## Inversion of Configuration in the Bromination of Vinylic Mercurials<sup>1</sup>

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The bromination of cis- and trans-1-propenyl- and 2-butenylmercury(II) bromide in carbon disulfide occurs with predominant inversion of configuration at the double bond. The rate of bromination of trans-2-butenylmercury(II) bromide in carbon disulfide is 75 times as fast as that of n-propylmercury(II) bromide. The observed inversion of configuration is consistent with a trans addition of bromine to the carbon-carbon double bond of the vinylic mercurials followed by a trans elimination of mercury (II) bromide. In contrast to the results in carbon disulfide, the bromination of the vinylic mercurials takes place with retention of configuration in pyridine.

A study of the bromination of cis- and trans-2butenyl- and 1-propenylmercury(II) bromides (compounds 1, 2, 3, and 4) in carbon disulfide was initiated in an attempt to generate free propenyl radicals of known stereochemistry and to investigate their stereochemical fate.2 A free-radical pathway for the bromination of these compounds in carbon disulfide was

anticipated at the outset, since Jensen had shown that the bromination of either cis- or trans-4-methyleyeloproduced the same ratio of cis- and trans-1-bromo-4methylcyclohexane and had interpreted the loss of of stereochemistry in terms of a free-radical reaction.3 The possibility of an alternative pathway involving retention of stereochemistry in the bromination of the

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vinylic mercurials was also considered, since Jensen<sup>3</sup> had shown that the bromination of the isomeric methylcyclohexyl mercurials in methanol proceeds with 85% retention of stereochemistry. Here, we report the surprising finding that bromination of 1-propenyl and 2-butenyl mercurials leads to propenyl and butenyl bromides of inverted stereochemistry.4

## Results

Synthesis and Stereochemistry of Vinylic Mercurials. -cis- and trans-1-properly- and 2-butenylmercury(II)

hexylmercury(II) bromide in degassed carbon disulfide

<sup>(1)</sup> Supported in part by the National Science Foundation, Grants GP- $28586\,\mathrm{X}$  and GP-2018.

<sup>(2)</sup> L. A. Singer in "Selective Organic Transformations," Vol. II. B. S. Thyagarajan, Ed., Wiley, New York, N. Y., 1972, p 239.

<sup>(3)</sup> F. R. Jensen and L. H. Gale, J. Amer. Chem. Soc., 82, 148 (1960).

<sup>(4)</sup> Although the chemistry of cis- and trans-1-propenylmercury(II) bromides has been studied in detail.5 the halogenation of these compounds has not been reported.

<sup>(5)</sup> A. N. Nesmeyanov, A. E. Borisov, and N. V. Novikova, Izv. Akad. Nauk SSSR, Otd. Khim. Nauk, 1216 (1959); Bull. Acad. Sci. USSR. Div Chem. Sci., 1174 (1959).

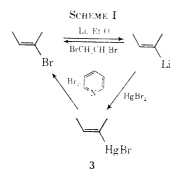
TABLE I Bromination of Vinylic Mercurials in Pyridine and Carbon Disulfide

Compd	Con Par	RHgBr, mmol	Br <sub>2</sub> , mmol	Vinyl bromides		
•	Conditions			——————————————————————————————————————	trans	Yield, "
Cis, <b>3</b>	Pyridine, air, $0.035 M$	0.25	0.25	99.6	0.4	92
Trans, 4	Pyridine, air, $0.035 M$	0.25	0.25	8.9	91.9	95
Cis, 1	$CS_2$ , air, $0.05 M$	0.11	0.11	21.8	78.2	.,,,
Trans, 2	$CS_2$ , air, $0.05 M$	0.10	0.10	73.5	26.5	
Cis, <b>3</b>	$CS_2$ , air, $0.03 M$	0.27	0.27	11.6	88.4	100
Trans, 4	$CS_2$ , air, $0.06~M$	0.20	0.20	92.9	7.1	102
Trans, 4	$CS_2$ , degassed, 0.06 M	0.12	0.12	91.8	8.2	11/2
Trans, 4	MeOH, air, $0.02 M$	0.02	0.02	77.8	$\frac{3.2}{22.2}$	

bromide were synthesized stereospecifically by reaction of mercury(II) bromide with the corresponding vinylic lithium reagents<sup>6</sup> prepared from stereochemically pure vinylic bromides. The isomeric 1-propenyl- and 2-butenyllithium reagents had been prepared previously and demonstrated to be configurationally stable, once formed.7-9 The stereochemistry of cis- and trans-1propenylmercury(II) bromide, 1 and 2, was originally assigned by Nesmeyanov<sup>5</sup> on the basis of the method of synthesis from the corresponding lithium reagents. Jensen<sup>10</sup> has recently demonstrated that exo- and endo-2-norbornylmagnesium bromide react with mercury (II)bromide to give the corresponding alkylmercury(II) bromides with complete retention of configuration. The stereochemistry of di-cis- and di-trans-propenylmercury(II), 5 and 6, respectively, has been assigned unambiguously by nmr. 11

The assignment of stereochemistry of cis- and trans-2-butenylmercury(II) bromide is based on the method of synthesis and on the nmr spectra of the compounds. Owing to deshielding by the cis mercury atom, " the chemical shift of the vinyl proton cis to mercury in 3 is à 5.56 while the chemical shift of the vinyl proton trans to mercury in 4 is  $\delta$  6.25.

The reaction of alkyl mercurials with bromine in pyridine is stereospecific.3 We have found that the reaction of the vinylic mercurials 3 and 4 with bromine in pyridine produces the corresponding vinylic bromides with retention of configuration (Table I). The chemical evidence for the assignment of configuration of cis-2-butenylmercury(II) bromide, 3, is shown in Scheme I.



<sup>(6)</sup> G. M. Whitesides, C. P. Casey, and J. K. Krieger, J. Amer. Chem. Soc., 93, 1379 (1971).

Bromination of Vinylic Mercurials in Carbon Disulfide.—The bromination of the vinylic mercurials 1, 2, 3, and 4 in carbon disulfide was highly stereoselectors but gave vinyl bromides having opposite stereochemistry from the mercury compounds from which they were derived (Table I). Similar results were obtained in degassed solutions or in the presence of air. which normally retards free-radical brommation of mercurials.3 Bromination of 4 in methanol also great predominantly the vinylic bromide of opposits and figuration.

Relative Rates of Bromination of Alkyl and Vinylic Mercurials.—To aid in the determination of the mech anism of the bromination of the vinylic mercurials. it was of interest to determine the relative rates of busmination of 1-hexene, trans-2-butenylmercury(II) bromide, 4, and n-propylmercury(II) bromide. Since both vinylic and alkylmercury(II) compounds decolorize carbon disulfide or pyridine solutions of bromine immediately, competition techniques were used to measure relative rates. Dilute carbon disulfide or pyridine solutions of pairs of the substrates were treated with a deficiency of a dilute solution of bromine. The relative yields of the two products were determined by gas chromatography and used to establish the relative rates of bromination shown in Table II.

## Discussion

Two mechanisms for the halogenation of organomercurials have been demonstrated. In polar solvents in the presence of air (a free-radical inhibitor). haolgenation of alkylmercury(II) halides proceeds by a stereospecific four-center mechanism leading to retention of configuration. In nonpolar degassed solvents, the halogenation proceeds by a completely nonstereospecific free-radical mechanism. The inversion

(12) F. R. Jensen and B. Rickborn, "Electrophilic Substitution of Organomercurials," McGraw-Hill, New York, N. Y., 1968, Chapter 4, and references cited therein.

<sup>7)</sup> D. Seyferth and L. G. Vaughan, J. Amer. Chem. Soc., 86, 883 (1964).

<sup>(8)</sup> A. S. Dreiding and R. J. Pratt, J. Amer. Chem. Soc., 76, 1902 (1954).

<sup>(9)</sup> The stereochemistry of 1-propenyllithium reagents has been determined directly by nmr spectroscopy: D. Seyferth and L. G. Vaughan, J. Organometal, Chem., 1, 201 (1963).

<sup>(10)</sup> F. R. Jensen and K. L. Nakamaye, J. Amer. Chem. Soc., 88, 3437

<sup>(11)</sup> D. Moy, M. Emerson, and J. P. Oliver, Inorg. Chem., 2, 1261 (1963).

TABLE II
RELATIVE RATES OF BROMINATION IN
CARBON DISULFIDE AND PYRIDINE

Registry no. 25264-93-1	Compd	Relative rate in CS <sub>2</sub>	Relative rate in C <sub>b</sub> H <sub>b</sub> N I.0
	$/=\langle HgBr $ (4)	2.2	34
18257-68-6	∕ HgBr	0.029	0.9

of configuration observed in the bromination of the vinylic mercurials is not compatible with either of the two above mechanisms and must be occurring by a different mechanistic pathway.

The predominant inversion of configuration observed in carbon disulfide can best be explained by a trans addition of bromine to the carbon-carbon double bond of the vinylic mercurial followed by a trans elimination of mercury(II) bromide from the resulting adduct. This addition-elimination mechanism implies that the carbon-carbon double bond is more reactive toward

bromine than is a carbon-mercury bond. If the addition-elimination mechanism is correct, then, by extension, one would expect a vinylic mercurial to be more reactive than an alkyl mercurial in carbon disulfide. The relative rate studies reported in Table II are in agreement with the proposed addition-elimination mechanism. Thus, trans-2-butenylmercury(II) bromide, 4, is about 75 times more reactive toward bromine in carbon disulfide than is n-propylmercury(II) bromide. The vinylic mercurial is about 2.2 times more reactive than 1-hexene.

The retention of configuration observed in the bromination of vinylic mercurials in pyridine must be due to direct electrophilic attack of bromine on the carbon-mercury bond. Pyridine interacts strongly both with organomercurials and with bromine; either interaction could lead to the change in mechanism upon changing solvent from carbon disulfide to pyridine. Bromine reacts with pyridine to produce a bromine-pyridine complex<sup>12</sup> which is a more electrophilic brominating agent than bromine. However, the pyridine-bromine complex should be more reactive toward both the

carbon-carbon double bond and the carbon-mercury bond of a vinyl mercurial. Consequently, a change in mechanism would not necessarily be expected upon changing the nature of the electrophile. Pyridine forms complexes with alkylmercury compounds. The electron donation from pyridine to mercury would be expected to activate selectively the carbon-mercury bond of a vinylic mercurial toward reaction with electrophiles. Thus, although in carbon disulfide the car-

bon-carbon double bond of 1-hexene is 34 times more reactive toward bromine than the carbon an energy being of n-propylmercury(II) bromide, the two companies have the same relative reactivity towards brome a pyridine suggesting an increased relative reactivity of trans-2-butenylmercury II. 15 mide compared with 1-hexene increases from 100 mag on going from carbon disulfide to pyridine.

The inversion of configuration reported here for  $\psi_{\alpha}$ bromination in carbon disulfide of the vinylic more curials 1, 2, 3, and 4 is surprising, since the extent research on the halogenation of vinytic service at a ried out in the research groups of Neshawayayay was Reutov had previously indicated that halogrees occurred either by a four-center mellsown as ag complete retention of stereochemistry or last to nonstereospecific free-radical mechanism in the contraction of the con genation of vinylic mercurials is reconsidcific in polar selvents and nenstereospecific solvents. Examples of balegoration or the s retention of stereochemister include disof cis- and trans-stillerylmereury (H) Let a hogy if oxane,14 the indination of rise and franciscoses mercury(II) chloride in methanol, dioxecformamide. 15 and the bromingation of zero a lowers styrylmercury(II) bromide in methersulfoxide. In contrast, the same next... trans vinylie halides is obtained in the lear iodination of cis- and trans-B-streviews mide in carbon tetraebloride or in large study of the oxymercuration products of eller reported that the iodination of classified ? mercuri-4-methoxy-2-pentene in carbon and the bromination of cis- and trans-decision. 4-methoxy-2-pentene in pyridine both sake pla retention of configuration. Thus, the last trees in carbon disulfide of compounds 1, 2, 3 pm 2 stitutes the first example so far reported of the onverse, a of configuration in the halogenation of virialic curials.

The only other examples of inversion of configuration in the halogenation of vinyl metal constraints from found in the chemistry of boron. Mathematical ported that the bromination of cis- or the second configuration. The reaction proceeds by the times were figuration.

<sup>(13)</sup> For a recent review of halogenation of vinytic mercurans congluss on the work of Russian chemists, see L. G. Makarova in "Organizations," Vol. 1, E. I. Becker and M. Tsutsui, Ed., Wijev, New York, 2011, 1970, pp 325-345.

 <sup>(14)</sup> A. N. Nesmeyanov and A. E. Borisov, Tetrahedian 1, 1, 2, 1997.
 (15) I. P. Beletskava, V. I. Karpov, and O. A. Regtow 2x 2000 (1988).
 SSSR, Otd. Khim. Nauk, 1707 (1964); Bull. Acad. Sci. USSE, 160 (1988).

<sup>(16)</sup> W. L. Waters, W. S. Linn, and M. C. Caserio, J. Apple 1915 of St. 90, 6741 (1968).

<sup>(17)</sup> The assignment of stereochemistry of the mercury composition a made on the assumption that the trans <sup>138</sup>Hg-H coupling consequence than the cis <sup>139</sup>Hg-H coupling constant; this assumption, a second studies of model compounds it. Nonetheless, the assumption, as second error, since the chemical shift of the vinyi hydrogen in the compounds it signed the structure cis-3-chloromercur 4-methox, 2 neutron corrections ppm downfield from the vinyl hydrogen in the compound assumption as structure, although an opposite effect would be expected from the ways at Oliver, it

 <sup>(18)</sup> W. L. Waters and E. F. Kiefer, J. Amer. Chem. Soc., 39 (19); 1
 (19) D. S. Matteson and J. D. Leidtke, J. Amer. Chem. Soc., 37, 129
 965).

Table III

Competition Reactions for Bromine in Carbon Discipline and Pyridine

	Initial concentrations of reactants, 10 <sup>2</sup> M			Molar ratio of products				
Solvent	/ <del>-</del> HgBr	HgBr	<b>/</b>	$\mathrm{Br}_2$	2/Bromo- 2/butene-	Br	Pr Br	
$CS_2$		0.56	0.56	0.062		1		Rate site
		0.00				1	32	- 1
$\mathrm{CS}_2$	4.1		5.5	2.2	1.42		1 0	2 2 3
$\mathrm{C_5H_5N}$	7.6	8.5		7.1	29 - 4	1 ()		314
$\mathrm{C_{5}H_{5}N}$		8.3	9.5	10.4		1.0	1 (10	

<sup>a</sup> Determined by disappearance of 1-hexene.

tion of bromine to the carbon-carbon double bond to give an isolated intermediate dibromide, which then undergoes a base-catalyzed trans elimination. The

carbon-boron bond is relatively inert to bromine<sup>20</sup> and it is not surprising that the carbon-carbon double bond is attacked preferentially in vinyl boron compounds. Brown<sup>21</sup> has recently shown that the dibromide prepared from a vinylborane decomposes in the presence of base by a trans elimination to give a vinyl bromide of inverted configuration; however, thermal decomposition of the dibromide in carbon tetrachloride proceeded by a cis elimination to give a vinyl bromide of the same configuration as the vinylborane.

## **Experimental Section**

trans-2-Butenylmercury(II) Bromide.—Mercury(II) bromide (17.0 g, 0.047 mol) was added to an ether solution of trans-2-butenyllithium<sup>6</sup> prepared from trans-2-bromo-2-butene (6.4 g, 0.047 mol). The reaction mixture was poured into water and the ether layer was separated, washed with water, dried (MgSO<sub>4</sub>) and concentrated to 100 ml. Cooling to  $-78^{\circ}$  gave white, lustrous crystals of trans-2-butenylmercury(II) bromide (3.5 g, 22°C, yield): mp 110–113°:  $\delta_{\text{TMS}}^{\text{CH-SC}_2}$  6.25 (quartet of multiplets, J=6.5 Hz, 1 H, H C=C), 2.01 (quintet, J=1.5, Hz, 3 H, C=CCH<sub>3</sub>Hg), and 1.84 (doublet of quartets, J=6.5, J'=1.5 Hz, 3 H, CH<sub>3</sub>C=CHg). Coupling due to <sup>199</sup>Hg was also evident:  $J_{\text{Hg,}\Theta\text{-}\text{CH}_3}=188\,\text{Hz}; J_{\text{Hg,}B\text{-}\text{CH}_3}=44\,\text{Hz}.$ 

Anal. Calcd for C<sub>4</sub>H<sub>7</sub>BrHg: C, 14.32; H, 2.10; Br, 23.81. Found: C, 14.23; H, 2.02; Br, 23.48.

c/s-2-Butenylmercury(II) Bromide.—Reaction of mercury(II) bromide (9.0 g, 0.024 mol) with an ether solution of c/s-2-butenyllithium<sup>6</sup> (35 ml, 0.61 N, 0.021 mol, 85% cis) at room temperature for 2 days gave a solution containing a gray, floculent precipitate. The precipitate was collected and dissolved in 200 ml of methylene chloride. The methylene chloride solution was filtered, dried (MgSO<sub>4</sub>), and cooled to  $-78^{\circ}$  to give white crystals of c/s-2-butenylmercury(II) bromide (2.4 g), mp 170–172°. Additional c/s-2-butenylmercury(II) bromide (1.7 g, 4.1 g total, 57% yield) was isolated from the ether solution: nmr  $\delta_{\rm TMS}^{\rm CDCO}$  5.56 (quartet of multiplets, J=6.5 Hz, 1 H, HC=C), 2.00 (m, 3 H, C=CCH<sub>2</sub>Hg), and 1.79 (doublet of quartets, J=6.5, J'=1 Hz, CH\_CH=CHg). Coupling due to <sup>199</sup>Hg was evident:  $J_{\rm Hg, G-CH_3}=202$ ,  $J_{\rm Hg, \beta-CH_3}=40$  Hz.

Anal. Caled: C, 14.32; H, 2.10; Br, 23.81. Found: C, 14.23; H, 2.13; Br, 23.52.

trans-1-Propenylmercury(II) Bromide.—Reaction of the (II) bromide (17.5 g. 0.048 mol) with an other solution of propenyllithium<sup>6</sup> (48 ml, 1.25 N, 0.047 mol, 87) temperature for 1 hr give a gray precipitate and the fitter tion. The reaction mixture was poured into 100 h of methylene chloridate solid was dissolved in 100 ml of methylene chloridate resulting yellow solution was dried (MgSO, 44) and centrated to 40 ml. Cooling the solution to a cooling crystals of trans-1-propenylmercury(II) bromade yield), mp 120–123° (lit. 22 mp 120–121.5°).

cis-1-Propenylmercury(II) Bromide a Reaction at (II) bromide (12.5 g, 0.035 mol) with an ether sade a propenyllithiums 25 ml, 1.32 N, 0.33 mol, 94 temperature gave a dark solution. The ether sade appound into water, separated, washed with water, and appound into water, separated washed with water, and appointed filtered, and cooled to  $-78^{\circ}$  to give cis-1-properate bromide (3.0 g, 29°, yield), mp 60 64 at the effect of from 10 ml of other gave 2.4 g of cooleagues bromide, mp 63 67° (ht 6) ap 62.5 60 5 s.

n-Propylmercury(II) bromide was prepared by the constraint Slotta and Jacob 2°. Reaction of n-propylmantescent test (0.31 mol) with mercury II bromide at 40 mol process propylmercury(II) bromide (58.1 g, 0.19 mol, 596), and the clitter of the constraint of the con

Reaction of Vinylic Mercury II: Bromides with Browning When a 0.1 N solution of the vinylic mercury II: be made as 0.1 N solution of the vinylic mercury II: be made as 0.1 N solution of bromine in either carbon disufficie, preface or pyridine were mixed in the presence of air, the browning of disappeared immediately. The cistrans ratio of the virbromides formed was determined by glpc on a 12-ft V7-TC column at 30°. In several cases, yields were determined decane as an internal vpc standard. The vinylic highest identified by comparison of glpc retention times with according samples. In addition, trans-2-bromo-2-buttene from the mination of cis-2-bromo-2-buttene from the bromination of the butteryllar bromide were collected from the content matograph and were identified by comparison of their confection with their spectra of authentic samples.

In one case, a solution of trans-2-butenylmercury (1) proposed and a solution of bromine in earbon disulfide were placed as opposite arms of an inverted U tube. The solutions were degassed by freezing the solutions with liquid nitrogen, evacuating the apparatus with a vacuum pump, closing the system off itset the vacuum line, and melting the solutions. This procedure was repeated five times before the solutions were warned to over temperature and mixed.

The reactions of bromine with the vinylic mercur  $\mathbb{N}(W)$  by an example are summarized in Table I.

Competition Experiments.—The relative rates of bromanate is of 1-hexene, n-propylmercury(II) bromide, and trans-Admicration mercury(II) bromide, 4, were determined by competition of the substrates for a deficiency of bromine. Dilute curbon as illustrates for a deficiency of bromine. Dilute curbon as illustrated in pyridine solutions of pairs of the substrates were strong rapidly at room temperature while a dilute carbon destified a pyridine solution of bromine was added slowly by syntage. The relative amounts of 1,2-dibromohexane, n-bromopropulae and n-octane or n-nonane as an internal standard (Table 11). The yields of all three products were found to be nearly quantitative in separate bromination experiments.

(23) K. H. Slotta and K. R. Jacob, J. Prakt. Chem., 120, 249 (1925)

<sup>(20)</sup> Tri-n-butylborane is not attacked by bromine or iodine in carbon tetrachloride; however, neat bromine attacks both the carbon-boron and the carbon-hydrogen bonds. J. R. Johnson, H. R. Snyder, and M. G. Van-Campen, Jr., J. Amer. Chem. Soc., **60**, 115 (1938).

<sup>(21)</sup> H. C. Brown, D. H. Bowman, S. Misumi, and M. K. Unni, J. Amer. Chem. Soc., 89, 4531 (1967).

<sup>(22)</sup> D. Seyferth and L. G. Vaughan, J. Organometal, Chem. 5, 780 (1966)

The relative rates were calculated using the following formula the sky into account differences in initial and final concentrations as desirate.

$$\frac{k_A}{k_B} = \frac{\ln |A| - \ln |A_0|}{\ln |B| - \ln |B_0|}$$

Registry No.--1, 6727-46-4; 2, 6727-44-2; 3, 46782-37-4; 4, 40782-38-5; mercury (H) bromide, 7789-47-; trans-2-basery lithium, 28944-86-7 crs-2-batenyllithium, 28944-85-6 crgrs-1-propenyllithium, 6386-72-7; crs-1-propenyllithium, 6386-72-7 bromine, 7726-95-6.