## **161.** A Convenient Method of Preparation of Certain Primary Amines.

By R. Brown and W. E. Jones.

Various primary amines, uncontaminated by the corresponding secondary and tertiary compounds, are readily obtained by condensation of O-methylhydroxylamine (1 mol.) with 2 mols. of alkylmagnesium chlorides or preferably bromides, but not with iodides. The yields vary from 40 to 90%. The reaction is also applicable to aliphatic dibromides; e.g., cadaverine is obtainable in 68% yield from pentamethylene bromide.

An investigation requiring considerable quantities of tert.-butylamine led to an examination of its preparation on a moderately large scale. Van Erp (Rec. Trav. chim., 1895, 14, 16) and Brander (ibid., 1918, 37, 67) prepared it respectively by the Hofmann degradation of pivalic amide and by heating tert.-butyl chloride with ammonia, but in both cases the yields were extremely small. Coleman and Yager (J. Amer. Chem. Soc., 1929, 51, 567) obtained it in 60% yield by condensation of tert.-butylmagnesium chloride with chloroamine, but the instability of the latter rendered this method unsuitable for our purpose. Sheverdina and Kocheshkov (J. Gen. Chem. Russia, 1938, 8, 1825), however, overcame this difficulty by replacing the chloroamine by the stable O-methylhydroxylamine, two mols. of Grignard compound being required.

$$2RMgCl + MeO\cdot NH_2 = RNH\cdot MgCl + RH + MeOMgCl$$

The experiments of Sheverdina and Kocheshkov (loc. cit.) were carried out on a very small scale (0.03 molar) and at such great dilution as to make larger-scale preparation impracticable, but suitable modifications have permitted an increase to 3-molar scale, the yield being 70% calculated on O-methylhydroxylamine.

A number of other halides have also been examined under the same conditions, the results being shown in the Table, together with comparative yields obtained by Sheverdina and Kocheshkov (loc. cit.) and by Coleman et al. (loc. cit. and J. Amer. Chem. Soc., 1928, 50, 1193; 1933, 55, 3669; 1936, 58, 27), all the work of the latter authors involving the use of chloroamine. In contrast to the experience of Coleman and his colleagues with chloroamine the yields with bromides were slightly better than when using the corresponding chlorides, but the result with isoamyl iodide confirms the conclusions of all previous investigators that the use of iodides in this type of reaction is to be avoided. No explanation of this phenomenon could be found, all attempts, for example, to correlate the yield of amine with the relative proportions of RMgX and R<sub>2</sub>Mg in the Grignard solution being unavailing.

The method finally adopted and exemplified by *tert*.-butylamine in the experimental section is rapid and easy to operate, and permits the facile preparation of a number of primary amines which are otherwise difficult to prepare. A further advantage is that the products are free from secondary and tertiary amines, although ammonia always appears to be formed.

	9	% Yield R·NH	2.		% Yield R·NH <sub>2</sub> .		
	Sheverdina				Sheverdina		
Halide.	Present authors.	and Kocheshkov.	Coleman $et al.$	Halide.	Present authors.	and Kocheshkov.	Coleman et al.
Ethyl bromide	81 *	66.6 *	27.7 *	isoAmvl chloride	60	80.1 *	$55 \cdot 2$
<i>n</i> -Propyl bromide	85 *		27.0 *	isoAmyl bromide	71		
<i>n</i> -Butyl chloride	58	-	58.9	isoAmyl iodide	Trace	5·3 *	10.9
<i>n</i> -Butyl bromide	63		$27 \cdot 2$	tertAmyl chloride	48		$66 \cdot 2$
isoButyl bromide	90			Allyl bromide	40		
tertButyl chloride	70	73.6 *	$60 \cdot 2$	Benzyl chloride	57	_	92
n-Amyl chloride	46	****		2-Phenylethyl chloride	68		74.0
<i>n</i> -Amyl bromide	65			Cinnamyl chloride	Trace		14

<sup>\*</sup> Isolated as, and yields calculated on, hydrochlorides.

In addition to the mono-halides listed in the Table experiments were carried out with pentamethylene bromide, hexamethylene bromide, and decamethylene bromide. In agreement with the work of Hilper and Grüttner (Ber., 1914, 47, 178) and of Bygdén (Ber., 1915, 48, 1238) it was found that attempts to form the Grignard compounds under normal anhydrous conditions resulted in the formation of highly polymerised compounds

completely unreactive towards O-methylhydroxylamine which was recovered unchanged. When, however, a small quantity of water was present, as suggested by Bygdén (loc. cit.), the normal Grignard compounds were formed, and these reacted readily with O-methylhydroxylamine. The optimum quantity of water required in the ether was found to be 01%. Under these conditions the yields of cadaverine, hexamethylenediamine, and decamethylenediamine were respectively 68%, 51%, and 53% calculated on O-methylhydroxylamine; the method provides a convenient means of preparation of this type of compound.

## EXPERIMENTAL.

tert.-Butylamine.—A solution of O-methylhydroxylamine (Traube, Ohlendorf, and Zander, Ber., 1920, 53, 1477) (141 g., 3 mols.) in anhydrous ether (300 c.c.) was added gradually with vigorous stirring to a solution of tert.-butylmagnesium chloride (from 609 g. of tert.-butyl chloride) in ether (31.), the internal temperature during the addition being maintained at  $-10^{\circ}$  to  $-15^{\circ}$ . After addition was complete the temperature was kept at  $-10^{\circ}$  for a further  $\frac{1}{2}$  hour, then allowed to rise slowly to room temperature, and the reaction mixture finally refluxed for 2 hours. It was then cooled and treated to with  $\frac{5}{2}$  hydroxhoric acid (21), the large transport and the reaction mixture of the support of the small property was expected to small allowed to rise slowly to room temperature, and the reaction mixture finally refluxed for 2 hours. It was then cooled and treated at 0° with 5N- hydrochloric acid (31.), the layers were separated, and the aqueous solution was evaporated to small bulk under reduced pressure. The residue, containing test.-butylamine hydrochloride, magnesium chloride, and a little ammonium chloride, was strongly basified with 50% aqueous potassium hydroxide and the test.-butylamine distilled out, the fraction b. p. 35-70° being collected. The crude amine was dried (KOH) and redistilled. Yield, 153 g. B. p. 40-45°. The amine after several distillations had b. p. 44.4°, a2° 0.6951, n2° 1.3789. The following were prepared: N-Benzoyl derivative, needles from alcohol, m. p. 134° (Found: N. 7-8. C.1H<sub>11</sub>, ON requires N. 7-8%). Picrate, prepared in benzene; needles from ethyl acetate-chloroform, m. p. 197-198° (Found: C. 39.7; H. 4+9; N. 18-4. C. 18-14, C. 18 treated at  $0^{\circ}$  with 5½- hydrochloric acid (3 l.), the layers were separated, and the aqueous solution was evaporated to small

N, 17·7%).

Allylamine.—Allyl bromide gave a 40% yield of the amine, b. p. 58°. It was characterised as its picrate, m. p. 140° (Gabriel and Eschenbach, Ber., 1897, 30, 1125, give m. p. 140—141°).

Benzylamine.—The yield of amine, b. p. 90°/12 mm., prepared from the chloride by the above method was 57%. No reason could be found for this yield being so much lower than that obtained with chloroamine (Coleman et al., loc. cit.). The amine was identified as its picrate, m. p. 194° (Moureu and Lazennec, Bull. Soc. chim., 1906, 35, 1183, give m. p.

2-Phenylethylamine.—The yield of this amine, b. p. 76—78°/10 mm., prepared from the chloride, was 68%. It was

characterised as the picrate, m. p. 167° (Kolshorn, Ber., 1904, 37, 2484, gives m. p. 167—168°).

Cadaverine.—Pentamethylene bromide (23 g.) was added gradually with stirring to magnesium turnings (7.6 g.) in ether (150 c.c.) containing 0.1% of water. The mixture was cooled so that the ether was just refluxing, and when the reaction had moderated the reaction mixture was kept overnight at room temperature and then refluxed for 4 hours. was then cooled to  $-15^{\circ}$  and a solution of O-methylhydroxylamine (4.7 g.) in dry ether (15c.c.) was added gradually with stirring, the temperature being maintained between  $-10^{\circ}$  and  $-15^{\circ}$ . After addition of the O-methylhydroxylamine was complete, the reaction mixture was allowed to attain room temperature and then refluxed for 2 hours. complex was decomposed with 5N-hydrochloric acid at 0°, the aqueous layer separated and made alkaline with 40% aqueous sodium hydroxide, and the diamine steam-distilled out. The distillate was made acid with hydrochloric acid and evaporated to dryness under reduced pressure, and the residue extracted with alcohol containing a trace of water. Evaporation of the alcohol gave the cadaverine hydrochloride; the free amine was liberated by means of 50% aqueous potassium hydroxide, dried over solid potassium hydroxide, and distilled. B. p. 178—180°. Yield 68%. The dipicrate crystallised from water in needles, m. p. 237° (decomp.) [Baumann and Udránszky, Z. physiol. Chem., 1889, 13, 570, give m. p. 220—222° (decomp.)].

Hexamethylenediamine.—Prepared from hexamethylene bromide as for cadaverine. B. p. 204°. Yield, 51%. The picrate crystallised from water in needles, m. p. 220° (decomp.) [Ssolonina, Bull. Soc. chim., 1896, 16, 1880, gives

m. p. ca. 220° (decomp.)].

Decamethylenediamine.—Prepared in 53% yield from decamethylene bromide in the same way as cadaverine. M. p. The picrate crystallised from water in needles, m. p. 134° (Found: C, 42.0; H, 4.5; N, 17.4. C<sub>22</sub>H<sub>30</sub>O<sub>14</sub>N<sub>8</sub> requires C, 41.9; H, 4.8; N, 17.7%).

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