Note on Ion Exchange with Mercury(II)

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Little is known about the ion exchange behaviour of mercury in spite of the great interest in tracing its fate in polluted waters, etc. We have started an investigation of the ion exchange properties of Hg(II) using ion exchangers with various types of ion exchanging sites. Since this work has to be discontinued at present we think the results obtained so far may be of some general interest.

Batch experiments were performed at 25°C, the ionic medium being 0.1 M HNO₃. The concentration of Hg(II) varied between 10⁻⁶ and 10⁻⁴ M. Mercury was determined radiometrically as ⁴⁰⁷Hg employing a Tracerlab Compumatic Scaler and a Tracerlab scintillation counter model Sc-57A for γ-rays. After equilibrium had been attained the two phases were separated by centrifugation and filtration through glass-wool. The filtrate was tested with a Tyndall lamp for colloids present. Only clear samples were accepted for analysis.

Table 1. Capacities (mequiv./g resin) of partially air dried ion exchange resins.

<table>
<thead>
<tr>
<th>Resin</th>
<th>Capacity</th>
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<tr>
<td>Dowex 50W X-8</td>
<td>3.78±0.01</td>
</tr>
<tr>
<td>Dowex A1</td>
<td>4.49±0.05</td>
</tr>
<tr>
<td>Amberlite IRC-50</td>
<td>6.96±0.1</td>
</tr>
<tr>
<td>Sephadex SH</td>
<td>0.17±0.01</td>
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</tbody>
</table>

The resins used were Dowex 50W X-8, Dowex A1, Amberlite IRC-50, and Sephadex SH. Table 1 gives the capacities of the resins. In the case of Dowex 50 this was determined by emf-titration; in all other cases by back titration with acid after the addition of excess sodium hydroxide solution to the acid form of the resin.

Fig. 1 gives log x against log [Hg²⁺]eq for the three resins: Dowex 50, Amberlite IRC-50, and Dowex A1; x is the equilibrium quotient for the reaction

\[ \text{Hg}^{2+} + 2\text{HR} \rightleftharpoons \text{HgR}_2 + 2\text{H}^+ \]  

and

\[ x = \frac{X_{\text{HgR}_2}}{X_{\text{H}^+}^2} \frac{[\text{H}^+]^2}{[\text{Hg}^{2+}]} \]

The results with Sephadex SH are not given in Fig. 1 because in this case the amount of mercury taken up by the resin always exceeded the experimentally determined capacity. A possible explanation is partial reduction of mercury by the resin, which is known to show electron exchange properties.

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Preparation of Methyl-substituted Trithienylphosphines and Their Sulfides

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In connection with current studies of nuclear magnetic resonance spectra of aromatic and heteroaromatic phosphine derivatives it was of interest to obtain information concerning the signs and magnitudes of the expected four-, five-, and six-bond methyl proton-ring phosphorus long-range spin-spin coupling constants in phosphorus substituted methylthiophenes. In this paper the preparations of the six hitherto unknown isomeric tri-(methylthienyl)-phosphines (Ia—VIa) and their sulfides (Ib—VIb) are described.

\[
\begin{align*}
\text{I} & \quad \text{II} & \quad \text{III} \\
\text{IV} & \quad \text{V} & \quad \text{VI}
\end{align*}
\]

\(a: X = \text{lone pair}\)
\(b: X = 8\)

The phosphines were prepared in 28—66 % yields from the corresponding methylthienylthium derivatives, obtained either by low-temperature metal-halogen interconversion of the methylbromothiophenes (compounds Ia—IVA) or by direct metallation of the methylthiophenes (compounds Va and VIa) with butyllithium, followed by reaction with phosphorus tribromide. The phosphines were converted to their sulfides in quantitative yields by reaction with elementary sulfur in solutions of methanol or chloroform. Structures and purities of the isolated compounds were checked by \(^1\)H NMR spectroscopy, mass spectrometry, or by microanalyses. All compounds were obtained in a pure state, with the exception of the mixture of 4-methyl-2-thienyllithium (ca. 83 %) and 3-methyl-2-thienyllithium (ca. 17 %) resulting from metallation of 3-methylthiophene with butyllithium which only allowed the isolation of tri-(4-methyl-2-thienyl)-phosphine (Va) as an oily mixture (ca. 54 %) with some of the isomers di-(4-methyl-2-thienyl)-3-methyl-2-thienyl phosphine (ca. 40 %) and di-(3-methyl-2-thienyl)-4-methyl-2-thienyl phosphine (ca. 6 %) as deduced from \(^1\)H NMR spectra. Separation by means of gas chromatography or thin layer chromatography could not be achieved. Attempts to prepare (Va) in a pure state by other synthetic routes were not carried out as the complete analysis of its \(^1\)H NMR spectrum could be performed on the mixture of isomers. However, the corresponding sulfide (Vb), prepared from the isomeric mixture, could easily be obtained in a pure crystalline state by fractional recrystallization. In the preparations of the phosphines (Ia—IIIa) low-boiling products, later shown to result from a thiophene ring-opening reaction, were obtained.

Experimental. Boiling and melting points are uncorrected.

Starting materials. 2- and 3-Methylthiophene were commercially available (Schuchardt, Munich) and used without further purification. 2-Methyl-3-bromothiophene was prepared according to Gronowitz et al., using dimethyl sulfate instead of methyl p-toluenesulfonate, b.p. 55/9 mmHg (b.p. 56—58/10 mmHg), yield 88 % (84 %). 2-Methyl-3-bromothiophene was obtained from 4-bromo-3-thienyllithium and dimethyl sulfate, b.p. 58—60/10 mmHg (b.p. 181—184/770 mmHg), yield 90 %.

2-Methyl-4-bromothiophene was obtained from 4-bromo-2-thienyllithium and dimethyl sulfide (at —70 °C) as an isomer-pure product (> 95 %) which is consistent with the observations of Gronowitz and Frosting: b.p. 58—59/9 mmHg (b.p. 61—62/11 mmHg), yield 81 %. 3-Methyl-2-bromothiophene was prepared by the method of Steinkopf and Jacob.