

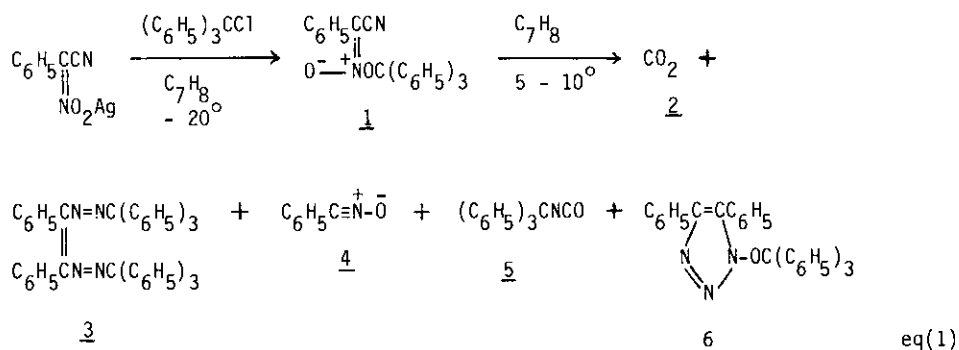
THE FORMATION AND FACILE CONVERSION OF TRITYL PHENYLCYANOMETHANE NITRONATE

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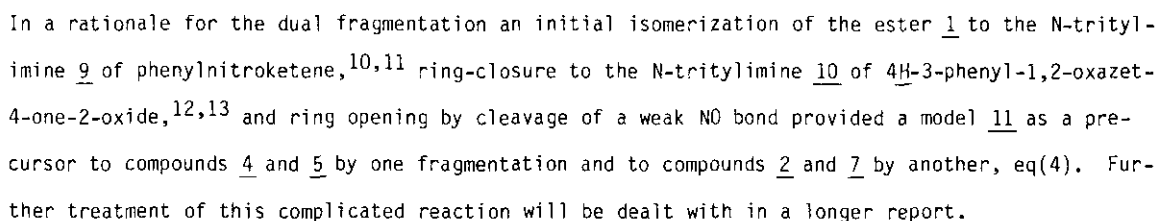
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Abstract - Trityl chloride and silver phenylcyanomethane nitronate in toluene at -20°C gave trityl phenylcyanomethane nitronate 1, an unstable ester that rearranged and fragmented at $5-10^{\circ}\text{C}$ to 1-triphenylmethoxy-4,5-diphenyl-1,2,3-triazole 6 with coproducts carbon dioxide, α,α' -bis(triphenylmethaneazo)stilbene 3, benzonitrile-N-oxide 4, and trityl isocyanate 5.

A product $\text{C}_{33}\text{H}_{25}\text{N}_3\text{O}$ obtained from an unexplained reaction¹ between silver phenylcyanomethane nitronate and trityl chloride in toluene at 0°C has been identified by X-ray crystallographic analysis as 1-triphenylmethoxy-4,5-diphenyl-1,2,3-triazole 6². Coproducts included carbon dioxide 2, α,α' -bis(triphenylmethaneazo)stilbene 3, benzonitrile-N-oxide 4, and trityl isocyanate 5, eq(1)¹. Trityl phenylcyanomethane nitronate 1 has now been identified as the initial product of the reaction at -20°C by comparative infra-red spectroscopic analysis (-20°C) of the ester 1 and the related benzyl³ and benzhydryl⁴ phenylcyanomethane nitronates (decomposition at 25°C)⁵. The conversion, eq(1), is important as a rare example of a transfer at low temperature of the oxygen atoms in a nitronate group to a carbon atom and is pertinent to explosive reactions of nitro compounds in which carbon dioxide may be similarly produced.



Product analysis revealed two fragmentation modes for the ester 1: one afforded benzonitrile-N-oxide 4 and trityl isocyanate 5, while the other differed by an interchange of oxo and tritylimino substituents (X, Y) to afford carbon dioxide 2 and benzonitrile-N-tritylimine 7, eq(2)^{6,7}. The

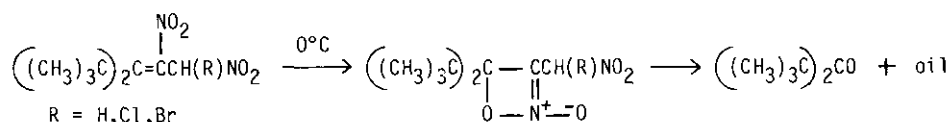
$$\begin{array}{ccc} \text{C}_6\text{H}_5\text{CCN} & \longrightarrow & \text{C}_6\text{H}_5\text{C}\equiv\text{NX} + \text{Y}=\text{C}=\text{O} \\ \text{O} \parallel & & \\ \text{O}-\text{NOC}(\text{C}_6\text{H}_5)_3 & & \\ \underline{1} & \underline{4} \text{ X} = \text{O} & \underline{5} \text{ Y} = (\text{C}_6\text{H}_5)_3\text{CN} \\ & \underline{7} \text{ X} = \text{NC}(\text{C}_6\text{H}_5)_3 & \underline{2} \text{ Y} = \text{O} \end{array} \quad \text{eq(2)}$$


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REFERENCES

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4. R. L. Shriner and G. B. Brown, *J. Org. Chem.*, 1938, 2, 561.
5. Since dual absorption bands near 1550 and 1350 cm^{-1} ($-\text{NO}_2$) were absent at -20°C the formation of the nitronate esters predominated over the formation of the isomeric nitro compounds to be derived from C- and N-alkylation.
6. The nitrilimine 7 was previously abandoned as a precursor to the bisazostilbene 3 when benzonitrile-N-oxide 4 and trityl isocyanate 5 failed to give an adduct that might have fragmented to the nitrilimine 7 and carbon dioxide⁷.
7. N. E. Alexandrou, *J. Org. Chem.*, 1965, 30, 1335.
8. Attempts to realize independently a dipolar addition between benzonitrile-N-tritylimine 7 and benzonitrile-N-oxide 4 were abandoned when (a) tritylhydrazine and benzaldehyde failed to convert to the expected hydrazone (to be oxidized to the imine 7) and (b) N-tritylbenzohydrazidoyl chloride ($\text{C}_6\text{H}_5\text{C}(\text{Cl})=\text{NNHC}(\text{C}_6\text{H}_5)_3$), to be dehydrochlorinated to the imine 7, was not obtained from β -N-tritylbenzohydrazide ($\text{C}_6\text{H}_5\text{CONHNHC}(\text{C}_6\text{H}_5)_3$) upon treatment with either thionyl chloride or phosphorus pentachloride.
9. The combined yield (42%) of compounds 3 and 6 is in good qualitative agreement with the yield (50%) reported for carbon dioxide¹.
10. The 1-5 sigmatropic isomerization 1 $\rightarrow$ 8 represented a formal change from O- to N-tritylation of a cyanomethane nitronate anion.
11. Attempts to prepare compound 8 by a reaction between trityl isocyanide and the sodium salt of phenylchloronitromethane were unsuccessful⁷.
12. The formation of 4,4-di-*t*-butyl-3-methyl-4H-1,2-oxazete-2-oxide from 3-*t*-butyl-4,4-dimethyl-2-nitro-2-pentene on standing at 25°C was the first recognized example of an isomerization of an α,β -unsaturated nitro compound to a four-membered heterocycle. It underwent facile dissociation to give di-*t*-butyl ketone and an oil¹³.



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