81. Stereospecific, R,AlCl-Promoted Intramolecular €he Reaction of a 1,6-Dienoate: Evidence for a Concerted Mechanism

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Summary

Treatment of the 83%-trans-¹³CH₃-labelled 1,6-dienoate 7 with Et₂AlCl at -78° provided in high yield the ene product 9 containing 83% ¹³C localized in the olefinic $C(8)$ -methylene group. Accordingly, H-transfer occurs exclusively from the *trans*methyl group of **7**, consistent with a concerted ene process $7 \rightarrow 9$ thereby ruling out an intermediate cation **8** *(Scheme 4).*

Introduction. – Recently we have reported direct, efficient, regio-, diastereo- and enantioselective syntheses of the neurophysiologically interesting algae constituents *a*allokainic acid **(1)** [1][2] and a-kainic acid **(2)** [3][4].

Each of these syntheses features an intramolecular ene reaction¹) $\mathbf{I} \rightarrow \mathbf{II}$ following one of the two stereochemically different strategies. Thus, either the configurationally pure center C(2) of **I** dictates the chirality at C(3) and C(4) in **I1** or, alternatively, the chiral ester substituent \mathbb{R}^2 in I first induces centers $C(3)$ and $C(4)$ which then in turn control the configuration at C(2) in **1.** A prerequisite for the latter alternative was the ability to control the relative configuration of centers $C(3)/C(4)$ in the process **l** \rightarrow **H** which was achieved by modifying the masked carboxyl equivajent X and the enophile

¹) For a review see [5].

geometry in **I.** On thermal cyclization of the **bis(ethoxycarbony1)-substituted** *(Z)* enoate **3,** we [1][6] and others [7] observed an unusual diastereoselectivity in favor of the *trans*-product **5** (ratio $4/5 = 1:3$ irrespective of the reaction temperature).

An even more pronounced stereoselectivity was achieved when **3** was cyclized in the presence of Et,AICI **[612).** Thus, treatment of **3** with Et,AICI **(3** mol-equiv.) in dry CH₂Cl₂ at -78° for 8 h or at -35° for 30 min yielded exclusively the *trans*-product 5 in 86% yield. No trace of the cis-isomer **4** was found in the reaction mixture.

Apart from the relevance of this result for the synthesis of racemic and enantiomerically pure *(+)-a* -allokainic acid [1][2], we were interested in the mechanistic origin of this spectacular Lewis-acid effect.

Results and Discussion. - Coordination between Lewis Acid and 1,6-Diene. Given the number of basic functionalities in **3** it was not surprising that the cyclization rate at -78° depended on the excess of Et₂A₁C₁ present (requiring for completion 8 h using 3 mol-equiv. and only *5* min using 20 mol-equiv. of the Lewis acid). To study the relevant coordination sites in **3** the 'H-NMR spectra of enoate **6** were monitored in relation to the molarity of Et₂AlCl (*Table*). On increasing the latter, the signals of the olefinic protons $H-C_a$ and $H-C_b$ are shifted down-field; this indicates that the extent of enoate/Et,AlCl coordination in **3** parallels the rate increase for the reaction $3 \rightarrow 5^3$).

Stereochemical Working Hypothesis. The question arose whether the Et,AlCI-promoted cyclization $3\rightarrow 5$ is in fact a concerted ene reaction or rather proceeds *via* a cationic intermediate resulting from electrophilic attack of the coordinated enoate at

which indicates the importance of CH₃-substitution in the 'ene' unit.
\n**i**
$$
R^1 = R^2 = H
$$
, $R^3 = Me$, $X = N - COCF_3$
\n**ii** $R^1 = COOEt$, $R^2 = H$, $R^3 = Me$, $X = CH_2$
\n**iii** $R^1 = H$, $R^2 = COOEt$, $R^3 = H$, $X = CH_2$
\n**iv** $R^1 = H$, $R^2 = COOEt$, $R^3 = Me$, $X = CH_2$
\n**v** $R^1 = R^2 = COOEt$, $R^3 = Me$, $X = CH_2$

^{2,} For RA ICI₂- and R_2 AICI-catalyzed bimolecular ene reactions see [8]; for a review see [9].

^{3,} The significance of enoate coordination and 'ene' substitution on the rate of the Et,AICl-inducd cyclization **3+5** is also illustrated by the following observations [lo]. All 1,5-dienes **i** to **v** underwent thermal ene reactions to give 5-membered ring systems. However, *only* the enoates **iv** and **v** cyclized at room temperature in the presence of an excess of Et₂AICI in dry CH₂Cl₂ (50 to 100 h). The relative rates of the thermal reactions (iv: 170°, 20 h/ v: 220°, 16 h/ 3: 70°, 80 h) reflect those of the Et₂AlCl-induced cyclizations; under all reaction conditions **3** cyclized most rapidly, probably owing to entropic reasons. Furthermore, in contrast to diene **iv,** no Et,AlCI-promoted cyclization of diene **iii** was observed under identical conditions

| Mol-equiv. Et ₂ AlCl | CH_3-C^2 | CH_3-N^2 | $CH2-O3$ | $H-C_{\star}$ | $H-C_R$ |
|-----------------------------------|------------|------------|----------|---------------|---------|
| $\overline{}$ | 1.24 | 3.33 | 4.17 | 6.12 | 6.51 |
| $\overline{2}$ | 1.28 | 3.33 | 4.24 | 6.20 | 6.55 |
| 3 | 1.29 | 3.33 | 4.28 | 6.28 | 6.61 |
| 6 | 1.32 | 3.33 | 4.36 | 6.44 | 6.69 |
| a) Center of the signal group. | | | | | |

Table. ^{*'H-NMR Signal Shifts* (in ppm, CDCl₃) *of* (Z)- $H_3C-N(COCF_3)-C(COOEt_2)_2-gCH=G(H-COOEt$ (6)} *in Relation to Et₂AlCl-Molarity*

the isolated olefinic bond4). This mechanistic problem may be particularly relevant in terms of the observed stereoselection which is kinetically controlled in both the thermal and Lewis-acid-induced cyclizations of **3').** Assuming the operation of a concerted ene process, we thus attempted to rationalize the observed stereochemistry by examination of the transition states *(Scheme 3).* This analysis accounts for H-transfer from both

the allylic trans-methyl **(A** and **B)** and cis-methyl group **(C** and **D).** Transition states **B** and **C** are readily excluded: **B** on the basis of steric repulsion (malonate/olefinic methyl) and **C** due to angle strain'). Orientation **A** seems to be favored over **D** which suffers from 1,3-diaxial perturbation; this steric repulsion should increase on coordi-

C E = COOEt **D**

^{4,} So far, the dichotomy between concerted and cationic mechanisms of intermolecular Lewis-acid-mediated ene reactions has been studied only by means of H/D-isotope effects and product distributions [9].

 5 **A** 1 :I mixture of **4** and *5* remained virtually unchanged either on heating at 180" for **10** min or on treatment with Et₂AlCl (30 mol-equiv.) in CH₂Cl₂ at 25° for 10 min.

 6 This angle strain argument is in accord with the stereochemistry of numerous other intramolecular ene reactions [5] [Ill [12].

nation of the ester and amide units with the *Lewis* acid. Consequently, trans-product *5* is formed exclusively in the presence of Et,AICI *via* **A.** We may thus predict that in a concerted Et₁AlCl-promoted ene process $3 \rightarrow 5$, H-atom is transferred selectively from the *trans*-positioned allylic CH_3 -group⁷). Its specific labelling with ¹³ $C⁸$) should lead to 9 with all **I3C** localized in the olefinic methylene C(8)-atom *(Scheme 4).* Alternatively, if C,C-bond closure and H-transfer are non-concerted such as in the formation of carbocation 8 the ¹³C-label would be scrambled between $C(8)$ and $C(9)$ in the cyclization product.

Synthesis and Et,AlCl-Promoted Cyclization of *"C-Labelled Diene* **7.** The essential problem thus boiled down to a stereospecific preparation of the trans-¹³C-methyllabelled bromide **14.** To establish selectively the desired alkene geometry, alkyne carbometallation') appeared to be the method of choice. Esterification [16] of 2-butynoic acid **(10)** with (+)-borneol/DCC/DMAP furnished, after chromatography and sublimation, the crystalline butynoate 11 in 98% yield. Stereospecific syn-addition [17] of ¹³Cdimethylcopperlithium (prepared *in situ* from 99% isotopically pure [¹³C]MeI) to butynoate 11 in THF at -78° afforded after chromatography olefin 12 in 99% yield. The

As a working model we postulate a chair-like transition state for the ene reaction. The above-mentioned \mathcal{D} prediction that the H-atom should be transferred exclusively from the allylic trans-methyl group of **3** holds also for the traditional model [13] which assumes that the migrating H-atom lies on the axis which joins the termini of the ene and enophile units.

To avoid the interference with kinetic **H/D** isotope effects [9] and for reasons of NMR-analytical conve- 8 nience $[14]$ we preferred ¹³C- over ²H-labelling.

ን For a review see **115).**

¹³C-NMR spectrum of 12 shows a single peak at $\delta = 27.2$ ppm, assigned to be labelled trans-methyl group. Its corresponding proton signal appears in the 'H-NMR spectrum as a doublet centered at $\delta = 1.90$ ppm (¹³C, H-coupling constant $^1J_{(CH)} = 126$ Hz). The cis-methyl signal at $\delta = 2.17$ ppm is split into a doublet with ${}^{3}J_{(C,H)} = 4.5$ Hz. Not a trace of a signal is visible at $\delta = 1.90$ ppm (unlabelled *trans-CH*,) consistent with virtually 100% stereospecific incorporation of the ¹³C-label. Reduction of the bornyl ester **12** with (prepared in *situ)* AlH, in THF at 0" provided the labelled methyl-butenol **13** in 83% yield after bulb-to-bulb distillation ($25^{\circ}/2$ Torr). Inspection of the ¹³C-NMR spectrum of **13** reveals a major peak at $\delta = 25.5$ (trans-¹³CH₃) and a minor peak at $\delta = 16.8$ ppm¹⁰) (integral ratio 20:1). The ¹H-NMR spectrum shows the *trans*-methyl signals as two doublets centered at $\delta = 1.72$ ppm $(^1J_{(C,H)} = 124$ Hz, 2.54 H) and $(^3J_{(C,H)} = 4$ Hz, 0.46 H) and the complementary cis-methyl doublets centered at $\delta = 1.68$ ppm $(^1J_{\text{C,H}} = 124$ Hz, 0.48 H) and $(^3J_{\text{C,H}} = 4$ Hz, 2.52 H). Accordingly, the ¹H-NMR integrals indicate some scrambling of the ¹³C-label on reduction of 12 resulting in a 5:1 mixture of **13/16'").** Conversion of the allylic alcohol **13/16** to the bromide **14/17** was successfully accomplished¹¹) using hexabromoacetone/Ph,P in sulfolane, analogous to

¹⁰) Since the ¹³C-signals of the *trans*- and cis-methyl groups in unlabelled 3-methyl-2-butenol appear at δ = 25.3 and δ = 17.4 ppm (integral ratio 1:1), the minor peak at δ = 16.8 ppm could not be clearly assigned.

¹¹) The desired N-alkylation $15\rightarrow 7$ could not be achieved using the corresponding tosylate or chloride.

the preparation of volatile chlorides by means of hexachloroacetone/ Ph , Pl [18]. The *trans-* and *cis-methyl groups of 14/17 exhibit two* ¹³C-NMR peaks at $\delta = 25.7$, 17.4 ppm (integral ratio 7:1) and four 'H-NMR doublets at $\delta = 1.78$, 1.74 ppm, respectively. Integration of both 'H-NMR signal pairs $(^1J_{\text{C-H}} = 125 \text{ Hz } vs. \ ^3J_{\text{C-H}} = 4 \text{ Hz})$ centered at $\delta = 1.78$ and 1.74 ppm showed the same stereoisomer ratio $14/17 = 5.2$:1.

N-Alkylation of amide **15** with the thus obtained bromide **14/17** proceeded most reliably after deprotonation of **15** with KH/18-crown-6 in HMPA to afford the crystalline diene 7/18 in 52% yield. ¹³C-NMR analysis of 7/18 shows two peaks at $\delta = 25.5$ and 18.0 ppm (integral ratio 8:1). Integration of the $H\text{-NMR}$ methyl signals centered at $\delta = 1.70$ and 1.53 ppm reveals a 5.1:1 ratio for $7/18$.

Having thus prepared the 13C-labelled 1,6-diene **7** the stage was set for the crucial cyclization. Treatment of the above $7/18$ mixture with Et,AlCl (3 mol-equiv.) at -78° in CH,Cl, for **8** h provided the trans-substituted pyrrolidine **9/19** in 86% yield. The olefinic methylene $(C(8))$ and the allylic methyl $(C(9))$ groups of the cyclization product **9/19** show two ¹³C-NMR peaks at $\delta = 116.6$, 18.0 ppm (integral ratio 6:1) and two pairs of 'H-NMR signals centered at $\delta = 4.93, 1.70$ ppm, respectively. The 'H-NMR integrals of both signal pairs and ${}^{13}CH_2(^1J_{(CH)} = 154 \text{ Hz})/{}^{12}CH_2(^3J_{(CH)} = 6 \text{ Hz})/$ ¹²CH₃(³J_(CH) = 6 Hz)/¹³CH₃(¹J_(CH) = 126 Hz) exhibit the same ratio of 5:1 for **9/19**.

Conclusion. – The localization of ¹³C-label in the precursors 14 and 7, as well as in the ene product **9** has been measured by reliable integration of the relevant 'H-NMR signals (\approx 5% precision). The corresponding ¹³C-NMR spectra confirm these findings in a qualitative sense. Accordingly, the 5:1 stereoisomer ratio of *trans/cis*-¹³C-labelled olefinic precursors **14/17** and **7/18** corresponds well to the 5:l ratio of "C-labelled **C(8)-methylene/C(9)-methyl** groups in the pyrrolidine products **9/19.** Thus, *only* the trans-methyl group in diene **7** is transformed to the olefinic methylene group in **9** during the cyclization. This result strongly supports a concerted mechanism for the ene process $7\rightarrow 9$ where (for geometrical reasons) hydrogen is transferred exclusively from the trans -positioned allylic methyl group of **7.** Within experimental error the intermediacy of cation **8** can be excluded. We thus propose a transition state 7^* where coordination of $R₂$ AlCl with the conjugated ester carbonyl is mainly responsible for the acceleration of the ene reaction by lowering the enophile LUMO energy [9].

The mechanistic understanding achieved by this study may help to stimulate further rational applications of diastereoselective and π -facial-selective ene reactions in synthesis").

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 $\bar{\mathcal{A}}$

¹²) For recent examples of π -face-selective Lewis-acid-promoted ene reactions see [6] [19] [20].

Experimental Part

General. All reactions were carried out under Ar-atmosphere with magnetic stirring. Solvents were dried by distillation from drying agents as follows: diethylether (Et,O, KH) tetrahydrofurme (THF, K-metal), hexamethylphosphoramide (HMPA, CaH₂), dichloromethane (CH₂Cl₂, P₂O₅). 'Workup' denotes washing of the org. phase with sat. aq. NaCl, drying over solid Na₂SO₄ and removal of solvent by distillation *in vacuo* using a rotatory evaporator. Column chromatography was carried out on *SO, (Merck.* Kieselgel 60) using hexane/ Et,O (ratio in parentheses). Melting points (m.p.) were determined on a *Kofler* hot stage and are uncorrected. IR spectra in CCl₄, unless otherwise specified, \tilde{v}_{max} in cm⁻¹. NMR spectra in CDCl₃, ¹H-NMR spectra at 360 MHz, unless otherwise specified, ¹³C-NMR spectra at 90.561 MHz, standard TMS (ppm) = 0.

Ethyl (Z)-4,4-bis(ethoxycarbonyl)-4-(N-methyltrifluoroacetamido)-2-butenoate (6). KH (washed with pentane, 28 mg, 0.7 mmol) was added to a solution of *ethyl* (Z)-4,4-bis(ethoxycarbonyl)-4-(trifluoroaceta*rnido)-bbuten~afe* **(15)** (185 mg, **0.5** mmol), 18-crown-6 (291 mg, 1.1 mmol) in HMPA (1.5 ml) at *0".* After 15 min at 0° CH₃I (44 μ 1, 0.7 mmol) was added to the mixture which then was stirred at r.t. for 24 h and finally poured into 10% aq. citric acid. Workup and chromatography (5:l) furnished *6* (oil, 184 mg, 96%). **IR** (film): 1757, 1733, 1703, 1650, 1192, 1054, 1022. 'H-NMR (100 MHz): 1.1-1.4 (9H); 3.33 *(m,* 3H); 3.94.4 (6H); 6.12 *(d, J* = 12, 1H); 6.51 *(d, J* = 12, 1H). MS: 383 (49, C₁₅H₂₀F₃NO₇⁺), 338 (44), 311 (14), 310 (100), 282 (24), 265 (43).

I,7,7-TrimethyIbicycl0[2.2.1 Jhept-Z-yl2-butynoate **(1 1).** A solution of dicyclohexylcarbodiimide (4.9 g, 23.8 mmol) in CH,CI, (20 ml) was added dropwise to a mixture of 2-butynoic acid **(18,** 2.0 g, 23.8 mmol), (-)-barneol (4.0 g, 25.97 mmol) and dimethylaminopyridine (30 mg, 0.25 mmol) in CH₂Cl₂ (20 ml) at 0°. Subsequent stirring of the mixture at *0"* for 24 h, dilution with CH,Cl,, filtration through *Celite,* evaporation of the filtrate and chromatography (9:l) of the residue gave the crystalline ester **11** (5.12 g, 97.7%) which was sublimed (150" (hath)/0.2 Torr), m.p. 59". IR (CHCI,): 2242, 1697. 'H-NMR (100 MHz): 0.87 **(s,** 3H); 0.89 (s, **3H);** 0.91 (s, 3H); 0.9-2.1 (6H); 2.0 (s, 3H); 2.35 *(m, 1H)*; 4.97 *(m, 1H)*. MS: 220 (34, C₁₄H₂₀O₂⁺), 136 (100), 121 (46), 111 (31), 110 (46), 109 (31), 95 (89).

["C/MethyNithium. [13C]CH31 (99%, I3C, 2.2 ml, 35 mmol) was added to a suspension of Li-powder (30% suspension in mineral oil, 2.26 g, 96.9 mmol, washed with pentane) in Et₂O at such a rate $(ca. 1 h)$ that the mixture is kept at gentle reflux. Stirring of the mixture at 25° for 24 h, filtration, washing of the solid with Et₇O (10 ml) and combination of the filtrates gave a solution of [I3C]CH,Li which was analyzed by *Gilman's* titration $[21]$ (40 ml, 0.67_N, 26.8 mmol, 77%).

1,7,7-Trim~thylhicyclo[2.2.l]hept-2-yl jE)-3-methyl[4-"C]-Z-butenoate **(12).** 0.67~ [I3C]CH,Li (Et,O, 35 ml, 22.4 mmol) was added dropwise to a suspension of **CuI** (2.45 g, 12.89 mmol) in dry THF (48 ml) at 0". After stirring the mixture at 0" for 15 min a solution of **11** (2.45 g, 14 mmol) in THF (12 ml) was added dropwise over 1 h at -78° . Stirring of the mixture at -78° for 3 h, followed by addition of MeOH (2.5 ml) and sat. aq. NH₄Cl (25 ml) at -78° , workup and chromatography (19:1) furnished the labelled ester 12 (oil, 2.6 g, 99%). 'H-NMR: 0.85 **(s.** 3H): 0.88 **(s,** 3H); 0.91 **(s,** 3H); 0.98 *(m,* 1H); 1.18-1.38 (2H); 1.6-1.83 (2H); 1.90 *(d, J* = 126, 3H); 1.96 *(m,* IH); 2.17 *(d, J* =4.5, 3H); 2.38 *(m,* IH); 4.91 *(m,* IH); 5.72 **(br.** *d. J* = 8.5. IH). ${}^{13}C$ -NMR: 27.3.

1.7,7-Trimethylbicyclo/2.2.l/hept-d-yl3-methyl-2-hutenoate. Butynoate **11** (400 mg, 1.82 mmol) was treated with dimethylcopperlithium as described above to give the unlabelled ester (oil, 382 mg, 89%), IR (film): 1716, 1650, 1229, 1146, 851. 'H-NMR: 0.85 (s, 3H); 0.88 (5, 3H); 0.91 **(s,** 3H); 0.98 *(m,* 11H); 1.18-1.4 (2H); 1.62 -1.85 (2H); 1.90 **(s,** 3H); 1.96 *(m,* 1H); 2.17 **(s,** 3H); 2.38 *(m,* 1H); 4.91 *(m,* IH); 5.72 **(s,** IH). '?C-NMR: 167.0 **(s);** 155.5 **(s);** 116.7 *(d);* 78.9 *(d);* 48.7 *(3);* 47.7 **(s);** 45 *(d);* 36.9 (4); 28.0 *(t);* 27.2 *(t* + **y);** 20.1 *(t);* 19.7 *(4);* 18.8 *(4);* 13.5 (q). MS: 236 (13, $C_{15}H_{24}O_2^+$), 153 (4), 137 (8), 136 (26), 110 (23), 83 (100).

 (E) -3-*Methyl[4-¹³C]-2-butenol* (13). Solid LiAlH₄ (57 mg, 15 mmol) was added in one portion to a solution of anh. AICl, (665 mg, 5 mmol) in THF (50 ml) at 0". After stirring the mixture at 0" for 30 min a solution of **12** (2.71 g, 11.43 mmol) in THF (15 ml) was added dropwise at 0". Stirring of the mixture at *0"* for 2 h, slow addition of sat. aq. Na₂SO₄, filtration, concentration of the filtrate by evaporation of the solvent at 1 atm through a *Vigrrux* column followed by distillation of the residue (25"/2 Torr) gave **13** (oil, 820 mg, 83%). 'H-NMR: 1.68 *(d, J* = 124, 0.48H): 1.68 *(d, J* = 4, 2.52H); 1.72 *(d, J* = 4, 0.46H); 1.72 *(d, ^J*= 124, 2.54H); 4.1 $(d, J = 7, 2H)$; 5.38 *(m, 1H)*. ¹³C-NMR: 25.5, 16.8 *(integral ratio 20:1)*.

3-Methyl-2-butenoL **1,7.7-Trimethylbicyclo[2.2.1]hept-2-yl** 3-methyl-2-butenoate (1.18 g, 5 mmol) was reduced with AlH₃ as described above to give unlabelled 3-methyl-2-butenol (oil, 300 mg, 70%). ¹H-NMR (100 MHz): 1.70 **(9,** 3H); 1.76 **(s,** 3H); 4.14 *(d, J* = 7, 2H); 5.44 *(m,* 1H). I3C-NMR: 135.0 **(s);** 123.7 *(d);* 58.6 *(t);* 25.3 *(4);* 17.4 *(q).*

 (E) -1-Bromo-3-methyl[4- ^{13}C]-2-butene (14). Hexabromoacetone [22] (3.69 g, 6.93 mmol) was added portionwise over 20 min to a stirred mixture of **12** (0.8 g, 9.2 mmol), Ph3P (3.26 g, 12.4 mmol) in sulfolane (10 ml) at 0". Then the mixture was allowed to warm to r.t. and to give after distillation at 25"/2 Torr the bromide **14** (oil, 680 mg, 49%). 'H-NMR: 1.74 *(d, J* = 125, 0.49 H); 1.74 *(d, J* = 4, 2.51 H); 1.78 *(d, J* = 4, 0.48 H); 1.78 *(d, J* = 125, 2.52 H); 4.02 *(d, J* = 8.4, 2H); 5.53 *(m, 1H)*. ¹³C-NMR: 25.7, 17.4 *(integral ratio 7:1)*.

Z-Bromo-3-methyl-2-butene. 3-Methyl-2-butenol (280 mg, 3.26 mmol) was treated with hexabromoacetone/ Ph,P as described above to give unlabelled **I-bromo-3-methyl-2-butene** (oil, 195 mg, 40%). IR: 1664, 860, 693. 'H-NMR (100 MHz): 1.73 (s, 3H); 1.78 **(s, 3H);** 3.88 *(d, J* = 8.4, 2H); 5.47 *(m,* IH). I3C-NMR: 140 (s); 120.9 *(d);* 29.6 (t); 25.7 (y); 17.5 *(9).*

Ethyl (Z)-4,4-bis(ethoxycarbonyl)-4-[N-((E)-3-methyl[4-¹³C]-2-butenyl)trifluoroacetamido]-2-butenoate **(7).** KH (washed with pentane, 100 mg, 2.5 mmol) was added portionwise at 0" to a solution of **15** [23] (720 mg, 1.95 mmol) and 18-crown-6 (1.14 g, 4.32 mmol) in HMPA (5 ml). After 15 min at *0",* **14** (550 mg, 3.67 mmol) was added dropwise to the mixture, which then was stirred at r.t. for 24 h and finally poured into 10% aq. citric acid. Workup and chromatography (7:3) furnished unchanged amide **15** (325 mg) followed by the more polar diene **7** (428 mg, 50% or 91% based on recovered 15) which was recrystallized (Et₂O/pentane, - 23°), m.p. 4849". 'H-NMRI3): 1.2-1.4 (9H); 1.53 *(d, J* = 124, 0.49 H); 1.53 *(d, J* = 2, 2.51 H); 1.70 *(m, J 5* 4, 0.50 H); 1.70 *(d, J* = 124, 2.5 H); 4.1-4.4 *(8H)*; 5.25 *(m, 1H)*; 6.13 *(d, J* = 12, 1H); 6.31 *(d, J* = 12, 1H). ¹³C-NMR: 25.5, 18.0 (integral ratio 8:l).

Ethyl *(Z)-4.4-bis(ethoxycurbonyl)-4-[N-(3-methyl-bhutenyl)tr1~uorouceiumido]-2-buienoate.* Treatment of **15** (152 mg, 0.41 mmol) with KH, 18-crown-6 and unlabelled l-bromo-3-methyl-2-butene as described above furnished unchanged **15** (61 mg) and unlabelled ethyl **(Z)-4,4-bis(ethoxycarbonyl)-4-[N-(3-methyl-2-butenyl)trifluoroacetamido]-2-butenoate** (93 mg, 52 % or 86 % based on recovered **15),** m.p. 48-49" (Et,O/pentane, - 10"). IR: 1747, 1710, 1292, 1260, 1210, 1145, 1100, 1030. 'H-NMR: 1.2-1.4 (9H); 1.51 (s, 3H); 1.70(d, *J* = **1,** 3H); 4.14.4 **(8H);** 5.24 *(m.* IH); 6.13 *(d, J* = 12, IH); 6.31 *(d, J* = 12, IH). 13C-NMR: 165.1 **(s);** 165.0 (s); 136.5 *(d);* 133.9 **(s);** 126.0 *(d);* 121.4 *(d);* 72.9 **(s);** 62.8 *(t);* 62.4 **(s);** 60.8 (t); 45.5 *(t);* 45.3 (t); 25.4 *(q);* 17.9 *(9);* 14.0(q); 13.6 *(4).* MS: 437 (5, C,gH26F3N07'), 392 (S), 324 (29), 258 (57), 251 (20), 212 (99, 184 (32), 180 (IOO), 156 (22).

Ethyl *trans-[2,2-bis(ethoxyc~rbonyl)-4-(i-methyl(2-'~C]vinyl)-i-(trifluoroacetyf)pyrroiidin-3-yl]acetate* **(9).** Et2AICI (2.08~ in hexane, 0.33 ml, 0.69 mmol) was added dropwise to a solution of diene **7** (100 mg, 0.23 mmol) in CH₂Cl₂ (1 ml) at -78° . The mixture was stirred at -78° for 8 h, then quenched at -78° with sat. aq. Na₂SO₄, diluted with CH₂Cl₂ and dried with anh. Na₂SO₄. Evaporation and chromatography (3:1) afforded **9(oil,86mg,86%).'H-NMR:1.16-1.4(9H);1.70(d,J=6,2.5H);1.70(d,J=126,0.5H);2.31(dd,J=16.6** and 7, 1H); 2.62 *(dd, J* = 16.6 and 7, 1H); 2.85 *(m,* 1H); 3.0 *(m,* 1H); 3.62 *(t, J* = 11, 1H); 4.0 *(t, J* = 9, 1H); **4.08(q,J=7.3,2H);4.2-4.3(4H);4.93(r,J=6,0.34H);4.93(br.d,J=154,** 1.66H).13C-NMR: 116.6,18.0 (integral ratio 6:1).

Ethyl *trdns-[2,2-bis(ethoxycarbonyl)-4-isopropenyl-l-(trifuoroaceiyl)pyrrolidin-3-yl]aceiuie.* Treatment of unlabelled ethyl (Z)-4,4-bis(ethoxycarbonyl)-4-[N-(3-methyl-2-butenyl)trifluoroacetamido]-2-butenoate (100 mg, 0.23 mmol) with Et₂AlCl at -78° as described above furnished the unlabelled ethyl trans-[2,2-bis(ethoxycarbonyl)-4-isopropenyl-1-trifluoroacetylpyrrolidin-3-yl]acetate (oil, 89 mg, 89%). IR: 1744, 1713, 1445, 1244, 1150, 1095, 1069, 1040, 912. 'H-NMR: 1.16-1.4 (9H); 1.72 **(s,** 3H); 2.33 *(dd, J* = 16.6 and 7, iH); 2.63 *(dd, J* = 16.6 and 7, 1H); 2.85 *(m, 1H)*; 3.07 *(m, 1H)*; 3.63 *(t, J* = 11, 1H); 4.02 *(t, J* = 9, 1H); 4.09 *(g, J* = 7.3, 2H); 4.24.3 (4H); 4.95 (br. s, 2H). I3C-NMR: 170.6 (s); 166.2 (s); 165.2 **(3);** 139.2 (s); 116.6 (t); 114.4 (s); 74.5 **(s);** 62.4 *(t);* 60.7 *(t);* 51.0 *(d);* 50.3 (t); 45.5 *(d);* 33.7 *(t);* 18.0 *(9);* 14.0 *(4);* 13.9 (y); 13.8 *(4).* MS: 437 (21, CIYH,,F,NO7+), 392 (7), 364 (37), 318 (IOO), 290 (74), 272 (21), 262 **(15),** 259 **(18),** 216 (12).

¹³⁾ The integrals of overlapping satellite bands were assigned as being equal to those of the corresponding isolated satellite bands.

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