Thermal Stability and Fluxional Behaviour of Bicyclic Sulphur-Nitrogen Structures of Formula EN_5S_3 ($E = CR$, PR_2 , or SO_2^-)

René T. Boeré, Richard T. Oakley,* and Maurice Shevalier

Guelph Waterloo Centre for Graduate Work in Chemistry, Guelph Campus, Department of Chemistry and Biochemistry, University of Guelph, Guelph, Ontario N 1G 2W1, Canada

Two thermally induced N-S bond cleavage processes are available to EN₅S₃ heterocycles: (i) an irreversible loss of NSN to form EN_3S_2 (for $E = PR_2$ and SO_2^-) and (ii) a pseudo-degenerate 1,3-nitrogen shift which exchanges all nitrogen atoms (for $E = CPh$).

The often explosive thermal instability of binary sulphurnitrogen compounds is a well recognised but poorly understood feature of their chemistry. In order to gain a deeper insight into the mechanisms of these decompositions, we are currently exploring the relationships between the thermal stabilities and the molecular and electronic structures of closely related compounds. Herein we compare the kinetic stabilities of heterocyclic derivatives based on the bicyclic

framework EN_5S_3 . We also report ¹⁵N n.m.r. spectroscopic evidence for nitrogen atom scrambling in $PhCN₅S₃$, an effect hitherto unobserved in sulphur-nitrogen chemistry.

Recently we reported a kinetic analysis of the thermal decomposition of $\overline{Ph}_2 \overline{PN}_5 S_3 (1)$ (E = PPh₂) to the corresponding dithiatriazine $Ph_2PN_3S_2$ (2) $(E = PPh_2)$ (Scheme 1).¹ An analogous decomposition pathway is observed for the anion $SO_2N_5S_3$ ⁻ (1) $(E = SO_2$ ⁻), which is easily prepared as its

 $(E = CR or PR₂)$

Scheme 2

Figure 1. Time dependence of the ¹⁵N n.m.r. spectrum of PhCN₂*N₃S₃ (in CDCl_c). The $t = 0$ case refers to data collected at -20 °C (where fluxional behaviour is frozen out).

yellow $(\text{Ph}_3\text{P})_2\text{N}^+$ salt \dagger by oxidation of $(\text{Ph}_3\text{P})_2\text{N}^+$ SO₂N₃S₂ with PhICl₂ in acetonitrile to yield $(\text{Ph}_3\text{P})_2\text{N}$ +SO₂N₃S₂Cl₂-,[†] followed by treatment with $(Me_3SiN)_2S$ (overall yield 59%). Thus, gentle warming of solutions of $(\text{Ph}_3\text{P})_2\text{N}$ +SO₂N₅S₃- in acetonitrile effects the reversion of the bicyclic anion to the monocyclic $SO_2N_3S_2$ species,² which can be isolated in 60% yield.

In comparison to the above EN_5S_3 derivatives, the isoelectronic species PhCN₅S₃³ (1) $(E = CPh)$ is more thermally stable. It can be sublimed without decomposition at 80 "C and 0.1 Torr , and although it does slowly decompose on prolonged reflux in acetonitrile, the dithiatriazine PhCN₃S₂ (2) $(E =$ CPh) is not formed; instead the major isolable products are

Figure 2. ¹⁵N N.m.r. spectrum (in CDCl₃) of $Ph_2PN_2^*NS_2$ prepared by the thermolysis of $Ph_2PN_2*N_3S_3$.

 S_4N_4 and benzonitrile. Similar results have been noted when E $= CNR₂$.⁴ The resistance of RCN₅S₃ derivatives to thermolysis is surprising in view of the long SN bonds to the bridging NSN unit, $3-5$ but may be related to the differences in the electronic structures of dithiatriazines where $E = CR$ and $E =$ $PR₂$ or $SO₂$ ⁻. While the latter are known to possess singlet ground states,^{2,6} π ground states have been predicted for a variety of $RCN₃S₂$ structures.^{4,7,8} Accordingly formation of $RCN₃S₂$ *via* the concerted elimination of a singlet⁹ NSN fragment from the RCN_5S_3 structure would be symmetry forbidden.

In order to examine more carefully the structural stability **of** PhCN₅S₃ we prepared the specifically ¹⁵N labelled¹⁰ derivative PhCN₂*N₃S₃ (*N = 99% ¹⁵N) by the method illustrated in Scheme 2. Consistently with the expected labelling pattern the ¹⁵N n.m.r. spectrum \ddagger (at -20° C) of the material prepared at -20°C shows two signals at **6** 330.1 (2N) and 54.9 (1N). However, when the solution is warmed to room temperature a third signal (at **6** 176.1), generated by the slow incorporation

t The elemental composition of this compound has been confirmed **by** elemental analysis.

^{\$} **15N** N.m.r. spectra were recorded at **40.5** MHz on a Bruker **WH-400** spectrometer, using 30 s pulse delays and a pulse width of 33 μ s. Chemical shifts are reported with reference to liquid ammonia at 25° C.

of 15N into the carbon-bound nitrogen sites, slowly emerges from the base line. Figure 1 illustrates the progress of this nitrogen exchange over a **14** h period. The limiting spectrum exhibits the $2:2:1$ intensity pattern expected for complete scrambling of ¹⁵N and ¹⁴N nuclei. In contrast to the above behaviour $Ph_2PN_2*N_3S_3$ (also prepared according to Scheme 2) undergoes no observable nitrogen exchange at room temperature; only two signals at **8** 322.0 (2N) and 79.2 (1N) are present, although after 16 h some decomposition to $Ph₂PN₂[*]NS₂$ is apparent. Furthermore, ¹⁵N n.m.r. analysis of the phosphadithiatriazine isolated from the thermolysis of $Ph_2PN_2*N_3S_3$ in refluxing toluene shows (by comparison with the known ¹⁵N n.m.r. spectrum of $Ph_2P^*N_3S_2$ ⁶ only small amounts of 15N incorporation into the phosphorus-bound nitrogen sites (Figure 2).

While the observation of nitrogen scrambling in $PhCN₅S₃$ is without precedent in sulphur-nitrogen chemistry, the possibility of such a phenomenon was recognized some years ago by Bartetzko and Gleiter for the isoelectronic S_4N_5 ⁺ cation.¹¹ Accordingly the fluxional process that leads to interchange of all nitrogen atoms in PhCN₅S₃ can be viewed as involving a series of pseudo-degenerate 1,3-nitrogen shifts, as depicted in Scheme **3.** While further theoretical and experimental studies are required to establish the activation barriers for such 1,3-shifts in EN_5S_3 structures, and the relative rates of the nitrogen exchange and NSN elimination pathways, the present results indicate that the scrambling process predominates when $E = CR$. When the NSN elimination channel is allowed, and when the 1,3-shift mechanism involves a high

energy intermediate *(e.g.* a five-co-ordinate phosphorus or sulphur), only thermal decomposition to EN_3S_2 is observed.

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