

A New Strategy for the Synthesis of Cinnoline Derivatives

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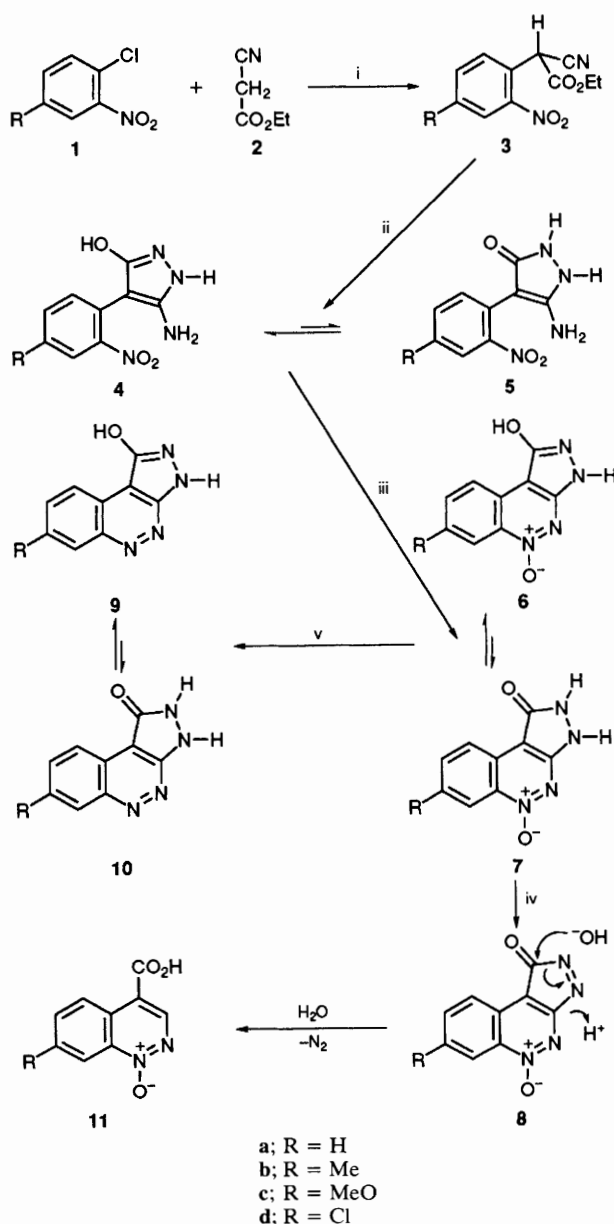
Readily accessible 3-amino-5-hydroxy-4-(2-nitroaryl)pyrazoles undergo efficient base-catalysed cyclisation to afford 7-substituted 1-hydroxy-3*H*-pyrazolo[3,4-*c*]cinnoline 5-*N*-oxides, hypochlorite oxidation of which provides a viable synthetic route to hitherto inaccessible cinnoline-4-carboxylic acid 1-*N*-oxides.

As part of an investigation concerned with the design of new radical scavenging agents having anti-inflammatory activity we required efficient and unambiguous synthetic access to cinnoline 1-*N*-oxides containing exploitable functionality at the 3- and/or 4-positions. There is no simple method for the synthesis of cinnoline *N*-oxides other than direct peracid oxidation of the parent heterocycles.¹ Moreover this approach in practice is often ambiguous in regard to the site of mono-oxidation, inefficient owing to the formation of mixtures of 1- and 2-*N*-oxides, and incompatible with existing

functionality sensitive to oxidation. These potentially undesirable features precluded the use of direct peracid oxidation for the synthesis of the cinnoline *N*-oxides in question and prompted the development of an alternative synthetic route based on a new strategy for the construction of the cinnoline ring system. This incorporates two novel synthetic procedures. Firstly the regiospecific formation of a cyclic azoxy nucleus using the known² but little explored base-catalysed aldol-type condensation of an amino substituent with a suitably positioned aromatic nitro group, and secondly the previously unexploited oxidative transformation of a fused pyrazolone ring into a carboxy substituent.³ Application of the new strategy is now shown to provide a viable synthetic route to otherwise inaccessible cinnoline-4-carboxylic acid 1-*N*-oxides.

The key starting materials for the new synthetic route to cinnoline 1-*N*-oxide derivatives (Scheme 1) were 3-amino-5-hydroxy-4-(2-nitroaryl)-1*H*-pyrazoles **4a-d**. These previously undescribed pyrazole derivatives were readily obtained (Scheme 1) by the condensation of easily accessible (Table 1) ethyl 2-cyano-2-(2-nitroaryl)acetates **3a-d** with hydrazine under standard conditions. The lack of carbonyl absorption in the IR spectra of the products **4a-d** supports their formulation as hydroxypyrazoles rather than the alternative tautomeric pyrazolinones **5a-d**.⁴ However the unusual behaviour of the hydroxypyrazole derivatives **4a-d** at their melting points (Table 1), the reason for which is not yet clear, may indicate their tautomeric conversion into the pyrazolinones **5a-d** at elevated temperature.

Heating the amino-nitropyrazoles **4a-d** with 2 mol l⁻¹ aqueous sodium hydroxide under reflux (2 h) resulted in their clean cyclisation to the expected pyrazolo[3,4-*c*]cinnoline *N*-oxides **6a-d** in good to excellent yields (Table 1). The gross structures of these derivatives of the previously undescribed pyrazolo[3,4-*c*]cinnoline ring system are supported by their simple reduction to the parent heterocycles **9a-d** in high yield (Table 1) using sodium dithionite in aqueous dimethylformamide (DMF). The existence of the pyrazolocinnoline derivatives **6a-d** and **9a-d** predominantly in the hydroxy rather than the keto tautomeric forms **7a-d** and **10a-d** respectively is, as for the hydroxypyrazoles **4a-d**, indicated by the absence of carbonyl absorption in their IR spectra.



Scheme 1 Reagents and conditions: i, NaH, DMF, 100 °C; ii, NH₂NH₂·H₂O, EtOH, reflux; iii, 2 mol l⁻¹ NaOH, reflux; iv, 14% NaOCl aq., 2 mol l⁻¹ NaOH, room temp.; v, Na₂S₂O₄, DMF, H₂O, reflux

Table 1

Compound ^a	Yield (%) ^b	Mp (t ^o C)	Compound ^a	Yield (%) ^b	Mp (t ^o C)
3a	92	60	6c	53	>320
3b	83	oil	6d	75	>320
3c	62	oil	9a	97	330 ^f
3d	97	82	9b	77	>320
4a	64	178 ^c	9c	88	>320
4b	53	124 ^d	9d	63	>320
4c	81	224	11a	100	222
4d	22	135 ^e	11b	83	250
6a	80	320 ^f	11c	90	320 ^f
6b	56	>320	11d	73	252

^a Satisfactory elemental combustion analyses and mass, IR and ¹H NMR spectral data were obtained for all new compounds. ^b Yields are unoptimised. ^c Monohydrate. ^d With resolidification and remelting at 213 °C. ^e With resolidification and remelting at 221 °C. ^f Decomp.

Exposure of the pyrazolocinnoline *N*-oxides **6a-d** to aqueous alkaline sodium hypochlorite at room temperature for 0.5 h resulted in their smooth conversion in high yield (Table 1) into the corresponding previously unknown cinnoline-4-carboxylic acid 1-*N*-oxides **11a-d**. These processes are readily explained (Scheme 1) in terms of the oxidative transformation of the pyrazolocinnoline *N*-oxides **6a-d** (presumably reacting in the pyrazolinone tautomeric forms **7a-d**) into transient pyrazolone structures **8a-d** which suffer spontaneous hydrolytic ring cleavage with loss of nitrogen, in the alkaline medium. The efficiency of these transformations demonstrates that the apparent existence of the pyrazolocinnoline *N*-oxides largely as the hydroxy tautomers **6a-d** does not preclude their ability to react in the pyrazolinone tautomeric forms **7a-d** when the occasion demands. Preliminary investigations indicate that the analogous oxidative cleavage reactions of the parent pyrazolocinnolines **9a-d** are not so straightforward.

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