by evaporation under reduced pressure, and the residue was **dried** in vacuo. This yielded a sample of [2-methyl-2-[(diiodophenylstanny1)methyll- 1,3-propanediyl] bis [diiodophenylstannane] **(6)** that was sufficiently pure for immediate use in the following reaction: 'H NMR **(90** MHz, CDC13) **6** 1.47 **(8,** 3 H), 2.71 (a, 6 H), 7.4-7.7 (m, 15 H).

7-Methyl-l,3,5-triphenyl- 1,3,5-tristanna-2,4,9-trithiatricyclo[3.3.1.1^{3,7}]decane (1-CH₃). A sample of [2-methyl-2-[(diiodophenylstannyl)methyl] -1,3-propanediyl] bis[diiodophenylstannane] **(6)** derived from [2-methyl-2-[(triphenylstanny1) methyl] - 1,3-propanediyl] bis [triphenylstannane] **(5;** 80.3 mg, 0.0717 mmol) was treated with a solution of **hexaphenyldistannathiane** $(317 \text{ mg}, 0.433 \text{ mmol})^{15}$ in dichloromethane (10 mL) , and the mixture was stirred at 25 "C for 2 h. Solvent was then removed by evaporation under reduced pressure. Flash chromatography²⁷ of the residue (silica, hexane (60%)/dichloromethane (40%)) provided an analytically pure sample of 7-methyl-1,3,5-tri**phenyl-1,3,5-tristanna-2,4,9-trithiatricycl0[3.3.l.l~~~]decane** (1-CH,; 45.9 mg, 0.0610 mmol, 85.1%) as a white solid: mp 240-243 °C; IR **(KBr)** 1480,1430,1070,725,695 cm-'; 'H **NMR** (400.13 MHz, CDCl₃) δ 1.46 (s, average $^{4}J(^{117,119}Sn,^{1}H) = 26.1$ Hz, 3 H), 1.79 (s, average $^{2}J^{(117,119}\text{Sn},^{1}\text{H}) = 42.4 \text{ Hz}, 6 \text{ H}$), 7.5 (m, 9 H), 7.7 (m, 6 H); ¹³C NMR (100.62 MHz, 5% CD_2Cl_2 in CH_2Cl_2 , gated proton decoupling) δ 34.5 (s, $^1J(^{117}Sn,^{13}C) = 381$ Hz, $^1J(^{119}Sn,^{13}C) = 398$ Hz), 45.2 (s, average 3J(117,119Sn,13C) = 88.4 Hz), 45.8 **(8,** average $^{2}J^{(117,119}Sn,^{13}C) = 26.7 \text{ Hz}$), 128.9 (s), 130.2 (s), 134.7 (s), 139.5 (s); maas spectrum (FAB, 18-crown-6), *m/e* 756,679,265. **Anal.** Calcd for $C_{23}H_{24}S_3Sn_3$: C, 36.70; H, 3.22. Found: C, 36.85; H, 3.58.

Reaction **of Hexaphenyldistannathiane** and Tetramethylstannane with Iodine. **A** solution of hexaphenyldistannathiane (107 mg, 0.146 mmol)¹⁵ and tetramethylstannane (25.8 mg, 0.144 mmol) in chloroform (15 mL) was treated with a solution of iodine (37.2 mg, 0.147 mmol) in chloroform (10 mL), and the mixture was stirred at 25 "C in the dark for 13 h. Volatiles were then removed by evaporation under reduced pressure, and

the residue was extracted with acetonitrile. The insoluble fraction, a pale yellow solid, was further washed with ether, dried, and identified by its characteristic solubility and chromatographic properties as elemental sulfur (3.1 mg, 0.097 mmol, 66%). Concentration of the acetonitrile solution of the soluble fraction produced colorless crystals of iodotriphenylstannane (101 mg, 0.212 mmol, 72.6%): mp 119-120 °C (lit.²⁸ mp 121 °C).

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1-H, 87922-35-8; 1-CH3, 111769-98-3; 2-H, Registry **No.** 88212-59-3; 2-CH3, 111769-99-4; 3,96504-44-8; **4,** 111770-00-4; **5,** 111770-01-5; **6,** 111770-02-6; **7,** 111770-03-7; 12, 639-58-7; bis- (triphenyltin) sulfide, 77-80-5; tetramethylstannane, 594-27-4; iodotriphenylstannane, 894-09-7; bromotriphenylstannane, 962- 89-0; 4-nitrobenzoyl chloride, 122-04-3; 4-nitroacetophenone, 100-19-6; **2-(hydroxyme~hyl)-2-methyl-l,3-propanediol,** 77-85-0; triphenylstannane, 892-20-6.

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Synthesis and Reactlons of (Trimethylgermy1)acetates

Sumie Inoue, Yoshiro Sato,* and Tomoko Suzuki

Faculty of Pharmaceutical Sciences, Nagoya City University, Tanabe-dori, Mizuho-ku, Nagoya 467, Japan

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Base-assisted trimethylgermylation of ethyl acetate gave high yields of ethyl **(trimethylgermy1)acetate (2), bis(trimethylgermy1)acetate** (5), and **tris(trimethylgermy1)acetate (6).** The Peterson reaction of (trimethylgermyl)(trimethylsilyl)acetate (4), prepared from ethyl (trimethylsilyl)acetate (1), with aldehydes selectively yielded α -(trimethylgermyl)- α , β -unsaturated alkenoates (15). The addition of fluoride anion to **4** successfully induced selective desilylation. Some chemical properties of **6** were also examined.

Although the synthetic utility of α -trimethylsilyl esters has been amply demonstrated,^{1,2} there is no direct method to prepare them from the corresponding esters. **All** reports to date on the direct trimethylsilylation of esters have shown that the predominant product is that due to 0-silylation and not C-silylation.^{1,3} One exception to this

general observation is that of tert-butyl acetate, which can be attributed to a larger steric effect in the 0-silylated material than in the C-silylated isomer.⁴ However, the trimethylsilylation of tert-butyl (trimethylsily1)acetate gave a mixture of both the C-silylated and O-silylated products.⁵ Larson reported that methyldiphenylsilylation of esters favors the formation of the C-silylation product.6 Previously we reported that the trimethylgermyl group tended to be introduced on the α -carbon rather than the carbonyl oxygen of ketones, whereas the trimethylsilyl group was usually introduced on the $oxygen.⁷$ This paper describes

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Table I. Reaction of the Germylacetate Anions with Aldehydes

entry	acetate			aldehyde		product		
		м			reactn conditns ^a		% yield ^b	ratio ^c $(E:Z)$
		Si	н	Ph		14a	84 ^d	$3 - 8:1$
		Ge	н	Ph		14a	65	>95 _i < 5
		Si		n -Oct		14b	81 ^d	1:1
		Ge		n -Oct		14b	57	95:5
		Si	Me ₃ Ge	Ph		15a	49	>95<5
		Ge	$Me_{3}Ge$	Ph		15a	52	>95<5
		Si	Me ₃ Ge	n -Oct		15b	47	53:47
		Ge	Me ₃ Ge	n -Oct		15b	43	72:28
		Si	Me ₃ Ge	n -Bu(Et)CH		15c	43	69:31
10		Ge	Me ₃ Ge	$n\text{-}Bu(Et)CH$		15с	62	75:25

^a Reaction conditions: A, -78 °C/1 h, -25 °C/1 h, and then 25 °C/1 h; B, -78 °C/1 h and then 0 °C/1 h. ^b Isolated yield. ^c Determined from the integrated values of GLC analyses. dReference 10.

^a Oven temperature of Kugelrohr distillation apparatus.

the synthesis of mono-, bis-, and tris(trimethylgermyl)acetates from ethyl acetate and the reactions of the related compounds.

Results and Discussion

Ethyl (trimethylgermyl)acetate (2) was conveniently obtained in high yield by trimethylgermylation of ethyl acetate with the aid of lithium diisopropylamide (LDA), even though it had been previously prepared by the Reformatskii reaction of ethyl bromoacetate with chlorotrimethylgermane.⁸ Trimethylgermylation of lithium salts of ethyl (trimethylsilyl) acetate (1) and 2 gave high yields of ethyl (trimethylgermyl)(trimethylsilyl)acetate (4) and ethyl bis(trimethylgermyl)acetate (5), respectively, whereas trimethylsilvlation of 1 afforded only O-ethyl O-trimethylsilyl ketene acetal (3) (Scheme I). The use of excess amounts of LDA and chlorotrimethylgermane with 2 gave ethyl tris(trimethylgermyl)acetate (6) in a high yield. We previously showed that the trimethylgermylation of (trimethylgermyl) acetonitrile in the presence of 2 equiv of a base formed stable crystals of tris(trimethylgermyl)acetonitrile.⁹ Compound 6 is also in the form of stable crystals and can be kept without detectable decomposition at room temperature.

Some reactivities of 6 were examined (Scheme II). The lithium aluminum hydride reduction gave a 78% yield of 2,2,2-tris(trimethylgermyl)ethanol (7) in ether, and the treatment with *n*-butyllithium at room temperature afforded an 86% yield of 1.1.1-tris(trimethylgermyl)-2-hexanone (9). Thus, the tris(trimethylgermyl)methyl group was not affected by the reduction or alkyllithium. Both tris(trimethylgermyl)-substituted compounds 7 and 9 were also stable in a moist atmosphere, and the former could be easily purified by distillation. However, the distillate of the latter at 100 °C (3 mmHg) was contaminated with 1,1-bis(trimethylgermyl)-2-hexanone (11), which was presumed to be the hydrolyzed form of the thermal rearrangement product, 1,1-bis(trimethylgermyl)-2-(tri-

methylgermoxy)-1-hexene (10). The isomerization from 9 to 10 was accelerated in the presence of silica gel in an ether solution.

When 7 was treated with boron trifluoride etherate, an 83% yield of 1.1-bis(trimethylgermyl)ethylene (8) was obtained with hydroxytrimethylgermane being generated as a byproduct. Treatment of 6 with phenyllithium gave only a small amount of α, α -bis(trimethylgermyl)acetophenone (12), although there was no reaction with either n -butylmagnesium bromide or phenylmagnesium bromide.

It is well-known that lithio(trimethylsilyl)acetates react with aldehydes to give α,β -unsaturated acid esters (14)^{10,11}

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and that α -(trimethylsilyl)- α , β -unsaturated esters are obtained from **lithio[bis(trimethylsilyl)]acetate** (Peterson reaction).^{2f,5} A similar reaction using the germyl analogue **2, 4,** or **5** with some aldehydes was carried out (Scheme 111), and the results are summarized in Tables I and 11. The results of entries **2** and **4** are comparable with previously reported data (entries **1, 3).**

The yields of ethyl α , β -unsaturated alkenoates 14a,b from **2** were somewhat lower than those from **1** although the stereoselectivities were higher (entires **1-4** in Table I). The reaction of the aldehydes with the anion of **4** or **5** gave the same product, ethyl α -(trimethylgermyl)- α , β -unsaturated alkenoates **(15a-4,** respectively (entries **5-10).** We previously revealed that the reaction of the (trimethylgermyl) (trimethylsily1)acetonitrile anion with aldehydes formed α -(trimethylgermyl)- α , β -unsaturated alkenenitriles as the sole reaction product. $\frac{9}{7}$ Thus, the trimethylsiloxy group is selectively eliminated from the Peterson-type reaction intermediate (e.g., 13 , $M = Si$, $Y = Me₃Ge$, in Scheme 111) in the reaction of trimethylgermyl, trimethylsilyl-substituted carbanion with carbonyl compounds.

The yields of **15a-c** were unexpectedly low with **34-56% of** used **4** or **5** remaining at the end of the reaction (entries **5-10).** When the temperature of the reaction mixture rose to room temperature after the addition of the aldehyde, **(trimethylgermyl)(trimethylsilyl)ketene (16)** or bis(trimethylgermy1)ketene **(17)** was notably produced. **A** high yield of either **16** or **17** was isolated from the solution of **4'** or **5'** after being at 0 "C for **1** h. There may be an equilibrium between the acetate anion **(4'** or **5')** and the adduct **(13)** which makes it difficult to complete the reaction. The analogous formation **of** bis(trimethylsily1) ketene from tert-butyl **lithiobis(trimethylsily1)acetate** has been reported by Rathke.¹²

The fluoride ion induced reaction of organosilicon compounds is an important synthetic method. Degermylation also occurs quickly by the addition of fluoride ion, e.g. **a-(trimethylgermy1)acetophenone** was degermylated at **-78 OC.'** However, selective desilylation of silyl, germyl-substituted compounds may be possible because there is a considerable difference between the bond dissociation

energies of $Si-F$ $(143 \text{ kcal/mol}^{-1})$ and $Ge-F$ (113 kcal) $mol⁻¹$ bonds.

The addition of tetra-n-butylammonium fluoride (TBAF) followed by water to **4** gave a mixture of **2** and It has been reported that the trimethylgermyl group of **(trimethylgermy1)acetonitrile** rearranges quickly to give **bis(trimethylgermy1)acetonitrile** in the presence of a base.13 The formation of **5** may also result from the intermolecular anionic rearrangement of the trimethylgermyl group to the (trimethylgermy1)acetate anion produced by the desilylation of **4.** In the presence of non-enolizable aldehydes such as benzaldehyde and trimethylacetaldehyde, the corresponding ethyl α , β -unsaturated carboxylates **(14a,d)** were obtained. (Trimethylsily1)acetic acid ester with the non-enolizable aldehydes gave the Reformatskii-type products under a similar reaction condition.¹⁴ The formation of 14 may proceed via the intermediate **18** with the elimination of **trimethyl(trimethylsi1oxy)germane.** Small amounts of ethyl **2-(trimethylgermy1)propionate (19)** were isolated from the desilylation of **4** in the presence of methyl iodide.

Experimental Section

All reactions were carried out under a nitrogen atmosphere. Diethyl ether and tetrahydrofuran were dried by distillation from sodium benzophenone ketyl prior to use. 'H NMR spectra were recorded on a JEOL JNM-PMX 60 spectrometer using Me₄Si as internal standard. IR spectra were recorded on a JASCO IRA-2 spectrometer. All melting points and boiling points are uncorrected. n-Butyllithium, 15% in hexane, was purchased from Merck & Co., Inc, Schuchardt.

Ethyl (Trimethylgermy1)acetate (2). A. A solution of n-BuLi in hexane (21 mL, 33 mmol) was added to a solution of diisopropylamine (3.33 g, 33 mmol) in ether (15 mL) at 0 $^{\circ}$ C with continuous stirring for 0.5 h. The resulting LDA solution was cooled at -78 °C, and a solution of ethyl acetate $(2.64 g, 30 mmol)$ in ether (15 mL) was added dropwise. After 1 h, a solution of chlorotrimethylgermane (5.05 g, 33 mmol) in ether (15 mL) was added with additional stirring for 1 h. Saturated aqueous NH4Cl was added to the mixture. The organic layer was separated, and the aqueous layer was extracted with ether. The combined extracts were dried over anhydrous MgSO,, concentrated, and distilled under reduced pressure to give **2** (4.80 g, 78%): bp 76-78 "C (30 "Hg) [lit? bp 83-84 "C (37 mmHg)]; IR **(film)** 1720 (CO), 830, 610, 580 cm⁻¹ (Me₃Ge); ¹H NMR (CDCl₃) δ 0.26 (9 H, s, $Me₃Ge$, 1.26 (3 H, t, $J = 8$ Hz, CH₃), 1.96 (2 H, s, CH₂CO), 4.20 $(2 \text{ H}, \text{ q}, J = 8 \text{ Hz}, \text{ CH}_2).$

B. A solution of ethyl bromoacetate (43.25 g, 259 mmol) in benzene⁻(25 mL) was added to a mixture of chlorotrimethylgermane (30.83 g, 201 mmol), activated granular zinc (26.28 g, 402 mmol), benzene (50 mL), and THF (80 mL). After 20 h of stirring at 30 °C, saturated aqueous $NH₄Cl$ was added to the

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mixture. The organic layer was separated, and the aqueous layer was extracted with benzene. The combined extracts were dried $(MgSO₄)$, concentrated, and distilled to give 2 (29.41 g, 71%) (lit.⁸

47%).
Ethyl (Trimethylgermyl)(trimethylsilyl)acetate (4). To **a** solution of LDA, prepared from diisopropylamine $(3.09 \text{ g}, 30.5 \text{ m})$ mmol) and n-BuLi **(20** mL, **31.2** mmol) in ether **(15** mL), was added dropwise a solution of ethyl (trimethylsily1)acetate **(4.83** g, **30.2** mmol) in ether **(10** mL) at **-78** "C. After **1** h of stirring, a solution of chlorotrimethylgermane (5.82 g, 37.9 mmol) in ether **(5** mL) was added with additional stirring for **3.5** h at the same temperature. After the addition of a saturated aqueous NH_4Cl solution, the organic layer was separated and the aqueous layer was extracted with benzene. The combined extracts were washed with water and saturated aqueous NaC1, dried over anhydrous $MgSO₄$, and concentrated. Distillation of the residual oil gave 1250, 820, 600 cm^{-1} (Me₃Si and Me₃Ge); ¹H NMR (CDCl₃) δ 0.13 **(9** H, **s,** Me3Si), **0.29 (9** H, **s,** Me3Ge), **1.23 (3** H, t, *J* = 8 Hz, CH3), **1.59 (1** H, **s,** CH), **4.13 (2** H, q, *J* = **8** Hz, CH2). Anal. Calcd for CloHz4Ge02Si: C, **43.35;** H, **8.75.** Found: C, **43.42;** H, **8.73. 4** (6.59 g, 79%): bp 101-105 °C (15 mmHg); IR (film) 1690 (CO),

Ethyl Bis(trimethylgermy1)acetate (5). In a manner similar to that described for **4,** diisopropylamine **(3.09** g, **30.6** mmol), n-BuLi **(19.5** mL, **30.5** mmol), **2 (6.14** g, **30** mmol), and chlorotrimethylgermane **(5.05** g, **33** mmol) were allowed to react and worked up to give **5 (8.92** g, **93%):** bp **121-122** "C **(28** mmHg); IR (film) **1695 (CO)**, **820**, **595**, **565** cm⁻¹ (Me₃Ge); ¹H *NMR* (CDCl₃) δ 0.25 (18 H, s, Me₃Ge), 1.22 (3 H, t, $J = 6.5$ Hz, CH₃), 1.60 (1) H, **s,** CH), **4.05 (2** H, q, *J* = **6.5** Hz, CH2). Anal. Calcd for $C_{10}H_{24}Ge_2O_2$: C, 37.36; **H**, 7.52. Found: C, 37.30; **H**, 7.47.

Ethyl Tris(trimethy1germyl)acetate (6). A solution of **2 (6.16** g, **30** mmol) in THF **(8** mL) was added dropwise at **-78** "C to a LDA solution, prepared from diisopropylamine **(12.04** g, **119** mmol) and n-BuLi **(77** mL, **120** mmol) in THF **(40** mL). After **1** h of stirring, a solution of chlorotrimethylgermane **(18.48** g, **120** mmol) in THF **(18** mL) was added dropwise with additional stirring for **4** h at the same temperature. Saturated aqueous NH4Cl was added to the mixture, and it was extracted with benzene. The organic layer **was** washed with water and saturated aqueous NaCl, dried over MgSO,, and concentrated. Distillation of the residue gave **6 (10.78** g, **82%):** bp **14&150** "C **(20** mmHg); mp **160-171** "C; IR (Nujol) **1675** (CO), **820,595,565** cm-' (Me3Ge); ¹H NMR (CDCl₃) δ 0.30 (27 H, s, Me₃Ge), 1.23 (3 H, t, $J = 7$ Hz, CH₃), 4.10 (2 H, q, $J = 7$ Hz, CH₂). Anal. Calcd for C₁₃H₃₂Ge₃O₂: C, **35.64;** H, **7.36.** Found: C, **35.75;** H, **7.25.**

2,2,2-Tris(trimethylgermyl)ethanol(7). A mixture of **6 (2.21** g, **5** mmol) and LiAlH, **(387** mg, 10 mmol) in ether **(30** mL) was heated at reflux for **8** h. The reaction mixture was cooled in an ice bath and worked up by the addition of ethyl acetate (0.5 mL), **10%** NaOH (0.5 mL), and water **(1.5** mL). The mixture was filtered, and the filter cake was washed with ether. The filtrate and washings were combined and washed with saturated aqueous NaCI, dried over MgSO,, and concentrated. Distillation of the residue gave **7 (1.55** g, **78%):** bp **120-130 "C (10** mmHg, oven temperature of a Kugelrohr distillation apparatus); mp **181-185** "C; IR (Nujol) **3450** (OH), **810,580,560** cm-' (Me3Ge); 'H NMR (CDCI,) 6 **0.28 (27** H, **s,** Me3Ge), **2.68 (2** H, **s,** CH2), **4.06 (1** H, **s,** OH). Anal. Calcd for CllH30Ge0: C, **33.35;** H, **7.63.** Found: C, **33.06;** H, **7.53.**

1,l-Bis(trimethylgermy1)ethylene (8). A solution of boron trifluoride etherate **(188** mg, **1.3 "01)** in dichloromethane **(3** mL) was added dropwise to a solution of **7 (396** mg, **1.0** mmol) in dichloromethane at 0 $^{\circ}$ C. After 1 h of stirring, saturated aqueous NH4CI was added and the mixture was extracted with ether. The ethereal extract was washed with water, saturated aqueous NaCl, dried over MgSO₄, and concentrated under reduced pressure to give **8 (216** mg, **83%):** bp **70** "C **(77** mmHg, Kugelrohr); IR (film) **1230,950,815,590,560** cm-'; 'H NMR (CDCl,) **d 0.22 (18** H, **s,** $Me₃Ge$, 6.13 (2 H, s, CH₂). Anal. Calcd for $C_8H_{20}Ge_2$: C, 36.76; H, **7.71.** Found: C, **36.49;** H, **7.56.**

l,l,l-Tris(trimethylgermyl)-2-hexanone (9). To a solution of **6 (448** mg, **1.1** mmol) in ether **(5** mL) was added dropwise n-BuLi **(2.1** mL, **3.3** mmol) at room temperature with continuous stirring for **1** h. The mixture was cooled in an ice bath, and saturated aqueous NH₄Cl was added. The ethereal extract of the reaction mixture was washed with water and saturated aqueous

NaCl and dried over anhydrous $MgSO_4$. Evaporation of the solvent gave 470 mg of the crude oil that was chromatographed on a silica gel column (hexane/ethyl acetate, 80:1) to give 9 (388 mg, **86%):** IR (film) **1640** (CO), **820,660,570** cm-' (Me,Ge); 'H NMR (CDCl₃) δ 0.36 (27 H, s, Me₃Ge), 0.86 (3 H, t, $J = 8$ Hz, CH₃), **1.12-1.71 (4 H, m, CH₂CH₂), 2.33 (2 H, t,** $J = 8$ **Hz, CH₂CO). Anal.** Calcd for C15H36Ge30: C, **40.02;** H, **8.06.** Found: C, **40.20:** H, **8.05.**

1,1-Bis(trimethylgermyl)-2-hexanone (11). Distillation of **9 (260** mg, **0.78** mmol) in a Kugelrohr distillation apparatus at **100** "C under **3** mmHg gave **235** mg of a mixture of **9** and **11** (GLC ratio, 83:17). The ratio of 9 to 11 was changed to 17:83 after being stirred with silica gel $(1 g)$ in ether $(5 mL)$ for 1 day at room temperature. The silica gel was filtered off, and the filtrate was distilled to give **186** mg of **11:** bp **115** "C (9 mmHg, Kugelrohr); *JR* (film) **1665** (CO), **820,600,570** cm-' (Me3Ge); 'H **NMR** (CDCl,) 6 **0.25 (18** H, **s,** Me3Ge), **0.90 (3** H, t, *J* = **6** Hz, CH,), **1.14-1.66 (4** H, m, CH,CH,), **2.18** (1 H, **s,** CHCO), **2.31 (2** H, t, *J* = **7.4** Hz, CH₂CO). Anal. Calcd for $C_{12}H_{28}Ge_2O$: C, 43.21; H, 8.46. Found: C, **43.07;** H, **8.34.**

a,a-Bis(trimethylgermy1)acetophenone (12). A solution of phenyllithium in cyclohexane/diethyl ether **(7:3) (2.0** M, **3** mL, **6 mmol) was added dropwise to a solution of 6 (438 mg, 1.0 mmol)** in ether **(5** mL) at room temperature with continuous stirring for **1** h. The reaction mixture was worked up in a manner similar to that described for **9.** Chromatography of the crude oil **(555** mg) on a silica gel column (hexane/ether, **401)** gave **6 (310** mg, **71%)** and **12 (36** mg, **10%):** bp **130** "C **(3** mmHg, Kugelrohr); *JR* **(film) 1635** (CO), **820,600,565** cm-' (Me,Ge); 'H *NMR* (CDCI,) **0.27 (18** H, **s,** Me3Ge), **3.10** (1 H, **s,** CH), **7.20-7.61** and **7.78-8.05** (each 3 H and 2 H, m, ArH). Anal. Calcd for $C_{14}H_{24}Ge_2O$: C, **48.95; H, 7.04. Found: C, 48.95; H, 7.06.**

Ethyl 2-Alkenoate (14) and Ethyl 2-(Trimethylgermy1)- 2-alkenoate (15) (General Procedure). A solution of 2.4 mmol of 2, 4, or 5 in THF (3 mL) was added dropwise at -78 °C to a solution of LDA, prepared from n-BuLi $(1.5 \text{ mL}, 2.4 \text{ mmol})$ and diisopropylamine **(243** mg, **2.4** mmol) in THF **(4** mL). After **1** h of stirring, a solution of an aldehyde **(2** mmol) in THF **(3** mL) was added dropwise, and stirring was continued at **-78** "C for 1 h, **-25** "C for **1** h, and **25** "C for **1** h (entries **1-4** in Table **I)** or **-78** "C for **1** h and **0** "C for **1** h (entries **5-10).** Saturated aqueous NH4Cl was added to the reaction mixture. The ethereal extract of the mixture was washed with water and saturated aqueous NaCl, dried over anhydrous MgS04, and concentrated under reduced pressure. The residue was distilled (entries **5, 6)** or chromatographed on a silica gel column (hexane/ether) (entries **2,4, 7-10)** to give **14-E** and **14-2** or **15-E** and **15-2.**

The *E,Z* assignment of two geometrical isomers of **15** was performed, according to the reported method, by the comparison of the chemical shifts of olefinic protons and trimethylgermyl groups in their 'H NMR spectra and the retention times of $GLC^{9,15}$ The characteristic data of $15-E$, Z are summarized in Table 11.

(Trimethylgermyl)(trimethylsilyl)ketene (16). A solution of **4 (554** mg, **2** mmol) in ether **(3** mL) was added dropwise at 0 "C to a solution of LDA, prepared from n-BuLi **(1.3** mL, **2** mmol) and diisopropylamine **(202** mg, **2** mmol) in ether **(4** mL). After 1 h of stirring at 0 °C, saturated aqueous NH₄Cl was added. The ethereal extract of the mixture was washed with water and saturated aqueous NaCl and dried over MgSO,. Evaporation of the solvent gave crude **16 (391** mg, **85%).** Kugelrohr distillation of the crude product gave **295** mg **(64%) of** the pure product: bp **75 "C (21** mmHg) [lit.16 bp **60** "C **(12** mmHg)]; 'H **NMR** (CDCl,) δ 0.18 (9 H, s, \overline{Me}_3 Si), 0.38 (9 H, s, \overline{Me}_3 Ge).

Bis(trimethylgermy1)ketene (17). In a manner similar to that described for **16,5 (772** mg, **2.4** mmol), n-BuLi **(1.5** mL, **2.4** mmol), and diisopropylamine **(243** mg, **2.4** mmol) were treated to give crude **17 (609** mg, **92%).** Distillation of the crude product gave **505** mg **(76%)** of pure product: bp **80** "C **(12** mmHg) [lit.17

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bp **80-81** OC **(20** ~nmHg)]; 'H NMR (CDC13) *6* **0.39 (8)** (lit." 6 **0.29). Desilylation of 4 with Tetra-n-butylammonium Fluoride (TBAF).** To a solution of 4 (554 mg, 2.0 mmol) in THF (10 mL) was added dropwise a solution of TBAF $(0.2 \text{ M} \text{ in } THF, \text{dried})$ over molecular sieves **4A,** 10 mL, **2.0** mmol) at **-78** "C. After **5** s, the reaction mixture was mixed with chilled ether **(50** mL). The mixture was washed with *5%* HCl, water, and saturated aqueous NaC1. Concentration of the organic layer on a rotary evaporator, after drying over anhydrous $\overline{MgSO_4}$, gave 342 mg of a mixture of 2 (74%) and 5 (6%). Identification of the products was accomplished by the comparison with authentic samples on GLC and NMR. Yield of each compound was calculated from the mole ratio based on the proton ratios in 'H NMR of the mixture.

Ethyl Cinnamate (14a). To a mixture of benzaldehyde **(212** mg, **2.0** mmol) and TBAF **(0.2** M in THF, **5** mL, **1.0** mmol) was added dropwise a solution of **4 (277** *mg,* **1.0** mmol) in THF **(3** mL) at **0** "C. After 1 h of stirring, the reaction mixture was mixed with ether **(50** mL). The mixture was washed with saturated aqueous NaC1, dried over MgS04, concentrated, and distilled to give **102** mg (58%) of **14a.**

Ethyl 4,4-Dimethyl-2-pentenoate (14d).'* To a solution of **4 (554** mg, **2.0** mmol) and trimethylacetaldehyde **(345** mg, 4.0 mmol) in THF (5 mL) was added dropwise TBAF (0.2 M in THF) , **2** mL, **0.4** mmol) at **-78** "C with continuous stirring for **1** h. After the addition of ether **(50** mL), the mixture was washed with

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saturated aqueous NaCl, dried over MgSO_4 , and concentrated under reduced pressure. Silica gel column chromatography (hexane/ether) of the residue gave 14d (188 mg, 60%).

Ethyl 2-(Trimethylgermy1)propionate (19). To a solution of **4 (554** mg, **2.0** mmol) and methyl iodide **(568** mg, **4.0** mmol) in THF **(5** mL) was added dropwise TBAF **(0.2** M in THF, **5** mL, 1.0 mmol) at -78 °C with continuous stirring for 1 h. After the addition of ether **(50** mL), precipitated tetra-n-butylammonium iodide was filtered. The filtrate was washed with saturated aqueous NaCl, dried over MgS04, and concentrated under reduced pressure. The residue was chromatographed on a silica gel column (hexanelether, **401)** to give **2** (181 mg, **45%)** and **19 (55** mg, **13%):** bp **65** "C **(13** mmHg, Kugelrohr); IR (film) **1715** (CO), **830,610,** 575 cm^{-1} (Me₃Ge); ¹H NMR (CDCl₃) δ 0.22 (9 H, s, Me₃Ge), 1.20 $(1 H, q, J = 6.4 Hz, CH), 4.10 (2 H, q, J = 7.2 Hz, CH₂).$ Anal. Calcd for C₈H₁₈GeO₂: C, 43.91; H, 8.29. Found: C, 43.70; H, 8.18. $(3 H, d, J = 6.4 Hz, CH₃CH), 1.23 (3 H, t, J = 7.2 Hz, CH₃), 2.16$

Registry No. 1, 4071-88-9; 2, 32583-34-9; 4, 111999-58-7; 5, 111999-62-3; 11, 27394-64-5; 12,38860-13-8; (E)-14a, 4192-77-2; (Z)-14a, 4610-69-9; (E)-14b, 51577-37-8; (Z)-14b, 51577-38-9; 14d, **87995-20-8; (E)-15a, 111999-64-5; (Z)-15a, 111999-65-6; (E)-15b, 111999-69-0; 16,38860-05-8; 17,32329-69-4; 19,111999-63-4;** ethyl acetate, **141-78-6;** ethyl bromoacetate, **105-36-2;** benzaldehyde, **100-52-7;** trimethylacetaldehyde, **630-19-3;** nonanal, **124-19-6;** 2-ethylhexanal, **123-05-7. 111999-59-8; 6, 111999-60-1; 7, 111999-61-2; 8, 7428-06-0; 9, 111999-66-7; (Z)-15b, 111999-67-8; (E)-15~, 111999-68-9; (Z)-15~,**

Lanthanoid-Olefin Complex Formation: A Matrix- Isolation Ultraviolet-Visible and Infrared Study of the Direct Synthesis of (C₂H₄)_nEu Using Europium Atoms[†]

Mark P. Andrews^{*} and Andrea L. Wayda*

AT&T Bell Laboratories, 600 Mountain Avenue, Murray Hill, New Jersey 07974

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 $Eu(C_2H_4)$ _n complexes have been synthesized at 12 K by codepositing Eu atoms with ethylene, neat or doped in **Ar** or Xe. UV-visible absorption spectroscopy of the highest stoichiometry product revealed a broad, intense visible transition centered at **537** nm. The complex is unstable above **50** K, consistent with the absence of any evidence for reaction in *77* K metal vapor reactions. Optical spectroscopy of the products formed in dilute matrices indicated the formation of a lower stoichiometry $E\hat{u}$ -C₂H₄ complex. This was confirmed by infrared studies of absorptions in the coupled ν (C=C)/ δ (CH₂) region. Under high dilution conditions, a matrix split $1198/1182 \text{ cm}^{-1}$ doublet is assigned to a monoethylene europium π complex. Matrix-annealing experiments combined with Eu/C $_2\rm{H}_4$ reactions in C $_2\rm{H}_4$ /rare-gas matrices of increasing C_2H_4 concentration trace the conversion of $Eu(C_2H_4)$ to the highest stoichiometry species absorbing at 1233 cm⁻¹. In general, the Eu(C₂H₄)_n IR spectra are similar to those of matrix-isolated group VIII $(8-10)$ and IB (11) ethylene π complexes. The optical spectra of the odd-electron lanthanoid-olefin molecules, however, most closely resemble those of the coinage metals. The latter show intense visible metal-to-ligand charge-transfer transitions. Similarly, the green absorption of the europium olefin complexes may be associated with excitations of the Eu f electrons into the olefin. The comparatively low thermal stability of the Eu and group IB **(11)** olefin complexes is rationalized with the Dewar-Chatt-Duncanson bonding description. Molecular orbital calculations indicate that a model $C_{2\nu} \, {\rm Eu(C_2H_4)}$ species is only weakly bonded by donation from a europium f orbital into the **a*** system of ethylene, with little "forward" donation from the olefin σ or π orbitals.

Although common in organotransition-metal chemistry, isolable, formally zerovalent complexes are extremely rare

Introduction in organolanthanoid chemistry.¹ This dichotomy in chemical stability is undoubtedly linked to several factors, chief among them being **(1)** the electropositive character of the lanthanoid metals which favors di- and trivalent oxidation states of these metals and **(2)** the relative inability of the corelike lanthanoid metal valence electrons

t In this paper the periodic group notation in parentheses is in accord with recent actions by IUPAC and ACS nomenclature cornmittees. A and B notation is eliminated because of wide confusion. Groups IA and IIA become groups 1 and **2.** The d-transition elements comprise groups through l2> and the Pblock comprise groups 13-18. (Note that the former Roman number designation is preserved in the last digit of the new numbering: e.g., III \longrightarrow N ; d. \longrightarrow 3 and 13.) 1344.

⁽¹⁾ The first well-characterized examples **of** formally zerovalent, thermally stable, isolable species of these metals, the lanthanoid diazadiene complexes, have only recently been reported. **See** Cloke, F. G. N.; de Lema, H. C.; Smeh, **A. A.** *J. Chem. SOC., Chem. Commun.* **1986,**