sequence of reactions: (1) lithium aluminum hydride reduction (95%), (2) selective tosylation (86%), and (3) treatment of the primary tosylate with potassium *tert*-butoxide in tetrahydrofuran (87%). The oxetane 11 was obtained as a colorless liquid: bp 110 °C (bath temp, 1 mm); NMR (CDCl₃) 1.23 (s, 3 H), 3.87–4.23 (m, 3 H); IR (liquid film) 1370, 990–980 cm $^{-1}$; mass m/e 210 (M $^+$).

(28) For the oxetane rearrangement reactions, diethylaluminum N-methylanilide was found to be a superior reagent to diethylaluminum 2,2,6,6-tetramethylpyperidide, with which the rearrangement required the longer period of reflux. see ref 31.

of reflux, see ref 31.

(29) Bp 50 °C (bath temp, 1 mm); NMR (CDCl₃) 1.00 (s, 6 H), 3.27 (s, 2 H), 5.35 °S. 46 (m, 2 H); IR (liquid film) 3350, 1040, 970 cm⁻¹; mass *ml* e 142 (M⁺); homogeneous by TLC, AgNO₃-silica gel TLC, and GLC (AgNO₃-Carbowax 20M) (>99 % pure). The *E,Z*-mixture of the alcohol 12 was prepared independently by the Wittig reaction (butylidenetriphenylphosphorane and 2.2-dimethyl-3-hydroxypropanal).

phorane and 2,2-dimethyl-3-hydroxypropanal).
(30) Bp 145 °C (bath temp, 1 mm); NMR (CDCl₃) 1.01 (s, 3 H), 3.17–3.53 (m, 2 H), 5.30–5.44 (m, 2 H); IR (liquid film) 3350, 1035, 980 cm⁻¹; mass *m/e*

210 (M+).

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Cyclizations via Organopalladium Intermediates. Macrolide Formation

Sir

Formation of C-C bonds in a cyclization reaction varies in efficiency as a function of the ring being formed among other factors. While great strides have been made in developing methods for formation of three- to seven-membered rings, larger rings such as those of 14 and 16 members are not very accessible by C-C bond formation. For example, while progress in macrolactonization has been forthcoming, formation of macrolides by C-C bond formation is very limited. The importance of the macrolide antibiotics led us to identify such ring systems as our target. We wish to report a new approach to cyclizations that includes access to large rings and is exemplified by a synthesis of exaltolide.

Initial attention focused on allylic acetates 1⁴ and 2⁴ (Scheme I) which are available from the Diels-Alder adduct of 1-acetoxybutadiene and acrolein⁵ by straightforward methods as outlined in Scheme I. Conversion of 1 (X = PhSO₂) to its anion in THF and addition of the resultant solution to a refluxing THF solution of 2-6 mole % of tetrakis(triphenylphosphine)palladium led to a 75% isolated yield of the desired cyclized product 3⁴ as a 4:1 mixture which is isomeric at C(9).⁶ While the addition of the substrate to the catalyst could be performed by rapid addition, improvement in yield was observed by slow addition utilizing a mechanically driven syringe pump. Cyclization times were on the order of 4-10 h. It is important that if any excess sodium hydride is used to generate the anions, it must be removed by filtration prior to the addition to the palladium catalyst.

Cyclization of 2 (X = PhSO₂) proved to be most interesting in light of our previous results in which a nitrogen nucleophile was employed.^{6a} In contrast to that case, the major product was the [4.2.0] compound 4. For characterization, the sample was desulfonylated (6% Na(Hg), Na₂HPO₄, CH₃OH) and analyzed by VPC. The minor product (20%) was identified as the bicyclo[2.2.2] octene by comparison to an authentic sample prepared by the Diels-Alder reaction between 1,3-cyclohexadiene and ethyl acrylate followed by transesterification. The major product (80%) was identified as 6. Most noteworthy is the NMR spectrum which establishes the structural rela-

Scheme I. Cyclizations of Allylic Acetates 1 and 2

OTs

$$X = PhSO_2$$

OAc

 A, b

CHO

CHO

 A, b

CHO

 A, b

OAc

 A, b

OAc

^a NaBH₄, CH₃OH, or C₂H₅OH, 0 °C, 95%. ^bTsCl, pyridine, 0 °C, 73%. ^c KOH(SO₂Ph)CO₂CH₃, NaI, HMPA, 55 °C, 62%. ^aNaH, THF, (Ph₃P)₄Pd, reflux, 67–75%. ^ePh₃P⁺CH₂OCH₃Cl⁻, t-C₄H₉Li, THF, 0 °C→ room temperature, 87%. ^f (CO₂H)₂, THF, H₂O, room temperature, 94%. ^g NaCH(SO₂Ph)CO₂CH₃, NaI, HMPA, 50 °C, 95%.

tionship of protons a-d at δ 5.81, 5.55, 3.31, and 3.03 with J_{ab} = 10 Hz, J_{ad} = J_{bd} = 4.5 Hz, and J_{cd} = 9 Hz, J_{ae} = 5.5 Hz,

 $J_{\rm ae'}=2.5$ Hz, and $J_{\rm cf}=9$ Hz. The contrast in regiochemistry between the case of carbon and nitrogen nucleophiles may reflect kinetic and thermodynamic control. Conformational considerations suggest that cyclization via 7 (bulky group pseudoequatorial on a half-chair ring) should be kinetically preferred over that via 8 (bulky group pseudoaxial on a twist boat ring). With carbon as the nucleophile, cyclization is irreversible. When the nucleophile is nitrogen, the initial azetine may undergo palladium catalyzed isomerization to the thermodynamically more stable isoquinuclidine.

Utilizing a similar scheme, allylic acetates 1 and 2 ($X = CO_2CH_3$) are also available. Interestingly, cyclization of these compounds led to very poor yields of products.

Scheme II outlines the formation of 14- and 16-member macrolides and the conversion of the latter to exaltolide. The preparation of the carboxylic acid 9 was achieved from ethyl adipate as outlined. Tetrahydrofuran and 1,6-hexanediol served as the precursors to the requisite alcohols 10 and 11. Conversion of acid 9 to its acid chloride and condensation with the alcohols led to the desired cyclization substrates 12⁴ and 13.⁴ Conversion to the anion with sodium hydride in THF and addition of the resultant solution via a syringe pump to a refluxing solution of 2-6 mole % of Pd(0) catalyst produced regiospecifically the medium ring compounds 14⁴ and 15⁴ (mp

Scheme II. Macrolide Formation
$$CO_{2}C_{2}H_{5} \xrightarrow{a,b} CO_{2}C_{2}H_{5}$$

$$CO_{2}H_{5} \xrightarrow{a,b} CO_{2}C_{2}H_{5}$$

$$CO_{2}H_{5} \xrightarrow{a,b} CO_{2}C_{2}H_{5}$$

$$CO_{2}H_{5} \xrightarrow{a,b} CO_{2}C_{2}H_{5}$$

$$CO_{2}H_{5} \xrightarrow{a,b} CO_{2}CH_{5}$$

$$OAc$$

$$OCC$$

$$OAC$$

$$OAC$$

$$OCC$$

$$OAC$$

$$OCC$$

$$OAC$$

$$OAC$$

$$OAC$$

$$OCC$$

$$OAC$$

$$OCC$$

$$OCC$$

$$OCC$$

$$OAC$$

$$OCC$$

 $^{a}BH_{3}$, THF, -15 °C \rightarrow room temperature, 93%, ref 9. $^{b}C_{5}H_{5}N^{+}$ HClCrO₃, NaOAc, room temperature, 75%. CCH₂=CHLi, ether, $-78 \rightarrow -20$ °C, 78%. ^d KOH, 50% aqueous C₂H₅OH, ^eC₅H₅N, Ac₂O, room temperature, 70%. ^f References 10.8 DHP, ¹¹ CH₂Cl₂, TsOH, 40-83%. h NaCH(SO₂Ph)CO₂CH₃, NaI, DMF, 60 °C. HCl, H₂O, THF, 50-69%. i TsCl, C₅H₅N, 0 °C, 93%. k SOCl₂, ether, reflux, ¹10 or 11, (C₂H₅)₃N, ether, 50 °C, 76 or 95%. mNaH, THF [Ph₃ P] $_4$ Pd, reflux, 49-69%. n (CH₃) $_4$ NOAc, HMPA, 90-95 °C, 76-81%. 06% Na(Hg), Na₂HPO₄, THF, C₂H₅OH, -20 °C, 70-89%. PH₂, 5% Pd/BaCO₃, 2 atm, room temperature, 95-99%.

105-106 °C) in 49-69% yield. 12 NMR examination of the crude material did not reveal the presence of any terminal vinyl group which would have resulted from reaction at the allylically related carbon. Further characterization was achieved by decarboxylation to 16 (mp 110-112.5 °C) and 17 (mp 91.5-92 °C) and desulfonylation to 18 (mp 25-33 °C) and 19 (mp 46-48 °C). Observation of a 15 Hz coupling constant between the vinyl protons in 14-17 indicates the major isomers in both rings have the E configuration. Detection of the Zisomer as a minor product in 18 was confirmed by irradiating the allylic protons at δ 2 and observing small peaks with the 10.8 Hz coupling constant in the vinyl region. Unfortunately, overlap of these absorptions with those for the E isomer precluded accurate determination of the isomeric ratio. The high regioselectivity and stereoselectivity make the cyclization particularly attractive.

Hydrogenation of 18 gives tridecanolide 20 (mp 24-6 °C)¹³ and of 19 gives exaltolide 21 (mp 33-35 °C). The flexibility of the sulfone ester moiety, in part illustrated by the transformations herein, makes the use of this group especially noteworthy. In contrast to the intermolecular versions, 6b-e the malonates are considerably poorer in the alkylations, a specific advantage of the sulfone ester in this case. The success of this macrocyclization may be a result of a template effect but such speculation is postponed for subsequent publications. Further aspects of this new cyclization method are under active investigation in our laboratories.

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- (11) DHP = dihydropyran.
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