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Pyrolysis of 2-Bis(methylthio)methylpyridine S-Oxides. Synthesis of Substituted Pyridinecarbaldehydes

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Summary A convenient preparation of pyridinecarbaldehydes by pyrolysis of 2-bis(methylthio)methylpyridine S-oxides is described.

In connection with other studies, we needed to prepare substituted pyridine-2-carbaldehydes. We herein report: (i) use of the carbanion of methyl methylsulphinylmethyl sulphide (1) to prepare heterocyclic aldehydes from the corresponding halide and (ii) the unexpected thermal rearrangement of the intermediate aldehyde dithioacetals. min followed by addition of 2-bromopyridine and stirring for 24 h at 30–35° gave (100%) the S-oxide (2; R = H), δ (CDCl₃) 2·28 and 2·32 (s, MeS), 2·51 (s, MeSO), 8·6 (d, [6-H, J 5 Hz), and 7·1–8·8 (m, py-H)]. Attempted purification of (2) by distillation did not afford the expected protected aldehyde but rather the free aldehyde (4; R = H) (74% overall yield), b.p. 181–182°, and dimethyl disulphide (70%), b.p. 116–117°, δ (CCl₄) 2·39 (s, MeS). Typical results are summarised in the Table.

Alternatively, thioacetals such as (2; R = H) can be

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2-Halogen R	opyrid ine X	Reaction t /h	ime Temp. /°C	Yield of (2) (%) ^a	Rearr. temp. /°C	Compound (4) (%) ^a m.p. (b.p./mmHg)
H H 6-Br	2-Br 2-Cl 2-Br	24 48 24	30 30 30	~100 b b	200 200 150	$\begin{array}{c} 74 \\ 12 \\ 45^{\circ} \\ 45^{\circ} \end{array} \right\} (181 - 182^{\circ} / 760)$
6-Br	2-Br 2-Br	${24 \\ 1}$	$\left. \begin{array}{c} 30\\ reflux \end{array} \right\}$	64	150	$\left. { { { { { { { { 5 } } } } \atop { { 6 1 } } } } } } \right\}$ 77—78°ª
6-C1	2-C1	$\left\{ egin{matrix} 24 \\ 1 \\ 48 \end{array} ight.$	$\begin{array}{c} 30\\ \text{reflux}\\ 30 \end{array}$	b	150	42 6970°d
н	3-Br		reflux	ь	240	10° (95—97°/15)
н	4-Cl	48	30	b	200	43 (77—78°/12)

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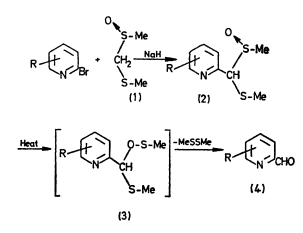
^a Yield data were not optimized. ^b Detected by n.m.r., but not isolated. ^c Isolation of aldehyde was by bisulphite extraction. ^d Satisfactory analyses were obtained. ^e Yield estimated by n.m.r. analysis of the mixture.

Ogura and Tsuchihashi¹ first described the use of reagent (1) for the preparation of alkyl aldehydes and Schlessinger, *et al.*,² have extended this work. Heterocyclic aldehydes have been prepared from the corresponding halide *via* an organometallic intermediate and ethyl orthoformate³ or *NN*-dimethylformamide,⁴ but this route is limited. Since attempts to prepare pyridine-2-carbaldehyde *via* 1,3-dithians⁵ failed, we investigated nucleophilic attack of the anion of (1) on simple 2-halogenopyridines. Reaction of (1) with sodium hydride in dimethoxyethane at 30° for 30

hydrolysed by treatment with concentrated hydrochloric acid in tetrahydrofuran at room temperature for several hours. Addition of $HgCl_2$ facilitates the work-up by eliminating disulphide contaminants. Thus, after 2 h under these conditions, the crude thioacetals afforded aldehydes (e.g. 4; R = H) in ca. 60—70% yield. Hydrolytic work-up procedures have no apparent advantage, unless the resultant aldehyde is thermally labile.

Nucleophilic substitution of the halogen of the 2-halogenopyridine with the protected formyl function proceeds

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smoothly at $25-85^{\circ}$ to give compound (2), which is stable under the mild reaction conditions. In general, we find that: (i) Br⁻ is displaced more rapidly than Cl⁻, (ii) only

one halogen per pyridyl ring is easily substituted, and (iii) the 2- and 4-positions are more reactive than the 3position. Upon heating to 150°-200°, compound (2cleanly rearranged to compound (4) and dimethyl disulphide in reasonable yields, which were not maximized. We suggest that a thermal sulphur to oxygen 1,2-migration occurs $[(2) \rightarrow (3)]$ probably proceeding via ionic decomposition followed by elimination of dimethyl disulphide.⁶ A similar rearrangement of methoxymethyl phenyl sulphoxide to give exclusively the OO-acetal has been recently reported.7

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