(m, 2 H, $-CH_2O_-$), 7.21 (d, 1 H, J = 8 Hz, Ar H), 7.52 (d, 1 H, J = 8 Hz, Ar H), 13: mp 127–128 °C; IR (CHOl₃) 3580, 3360, 1755, 1679 cm⁻¹; ¹H NMR (CDCl₃) δ 1.13 (s, 3 H, \Rightarrow C-CH₃), 1.23 (d, 3 H, J = 7 Hz, $-CHMe_2$), 1.27 (d, 3 H, J = 7 Hz, $-CHMe_2$), 1.5–3.2 (m, 7 H), 3.33 (septet, 1 H, J = 7 Hz, $-CHMe_2$), 1.5–3.2 (m, 1 H, J = 7 Hz, J = 7 H 6.87 (d, 1 H, J = 8 Hz, Ar H), 7.17 (d, 1 H, J = 8 Hz, Ar H). 14: mp 80-83 °C; IR (CHCl₃) 1755, 1680, 1660, 1637, 1582 cm⁻¹; UV (MeOH) 343 nm $(\epsilon 4954)$; ¹H NMR (CDCl₃) δ 1.10 (d, 6 H, J = 7 Hz, $-CHMe_2$), 1.15 (s, 3 H, $> C-CH_3$), 1.5-2.6 (m, 7 H), 2.93 (septet, 1 H, J = 7 Hz, $-CHMe_2$), 4.07 (m, 1 H, epoxy H), 4.70 (m, 2 H, $-CH_2-O_-$), 6.42 (d, 1 H, J=7 Hz, C_{11} H), 6.99 (d, 1 H, J = 7 Hz, C_{12} H).

Ketone 11 and the corresponding isomer with cis A-B ring fusion are each obtained in \sim 15% yield, and the cis isomer is the more stable isomer. Quinone resulting from oxidation of the aromatic ring is also produced (35%). Separation was effected by column chromatography on silica gel and recrystallization. Efforts to improve the oxidation procedure are in progress.

(10) Bis epoxidation of 14 with m-CPBA to afford 3 in one step (22%) has been observed and is under further study.

(11) The structure of racemic 1 (mp 255-256 °C) and 3 (mp 225-226 °C) was established by comparison of spectral data with those of the natural substances. We thank the late Professor S. M. Kupchan for providing IR and ¹H NMR spectral data.

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α -Alkoxyorganolithium Reagents. A New Class of Configurationally Stable Carbanions for Organic Synthesis

Recent preparations of stereochemically defined organolithium reagents have provided a most useful approach to the stereospecific construction of carbon-carbon bonds. These reagents have however been limited largely to vinyl- and cyclopropyllithiums.1 We report here the preparation of a new class of configurationally stable organolithiums which are sp³ hybridized, acyclic, and may be obtained as diastereomerically or enantiomerically pure reagents.

Early attempts to prepare chiral organometallics from metals and optically active alkyl halides led to extensively racemized products, a result presumably due to the intermediacy of free radicals on the reaction pathway.² Later investigations showed however that the exchange reaction of alkyllithiums with resolved sec-butylmercuric chloride proceeded with clean retention of configuration.3 Although the exchange reported is not a synthetically useful one owing to the presence of other lithium reagents in the product, the experiment did show that sp³ organolithiums should be configurationally stable once formed. Since α -alkoxyorganolithium reagents may be prepared by a fast, low-temperature exchange from the corresponding organostannanes, we felt that this route to organolithiums should be a stereospecific one.⁵

We therefore examined the stereochemistry of the exchange reaction in the following way. 2-Benzylpropanal (1) was first treated at -78 °C with tri-n-butylstannyllithium (from n-Bu₃SnH and LiNiPr₂) and then protected with chloromethyl methyl ether (i-Pr₂NEt, 0 °C, 1 h) to produce a 1:1 mixture of diastereomers 2a and 2b (75% yield). Although the compounds did not resolve on TLC, they could be cleanly separated on a preparative scale by medium-pressure liquid chromatography (MPLC) on silica gel. 6a Compounds 2a and 2b were

then individually treated with n-butyllithium (THF, -78 °C) and after 15 min the intermediate α -alkoxyorganolithium reagents were quenched with acetone. Careful high pressure liquid chromatographic (HPLC) examination of the products showed the reactions to be totally stereospecific. Thus 2a produced a single acetone adduct (90% yield) which was different from the single product afforded by 2b. These reactions were repeated at -30 °C (THF, 15 min) and again no loss of stereochemistry was observed.7 Analogous results were obtained with trapping by trimethylchlorosilane.

While the above experiments demonstrate the stereospecific nature of the exchange and trapping, they do not distinguish between retention and inversion. The expected net retention of configuration was verified in one case by trapping the intermediate α -alkoxyorganolithium reagent with a tin halide. Thus, while the lithium reagent prepared from 2a was unreactive with tri-n-butyltin chloride, it did add to tri-n-butyltin iodide at -50 °C. The resulting α -alkoxyorganostannane was shown to be the product of retention by its correlation with the starting material, 2a.

We have also prepared enantiomerically pure α -alkoxyorganolithium reagents from aldehydes by chromatographic resolution of suitable derivatives of the intermediate stannylcarbinols. An example of this operation starts with propanal. Addition of tri-*n*-butylstannyllithium and then esterification with (-)- α -methoxy- α -trifluoromethylphenylacetyl chloride [(-)-MTPA-Cl]⁸ gave quantitatively a pair of diastereomeric esters which could be separated by MPLC.6b,9 The more mobile R^{13} ester 4 was converted into the resolved stannylcarbinol by reduction (i-Bu₂AlH, PhCH₃, -78°C, 90% yield) and was then protected with benzyl chloromethyl ether (i-Pr₂NEt, 0 °C, >95% yield) to give 5. Finally the lithium reagent was prepared as usual (n-BuLi, THF, -78 °C) and was alkylated with dimethyl sulfate to yield 6. Hydrogenolysis (10% Pd/C,

Et₂O) gave optically active 2-butanol. Whereas the (-)-MTPA ester of racemic 2-butanol displayed the carbinol methyl resonances in the NMR (CDCl₃) as a pair of doublets at δ 1.25 and 1.33, the (-)-MTPA ester from 6 showed only the lower field doublet. 10 This ester was shown to be identical with that prepared from authentic (R)-(-)-2-butanol.

The results described above involve separations of diastereomeric organostannanes as a pathway to stereochemically defined organolithium reagents. Since stereoselective preparations of α -alkoxyorganostannanes would be of considerable value in this area, we briefly examined α induction in the addition of tributylstannyl nucleophiles to several α -substituted chiral aldehydes. Our results indicate that, for α induction based only on the relative sizes of α -substituents (Cram's rule),11 the tributylstannyl anion exhibits much the same stereoselectivity as unhindered Grignard reagents. Thus 2,3-dimethylbutanal (THF, -110 °C) gives essentially the same stereochemical product distribution with either tributylstannyllithium (3:1) or methylmagnesium bromide (2.5:1). In the case of the former addition, the product stannylcarbinol mixture was protected (BnOCH2Cl, i-Pr₂NEt), lithiated (BuLi, THF, - 78 °C), and methylated (Me₂SO₄) to give the same major methylcarbinol produced by the Grignard addition. This result would seem to indicate that methylation proceeds with retention unless steric α induction with methylmagnesium bromide is opposite that observed with tributylstannyllithium.

Stereoselectivity is somewhat improved with aldehydes substituted at the β position by oxygen. With α -asymmetric aldehydes of this type, the cyclic chelate mechanism¹² would presumably be operative and anti-Cram products would be predicted. When the β -alkoxy aldehyde 7 was treated with

tributylstannyllithium in THF, a 5:1 (-78 °C) or 8:1 (-110 °C) mixture of diastereomeric stannylcarbinols was produced. After protection (MeOCH₂Cl, i-Pr₂NEt), the major diastereomer was purified by MPLC on silica gel. Lithiation (BuLi, -78 °C, THF) and methylation (Me₂SO₄) then gave the anticipated¹³ threo product¹⁴ stereospecifically. For comparison, both methyllithium and methylmagnesium bromide add to 7 (THF, -78 °C) in an essentially stereorandom manner. Although the generality of stereoselection in tin anion additions remains to be established, these preliminary results suggest that tributylstannyllithium may be added to aldehydes with moderate stereoselectivity and that the direction of the addition is that predicted either by Cram's rule or by the cyclic chelation model.15

References and Notes

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- (6) A 25 \times 500 mm LiCroprep.Si60 (25–40 μ , E. Merck No. 9390) was used, 15 mL/min: (a) 2% ethyl acetate-petroleum ether; (b) 0.3% ethyl acetate-petroleum ether
- (7) A similar sequence with 2a at 0 °C gave mainly decomposition of the lithium

- reagent. The small portion of the reagent which did survive gave a 1:1 mixture of $\bf 3a$ and $\bf 3b$ [E = C(CH₃)₂OH] on trapping with acetone. It is not clear whether or not the isomerization is due to pyramidal inversion of the anion or due to some other process related to decomposition of the reagent.
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- (9) Optical resolution of the tributyltin adduct of propanal could also be effected via formation of a urethane with (-)- α -phenylethylamine [(a) COCl₂, i-Pr₂NEt; (b) (-)-PhCH(CH₃)NH₂]. With this derivative the MPLC separation was more difficult and the S urethane analogous to **4b** eluted first. Conversion to the stannyl carbinol was effected without loss of optical activity using HSiCl₃–EtY₃N; cf. W. H. Pirkle and J. R. Hauske, *J. Org. Chem.*, **42**, 1939 (1977); W. H. Pirkle and P. L. Rinaldi, *ibid.*, **43**, 3803 (1978).
- (10) Although no peaks resulting from the (-)-MTPA ester of the enantiomeric alcohol could be seen, proportions of that material as large as 5% could have escaped detection.
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- (13) Assuming retention of stereochemistry during methylation.
- Authentic threo material was prepared from tiglic acid as follows: (1) LiAIH4, Et₂O; (2) BnOCH₂CI, *i-*Pr₂NEt; (3) BH₃, THF; NaOH, H₂O₂; (4) MeOCH₂CI,
- (15) This work was supported by NSF Grant CHE 78-01769.
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Total Synthesis of Stemolide

Sir:

Falling in the same class as the potent cytotoxic agents triptolide, tripdiolide, and triptonide, the diterpenoid bisepoxide stemolide (1), possessing the novel $18(4\rightarrow 3)abeo$ abietane skeleton, was recently isolated and described by Manchand and Blount.² Herein we report a total synthesis of

this natural product, the first route to a representative of this structural type.3

To prepare for the later incorporation of the bis epoxide moiety, the starting material, methyl dehydroabietate, was first functionalized in the aromatic ring by treatment with acetyl chloride in CS2 in the presence of Al2Cl6, providing methyl 12-acetyldehydroabietate (80%). Baeyer-Villiger oxidation with 3,5-dinitroperbenzoic acid⁵-methanesulfonic acid (CH₂Cl₂, room temperature), saponification, and Oalkylation with MeI-NaH (THF, room temperature) led to methoxy ester 2,6 convertible by EtSLi7 (HMPA-THF, room temperature) into the corresponding acid 36 (76% from 2). Following the approach of Huffman and and Stockel,8 the substituted dehydroabietic acid 3 was transformed into the dehydroabietene 7 (mp 75-77 °C) by Curtius degradation to isocyanate 4, LiAlH₄ reduction followed by Eschweiler-Clarke methylation to 5, N-oxidation to 6, and Cope elimination (72% from 3). The α -epoxide resulting from m-chloroperbenzoic acid oxidation of 7, on treatment with Et₂Al-N-i-Pr₂⁹ $(C_6H_6/PE, 50 \, ^{\circ}C)$, generated allyl alcohol 8. After conversion (n-Bu₃P/CCl₄, 0 °C) of 8 to halide 9, displacement by lithium thiophenoxide (THF, room temperature) gave thioether 10 (81% from 7). The corresponding sulfonium fluoroborate 11 was converted by BuLi (THF, -78 °C) into ylide 12, which underwent in situ electrocyclic conversion at 0 °C into the