Synthesis of Biological Active Esters of the Isovaleric Acid by Isobutylene Hydroalkoxycarbonylation

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Abstract

Hydroalkoxycarbonylation of 2-methylpropene with carbon monoxide and alcohols under condition of homogeneous catalysis with transition metal complexes allows facile one-step synthesis of practically useful isovaleric acid esters. Many of them exhibit biological activity and are components of pharmaceuticals (Validolum, Corvalolum, etc.) or valuable intermediates in drug synthesis. Biological active isovaleric acid esters (1-methylisovalerate, ethylisovalerate, cyclohexylisovalerate, benzylisovalerate, α-monoglyceryde of isovaleric acid) were prepared by isobutylene hydroalkoxycarbonylation. New efficient technologies for preparation of drugs (Validolum, Ethyl ester of α-bromisovaleric acid and Corvalolum) are based on the isovaleric acid esters were worked out.

Keywords: Synthesis; Isovaleric acid; Esters; Drugs; Isobutylene; Hydroalkoxycarbonylation; Technologies

Introduction

Isobutylene as an accessible and inexpensive feedstock is of interest for synthesis of many practically useful compounds. Isobutylene carbonylation with carbon monoxide and alcohols under conditions of homogeneous catalysis with transition metal complexes allows facile one-step synthesis of practically useful isovaleric acid esters [1-5]. Many of them have biological activity and are components of pharmaceuticals (Validolum, Corvalolum, etc.) or valuable intermediates for their synthesis. Some isovlarate esters possess a characteristic odor and are used as fragrance compounds in the manufacture of perfumes, cosmetics and food essences [6].

We applied hydroalkoxycarbonylation of isobutylene with carbon monoxide and mono(poly)hydric alcohols in the presence of catalytic systems based on the phosphinopalladium complexes (Pd(PPh3)4), Pd(PPh3, TsOH and Pd(Acac)2)-(PPh3, TsOH) to prepare of biological active isovaleric acid esters: 1-methylisovalerate (possesses spasmodytic properties; it used as main active component of the spasmytic medicine Validolum), ethylisovalerate (possesses aromatic (fruit) odor; intermediate product for obtaining sedative and spasmytic medicines Ethyl ester of α-bromisovaleric acid and Corvalolum), cyclohexylisovalerate (bactericide activity (against Staphylococcus aureus, Escherichia coli, Pseudomonas aeruginosa)); antifungal activity (against Candida albicans)); benzylisovalerate (bactericide activity (against Escherichia coli, Staphylococcus aureus)); α-monoglyceryde of isovaleric acid (bactericide activity (against Escherichia coli, Pseudomonas aeruginosa)); antifungal activity (against Candida albicans)).

New efficient technologies for preparation of drugs are based on isovaleric acid esters – Validolum, Ethyl Ester of α-bromisovaleric acid (EEBIA) and Corvalolum – were worked out. Validolum – is a spasmytic (sedative) medicine. It has a sedative effect on the nervous system and a moderate reflex vaso-dilating effect. EEBIA possesses sedative and spasmytic properties. It is included in Corvalolum composition and may be used for producing other medicines. Corvalolum is a combined medicine and consists of EEBIA, phenobarbital, sodium hydroxide, peppermint oil, ethyl alcohol and water. Corvalolum possesses anetic and spasmytic properties.

Due to the more advanced technology of production the Medicines will have better qualitative characteristics. The cost of production of the Medicines with the use of new technologies is 2-3 times lower as compared to the medicines produced by existing at the present traditional technologies.

Experimental

The complexes Pd(PPh3)4 and Pd(PhAc)2 was obtained according to the known procedures [7]. p-Toluenesulfonic acid was recrystallized from 96% ethanol and dried until the composition TsOH*H2O. Triphenylphosphine was recrystallized from an ether-ethanol mixture to a constant melting point. Absolute ethanol, isobutylene of 99,5% purity, carbon monoxide of 99,8% purity, l-menthol of 99,7% purity, cyclohexanol of 99,5% purity, benzyl alcohol of 99,2% purity and glycerin of 99,5% purity were used. The experiments were carried out in the solvent-free mode in a laboratory stainless steel autoclave unit. The determination of the purity and the analysis of the products were carried out by means of a GLC technique, thin-layer chromatography, IR-spectroscopy and 'H NMR techniques. Gas chromatography was performed on a Hewlett-Packard 3890-II-Plus chromatograph with a flameionization detector; HP-Innowax cross-linked PEG capillary column (30000*0,25mm), film thickness 0,25 μm. Injector temperature 200°C, detector temperature 200°C, carrier gas nitrogen (25 ml/min). The oven temperature was programmed from 75 to 175°C at a rate of 10°C/min. The 'H NMR spectra were measured on a Varian Mercury-300 instrument (300 MHz) against internal TMS.

L-Menthyl Isovalerate

The 100 ml autoclave equipped with a stirrer and a carbon...
monoxide and isobutylene feeding device was charged with 0,133 g (1,15*10^4 mol) of Pd (PPh 3) 4, 0,263 g (1,38*10^4 mol) of TsOH, and 7,854 g (5,02*10^2 mol) of l-menthol. The autoclave was sealed, purged with CO to remove air, and filled with CO to a pressure of 1,0-1,1 MPa. Then 3,562 g (6,35*10^3 mol) of isobutylene was introduced and stirring and heating were switched on. The carbon monoxide pressure was brought to 2,0 MPa, the temperature was elevated within 1 h to 100°C, and the reaction mixture was agitated under these conditions for 4 h. On completion, the autoclave was cooled to room temperature and the reaction mixture was fractionated in a vacuum. 3,58 g (45,6% of the initial quantity) of unreacted l-menthol and 6,24 g (51,6%) (or 94,9% on converted l-menthol) of l-menthyl isovalerate were obtained. Bp 187°C / 30 mm Hg, nD 1,4480.

Isovaleric acid α-monoglyceride

A steel autoclave of 100 ml capacity was charged with 0,035 g (1,15*10^4 mol) of Pd (Acac) 2, 0,212 g (8,085*10^4 mol) of PPh 3, 0,263 g (1,386*10^4 mol) of TsOH, and 5,975 g (6,35*10^2 mol) of glycerin. The autoclave was sealed, purged twice with CO to remove air from the system, and then filled with CO to a pressure of 1,0-1,1 MPa. After that 7,125 g (12,7*10^3 mol) of isobutylene was introduced into the autoclave, and the carbon monoxide pressure was increased to 2,0 MPa. Stiring and heating were switched on. The reaction mixture was stirred within 4 h at a temperature of 100°C and a pressure of 2,0 MPa. Then the autoclave was allowed to cool down to a room temperature and left for a night. The next day, after the pressure was released to atmospheric, the reaction mixture was fractionated in a vacuum. The desired products were separated from the obtained mixture of products with unreacted glycerin by means of column adsorption chromatography on silica gel (0,005-0,04 mm). Chloroform and chloroform: methanol blend (9:1 by volume) were used as eluents. 1,81 g (30,8% of the initial quantity) of unreacted glycerin, 0,91 g (16,2%) (or 23,2 % on converted glycerin) of isovaleric acid α-monoglyceride and 1,08 (13,1 %) (or 18,7 % on converted glycerine) of isovaleric acid α,α' diglyceride were obtained. Isovaleric acid α-monoglyceride. Bp 187°C / 30 mm Hg, nD 1,4440. Isovaleric acid α,α'-diglyceride. Bp 198°C / 24 mm Hg, nD 1,4390.

Results and Discussion

Hydroalkoxycarbonylation reaction of isobutulene with carbon monoxide and monohydric alcohols (ethanol, cyclohexanol, l-menthol, benzy alcohol) in the presence Pd(PPh3)4-PPh3-TsOH system carried out at optimal conditions of isovaleric hydroxymethoxy carbonylation [8]: temperature 100°C; CO pressure 2,0 MPa; reaction time 4 h; reactants and catalyst components ratio [alcohol]:[isobutylene]:[Pd(PPh3)4]:[PPh3]:[TsOH]=435:550:1:3:12. The yields of the products were 71-95% (on converted alcohols).

Synthesis of isovaleric acid α-monoglyceride carried out at optimal conditions of hydroalkoxycarbonylation of isobutylene with carbon monoxide and glycerin in the presence of Pd(Acac)2-PPh3-TsOH system [9]: temperature 100°C; CO pressure 2,0 MPa; reaction time 3 h; reactants and catalyst components ratio [glycerin]:[isobutylene]:[Pd(Acac)2]:[PPh3]:[TsOH]= 550:1100:1:7:12. The yield of the isovaleric acid α-monoglyceride was 18,7% (on converted glycerin).

The selectivity in linear reaction products was 100%. Such a high regioselectivity is apparently provided both by the structure of the starting alkenes (isobutylene) and by the reaction mechanism. The most probable is a hydride mechanism. Evidence for this proposal comes from the observation of an exceptionally strong effect of the TsOH addition, which being a proton donor, facilitates formation of the primary active hydride complexes of the catalytic cycle.

L-Methylisovalerate is a main active ingredient of the drug Validolum. Validolum has a sedative effect on the nervous system and a moderate reflex vaso-dilating effect. It is used at light attacks of stenocardia, neurosis, hysteria. It is also used as anantihetic at sea sickness.

The existing industrial production of Validolum is based on the two stage scheme of the synthesis of L-methylisovalerate: 1) oxidation reaction of isomal alcohol to isovaleric acid; 2) esterification reaction of isovaleric acid by l-menthol. Such a technology of obtaining L-methylisovalerate is characterized by low technical-economic (duration of the esterification process is 48 h, yield of the target product no more than 75%) and low ecological characteristics (large amounts of waste waters at the stages of neutralization and washing) and low quality of products because of the presence of impurities. Validolum obtained by traditional technology contains 11 admixtures, the content of which reaches 8%.

The new technology developed by us makes possible to make the synthesis of L-methylisovalerate in one stage by reaction of hydroxymethoxy carbonylation of isobutylene with carbon monoxide and l-menthol in the presence of metalcomplex catalyst. The use of the more available raw materials and also the high effectiveness of the technology (duration of the process is 5 h, yield of the target product 95%) makes this process of obtaining L-methylisovalerate highly profitable. The product obtained with the new technology has higher quality and contains only 3-4 admixtures, the contents of all of which is not higher than 1-1,5%.

EEBIA possesses sedative and spasmylytic properties and in larger doses provides light soporific action. It is included in composition of the drug Corvalolum and may be used for producing other medicines. Corvalolum possesses anetic and spasmylytic properties. It is used for neurosis with increased irritability, for soft spasms of coronary vessels, tachycardia, anhypnosis, early stages of hypertension and bowel spasms.

The existing technology of EEBIA production is based on the four stage scheme of the synthesis. The first stage is obtaining of isovaleric acid by oxidation of isoamyl alcohol. Then follows the two staged bromination of isovaleric acid with bromine in the presence of PCl3. The obtained α-bromisovaleric acid is transferred into chloranhydride, which is subjected to esterification with ethanol. This method of EEBIA obtaining is characterized with complexity and is highly labor consuming process, has low technical, economic and ecological characteristics (use of expensive and rare raw materials, use and formation of aggressive starting products and secondary by-products: PCl3, HCl, H3PO4) and the low quality of the product due to the admixtures.

EEBIA contains only 3-4 admixtures, the content of all of which is not higher than 1-1,5%.
The proposed by us new technology of obtaining EEBIA (and Corvalolum production on its basis) is based on the new effective method of obtaining EEBIA from isobutylene, carbon monoxide, ethanol and bromine. The synthesis of ethylisovalerate is carried out by hydroethoxycarbonylation reaction of isobutylene by carbon monoxide and ethanol in the presence of metalcomplex catalyst. On the second stage the product (EEBIA) is synthesized by bromination of ethylisovalerate in the presence of the red phosphor. Quality of EEBIA obtained by the new technology is higher (contains less admixtures), the production costs are 2-3 times lower than the production costs of the existing four stage method of obtaining EEBIA.

1. $\text{CH}_3\text{CH} = \text{CH}_2 + \text{CO} + \text{C}_2\text{H}_5\text{OH} \rightarrow \text{CH}_3\text{CH} = \text{CH}_2\text{OC}_2\text{H}_5$

2. $\text{CH}_3\text{CH} = \text{CH}_2\text{OC}_2\text{H}_5 + \text{Br}_2 \rightarrow \text{CH}_3\text{CH} = \text{CH}_2\text{OC}_2\text{H}_5 + \text{HBr}$

$\alpha$-Monoglyceride of isovaleric acid was synthesized by carbonylation of isobutylene with carbon monoxide in the presence of glycerin and the catalyst system Pd(Acac)$_2$-PPh$_3$-TsOH. According to available data, the secondary hydroxyl group of glycerin reacts more slowly than a primary one. At ratios of [isobutylene]:[glycerin]=1:1 and 2:1, mono- and diglycerides are formed, and mono-, di- and triglycerides are formed when the ratio is 1:3. The maximum total yield of glycerides (23.3%) was obtained at a ratio of [isobutylene]:[glycerin]=2:1; the yields of monoglyceride and diglyceride make up 19.1 and 4.2%, respectively, in this case. The yields of mono- and diglycerides at [isobutylene]:[glycerin]=1:1 are 9.3 and 1.1%, respectively. At a ratio of [isobutylene]:[glycerin]=3:1, mono-, di-, and triglycerides were obtained with yields of 14.7, 3.0 and 0.4%, respectively.

Note that, in contrast to the known processes for the manufacture of glycerides of fatty acids by the direct esterification of the acids with glycerin and by transesterification of methyl (or ethyl) esters of fatty acids with glycerin when a mixture of $\alpha$- and $\beta$-isomers of monoglycerides is formed, the formation of the $\alpha$-isomer alone is observed during the hydroxycarbonylation of isobutylene with carbon monoxide and glycerin.

Thus, the feasibility of the synthesis of isovaleric acid esters with aliphatic polyols by means of alkoxy carbonylation of isobutylene in the presence of the catalytic system Pd(Acac)$_2$-PPh$_3$-TsOH was established. The reaction proceeds regioselectively at the terminal atom of isobutylene. Isovaleric acid monoglyceride is formed only in the form of the $\alpha$-isomer.

The proposed methods are highly economical and may be used for commercial production of the mentioned above biological active esters of the isovaleric acid. Optimal technological parameters for carrying out the processes were tested at the pilot plant. Technologically, organization of the productions being proposed does not present any great difficulties. Standard equipment may be used. It should be noted that all the proposed productions are based on the similar technology using one and the same equipment.

References