188. Epimeric Alcohols of the cycloHexane Series. Part V. The Optically Inactive 3-Methylcyclohexanols.

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The dl-cis- and dl-trans-3-methylcyclohexanols have been prepared in the pure state and been characterised by a number of derivatives. The trans-compound is the main product when m-cresol is hydrogenated under pressure with Raney nickel catalyst; the cis-compound is conveniently prepared by pressure hydrogenation of dl-3-methylcyclohexanone with the same catalyst. The isomers may readily be separated by forming the piperazine salts of their hydrogen phthalates, as the physical characteristics of these salts are such as allow a mechanical separation by flotation to be effectively applied.

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3-METHYL cycloHEXANOL is a very suitable subject for a stereochemical investigation, as the spatial configurations of the epimers are related in a fundamental way to those of the naturally-occurring terpene alcohols. As the molecule presents the features involved in cis-trans isomerism, and as the carbon atoms 1 and 3 are both asymmetric, the full study of the isomerism involves the preparation and examination of the four isomers written below, namely d- and l-cis- and trans-3-methylcyclohexanol, and the corresponding dl-compounds.

The 3-methylcyclohexanols have been investigated previously, but the data available show some serious discrepancies, and in no case has the examination been complete. Methods for preparing the dl-cis- (Ia and Ib) and the dl-trans-(IIa and IIb) compounds in quantity in the pure state are now described, together

with the characterisation of the pure substances by physical constants and by suitable derivatives. Knoevenagel (Annalen, 1896, 289, 142) first prepared dl-3-methylcyclohexanol by the reduction of dl-3-methylcyclohexanone and of dl-3-methyl- Δ^2 -cyclohexenone. He assigned the trans-configuration to the alcohol obtained by alkaline reduction, and assumed that the cis-form was the product of acid reduction. Present day views would support this classification but as his trans-form had the higher density and refractive index, at least one of his isomers was impure. Pickard and Littlebury (I., 1912, 101, 115) obtained a 3-methylcyclohexanol as a by-product from the catalytic hydrogenation of thymol, and this was probably mainly the trans-form. Auwers and Schmeltzer (Chem. Zentr., 1927, II, 1562) and Skita and his co-workers (Ber., 1931, 64B, 2878; Annalen, 1923, 431, 1) also isolated the 3-methylcyclohexanols as hydrogenation products, but their materials were not pure. The 3:5-dinitrobenzoates, for example, were used to purify the product of their hydrogenations of m-cresol, an unfortunate choice, as the dl-sis- and dl-trans-forms give 3:5-dinitrobenzoates which do not differ greatly either in melting point or solubility; and there is a difference of some 20° in m. p. between Skita and Faust's figure and that now recorded for dl-cis-3-methylcyclohexyl 3:5-dinitrobenzoate. Gough, Hunter, and Kenyon (I., 1926, 2052) submitted the p-nitrobenzoates to fractional crystallisation in order to separate and purify the dl-cis- and dl-trans-alcohols which they described, but we have found this method also unsatisfactory. Incomplete separation of the epimers probably explains the conflicting m. p.s given for solid derivatives of the isomers in the literature. The figures for the derivatives now described are given in the table, and the occurrence of dimorphism in the case of two derivatives was noted.

Derivatives of 3-Methylcyclohexanols.

| dl-cis | $n_{ m D}^{20}$ °. 1·4583 | d_{4}^{30} °. 0.9137 | Hydrogen phthalate. 72° 84 | p -Nitrobenzoate. 63° | 3:5-Dinitro- benzoate. 111° | Phenyl- urethane. 91° 103 | α-Naphthyl- urethane. 118° |
|----------|---------------------------|--------------------------|-------------------------------------|----------------------------------|-----------------------------------|------------------------------------|----------------------------------|
| dl-trans | 1.4573 | 0.9072 | 94 | 48 | 99 | 92 | 130 |

m-Cresol is the most convenient source of 3-methylcyclohexanol, but whereas previous workers generally used platinum catalysts or the less active forms of nickel it is now found that the phenol may be readily hydrogenated in large quantities in the homogeneous state by the use of Raney nickel and hydrogen under high pressure. The product is predominantly trans-3-methylcyclohexanol and, contrary to the statement of Gough, Hunter, and Kenyon (loc. cit.), it can be recovered as the hydrogen phthalate. This derivative solidified on standing and readily yielded pure dl-trans-3-methylcyclohexyl hydrogen phthalate on treatment with light petroleum.

The small quantity of the *cis*-compound could not be recovered economically from the residual material from the above preparation, but it was obtained (*ca.* 70%) by reduction of 3-methyl*cyclo*hexanone in acid media. Electrolytic reduction of alcoholic solutions containing sulphuric acid, and catalytic reduction using platinum oxide in glacial acetic acid, both gave yields rich in the *cis*-form, but optimum conditions were worked out for pressure hydrogenation of the ketone using Raney nickel, as this enabled work to be done conveniently and on a much larger scale. The rate of hydrogenation and the yield of the *cis*-isomer increase up to a temperature of about 140°; the rate of hydrogenation continues to increase with further rise of temperature, but the proportion of *cis*-form in the product becomes less. A reasonably exact method of assaying the relative proportion of the epimers in the hydrogenation products was based on density determinations after the preparation of pure specimens of the *cis*- and *trans*-compounds and determination of their physical constants. The difference in the refractive indices of the two forms is too small to be applied in this way, but the fraction *x* of the *cis*-form

present in a mixture whose $d_4^{30^\circ}$ value is a, is given by the expression $\frac{a - 0.9072}{x} = 0.9137 - 0.9072$.

A convenient method of separating mixtures of almost any proportions of the cis- and trans-forms consists in acting on the hydrogen phthalates with piperazine, in the ratio of one molecule of the base to two molecules of the acid phthalate. Acetone was found to be a suitable solvent, but traces of water must be present for

satisfactory operation, as then the salts formed may be separated mechanically by flotation on account of their very different physical properties. As the salts are somewhat unstable it proved best to recover the hydrogen phthalates from the crude separated piperazine salts. The recovery of purified hydrogen phthalates was effective, only relatively small amounts of piperazine being required and the base itself being readily recovered for further use.

In the course of preparation of pure derivatives for the characterisation of the epimeric alcohols it was found that $dl\text{-}cis\text{-}3\text{-}methylcyclohexanyl}$ phenylurethane and $dl\text{-}cis\text{-}3\text{-}methylcyclohexanyl}$ hydrogen phthalate exist in two modifications, a lower-melting (β) form which could in each case be converted into a higher melting (α) form on heating. The α - and β -phenylurethanes can be reproduced at will, the α -form (m. p. 103°) separating from light petroleum, and the β -form (m. p. 91°) crystallising from alcohol-water mixture. Skita and Auwers (loc. cit.) described the dl-cis-phenylurethane as melting at 90° whereas Gough, Hunter, and Kenyon (loc. cit.) gave its m. p. as 103° . The proof of the existence of the two modifications reconciles these apparently conflicting reports.

EXPERIMENTAL.

Hydrogenation of m-Cresol.—In a typical reduction m-cresol (90 g.) with Raney nickel (ca. 4 g. in 20 c.c. abs. alcohol) was shaken with hydrogen at an initial temperature of 150° and pressure 2,000 lb./sq. in. The reaction started immediately and the temperature soon rose to 200° at which the hydrogenation was allowed to proceed, the pressure being maintained above 1,000 lb. by passing in hydrogen as required. Reaction was complete in about 1.5 hours. After filtration from the catalyst, the product was stirred with dilute alkali to remove traces of phenol and then continuously extracted with ether. After drying the ethereal extract over anhydrous magnesium sulphate and removing the solvent, crude, pale-yellow 3-methylcyclohexanol (85 g.) was recovered. After distillation this had d_4^{30} 0.9082 indicating ca. 85% trans-epimer.

The crude material from such hydrogenations was esterified by heating overnight at 110—115° with 1·1 mol. phthalic anhydride, the syrupy product was stirred whilst hot into sodium carbonate solution (1·5 mols.; 5%) and agitated till dissolved. Traces of oil (possibly msthylcyclohexene) were removed by ether, the carbonate solution acidified with conc. HCl, and the precipitated syrup dissolved in chloroform to separate traces of phthalic acid. After removal of the chloroform (under reduced pressure at the end) the hydrogen phthalate was left standing with occasional stirring and, after some days, became semi-solid. The sticky solid was stirred with twice its volume of ligroin (b. p. 60—90°), filtered, and washed with the same solvent. Recrystallised four times from the ligroin it formed transparent prisms of dl-trans-3-methylcyclohexyl hydrogen phthalate, m. p. 94° (Found: C, 69·1; H, 7·1. Calc.: C, 68·7; H, 6·9%). Further quantities were recovered from the mother-liquors.

Attempts were made to separate the liquid hydrogen phthalate recovered from the light petroleum trituration by formation of metallic salts (Pickard and Littlebury, *loc. cit.*) but, although on the small scale the *trans*-compound could be recovered by fractional crystallisation of the calcium salt, no salt suitable for isolating the *cis*-form was discovered.

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dl-3-Methylcyclohexanone.—The conditions prescribed by Beckmann (Annalen, 1889, 250, 325) for the oxidation of l-menthol did not give good yields when applied to 3-methylcyclohexanol. Yields approaching 90% were obtained by the modified procedure described in a typical case, and never more than traces of unchanged alcohol were present.

dl-3-Methylcyclohexanol (70 g.) recovered from the above hydrogen phthalates was mixed with water (500 c.c.) and potassium dichromate (66.0 g., 10% excess) in a three-necked 2-l. flask fitted with a mechanical stirrer, thermometer and dropping funnel. Sulphuric acid (93%, 57 c.c.; 20% excess) was dropped in to maintain the temperature at 50—60°. Towards the end, the temperature fell and, after addition of all the acid and cooling, the mixture was extracted with ether, the extract washed thrice with sodium hydroxide (5%), then with water and dried over magnesium sulphate. After removal of the solvent the ketone was distilled, b. p. 64—65°/30 mm.

Hydrogenation of dl-3-Methylcyclohexanone.—(a) The ketone (25 g.) in glacial acetic acid (100 c.c.) in a 250 c.c. flask was shaken at laboratory temperature and pressure with Adams (platinum oxide) catalyst (0.3 g.) after removal of air removal of air removal of all the catalyst had frequently to be reactivated. After

Hydrogenation of dl-3-Methylcyclohexanone.—(a) The ketone (25 g.) in glacial acetic acid (100 c.c.) in a 250 c.c. flask was shaken at laboratory temperature and pressure with Adams (platinum oxide) catalyst (0·3 g.) after removal of air and introduction of hydrogen. Hydrogen uptake was slow and the catalyst had frequently to be reactivated. After 10 hours reaction had ceased and ca. 87% of the theoretical amount of hydrogen had been absorbed. After removal of the catalyst and dilution, the acetic acid was neutralised with strong cooling and the mixture continuously extracted with ether. Unchanged ketone was removed from the extract by sodium bisulphite, and the extract washed, dried and distilled under reduced pressure gave dl-3-methylcyclohexanol (14 g.), d₄₀³⁰·0·9117, indicating about 69% of the cis-epimer. (b) The method used by Gillespie, Macbeth, and Swanson (J., 1938, 1820) for the electrolytic reduction of cryptone to dihydrocryptol was used. Each reduction was carried out with 25 g. of the ketone, and was complete in 2 hours at a temperature of 30—35°. The alcoholic solution was diluted with water and continuously extracted with ether, and the extract worked up as above. 3-Methylcyclohexanol (16 g.) with d₄₀³⁰·0·9119, indicating about 72% of the cis-epimer, was usually obtained. (c) Cooke, Gillespie, and Macbeth (J., 1939, 518) obtained a good yield of cis-dihydrocryptol by hydrogenation of cryptone over Raney nickel, and the preparation of 3-methylcyclohexanol rich in the cis-epimer by hydrogenation at a temperature of about 140° was satisfactory. This method was adopted in all preparations of the cis-alcohol on account of its convenience and good yields. The ketone (50—60 g.) mixed with Raney nickel (3 g.) in abs. alcohol (20 c.c.) was hydrogenated in a glass-lined bomb at 140° the initial hydrogen pressure being 1,700—1,800 lb./sq. in. The hydrogen uptake extended over about 2 hours. After working up in the usual way 3-methylcyclohexanol, d₄₀³⁰ 0·9122 (indicating 77% cis-epimer), was obta

The al-trans-piperazine salt separated in masses of white needles when pure al-trans-3-methylcyclohexyl hydrogen phthalate (1 g.) and piperazine hydrate (0.35 g.) were dissolved separately in warm acetone and the solutions mixed. Recrystallisation from boiling acetone (100 c.c.) yielded white needles, m. p. 135—136°, which separated slowly. The salt is insoluble in ether and paraffin hydrocarbons, moderately soluble in alcohol, ethyl acetate and hot benzene. The dl-cis-salt was prepared similarly and recrystallised from hot acetone, in which it is easily soluble and from which it separates in shining granules having m. p. 122—123·5°. It is insoluble in ether, and more soluble than the trans-salt in other solvents, being readily soluble in cold alcohol and in chloroform. The salts are considered to be the neutral piperazine salts, $C_4H_8(NH)_2, 2C_7H_{13}O_2C\cdot C_6H_4CO_2H$, although it was impossible to get exact analytical figures for this formula as the salts appear to contain variable amounts of water.

The cis- and trans-salts could be separated when prepared from a mixture of the hydrogen phthalates by using about six times the weight of acetone. The following examples are typical. (a) The hydrogen phthalate (92 g.) formed from the alcohol prepared by the hydrogenation of 3-methylcyclohexanone was added to a solution of piperazine (36.5 g.) in warm acetone (600 c.c.). The reaction was exothermic and the acetone boiled. The hot solution was seeded with crystals of both the pure salts and left two days undisturbed at room temperature. A mixture of fine white needles and glistening prisms separated and after breaking up the crusts the whole mixture was agitated with a turbine stirrer. The

suspension of needles could then be poured from the sedimented heavy granules on to a filter. Both lots of crystals were separately stirred again with the filtrates and eventually a clean separation was effected into heavy crystals (53 g.) and needles of the trans-salt (18 g.). (b) Occasionally, particularly with mixtures containing predominantly trans-epimer, seeding of the hot acctone solution with the trans-salt alone caused the crystallisation of needles of the trans-salt without any cis-salt: the latter could be obtained after separating the precipitate and seeding the filtrate with the cis-salt. For example, the hydrogen phthalate (138 g.) obtained from the hydrogenation product of m-cresol was mixed with piperazine hydrate (55 g.) in hot acetone (900 c.c.). On seeding the hot solution, crystallisation started immediately and only needles appeared when left overnight. These were separated (79 g.) and the filtrate, when seeded with the cis-salt and left for two days, deposited a mixture of needles (6 g.) and prisms (24 g.) which were separated as above. On strong cooling a low-melting product (18 g.) of mixed cis- and trans-forms separated. Its formation during large scale separations is avoided by slow crystallisation without disturbance in a stoppered flask at room temperature.

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The salts remaining in the mother-liquors, after removal of the crystalline piperazine salts, are decomposed to the hydrogen phthalates for treatment again with piperazine. This is more satisfactory than attempted recovery by further concentration. For example, in the two separations described above, the mother-liquors, containing the low-melting mixture of salts (18 g.), were evaporated to small bulk and run into dilute hydrochloric acid. The hydrogen phthalate (105 g.) recovered as a brownish syrup after extraction with ether was again converted into the piperazine salt, and crystallisation and separation as above yielded the crude cis- (51·5 g.) and trans- (31·5 g.) salts. From the total crude hydrogen phthalates (230 g.) employed in (a) and (b) above these relatively simple operations gave good recoveries of the crude cis- (128·5 g.) and trans- (134·5 g.) salts. The fact that the separation of cis- and trans-salts was effected from mix-

tures rich in either form is also of interest.

It was found more convenient and effective not to attempt to purify the piperazine salts but to decompose them to hydrogen phthalates and purify the latter. (a) Crude dl-trans-piperazine salts 10 g.) was dissolved in boiling methanol (250 c.c.) and the hot solution stirred into cold HCl (600 c.c.; 5%). The hydrogen phthalate separated as an oil and was recovered by ether extraction (thrice). After removal of the solvent the crude hydrogen phthalate (92 g.; 95% recovery) was recrystallised thrice from light petroleum (b. p. 60—90°) giving the pure product (51 g.). The mother-liquors were used for crystallising further batches of the crude material. (b) The crude cis-salt (85 g.) when similarly decomposed gave the crude hydrogen phthalate (66 g.; 91% recovery) which yielded the pure product (26 g.) after three crystallisations from light potroleum (b. p. 40, 90°). crystallisations from light petroleum (b. p. 40—90°).

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The purified hydrogen phthalates were hydrolysed and the liberated alcohol separated by steam distillation of the solution containing aqueous sodium hydroxide (3 mols., 10%). The distillate was continuously extracted with ether, the extract dried with anhydrous magnesium sulphate and, after removal of the solvent, the alcohol was distilled under reduced pressure. This gave (93% yield) dl-trans-3-methylcyclohexanol, b. p. 60°/2 mm.; n½° 1-4573; d2° 0-9072. Similarly dl-cis-3-methylcyclohexanol was obtained having b. p. 72-73°/20 mm.; n½° 1-4583; d2° 0-9137.

Derivatives of the 3-Methylcyclohexanols.—dl-trans-3-Methylcyclohexyl p-nitrobenzoate crystallised from methanol-water in pale-yellow feathery crystals, m. p. 478° (Found; c. 63*9; H, 64.* Calc.; 63*85; H, 6.5%). dl-cis-3-Methylcyclohexyl p-nitrobenzoate crystallised from the same solvent had m. p. 62—63°. dl-trans-3-Methylcyclohexyl 3: 5-dintrobenzoate separated from ligroin or methanol-water in almost colourless granules. m. p. 99° (Found): c. 54*5; H, 5-2; N, 9-15. Calc.; C, 54*6; H, 5-4; N, 9-19%). The α-naphthylamine complex was precipitated by adding a 5% ethereal solution of the dintrobenzoate to a slight excess of α-naphthylamine in 85% alcohol. It recrystallised from light petroleum in long, brick-red needles having m. p. 142—143° (Found): c. 63*9; H, 5-6%). dl-cis-3-Methylcyclohexyl a-naphthyluriethane was prepared by mixing the alcohol and α-naphthyl isocyanate in equimolecular quantities in a little light petroleum. The solid which separated was recrystallised from light petroleum and had m. p. 128-5—129-5 (Found): c. 76·3; H, 7-5%). dl-cis-3-Methylcyclohexanyl α-naphthylurethane crystallised from light petroleum as prepared in a similar way and purified by recrystallised in small pearly plates, m. p. 91—92° (Found): c. 71·9; H, 8-2; N, 6-05. Calc. c. 7.2·05; H, 8-2; N, 6-0%)

melted and cooled the higher-melting, a_r form of hydrogen phthalate quickly separated in transparent irregular crystals having m. p. 82—83°. This form is identical with the dl-cis-hydrogen phthalate obtained from the piperazine salt separations described above (Found: C, 69·2; H, 7·0%).

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