

Isolation of N-acetylsialic acid from normal liver

As early as 1905 MANDEL AND LEVENE¹ showed the presence of a substance in the liver that gave a red-violet colour with orcinol-hydrochloric acid. This colour was later proved to be given by sialic acid² (*synonym* neuraminic acid³). It was therefore astonishing that KLENK AND FAILLARD⁴ found a negative orcinol-hydrochloric acid reaction in normal liver tissue. On the other hand, they isolated rather large amounts of the methyl glycoside of sialic acid (1.6% calculated on dried defatted material) in amyloid degeneration of liver. They postulated that sialic acid was a component characteristic of the pathological protein occurring in amyloid degeneration. As all proteins of globulin nature hitherto investigated contain sialic acids, and a large part of the liver proteins are globulins, we applied to normal livers the recently described method⁵ for the isolation of N-acetylsialic acid.

Normal human livers (checked by microscopical investigation) were removed as soon as possible after death (12–24 h) with intact blood vessels. The livers were perfused with physiological saline through the hepatic vein until the perfusion solution contained no visible amounts of blood cells or pigments. This was necessary because plasma proteins contain large amounts of sialic acid (about 600 mg N-acetylsialic acid/litre serum). After removal of blood, the livers were homogenized in a Turmix blender and extracted twice with 4 vol. ethanol. The defatted livers were then treated with 0.01 N H₂SO₄ at 0° C to remove cations and other low molecular substances. N-Acetyl sialic acid was liberated with 0.05 N H₂SO₄ at 80° C and the hydrolysates were chromatographed on Dowex-2 in formate form⁵. The fractions containing the sialic acid were lyophilized. The crystallization was performed by dissolving 200 mg of dried material in 1 ml of distilled water, and adding 10 vol. methanol-ethanol (2:1, v/v) and 20 vol. ethyl ether. A small amount of amorphous material was removed by filtration. To the filtrate, light petroleum (b.p. 30–40° C) was added to a faint opalescence and the flask was left for 24 h. The crystals were filtered off and additional ethyl ether-light petroleum was added to the filtrate, giving another crop of crystals. The two fractions were recrystallized from methanol-water by addition of ethyl ether.

The isolated sialic acid was compared with an authentic sample of N-acetylsialic acid (C₁₁H₁₉NO₉) isolated from human serum. X-ray diffraction pattern, optical rotation, *R_F*-values in paper chromatography⁵ and molar absorptancy indices in colorimetric methods⁶ were identical.

In one case a quantitative isolation of N-acetylsialic acid was performed. The whole liver (1850 g) contained 220 mg N-acetylsialic acid measured by the resorcinol method, *i.e.* 12 mg % protein-bound N-acetylsialic acid, calculated on fresh weight. The yield of recrystallized N-acetylsialic acid was 152 mg or 69 %.

Department of Medical Biochemistry, University of Gothenburg,
Gothenburg (Sweden)

ALF MARTINSSON
AINO RAAL
LARS SVENNERHOLM

¹ J. A. MANDEL AND P. A. LEVENE, *Z. physiol. Chem.*, 45 (1905) 386.

² G. BLIX, L. SVENNERHOLM AND I. WERNER, *Acta Chem. Scand.*, 6 (1952) 358.

³ E. KLENK, *Z. physiol. Chem.*, 273 (1942) 76.

⁴ E. KLENK AND H. FAILLARD, *Z. physiol. Chem.*, 299 (1955) 191.

⁵ L. SVENNERHOLM, *Acta Soc. Med. Uppsaliensis*, 61 (1956) 75.

⁶ L. SVENNERHOLM, *Biochem. J.*, 64 (1956) 11 P.

Received December 15th, 1956

The growth of cellulose microfibrils from *Acetobacter xylinum**

Extracellular production of cellulose microfibrils in cultures of *Acetobacter xylinum* is well established^{1,2} but the mode of their formation is obscure. An electron microscope study by MÜHLETHALER suggested that the microfibrils crystallize from a high-polymeric, amorphous precursor concentrated in a homogeneous region large compared to a bacterium². The preparations examined by MÜHLETHALER were pseudoreplicas of cellulose films withdrawn at intervals from active cultures grown in beer. Interpretation of the appearance of such preparations, however, may easily be complicated by debris from the culture, since insufficient washing of the films may leave residues from the beer, while too much washing may damage the specimen and remove pertinent material. The manner of formation of cellulose microfibrils has therefore been re-investigated in the electron microscope by incubating cultures in a synthetic medium of low molecular weight under conditions which minimized contamination and mechanical disturbance.

Cells were prepared free of cellulose as described by HESTRIN AND SCHRAMM¹ except that they were not freeze-dried but stored as a suspension in phosphate-citrate buffer, pH 6.0, 0.01 M in phosphate. Drops of an appropriate dilution of the suspension in 2 % glucose, 0.27 % Na₂HPO₄

* Issued as N.R.C. No. 4212.