pound I was prepared by the method of Farrar and Levine.<sup>14</sup> The product, which was an azeotrope, was found by gas phase chromatography to consist of 70% of I and 30% of the 2-isomer, in close agreement with the authors. Only chromatographically pure I was used to avoid contamination of the products by the Mannich bases of the 2-isomer. Except for II and III, which were prepared in isoamyl alcohol, each of the  $\beta$ -amino ketones was prepared from formaldehyde and the base in absolute ethanol by the usual procedure.<sup>11</sup>

Method of Testing for Antimicrobial Activity.— The test microorganisms consisted of stock laboratory strains of *Staphylococcus aureus*, *Escherichia coli*, and *Saccharomyces cerevisiae*. The minimal inhibitory concentration (MIC) of each compound, as well as of each of four antibiotics, for the test organisms was determined by the agar dilution method. Appropriate concentrations of each compound were incorporated in 15-ml. portions of liquified nutrient agar, the medium was then poured into Petri plates, and 0.05 ml. of 24-hr. nutrient broth cultures of the microbial species were spread on the solidified agar surfaces. For the yeast species, glucose yeast infusion agar and broth, instead of nutrient agar and broth, were employed.

The lowest concentrations of Mannich bases or of antibiotics that prevented the development of visible growth are listed in Table I. The range of values represents maximum fluctuation in a series of assays. The data were obtained from tests with solutions sterilized by filtration; autoclaving destroyed as much as 90% of the activity of the various bases. Additional tests carried out with the most active of the Mannich bases (VII) demonstrated that the substance is not germicidal at low concentrations nor is its antimicrobial potency inactivated by lecithin. Inasmuch as the compound suppresses the growth of gram-positive and gram-negative bacteria, and yeast cells at similar concentrations, the mechanism of action is probably not associated with suppression of cell wall synthesis. Further speculation concerning the mode of action must await additional observations of microorganisms exposed to the compound.

## Experimental

3-Acetylbenzo[b] thiophene (I).—To a rapidly stirred mixture of 134.2 g. (1.0 mole) of benzo[b] thiophene and 112 g. (1.1 moles) of acetic anhydride was added 10 g. of ferric chloride in one por-The temperature immediately rose to 96°, and the retion. action mixture became very dark. After stirring for 1.5 hr., 400 ml, of cold water was added, and the dark mixture was extracted with three 200-ml. portions of ether. The combined ether phase was washed with 10% sodium carbonate solution and dried with MgSO4. After removal of the solvent and fractionation, 105 g. (70%) of a clear colorless liquid boiling at 129-131° (1 mm.) was obtained. Upon cooling and seeding, a white crystalline mass formed which was recrystallized three times from ethanol to yield a product melting at  $64-65^{\circ}$ , which showed only one peak in the gas chromatograph. The reported melting point is  $64-65^{\circ}$ ,  $^{13} \lambda_{max}^{\text{KBr}} 6.05 \text{ (C=O)}$  and  $7.35 \mu \text{ (CH}_{s}-\text{C=O)}$ . The oxime was prepared as white platelets which melted at 123-124°.

Anal. Calcd. for  $C_{10}H_{9}NOS$ : C, 62.85; H, 4.75; N, 7.32. Found: C, 62.91; H, 4.69; N, 7.24.

Mannich Bases From I (II-IX).—In a 50-ml. flask containing 25 ml. of absolute ethanol (dry isoamyl alcohol for II and III) was added 0.05 mole of the respective amine, and the pH was adjusted to 3-4 with concentrated HCl. To this was added 8.8 g. (0.05 mole) of the ketone and 2.3 g. of paraformaldehyde. The reaction mixture was allowed to reflux for 4 hr. and was then poured into 100 ml. of dry acetone. After cooling in the refrigerator overnight, the white precipitate was collected and recrystallized from absolute ethanol.

## 4-(Alkoxystyryl)quinolines<sup>1</sup>

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The activity of 4-(4-aminostyryl)quinolines against Walker 256 tumors in rats and against KB tumor cells in cell culture<sup>2</sup> led us to prepare the series of 4-(methoxystyryl)quinolines listed in Table I for similar testing. In each case a mixture of lepidine hydrochloride and an equimolar quantity of the appropriate aldehyde was heated until reaction had taken place, the mixture was dissolved in hot methanol and neutralized by adding an excess of concentrated ammonium hydroxide, and the oil which separated upon addition of water was recrystallized from isohexane, isooctane, or ethanol until acceptably pure. Samples were submitted for antitumor tests in vivo and in vitro. None of the compounds tested showed clear-cut activity against the Walker tumor at a single 200 mg./kg. dose level, but several of them produced 50% reduction in growth rate of KB cells in tissue culture at concentrations below 4  $\gamma$ /ml., which placed them in the same range with 4-(4-dimethylaminostyryl)quinoline.

It has been suggested that there might be a correlation between the ultraviolet absorption maxima of such compounds and their cytotoxicity. Absorption spectra were examined with a Beckman DU spectrophotometer. Most of the peaks were not sharp but there were observable differences between the compounds. One interesting observation was that the wave length and intensity of absorption both diminished when a methand solution of the compound was allowed to stand 1 day, but the acetic acid solutions did not undergo this change. On the other hand, the acetic acid solutions had an additional peak at a longer wave length than the methanol solutions. The shift in the methanol solutions was least for the dimethoxy and trimethoxy compounds containing a methoxy group at the 2-position, but 4-(2-methoxystyryl)quinoline itself exhibited a decided shift. The greatest difference between log  $\epsilon$  for the acetic acid solution peak and the peak in methanol was exhibited by the three monomethoxy compounds and the 2,4-dimethoxy compound. The presence of a 2-methoxy group seemed to increase cytotoxicity except in the 2,3-dimethoxy compound.

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<sup>(2)</sup> C. T. Bahner, L. M. Rives, and C. Breder, J. Med. Chem., 7, 818 (1964); C. T. Bahner, D. Brotherton, W. H. Chapman, Jr., W. Longmire, H. B. Orr, L. M. Rives, E. B. Senter, and W. Yee, *ibid.*, submitted for publication.

<sup>(14)</sup> M. W. Farrar and R. Levine, J. Am. Chem. So ., 72, 4433 (1950).

TABLE I

4-(Алкохузтукил.)quinoline

- B
<u></u> /
0
H)
<pre>Z</pre>

													· λ <sub>max</sub> mµ (log	e)			Tr / Cr &
		71 W	Read	rtion term	Viold		, Caled	- 21	Found	1. C	Fresh	in methanol after	Change on	In acetic		$E10_{50}$ , $d$	200
No.	Alkoxy group	.0°.	hr.	С.	5	Formula	U	Ξ	0	Ξ	solution	standing	standing	acid	Difference <sup>c</sup>	γ∕ ml.	mg./kg.
Т	$2-0$ CH $_3$	1217.0	ĊJ	160	28	$\rm O_{18}H_{15}NO$	87. <u>7</u> 9	5.79	82,50	6 <del>1</del> .6	345 (4.3)	315(4.0)	-30(-0.3)	390(3.85)	+45(-0.4)	1.0	_
Ŷ	3-OCH.	0.54.9	+	14	≎: ₽	CisHeNO	82.15	5.70	1: 2: 2:	5.76 76	330 (4.3)	310(4.0)	-20(0.3)	320(4.0) 365(3.85)	+35(-0.4)	1-	
: .:	4-0CH3	<i>;</i> ,4[0	; +	991	<u>6</u>	C <sub>Is</sub> H <sub>is</sub> NO	82.75	õ. 79	82.58	5.53	350(4.3)	315 (4.0)	-35(-0.3)	320(4.0) 405(3.8)	+55(-0.5)	6	-
-	0.2 (OCH.)	116/4	9	150	:2	CHNO.	N N N	У. Х	+ - 	22	330(4.4)	320(4.3)	10(-0.1)	315(3.8) 380(4.3)	+50(-0.1)	Ē	_
h 10	2,0-(OCH3)2 9 4-(OCH3)2	$130^{f,i}$	2 21	145	; <u>;</u> ;	CraHisNO.	N S	N.S. (1	11,95	08.10	360(4,1)	350(3.8)	-10(-0.3)	430(3.7)	+70(-0.4)	17	
	9 5.(OCH.).	457.1	ı ÷	57	96	CHNO.			X0 - X1	6.15	360 (4.2)	360 (4.1)	0(-0.1)	315(3.8) $420(4.1)^{l}$	+60(-0.1)		
	-10-100-013/2		1		ì						325(4.2)	320(4.1)	$-\tilde{z}$ ( $+0.1$ )	370(4.1)			
1		1	01		ē			5 5 7	91 - 91 - 91	ע ס י	10-11-226	(N. 27012	15 0 - 15 L	330(4.0) 320(4.0)	(0, 0, -0.5)	17	) 2 0
-	$3,4-(0011_3)_2$	10 (1 + 3)	-	( NC)	1	2 N. 71 H 61 Y			01,01							:	
9	3.4-0.010	n-7071		17	5	C.H., VO.	le SE	SZ T	19,87	26° †	355 (4,2)	305(3,9)	50(0.3)	310(3.8) 418(4.0)	+65(-0.2)	X	` <b></b>
2	24403200 + <b>6</b> 6		-											315(3.9)			
6	3.4-(0C,H.),	$105^{j,a}$	+	150	91	$(2_{11}H_{21}NO_{2})$	[30, 81]	4.36	78. (6)	11.1	360(4.3)	330(3,8)	(-30(-0.5))	425(4.1)			
:														315(3.9)	+65(-0.2)	5	ľ,
10	$2.4.5-(OCH_3)_3$	$133^{1/2}$	ц. Г	145	44	$C_{20}H_{19}NO_3$	74.71	5.95	74.60	5.94	375(4.2)	370(3.9)	$-\tilde{a}(-0.3)$	460(4.3)	+85(+0.1)	-	
11	3.4.5-(OCH <sub>3</sub> ),	167 - 168		155	5	$C_{20}H_{19}NO_3$	11.11	5.95	74.88	5.84	345(4.5)	350 (3.6)	+5(-0.9)	410(4.3)	+65(-0.2)		
12	$3,4,5-(OCH_3)_3$	$134^{j,i}$	::	155	+	$\mathrm{C}_{20}\mathrm{H}_{19}\mathrm{NO}_3$	74.71	5.95	74.96	5.80	347(4.3)	315(4.0)	-30(-0.3)	410(4.2)	+65(-0.1)	21	1.7"
										:	-		-	315(4.0)	-		
and in	orrected for therm fresh methanol se	ometer ster alter $\frac{d}{d}$	n expost Results e	ne; det of the st	ermined w andard <i>in</i>	rith Thiele tub <i>ritr</i> ø KJB tume	e. <sup>6</sup> Ave ar cell inb	rage of t ibition to	wo analy ests carrie	ses by M ed out un	teller and St ider sponsor	rauss, Oxford, ship of the Ca	Fingland. 71 meer Chemoth	Atterence beta erapy Nation	xeen peak m ace al Service Cente	the actd r at the	solution Univer-
sity o	f Miami Cell Cult	ure Labora	fory and	Southe	m Resear	ch Institute.	- We are	grateful	to Profe	ssor Ale	xander Hade	low, Mr. J. E	. Everett, and	Mr. B. C. V.	Mitchley of the mission in A	: Cheste mubic o	r Beatty il an the
Resca lav fo	rch Institute for a flowing tumor im-	tata on tox plantation (	ar on the	a activi » first da	A agamst w of the t	the Watker 2 toxicity observ	ation. ´	r m rats Furnor-be	weigning earing an	nuz uuz mals we	v g. – rach ( re sacrificed	sompound was approximate	s administered ly S days later	as a single 1. and the aver	p. injection in A age weights of ti	u suum unors ii	n un unc i treated
md u	ntreated hosts are	coported a	us the r	atio T/(	🗄 / Rec	rystallized from	m ethanc	ol. • W1	ûte. 🤺 R	cerystal	lized from is	sohexanes.	Yellow. 'Re	erystallized fi	om isooctancs.	<sup>k</sup> The	peaks at
he lo	wer wave lengths <b>v</b>	ane very sma	dl elevat	tions on	a long pla	teau. ' Killed	none of :	s test rati	s at 400 n	ıg./kg.	" killed F.c	d 3 test rats <i>z</i>	it 400 mg./kg.				