Vol. 61

mated as cupric oxide, we are at present developing quantitative methods for these two drugs as well as sulfanilamide.

MEDICINAL CHEMISTRY LABORATORY W. A. LOTT THE SQUIBB INSTITUTE FOR MEDICAL RESEARCH NEW BRUNSWICK, NEW JERSEY FRANK H. BERGEIM **Received September 20, 1939** 

## BIOTIN AS A GROWTH FACTOR FOR THE BUTYL ALCOHOL PRODUCING ANAEROBES

Sir:

Recently the "growth factor" requirements of the butyl alcohol-producing clostridia has excited considerable interest.<sup>1,2,3</sup> It has been found that several species of these organisms require a widely distributed organic acid, which cannot be replaced by any known available growth factors.

Using the test conditions of McDaniel, et al.,<sup>3</sup> and a culture of Clostridium butylicum (Amer. Type Culture Coll. No. 6015), we have shown that biotin<sup>4</sup> is the only accessory substance required by this organism for luxuriant growth even on a synthetic medium.

The results of three tests are given in Table I, the first two on the basal medium of McDaniel, et al.,<sup>3</sup> the third using a basal medium exactly the same except that an additional 0.1% asparagin was substituted for the casein hydrolysate. Cultures 2 and 3 were subcultured from the unsupplemented tube of culture 1. Turbidities at the

	TABL	εI	
	Turbiditi	es (galvanomete	r readings)
Micrograms supplement per cc. of medium	Culture 1 (1-3 diln.) hydrolyzed casein medium	Culture 2 (1-2 diln.) hydrolyzed casein medium	Culture 3 (undiluted) asparagin medium
	Biotin Sup	plement	
0.000000	15.0	17.0	2.2
. 0000133	25.0	19.0	10.2
.0000266	32.0	24.0	29.0
. 000053	46.0	34.0	54.0
.00010	60.0	49.0	75.0
.00020	68.0	66.0	88.0
.00066	81.0	67.0	94.0
Liv	er Concentra	te Supplement	
.0133	33.0	28.0	39.0
.0333	52.0	39.0	60.0
.0666	65.0	58.0	79.0
.1332	6 <b>6</b> .0	66.0	94.0
. 2664	67.0	74.0	94.0
. 6660	70.0	68.0	96.0
(1) Weizmann el	al Biochem I	81 619 (1937)	

(2) Brown, Wood and Werkman, J. Bact., 36, 246 (1938).

(3) McDaniel, Woolley and Peterson, ibid., 37, 259 (1939).

(4) Kögl and Tonnis, Z. physiol. Chem., 242, 43 (1936).

end of the three-day incubation period were measured quantitatively with the thermoelectric turbidimeter described by Williams, et al.<sup>5</sup> The galvanometer scale was set to read zero with the uninoculated medium in the cell; a reading of 100 corresponds to complete opacity.

The "liver concentrate" was prepared in accordance with work of McDaniel, et al., and was kindly furnished by Dr. Woolley. Biotin proved to be approximately 500 times as active as this concentrate and it is evidently the only substance required in addition to the ordinary nutrients. Hydrolyzed casein evidently contains small amounts of some substance which is in this case physiologically equivalent to biotin. Whether the response which we have obtained is specific for biotin cannot at present be stated. In any event, these results supply further evidence for the great physiological activity of biotin.

We wish to express our thanks to Professor Kögl, who kindly furnished us with the sample of biotin which made this work possible.

(5) Williams, McAlister and Roehm, J. Biol. Chem., 83, 315 (1929).

DEPARTMENT OF CHEMISTRY	ESMOND E. SNELL
THE UNIVERSITY OF TEXAS	ROGER J. WILLIAMS
Austin, Texas	

**RECEIVED OCTOBER 13, 1939** 

## ON THE ABSORPTION SPECTRUM OF HYPERICIN Sir:

The pigments of Hypericum perforatum causing photosensitization in animals are spectroscopically closely related to irradiated oxypenicilliopsin from Penicilliopsis clavariaeformis [A. E. Oxford, Chem. and Ind., 57, 975 (1938); C. Dhéré and V. Castelli, C. R. Soc. Biol., 131, 669 (1939); C. Dhéré, ibid., 131, 672 (1939)].

Extracted hypericin was found chromatographically to consist of five components, and we have designated the two most abundant as X and Y. Samples were interchanged with Professor Raistrick, and component Y and irradiated oxypenicilliopsin appear very similar but not identical in chemical behavior. We wish at this juncture to supplement spectroscopic observations by Dhéré and Castelli: on these pigments. We have checked their absorption maxima in absolute alcohol within 10 Å. and are in fair agreement as to the intensities.

In Table I we show the effect of solvent composition on the absorption maxima. Values are