## Certain Diethers and Triethers<sup>1</sup>

By Lucile Liston and William M. Dehn

For the purpose of testing their solubility on cellulose esters, ethers of the types ROCH<sub>2</sub>CH<sub>2</sub>-OR' and ROCH2CH2OCH2CH2OR' were prepared from monoethyl and monobutyl ethers of glycol and diethylene glycol. During their formation, distillation and storage certain decompositions were observed, hence these ethers are considered good material for study of hydrolysis. In general, quantitative yields of the ethers were not obtained: lower and higher boiling fractions than the ether itself were always obtained and such fractions were increased by repeated distillation. Also whereas the first distillations of allyl diethers gave 40-67% yields, after some months of storage, being redistilled, they decomposed energetically into other products. Some of these decompositions were of explosive violence. These and other pyrolytic decompositions will be investigated.

The ethers of glycol and diethylene glycol were converted into their sodium compounds and then were refluxed with the alkyl chloride. Advantageously comminuted sodium was prepared by refluxing toluene containing sodium in a flask on an oil-bath. The flask was then stoppered and shaken vigorously while being held by two towels. The mixture of sodium and toluene was used directly, or the toluene was displaced by petroleum ether or the hydrocarbon was practically decanted. First the glycol monoether, then the alkyl halide were added from the top of a condenser at such rates that the reactions could be controlled. After heating in an oil-bath to complete the reactions, the mixtures were either distilled directly

Yield, Carbon, % Hydrogen, % % Calcd. Found Calcd. Found B. p., °C. Ethylene glycol diethers Allyl, ethyl 139 - 14260 64.59 64.65 10.84 10.83 n-Pentyl, ethyl 180-183 48 67.42 67.08 12.5812.37183-184 11.47Allyl, butyl 67 68.28 68.28 11.53 #-Pentyl, butyl 221-222 43 70.16 70.5912.8513.07 Butyl, ethyl 164 - 16590 65.69 65.44 12.43 12.39 Diethylene glycol diethers 200-203 Allvl. ethvl 62.00 62.38 10.41 10.29 40 Butyl, ethyl 218 - 21940 63.11 63.23 11.68 11.84 n-Pentyl, ethyl 121-12418 30 64.65 64.44 11.84 11.76

(1) Original manuscript received August 24, 1934:

or the contained salt was first removed by filtering or extracting with water.

Fractionations of these complex ethers gave indications of thermal decomposition, thus accounting for some of the low yields, especially in cases of the unsymmetrical complex ethers.

Ethylene glycol diethyl ether boiling at  $124^{\circ}$  was obtained in 70% yields; ethylene glycol dibutyl ether boiling at  $204^{\circ}$  was obtained in 75% yields.

**Summary.**—A number of new complex glycol ethers have been prepared and analyzed.<sup>2</sup>

(2) For other studies along this line, see Conn, Collett and Lazzell, THIS JOURNAL, 54, 4370 (1932); Cretcher and Pittenger, *ibid.*, 47, 163 (1925).

CHEMISTRY LABORATORY

UNIVERSITY OF WASHINGTON SEATTLE, WASH.

RECEIVED MARCH 7, 1938

## Some New Esters by Automatic Processes without Catalysts<sup>1</sup>

BY THOMAS R. LISTON AND WILLIAM M. DEHN

By use of a modification of the Betz-Holden automatic water-removing trap, new esters of the chloroacetic acids were prepared in very satisfactory yields. The modification of apparatus consisted merely of a stopcock at the bottom of the graduated tube so that the calculated volume to be occupied by the water, at the completion of the reaction, could be adjusted easily to reach exactly the overflow level. The advantages of this adjustment are that a visible indication of the progress of the reaction is afforded and the process can be continued to completion, or longer, without surveillance. The flask containing the acid and alcohol is connected with the water trap and the return condenser and the mixture is heated in an oil-bath for such time as is necessary to fill the trap with water to the level of overflow. At the beginning of the process, alcohol, acid, ester, and water will collect in the separator but progressively all but traces of acid and the water will flow back into the flask. With different reactions, the times necessary for heating were between one and eight hours. The products were washed with water containing sodium carbonate, dried with calcium chloride and fractionated, preferably in vacuo. The analyzed esters are (1) Original manuscript received June 19, 1934.

1265

Alcohol	Acid,	Alc.,	Time, hrs.	% ester	B. p. 756 mm. °C.	°C. <sup>B.</sup>	p. Mm.	<b>Sp. gr.</b> 20°/20°	Chlor Calcd.	ine, % Found
Esters of Acetic Acid										
n-Butyl	<b>24</b> 0	300	3	90	124-126					
Esters of Monochloroacetic Acid										
n-Butyl	47	50	1.5	97		94	38			
n-Amyl	100	120	3	88	196	105	35	1.0514	21.59	21.81
s-Amyl	47	50	3	89	185	93	<b>27</b>	1.0475	21.59	21.65
Diethyl methyl	94	100	4	85	184	93	30	1.0469	21.59	21.62
t-Amyl	94	100	2	65	168	88	43	1.0627	21.59	21.61
Esters of Dichloroacetic Acid										
n-Butyl	64	50	3	98		102	37			
n-Amyl	65	60	1.5	94	207	124	48	1.1455	35.63	35.52
s-Amyl	65	60	4	93	198	98	<b>20</b>	1.1210	35.63	35.29
Diethyl methyl	65	60	7	95	197ª	105	<b>4</b> 0	1.1225	35.63	35.35
t-Amyl	65	<b>60</b> °	2	75	180°	93	30	1.1230	35. <b>6</b> 3	35.72
Esters of Trichloroacetic Acid										
n-Butyl	82	50	3	98		111	40			
n-Amyl	61	38	3	91	218	118	30	1.2475	45.59	45.90
s-Amyl	81	60	2.5	95	206	108	30	1.2084	45.59	45.55
Diethyl methyl	163	100	5	80	<b>2</b> 01°	105	25	1.2081	45.59	45.71
t-Amyl	81	60	1.5	84	191*	105	30	1.2505	45.59	45.73
<sup>a</sup> These tend to dec	ompose part	lu into o	lafin and	anid						

" These tend to decompose partly into olefin and acid.

new; some old esters are given for comparisons of vields.

CHEMISTRY LABORATORY

UNIVERSITY OF WASHINGTON SEATTLE, WASHINGTON RECEIVED MARCH 7, 1938

# The Relative Partial Molal Heat Content of Sodium Bromide in Aqueous Solutions at 25°

#### By A. L. ROBINSON

Harned and Crawford<sup>1</sup> have measured recently the electromotive forces of the cells

Ag-AgBr/NaBr(m)/Na<sub>x</sub>Hg(m < 0.1)/NaBr(0.1)/AgBr-Ag at temperatures from 0 to 40° and from these have calculated values for the relative partial molal heat contents of sodium bromide. Their  $\overline{L}_2$  values at 25° may be compared with values determined from calorimetric measurements extending to 0.00016 m.<sup>2</sup>

m	<u>L</u> 2, e. m. f.	$\overline{L}_2$ , cal.
0.1	101	76
.2	79	56
.5	55	-32
1.0	-211	-208
1.5	-421	384
2.0	-587	- 540
2.5	-719	-656
3.0	-819	-753
3.5	-857	
4.0	887	

(1) H. S. Harned and C. C. Crawford, THIS JOURNAL, 59, 1903 (1937).

(2) H. Hammerschmid and A. L. Robinson, ibid., 54, 3120 (1932).

The agreement is fair although the differences are somewhat larger than the usual estimates of precision of the two methods.

DEPARTMENT OF CHEMISTRY RECEIVED APRIL 6, 1938 UNIVERSITY OF PITTSBURGH PTITSBURGH, PENNA.

## A Sensitive Manostat for Low Gas Pressures. A Correction

### By Theodore Soller, Seymore Goldwasser and Ralph A. Beebe

We are indebted to Professor Earl B. Working for calling our attention to the fact that the diagram of the photoelectric relay circuit given in our paper: "A Sensitive Manostat for Low Pressures and its Application to the Adsorption of Hydrogen and Deuterium on Copper,"<sup>1</sup> should contain a grid leak, that is, a high resistance of the order of 10<sup>8</sup> ohms, between the grid and the cathode of the 6C6 tube. With infinite resistance in the grid circuit, this tube would block.

We have found that under ordinary conditions of humidity, the leakage resistance of the condenser and the sockets is sufficient to prevent blocking, so it has not been necessary for us to introduce a separate resistor. But under conditions of extremely good insulation it would be necessary to introduce a separate high resistance of this order of magnitude.

(1) THIS JOURNAL, 58, 1703 (1936).