| Title | Disulphates of 16-oxygenated ketonic C ₁₉ steroids as biliary metabolites of androsterone sulphate in female rats |
|------------------|--|
| Sub Title | |
| Author | 松井, 道夫(Matsui, Michio) 箱崎, 美砂子(Hakozaki, Misako) |
| Publisher | 共立薬科大学 |
| Publication year | 1978 |
| Jtitle | 共立薬科大学研究年報 (The annual report of the Kyoritsu College of Pharmacy). No.23 (1978.) ,p.117- 119 |
| JaLC DOI | |
| Abstract | |
| Notes | 抄録 |
| Genre | Technical Report |
| URL | https://koara.lib.keio.ac.jp/xoonips/modules/xoonips/detail.php?koara_id=AN00062898-00000023- 0117 |

慶應義塾大学学術情報リポジトリ(KOARA)に掲載されているコンテンツの著作権は、それぞれの著作者、学会または出版社/発行者に帰属し、その権利は著作権法によって 保護されています。引用にあたっては、著作権法を遵守してご利用ください。

The copyrights of content available on the KeiO Associated Repository of Academic resources (KOARA) belong to the respective authors, academic societies, or publishers/issuers, and these rights are protected by the Japanese Copyright Act. When quoting the content, please follow the Japanese copyright act.

Disulphates of 16-Oxygenated Ketonic C₁₈ Steroids as Biliary Metabolites of Androsterone Sulphate in Female Rats *

MICHIO MATSUI and MISAKO HAKOZAKI

松井道夫, 箱崎美砂子

It is well established that the 16β -hydroxy-17-oxosteroid is readily rearranged to the stable epimer, the 17β -hydroxy-16-oxosteroid, by means of alkali or acid, and by isolation procedures from urine. In a paper dealing with biliary metabolites of androsterone conjugates in female rats, we reported that androsterone sulphate was metabolized to the disulphate of 3α , 17β -dihydroxy- 5α -androstan-16-one. But, we did not preclude the artifact formation by rearrangement of 16β -hydroxyandrosterone to 3α , 17β -dihydroxy- 5α -androstan-16-one during the isolation procedure, because 3β , 16β -dihydroxy-5-androsten-17-one, a model compound, was entirely converted into 3β , 17β dihyroxy-5-androsten-16-one by t.l.c. on silica gel plate. Very recently, Mattox *el al.* purified synthetic 3β , 16β -dihydroxy-5-androsten-17-one by partition chromatography on Celite 545. These studies prompted us to investigate the occurrence of 16β -hydroxyandrosterone in the 16-oxygenated ketonic steroid metabolites of androsterone sulphate by employing the partition chromatography to the separation of the solvolyzed metabolites.

In this report, we describe the synthesis of 16β -hydroxyandrosterone and the identification of 16β -hydroxyandrosterone disulphate as a biliary metabolite of androsterone sulphate in female rats.

The diacetate of 16β -hydroxyandrosterone was prepared by lead tetraacetate oxidation of 5α -androst-16-ene- 3α , 17-diol diacetate. Hydrolysis of the acetate group was done by formation of a C-17 semicarbazone, alkaline hydrolysis of the acetate group, and removal of the semicarbazone group. The configuration of the 16-hydroxy group was established by comparison of the sodium borohydride reduction product with 5α androstane- 3α , 16β , 17β -triol.

GC analyses of the TMS and MO-TMS derivatives of 16α -hydroxyandrosterone, its 16β -epimer, and 3α , 17β -dihydroxy- 5α -androstan-16-one demonstrated that the TMS ether of the 16β -hydroxysteroid was not separated from that of the 16-oxosteroid, while the MO-TMS derivatives provided good separation. Mass spectra of the MO-TMS derivatives showed the molecular ion at m/e 479, characteristic of the dihydroxymonooxo C₁₉ saturated steroid. The 16α -and 16β -hydroxysteroids gave base peak at m/e 448, whereas the 16-oxosteroid afforded base peak at m/e 158. Furthermore, sev-

^{*}本報告は Sreroids, 31, 219 (1978) に発表。

No.23 (1978)

eral ion abundance differences were observed among these spectra. The ions at m/e 433 and 343 were more abundant in the 16α -hydroxysteroid than in the other steroids, while the fragments at m/e 142 and m/e 133 were characteristic of the 16β -hydroxysteroid and the 16-oxosteroid, respectively.

The occurrence of epimerization of the 16β -hydroxy-17-oxosteroid to the 17β -hydroxy-16-oxosteroid is of crucial problem in the analysis of the 16β -hydroxy-17-oxosteroid. To check to what extent artifact formation occurred during the isolation procedure, 16β -hydroxyandrosterone was subjected to incubation in solvolytic medium, followed by partition chromatography. GC analysis revealed that only a few % of 16β -hydroxyandrosterone was epimerized to 3α , 17β -dihydroxy- 5α -androstan-16-one.

Following intraperitoneal injection of [3H]-androsterone sulphate into female rats with bile fistulas, the excretion of the radioactivity in the bile was about 76% during 24 hr. The biliary radioactivity consisted of monosulphate (65%) and diconjugate (35%) fractions. By solvolysis of the diconjugate fraction, 58% of the radioactivity appeared in the liberated steroid fraction. These figures were similar to those described in a previous paper. Partition chromatography of the liberated steroid fraction yielded 79, 2, and 4% of the radioactivity in 5α -androstane- 3α , 17β -diol, 3α , 17β -dihydroxy- 5α and rost an 16β -hydroxy and rost erone fractions, respectively. The production of large amounts of 5α -androstane- 3α , 17β -diol was a pronounced feature in this study. One explanation for this may be that we administered the increased amounts of androsterone sulphate into rats in this study. Eriksson et al. reported the change of metabolic patterns with different concentrations of progesterone in perfused rat liver. The each fraction obtained by partition chromatography was analyzed as the TMS and MO-TMS derivatives by GC and GC-MS. The identified metabolites provided relative retention times and mass spectra identical with those of the respective reference steroids. Details of the identification of 5α -androstane- 3α , 17β -diol were not given here, because it was already done previously. 16β -Hydroxyandrosterone and 3α , 17β dihydroxy- 5α -androstan-16-one were identified in the respective fractions, though the 16β -hydroxyandrosterone fraction was contaminated with small amounts of 3α , 17β -dihydroxy- 5α -androstan-16-one. No other steroid metabolites (16α -hydroxyandrosterone, etc.) were identified in these fractions. Although our results do not preclude the occurrence of the disulphate of 3α , 17β -dihydroxy- 5α -androstan-16-one, the production of 16β -hydroxyandrosterone disulphate as a biliary metabolite of androsterone sulphate in female rats is firmly established. To our knowledge, this is the first identification of 16β -hydroxyandrosterone as a disulphate in rat bile. Thus, the major portion of 3α , 17β -dihydroxy- 5α -androstan-16-one identified in the previous study must be the artifact formed from 16β -hydroxyandrosterone.

Several 17β -hydroxy-16-oxosteroids were isolated from the incubation of C₁₉ steroids with rat liver or human foetal liver, and from rat bile. The present study suggests

No.23 (1978)

the occurrence of the 16β -hydroxy-17-oxosteroids in these 16-oxygenated steroid metabolites. Recently, 16β -hydroxylating enzymes active on steroid sulphates were found in human foetal liver. Thus, androsterone sulphate must undergo direct metabolism by liver microsomal 16β -hydroxylase to 16β -hydroxyandrosterone 3-sulphate, which must be consequently conjugated with sulphuric acid to 16β -hydroxyandrosterone disulphate by the sulphotransferase located in the soluble fraction of the liver cell.