

SYNTHESIS AND REACTIONS OF 1,2,3-TRIAZINES

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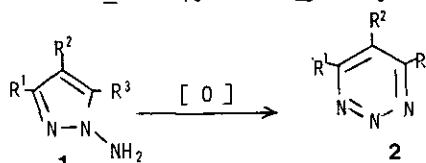
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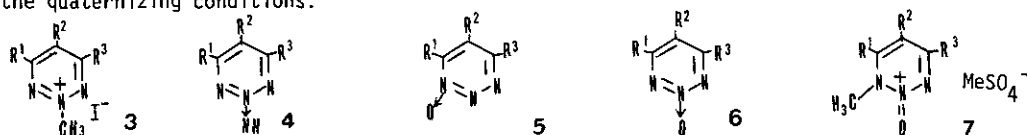
Monocyclic 1,2,3-triazines were synthesized and their properties were investigated.

Synthesis Oxidation of methyl- and/or phenyl-substituted *N*-aminopyrazoles(**1**) using $\text{Pb}(\text{AcO})_4$ afforded 1,2,3-triazines(**2**) although the method afforded only a trace amount of unsubstituted one (**2**, $\text{R}^1=\text{R}^2=\text{R}^3=\text{H}$). The latter was obtained by nickel peroxide-AcOH oxidation of **1** in 18% yield.

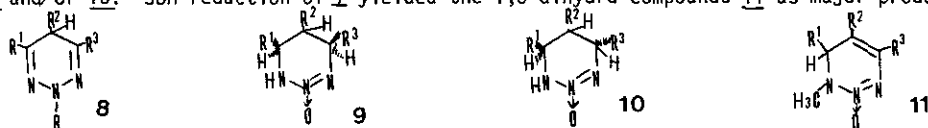


Isolated 1,2,3-triazines were moderately stable and the exact molecular structure of unsubstituted one was examined by X-ray crystallography.

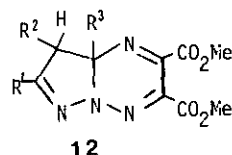
Quaternization Methylation of **2** with MeI afforded 2-methylated salts(**3**). *N*-Amination with *O*-mesitylenesulfonylhydroxylamine gave Δ -imino compounds(**4**) which existed in the monomeric form in a solution(CHCl_3). Although *N*-oxidation of **2** with *m*-chloroperbenzoic acid afforded the corresponding 1-oxides(**5**) and 2-oxides(**6**) when $\text{R}^1 \neq \text{Ph}$, only 2-oxides were obtained when $\text{R}^1=\text{R}^2=\text{Ph}$. The molecular structure of 4,5,6-triphenyl-1,2,3-triazine 2-oxide was confirmed by X-ray analysis. The methylation of the oxide **6** with Me_2SO_4 afforded *N*-methylated product(**7**) although **5** decomposed under the quaternizing conditions.



Reduction Catalytic reduction of **2**, **5**, and **6** gave the 2,5-dihydro compounds(**8**, $\text{R}=\text{H}$). Reduction of **2** and **3** with sodium borohydride(SBH) also yielded **8**($\text{R}=\text{H}$ and Me). The major products of SBH reduction of the oxide **5** were **2** and **8**, while the major products from **6** were 1,4,5,6-tetrahydro compounds **9** and/or **10**. SBH reduction of **7** yielded the 1,6-dihydro compounds **11** as major products.



Other Reactions 1,3-Dipolar cycloaddition of **4** with dimethyl acetylenedicarboxylate yielded the rearranged adducts(**12**). Results of other reactions will be stated.



References: JCS Chem.Comm.,1980, 1182; JCS Chem.Comm.,1981, 1174;
Heterocycles, 17, 317 (1982).