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Novel Cycloaddition Reactions of Isocyanides with Diphenylacetylene Using Transition-metal Complexes

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Summary Reaction of tetrakis(aryl isocyanide)nickel complexes (Ia—c) with diphenylacetylene gave the diiminocyclobutenes (IIIa—c); on pyrolysis, the acetylene complex (Va) gave (IIIa) as sole product, while with (Vd) the major product was iminocyclopentadiene (VId).

WE previously reported the cycloadditions of isocyanide to acetylenes substituted with electron-withdrawing groups to give tri-iminocyclopentene and pyrindine derivatives, whereas aryl and alkyl substituted acetylenes were unreactive, probably owing to the lower electron affinity of their triple bonds.¹ As the isocyano-group is isoelectronic with carbon monoxide and reactions of various metal carbonyls with carbon monoxide and reactions of various metal carbonyls with acetylenes are well known,² it was expected that the reactions of isocyanides with diphenylacetylene would give cyclized products via co-ordination to a transition metal.

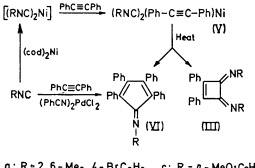
An equimolar mixture of (Ia)³ and diphenylacetylene (II) was refluxed in toluene for 1 h under nitrogen. Addition of n-hexane and chromatography of the filtrate yielded the yellow di-iminocyclobutene (IIIa), m.p. 205-206°, [30% based on (II)]. Its structure was established by elemental analysis, molecular weight determination, and its i.r., u.v.,

(III) (立)

a; $R = 2.6 - Me_2$, 4 - BrC₆H₂ b;R=Ph c;R=p-MeO·C₆H₄

and n.m.r. spectra. Hydrolysis of (IIIa) with aqueous HCl gave diphenylcyclobutenedione (IV), identical with an authentic sample.⁴ The tetrakis(aryl isocyanide)nickel complexes (Ib and c) reacted with acetylene (II) to give the diimino-cyclobutenes (IIIb), m.p. 158-160° (90%) and (IIIc), m.p. 169-171° (22%). Under similar conditions, tetrakis-(t-butyl isocyanide)nickel⁵ and tetrakis-(2,6-dimethylphenyl isocyanide)nickel were unreactive to (II).

A mixed ligand complex of isocyanide and acetylene formed initially is possibly an important intermediate in the formation of (III). Bis(cyclo-octa-1,5-diene)nickel [(cod),-Ni] reacted readily with 2 mol. equiv. of the isocyanide in toluene at room temperature and subsequent addition of I mol. equiv. of (II) gave yellow microcrystals of (Va) $[v(N=C) 2160 \text{ and } 2100 \text{ cm}^{-1}; v(C=C) 1830 \text{ cm}^{-1}]$ whose i.r. spectrum is similar to that of the known complex (Ve).6 Heating of complex (Va) at 80-100° in toluene gave di-iminocyclobutene (IIIa) (30%). However, pyrolysis of complex (Vd), prepared similarly, in toluene gave the cyclopentadiene (VId) as dark violet crystals, m.p. 190-



a; $R = 2, 6 - Me_2, 4 - BrC_6H_2$ c; R = p - MeO·C6H4 d; $R = 2, 6 - Me_2C_6H_3$ e; R = Bu^t

191° (65%), together with a small amount of (IIId), m.p. 175-177°. The structure of (VId) was supported by elemental analyses and hydrolysis with aqueous HCl to tetracyclone, as well as its i.r., u.v., and n.m.r. spectral data.

Reaction of the isocyanides R-NC (R as in a, c, and d) with (II) in the presence of bis(benzonitrile)palladium chloride (molar ratio 1:2:1) gave the iminocyclopentadienes (VIa), m.p. 186-187° (VIc), m.p. 206-207°, and (VId) in 21, 10, and 14% yield, respectively, in addition to the known palladium complex of tetraphenylcyclobutadiene, $Pd_{3}Cl_{3}(Ph_{4}C_{4})_{2}$,⁷ (30-40% based on Pd).

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