating in dry toluene gives the triazoline (7; $R^1 = R^2 = Me$), m.p. 165° (20%), presumably by C–N bond cleavage in the β -lactam leading to loss of isocyanate and formation of the 1,5-dipole (6) which cyclises. In this case C-C bond cleavage would give a zwitterion (4; $R^1 = R^2 = Me$) in which the carbanionic centre is less stabilised than in the diphenyl analogue and none of the imidazolinone (5; $R^1 = R^2 = Me$) was detected. On further heating in toluene the triazoline (7), a rare type of dihydro-1,2,3-triazole with two saturated nitrogen atoms, undergoes fragmentation to give benzocinnoline and benzonitrile. Predictably the methylphenylketen adducts show intermediate behaviour. Thus (2; $Ar = R^1$ = Ph, R^2 = Me) gave the imidazolinone (5) (30%) and the triazoline (10%) whereas (2; $Ar = p-NO_2 \cdot C_6H_4$, $R^1 = Ph$, $R^3 = Me$) gave only the triazoline (50%). As expected the more electron-withdrawing substituent on the terminal nitrogen favours cleavage of the C-N bond.

Formation of adducts (2) can be readily rationalised by a stepwise cycloaddition through the zwitterion (4) formed by nucleophilic attack of the terminal ylide nitrogen on the keten, although concerted $\pi^2_s + \pi^2_a$ addition, or $\pi^2_s + \pi^2_s$ addition favoured by configuration interaction,⁴ are also formal possibilities. Rare examples of 5 + 2 (possibly concerted $\pi^6_s + \pi^2_a$) additions of conformationally fixed

cisoid 1,5-dipoles to cumulenes have been observed;⁵ however, the preferred configuration and conformation of the acyclic 1,5-dipoles (1) are uncertain and may militate against formation of the 5 + 2 adduct by either stepwise or

concerted modes of addition.

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