

MOLECULAR COMBINATIONS OF β -METHYL-*D*-GLUCOSIDE, β -METHYL-*D*-XYLOSIDE AND β -METHYL-*L*-FUCOSIDE WITH POTASSIUM ACETATE¹

Sir:

While engaged in deacetylating tetraacetyl- β -methyl-*D*-glucoside with an alcoholic solution of potassium hydroxide, we observed the separation of a quantity of long silky needle-like crystals; these proved to represent a molecular combination of β -methyl-*D*-glucoside with potassium acetate ($C_7H_{14}O_6 \cdot CH_3COOK$). The same substance was prepared in quantity by mixing concentrated absolute alcohol solutions of the glycoside and of the anhydrous salt.² The α -methyl-*D*-glucoside when similarly treated in many experiments over a period of two years has never yielded any crystalline product. This specificity of the reaction constitutes its chief usefulness since it affords a convenient method of separating the beta form completely from its alpha isomer, a process which formerly was accomplished only by tedious fractionation. Application of this new double compound for such separations will be reported later.

The compound contains the sugar derivative and the salt in a molecular ratio of one to one as is indicated by the methoxyl and potassium analyses. It melts at 181–182° (corr.) and shows $[\alpha]_D^{20}$ in water -22.0° , which is the specific rotation that would be calculated on the assumption that the salt is inert and has no influence upon the rotation of the sugar derivative. The silky needles are rather hygroscopic.

The analogous compound of β -methyl-*L*-fucoside ($C_7H_{14}O_6 \cdot CH_3COOK$) is almost identical in superficial appearance, is similarly hygroscopic, melts at 208–212° (corr.) and shows $[\alpha]_D^{20}$ in water $+8.9^\circ$. It is also a one to one molecular combination as indicated by its analyses and its specific rotation. This substance has been particularly valuable in facilitating the separation of β -methyl-*L*-fucoside from the alpha isomer, which forms no analogous compound, since the separation by fractionation was especially difficult in this case.

The β -methyl-*D*-xyloside potassium acetate compound ($C_8H_{12}O_5 \cdot CH_3COOK$), which is also of

(1) Publication authorized by the Surgeon General, U. S. Public Health Service.

(2) The samples of commercial potassium acetate which we examined all contained a molecule of acetic acid of crystallization per molecule of salt, though the labels described the substance as CH_3COOK . The salt may be dried by heating at 150° but the same products are obtained from the methyl glycosides by using either the dried or the undried salt.

very similar appearance, is hygroscopic, melts at 171–172° (corr.), shows $[\alpha]_D^{20}$ in water -41.3° and is likewise a one to one combination.

The glycoside may be recovered in each case by precipitating the potassium as potassium acid tartrate from a solution containing 50% of alcohol. By treating the double compounds directly with acetic anhydride, the acetylated methylglycosides may be obtained almost quantitatively, the salt acting as catalyst.

Efforts to prepare similar derivatives of α - and β -methyl-*D*-arabinoside, α - and β -methyl-*D*-galactoside, α - and β -methyl-*D*-lyxoside, α -methyl-*D*-xyloside, α -methyl-*D*-glucoside and α -methyl-*L*-fucoside have been unsuccessful. Other communications will follow.

NATIONAL INSTITUTE OF HEALTH
WASHINGTON, D. C.

A. J. WATTERS
R. C. HOCKETT
C. S. HUDSON

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A NEW SYNTHESIS OF N-MONOPHENYLPIPERAZINE

Sir:

The only reported synthesis of N-monophenylpiperazine is that of V. Prelog and G. J. Driza [*Coll. Czechoslovak Chem. Comm.*, **5**, 497–502 (1933)].

The same synthesis had been accomplished by us in this Laboratory previous to the publication by the above authors. Since we had also noticed the physiological effects of N-monophenylpiperazine, it was decided to find, if possible, an easier and more convenient method for preparing it. This was desirable in view of the fact that we encountered considerable difficulty in preparing the intermediate bis-(β -haloethyl)-amines in sufficient quantities to make any large amounts of the N-monophenylpiperazine.

We have been able to prepare N-monophenylpiperazine in practically any amount by heating together aniline hydrochloride and diethanolamine hydrochloride at about 240° for from six to eight hours. In practice, the amines are mixed in molecular proportions and concd. hydrochloric acid added until the solution is neutral to litmus. The water is then boiled off and when the temperature has reached 220 to 240° it is kept there for from six to eight hours. Upon cooling the mass sets to a dark brown, gummy solid. The free base is liberated by treatment with concd. sodium hydroxide and the resulting oil is then fractionated