of asymmetry have been preserved. The details of these structure proofs, together with other transformations of the reserpine E ring, will be reported elsewhere.

Pharmacological evaluation of the ethers Id and II and their hydrochloride salts revealed a preponderance of sedative action in mice and dogs without any demonstrable hypotensive effect in dogs. The quieting action of the ethers differed from that caused by reserpine itself in several important respects: The onset of action was within minutes rather than hours. The duration of action was considerably shorter than that of reserpine. Cumulation was not evident upon repeated administration. Oral absorption was rapid and quite complete, since tranquilization of the same degree was achieved in 30 to 60 min following administration by either the oral or intravenous route. In contrast to the experience with reserpine, there was no evidence of drug action on the following day.

There were also distinct differences between the effects of reserpine and those of the ethers upon the gastrointestinal tract of the dog. The latter did not evoke increased motor activity in the small intestine of the dog as recorded from a Thiry-Vella intestinal loop. For this reason undoubtedly diarrhea, which was so common following reserpine in the dog, was not noted following sedative doses of the ethers. Their effect on gastric secretion was less than one-tenth that of reserpine.

A further interesting pharmacological property of the epiethers was the antagonism of the fibrillatory and cardiac depressant actions of aconitine upon the isolated cat heart. The epiethers were considerably more active than the normal ethers of methyl reserpate in this respect.

Zusammenfassung. Durch Umsetzung des Methyl-Reserpates (Ic) mit Diazomethan in Gegenwart von Fluoborsäure wird 18-O-Methylreserpsäure-methylester (Id) gewonnen. Reaktion des 18-O-p-Brombenzolsulfonylreserpsäure-methylesters (Ie) mit Methanol und Triäthylamin führt dagegen zum isomeren 18-epi-Methyläther (IIa). Diese Produkte, und besonders ihre wasserlöslichen Hydrochloride, besitzen ausgeprägte sedative Eigenschaften mit schnellem Wirkungseintritt; sie sind aber nicht hypotensiv wirksam.

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Uzarigenin and Desglucouzarin from Asclepias syriaca L.

ZECHNER¹ isolated from Asclepias syriaca L. of Croatian origin an amorphous glycoside with typical digitaloid effects², which was not chemically characterized in detail. Petricić et al.³ showed by paper chromatography the presence in Asclepias syriaca L. of five substances which gave a positive Baljet reaction. One of them was isolated in crystalline state; on hydrolysis it yielded rhamnose and glucose but the aglycone was not described.

We have examined Asclepias syriaca L. obtained from Western Slovakia. Paper chromatography showed the presence of nine substances, which gave positive reactions with m-dinitrobenzene 4 and with 3,5-dinitrobenzoic acid 5.

Ether, chloroform, and chloroform-ethanol-(2:1) extracts were prepared in the usual manner from the ethanol extract of the drug, after removal of impurities with lead acetate. From these extracts six substances were obtained

(two in crystalline form) by counter-current separation and chromatography on aluminium oxide.

The first crystalline substance isolated had the formula $C_{23}H_{34}O_4$, m.p. 243–250°C (from diluted ethanol), $[\alpha]_5^8l=+$ 14.9 \pm 2° (ethanol), $\lambda_{\rm max}$ 218 m μ in ethanol (log $\varepsilon=$ 4.24). The aforementioned UV-absorption is typical for an $\alpha\beta$ -unsaturated- γ -lactone. The substance gave a monoacetyl derivative $C_{25}H_{36}O_5$, m.p. 248–256°C (from acetone-ether), $[\alpha]_5^{8d}=+$ 3.5 \pm 2° (chloroform). The substance was compared with uzarigenin, using as criteria mixed m.p., IR-spectrum, and paper chromatographic behaviour, and was identical.

The second crystalline substance isolated had the formula $C_{29}H_{44}O_9 \cdot H_2O$, m.p. 260–272° (from diluted ethanol), $[\alpha]_B^{28} = -44.1 \pm 2^\circ$ (pyridine); on acetylation it gave a tetra-acetyl-derivative of the formula $C_{37}H_{52}O_{13}$, m.p. 174–176° (from acetone-ether); $[\alpha]_B^{89} = -8.85 \pm 3^\circ$ (chloroform); $\lambda_{\rm max}$ at 217 m μ in ethanol (log $\varepsilon=4.24$). On hydrolysis with 1% hydrochloric acid in dioxane, it gave uzarigenin and D-glucose. This evidence shows that the isolated substance is a monoglucoside of uzarigenin, viz. desglucouzarin, which has not as yet been obtained from a natural source.

From Asclepias curassavica L., a botanically differentiated species of Asclepias, Tschesche et al. ?, 8 were able to isolate uzarigenin besides six other digitaloid genins 9.

Zusammenfassung. In Asclepias syriaca L. wurden neun Cardenolide nachgewiesen. Zwei Substanzen konnten in kristallisiertem Zustand isoliert und eine davon mit Uzarigenin identifiziert werden. Das zweite kristallisierte Produkt war das β -D-Glucosid von Uzarigenin.

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Department of Chemistry of Natural Products, Slovak Academy of Science, Bratislava (CSR), September 19, 1960.

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- We thank Prof. R. TSCHESCHE, Hamburg, who kindly supplied us with a sample of uzarigenin. The UV spectra were measured in this Department by J. Suchý.

Zum Mechanismus der Immuncytolyse durch heterologe Antikörper und Komplement

Die Glykolyse von Zellen des Ehrlich-Aszitestumors (EAT) wird durch heterologe Antikörper und Komplement vollständig gehemmt¹. Dabei ist es unklar, ob die zu beobachtende Cytolyse die Ursache oder die Folge der Glykolysehemmung ist. Die Stoffwechseluntersuchungen und die Ergebnisse der Elektronenmikroskopie² gaben Hinweise, dass der Effekt auf die Glykolyse die Folge einer Störung im Stoffwechsel der Adenosintriphosphorsäure (ATP) ist.

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