

Table I. Effect of Lewis Acid in the Cyclization of $1a \rightarrow 4a/5a^a$

entry	reagent	temp, °C	% syn (4a) ^b	% anti (5a) ^b	yield, % ^{b,c} (mass recovery)
1	Me ₃ SiOTf	-70	96	4	100
2	TfOH	-70	96	4	62 (74)
3	$Ti(OiPr)_2Cl_2$	-20	87	13	21 (91)
4	AlCl ₃	-20	86	14	33 (78)
5	BCl ₃	-70	82	18	57 (73)
6	BF ₃ •OEt ₂	-20	77	23	95 (100)
7	TiCl₄	-90	47	53	55 (58)
8	$SnCl_4$ (1.0 equiv)	-70	45	55	35 (60)
9	SnCl ₄ (0.5 equiv)	-60	71	29	81 (81)

^aAll cyclizations were performed in CH₂Cl₂ (0.05 M) with 1.0 equiv of Lewis acid (except entry 9). At least 3 runs with each Lewis acid ($\pm 3\%$). ^bRatios and yields were calculated based on independently determined response factors vs cyclododecane. ^cYield is based on 4a + 5a vs cyclododecane; mass recovery is total integrated area including products of protiodesilylation.

Table II. Effect of Lewis Acid in the Cyclization of $2a \rightarrow 4a/5a^a$

entry	reagent	temp, °C	% syn (4a)	% anti (5a)	yield, % (mass recovery)
1	Me ₃ SiOTf	-70	93	7	100
2	BF ₃ OEt ₂	-60	94	6	63 (85)
3	TiČl₄	-70	41	59	44 (49)
4	SnCl ₄	-30	38	62	25 (25)

"See footnotes Table I.

Table III. Effect of Acetal Structure in the Cyclization of 1 and $2 \rightarrow 4/5$ with TMSOTf^a

substrate	ML'3	R	% syn (4) ^b	% anti (5) ^b	
1a	SiMe ₃	Me	96	4	
1b	SiMe ₃	Et	92	8	
1c	SiMe ₃	<i>i</i> Bu	90	10	
1 d	SiMe ₃	iPr	38	62	
2a	SnBu ₃	Me	93	7	
2b	SnBu ₃	Et	92	8	
2d	SnBu ₃	iPr	43	57	

^{*a*}All cyclizations were performed in CH₂Cl₂ (0.05 M) with 1.0 equiv of TMSOTf for 1 and 2.0 equiv for 2. At least 3 runs with each substrate ($\pm 3\%$). ^{*b*}Ratios were calculated on independently determined response factors vs cyclododecane. All yields were >95%.

Table IV. Effect of Enol Ether Structure in the Cyclization of $3 \rightarrow 4/5^{\circ}$

substrate	R	E/Z^b	temp, °C	% syn (4) ^c	% anti (5)°	yield, %'
(E)- 3a	Me	96/4	-50	60	40	25 ^d
(Z)-3a	Me	0/100	-70	53	47	79
(E)- 3 b	Et	100/0	-70	31	69	76
(Z)-3b	Et	23/77	-70	38	62	74
(E)-3d	iPr	100/0	-70	25	75	84
(Z)-3d	iPr	0/100	-70	27	73	76

^aAll cyclizations were performed in CH₂Cl₂ (0.05 M) with 0.95 equiv of TfOH. At least 3 runs with each substrate ($\pm 3\%$). ^bEstablished by capillary GC analysis. ^cRatios and yields determined by independently determined response factors vs cyclododecane. ^dThe major product resulted from protiodesilylation.

TMSOTf these methyl, ethyl, and probably isobutyl acetals react via an S_N2 -type mechanism, while the isopropyl acetals react via prior ionization to an oxocarbenium ion (i). Accordingly, these conclusions are supported by the observed Lewis acid dependences wherein the nature of the Lewis acid acetal complex is expected to influence the S_N2 -type reaction. Furthermore, the inherent strain in diisopropyl acetals should favor ready ionization compared to ethyl and methyl analogues.

Scheme II



Studies are in progress on the structure of Lewis acid acetal complexes and the stereochemical course of reactions with cyclic acetals.

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Supplementary Material Available: Full characterization for the models 1, 2, and 3 are provided along with representative cyclization procedures (8 pages). Ordering information is given on any current masthead page.

Metal-Promoted Carbon-Carbon Bond Formation in the Gas Phase: Reaction of Iron Carbonyl Cations with Allyl Chloride

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Gaseous transition-metal ions, bare¹ or bearing ligands,² have been shown to be extremely reactive with various organic compounds; for example, group 8 metal cations are able to cleave C-H and C-C bonds of alkanes.³ However, contrary to solution organometallic chemistry, gas-phase organometallic chemistry includes relatively few examples of carbon-carbon bond formation.⁴ We report here our first results concerning the Fe-(CO)_n⁺-allyl chloride (3-chloro-1-propene) system, in which a new type of C-C bond formation is encountered.

The reactivity of each $Fe(CO)_n^+$ ion (n = 0-5) with allyl chloride was studied by using a MS/MS/MS multiquadrupole spectrometer, described elsewhere,⁵ with the following configuration: (i) source, electron impact on $Fe(CO)_5$; (ii) first quadrupolar analyzer, selection of $Fe(CO)_n^+$ reagent ion; (iii) first collision cell, reaction with allyl chloride (ca. 1 mTorr); (iv) second quadrupolar analyzer, selection of a reaction product; (v) second collision cell, CAD of this product (collision gas; Argon, ca. 0.7 mTorr, collision energy 15–25 eV); (vi) third quadrupolar analyzer, scanning. The relatively high pressure in the first collision cell allows successive reactions.

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Table I. Primary Spectra from Reaction of $Fe(CO)_n^+$ Ions (from EI on $Fe(CO)_5$) with Allyl Chloride, Percent of Total Reaction for Each Important Product Ion,^a and Approximate Overall Rate Constants

	reactant ion						
product ion (m/z)	Fe ⁺	Fe ^{+ b}	FeCO ⁺	Fe(CO) ₂ ⁺	Fe(CO) ₃ ⁺	Fe(CO) ₄ + c	Fe(CO) ₅ ⁺
$C_3H_5^+$ and products ^d	83	89	53	23	7	2 (1)	1
$C_6H_{10}^{*+}$ (82), $C_5H_7^{+}$ (67) ^e	7	6	16	33	37	13 (9)	5
$Fe(CO)_{n=1}^{+}$			3	6	7	12 (15)	51
FeCl ⁺ (91.93)	4	3	2	0	0	0 (0)	0
$FeC_{3}H_{5}Cl^{+}$ (132, 134)	3	1	15	31	27	7 (11)	3
$FeC_4H_5ClO^+$ (160, 162)			0	0	8	24 (8)	11
$FeC_6H_9Cl^+$ (172, 174)	1	0	3	4	7	3(1)	I
$FeC_6H_{10}Cl_2^+$ (208, 210, 212)	0	0	0	0	5	34 (53)	16
rate constant (in ms ⁻¹ mTorr ⁻¹) ^f	6	g	14	17	7	35	g

^a Because of minor reaction pathways, not included here the sum of the values corresponding to a given reactant ion is generally slightly less than 100. ^b From FAB source. ^c In parentheses, values obtained when adjusting the energy of reacting ions so as to optimize the $FeC_6H_{10}Cl_2^+$ yield. ^dSum of the abundances of $C_3H_5^+$ (m/z 41) ion and its products of reaction with allyl chloride: m/z 55, 75, 79, 81, 89, 95, and 115 ions. ^eC_5H_7^+ ion arises from $C_6H_{10}^{\bullet+}$ cleavage (see text). ^fDetermined from kinetic studies: allyl chloride pressure 0.1-1 mTorr, confinement time 0.06-1.4 ms. ^gNot determined.

In the first step the second analyzer was left transparent and the second collision cell empty, in order to get the primary Fe- $(CO)_n^+$ -allyl chloride reaction spectra. These spectra, summarized in Table I, exhibited very different patterns depending on the number of CO ligands born by the reagent ion. The reactions were also performed at variable reaction time under ion confinement conditions,⁶ allowing us to determine their rate constants (Table I).

For n = 0 (bare Fe⁺ ion) the spectrum was very similar to the reaction spectrum of source-generated C₃H₅⁺ ion and allyl chloride; this indicates the following reaction scheme, which was also corroborated by ion rejection experiments.7

$$Fe^+ + C_3H_5Cl \xrightarrow{-FeCl} C_3H_5^+ \xrightarrow{C_3H_5Cl} C_6H_9^+, C_6H_7^+ \dots$$

$$FeCl^+ FeCl^+$$

This reactivity is not surprising considering the known reactions of alkyl halides with Fe⁺⁸ or other^{2c,8,9} transition-metal ions; apart from dehydrohalogenation, which when possible is the major reaction pathway, alkyl cation formation^{2c,8b,9} as well as MX⁺ formation^{8,9a} have been reported. Increasing n allows fixation of one and then two allyl chloride molecules to the metal atom, accompanied with departure of CO molecules. The major ironcontaining product ion is thus $FeC_3H_5Cl^+$ for n = 1-3 and $FeC_6H_{10}Cl_2^+$ for n = 4. According to the confinement experiments, the latter is formed through $Fe(CO)C_3H_5^+$ and Fe- $(CO)_2C_3H_5^+$ intermediates. Related "replacement" reactions of one or two CO ligands of metal carbonyl ions by an alkyl chloride molecule have been described.^{2a,c,8} $Fe(CO)_5^+$ ion is much less reactive^{2a} and undergoes mainly CO loss. Ion $C_6H_{10}^{++}$, along with its fragmentation product $C_5H_7^+$, was always found in variable proportions. This ion, which is not a product of C₃H₅⁺ reaction with C₃H₅Cl, should arise from an allyl coupling reaction in an iron complex ion. Its main precursors are likely $FeC_3H_5^+$ and $Fe(CO)C_3H_5^+$ ions.

The CAD spectra of the most significant products of the preceding reactions were then recorded, giving the following results.

$$C_{6}H_{10}^{++} (m/z \ 82) \rightarrow C_{5}H_{7}^{+} (m/z \ 67) \ 66\%$$

$$\rightarrow C_{4}H_{6}^{++} (m/z \ 54) \ 28\%$$

$$\rightarrow C_{3}H_{5}^{+} (m/z \ 41) \ 6\%$$

This spectrum is very similar to the fragmentation pattern of hexadiene radical ions,¹⁰ as expected if C_6H_{10} ⁺ is the coupling product of two C_3H_5 units. This pattern is unfortunately not sensitive to the position of the double bonds.

$$FeC_{3}H_{5}Cl^{+} (m/z \ 132, \ 134) \rightarrow FeCl^{+} (m/z \ 91, 93) + C_{3}H_{5}^{*} \ 28\%$$

$$\rightarrow FeC_{3}H_{4}^{+} (m/z \ 96) + HCl \ 62\%$$

$$\rightarrow C_{3}H_{5}^{+} (m/z \ 41) + FeCl \ 10\%$$

This spectrum is very different from that reported for Fe- $(propene)^+$ ion, in which the only fragmentation is propene loss.¹¹ Absence of C_3H_5Cl loss as well as C_3H_5 and FeCl losses indicate Fe⁺ insertion into the carbon-halogen bond, as observed in the case of methyl halides.⁸ The relative importance of HCl elimination decreases with increasing collision energy, as expected for a rearrangement reaction as opposed to simple cleavages.¹² $FeC_{3}H_{4}^{+}$ ion, possibly an allene complex, was also detected in primary reaction spectra as a minor product; its CAD showed C_3H_4 loss as only fragmentation. Thus structure 1 can reasonably be assigned to $FeC_3H_5Cl^+$ ion, since π -allyl complexes of iron¹³ and other metals^{13,14} are known in the gas phase.

$$FeC_4H_5ClO^+$$
 (m/z 160, 162) \rightarrow $FeC_3H_5Cl^+$ (m/z 132, 134) + CO

secondary fragmentation:identical as above

Formation of 1 shows that no CO insertion occurred in this complex, which can thus be represented by structure 2.

FeC₆H₁₀Cl₂⁺ → C₆H₁₀⁺ (
$$m/z$$
 82) + FeCl₂ 77%
(m/z 208, 210, 212) and fragments
→ FeC₃H₅Cl⁺ + C₃H₅Cl 23%
(m/z 132, 132–134, 134)

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Formation of a C_6 fragment indicates that a carbon-carbon bond has been formed between the two C₃ units incorporated in the $FeC_6H_{10}Cl_2^+$ complex, suggesting a (hexadiene)FeCl_2^+ structure (3 or double bond position isomer). However, this



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structure is not consistent with the observed allyl chloride loss, which may arise from structure 4. Thus we conclude that $FeC_6H_{10}Cl_2^+$ ion is actually a mixture of two complexes 3 and 4. In order to test this hypothesis, ligand exchange was attempted: FeC₆H₁₀Cl₂⁺ was submitted to reaction with acetonitrile or trideuterated acetonitrile in the second collision cell, at a variable reaction time. The following reaction sequences were observed.



 $FeC_3H_5Cl(CD_3CN)^+$ ion, deriving from rapid replacement of allyl chloride by acetonitrile in structure 4, leads successively to $FeCl(CD_3CN)_2^+$ and $FeCl(CD_3CN)_3^+$ ions. After a short time (ca. 0.2 ms) the sum of the relative intensities of these three ions remained constant and equal to approximately 40% of the total ion current. Therefore the m/z 208 ion mixture reacted with acetonitrile should contain ca. 40% of structure 4. The remaining slower reactions of m/z 208, involving either C₆H₁₀^{•+} formation or replacement of neutral C_6H_{10} with acetonitrile, are in good agreement with structure 3. $4 \rightarrow 3$ isomerization reactions will now be attempted.

The reactivity of methallyl chloride with iron complexes was also investigated: with $Fe(CO)_{n}^{+}$ it is the same as that of allyl chloride. With $FeC_3H_5Cl^+$ no significant ligand exchange was observed, as expected from structure 1; the major reaction was chloride abstraction.

$$FeC_{3}H_{5}Cl^{+} + C_{4}H_{7}Cl \rightarrow C_{4}H_{7}^{+} (m/z \ 55) + C_{3}H_{5}FeCl_{2}$$

$$C_{4}H_{7}Cl \rightarrow C_{8}H_{13}^{+} (m/z \ 109)$$

Ion $FeC_6H_{10}Cl_2^+$ gave the expected rapid but incomplete substitution of one allyl chloride ligand by methallyl chloride, followed however by complex reactions.

The coupling reaction occurring within the $FeC_6H_{10}Cl_2^+$ complex and leading after fragmentation to hexadiene radical cation is analogous to the transition-metal-promoted coupling of allylic halides in solution (Corey reaction).¹⁵ Interestingly, occurrence of this solution-like reaction requires the presence of several ligands on the starting iron complex ion. Further investigation is in progress to determine the scope of this reaction concerning the nature of the metal ion and ligands as well as of the organic halide.

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Registry No. 1, 119909-88-5; 3, 119909-89-6; 4, 119909-90-9; Fe+, 14067-02-8; $FeCO^+$, 35038-14-3; $Fe(CO)_2^+$, 35038-15-4; $Fe(CO)_3^+$, 35038-16-5; $Fe(CO)_4^+$, 35038-17-6; $Fe(CO)_5^+$, 59699-78-4; allyl chloride, 107-05-1.

Supplementary Material Available: Experimental spectra for all the processes mentioned in the text (primary reaction or CAD) (34 pages). Ordering information is given on any current masthead page.

The Unusually Robust P-H Bond in the Novel Cation HP(NMeCH₂CH₂)₃N⁺

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It was reported recently on the basis of NMR solution data that the phosphatrane cation 1 is unstable and was therefore not



isolated.¹ In earlier publications from our laboratories, we observed, on the other hand, that the unstable prophosphatrane 2 readily protonates to form the stable cation 3^{2-4} Here we show that in contrast to 2, compound 4 is stable and is capable of deprotonating a variety of weak acids, including water, to form the unusually stable cation 5.

Cation 5 is prepared as the chloride by adding a solution of (HMeNCH₂CH₂)₃N⁵ (1.67 g, 11.4 mmol) in CH₂Cl₂ (20 mL), over a period of 5 min to a stirred solution of ClP(NMe₂)₂ (1.76 g, 11.4 mmol) and Et₃N (1.5 g, 15 mmol) in CH₂Cl₂ (30 mL). Stirring at room temperature for 1 h followed by removal of the solvent and Et_3N afforded 5 as the chloride in quantitative yield. The salt was recrystallized from hexane/chloroform at -20 °C to give an 82% yield of the product as a colorless crystalline solid.6 Treatment of 5 (Cl⁻) with $AgBF_4$ in CH_2Cl_2 gave the BF_4^- salt in quantitative yield. The ³¹P NMR chemical shift of -10.6 ppm for 5 is indicative of five-coordinate phosphorus,^{2,7} as is its one-bond H-P coupling constant of 491 Hz $(3, 794 \text{ Hz};^2 1, 849 \text{ Hz}^1)$. The existence of an N_{ax} -P bond in 5 in solution is suggested by the observation of substantial $PN_{ax}CH_2$ coupling (4.7 Hz) and the observation of two-bond $PN_{ax}C$ coupling (6.1 Hz). These couplings are absent in prophosphatranes 2^3 and $4^{.5}$ The structure of 5 was confirmed for the BF₄ salt by X-ray crystallographic means (Figure 1).⁸ Although the H-P hydrogen in $5(BF_4)$ could not be located, the sum of the NPN angles in the equatorial plane (358.9 (4)°), the nearly right-angle relationship of the N_{ax} -P bond with the N_{eq} -P linkages (average 86.3 (4)°), the detection of the ${}^{1}H-{}^{31}P$ coupling in solution, and the directionality of the N_{ax} lone

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Allg. Chemie, accepted for publication. (6) NMR data: ³¹P (CDCl₃) δ -10.6; ¹H (CDCl₃) δ 2.61 (9 H, d, CH₃, $^{3}J_{PH} = 17.4 \text{ Hz}$), δ 3.03 (6 H, dt, $N_{ax}CH_2$, $^{3}J_{PH} = 11.0 \text{ Hz}$, $^{3}J_{HH} = 6.2 \text{ Hz}$), δ 3.58 (6 H, dt, $N_{ex}CH_2$, $^{3}J_{PH} = 4.7 \text{ Hz}$, $^{3}J_{HH} = 6.2 \text{ Hz}$), δ 5.20 (1 H, d, $^{1}J_{PH} = 491 \text{ Hz}$); ¹³C (CDCl₃), δ 34.4 (d, CH₃, $^{2}J_{PC} = 17.1 \text{ Hz}$), δ 41.3 (d, $N_{ax}CH_2$, $^{2}J_{PC} = 6.1 \text{ Hz}$), δ 47.3 (d, $N_{eq}CH_2$, $^{2}J_{PC} = 7.3 \text{ Hz}$). (7) Carpenter, L. E.; de Ruiter, B.; van Aken, D.; Buck, H. M.; Verkade, L. C. L. due Cham. Soc. 1065 (408)

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(8) Crystal data: space group Cmc^{2} , a = 8.7251 (5) Å; b = 12.3824 (7) Å, c = 13.1330 (7) Å, V = 1418.9 (2) Å³, Z = 4, $d_{calcd} = 1.423$ g/cm³, μ (MoK α) = 2.24 cm⁻¹; 897 unique reflections for $2 < \theta < 25^{\circ}$, 760 observed $(F_{o}^{2} > 3\sigma(F_{o}^{2}))$. The choice of the acentric space group was suggested by intensity statistics and was confirmed by successful refinement of the structure. The structure was solved by direct methods. Since the phosphatrane cage resides on a crystallographic mirror plane, a slight disorder arises from the possible orientations of atoms C(3), C(6), and C(7). The occupancies of the two orientations of the BF_4^- ion refined to 56.1 (5)% for the major orientation. Refinement of 113 parameters converged with agreement factors of the fol-lowing: $R_1 = \Sigma |F_o - F_c| / \Sigma F_o = 0.0887$ and $R_2 = \operatorname{sqrt} [\Sigma w (F_o - F_c)^2 / \Sigma w (F_o^2)] = 0.1189$. The refinement was carried out with the SHELX-76 package.

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