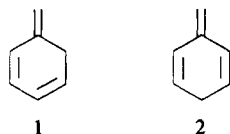


electron impact on the various precursors appear to be a mixture of benzyl and tropylium cations, at least on a millisecond time scale. The ratio of the two forms depends on the nature of the neutral precursor and the electron energy.^{4,5} The recent availability⁶ of two tautomers of toluene, **1** and **2**, has prompted an investigation of the energetics and reactivity of these new entries onto the $C_7H_8^+/C_7H_7^+$ surface.



The heats of formation of **1** and **2** have been determined by using ion cyclotron resonance spectrometry. We observe the onset for their deprotonation by anions of increasing base strength to give the 91^- ion. Tautomers which give a common anion on deprotonation should differ in acidity by the difference in their heats of formation, the higher energy forms being more acidic.⁷ Neither **1** nor **2** give any 91^- ion on treatment with HS^- or $t-BuS^-$. The bases were formed by electron impact on H_2S and $t-BuSS-t-Bu$, respectively. Both tautomers yield small amounts of 91^- ion when reacted with $i-PrS^-$ and appreciable amounts from CH_3S^- and stronger bases. Since the isotoluenes are both 23 ± 3 kcal/mol more acidic than toluene,^{7,8} the ΔH_f° for **1** and **2** must likewise be 23 ± 3 kcal/mol above that of toluene, or 35 ± 3 kcal/mol. MNDO calculations agree with these experimental results, giving ΔH_f° for **1** and **2** as 23 kcal/mol greater than that of toluene.⁹ These calculations also find the most stable structures for both tautomers to be within 1 degree of planarity for the carbon skeletons. This contrasts with an electron diffraction study of the 5,5-dimethyl derivative of **2**, which was found to exist in a boat conformation with approximately 7° puckering at each end.¹⁰ The MNDO method, however, often gives ring structures more planar than are experimentally observed.⁹ The 91^- ion formed from **2** by the stronger bases exchanges only two protons with D_2O and CH_3OD , indicative of the benzyl structure.

For **2**, the apparent ionization potential in the ion cyclotron resonance spectrometer by both the vanishing current method and the parallel slopes method relative to toluene under identical conditions is 8.6 ± 0.1 eV or 0.2 eV less than that of toluene. This agrees with the absolute value from MNDO calculations of 8.67 eV, although we note that such calculations do not give a good value for the toluene IP. These results put the M^+ of **2** at about 18 kcal/mol higher in energy than M^+ for toluene. The MNDO calculations also predict that the ionization potentials of **1** and **2** are identical. The 70-eV electron impact mass spectrum of **2**, obtained on a Hewlett-Packard 5992 GC/MS, resembles that of toluene, save that the proportion of fragment ions relative to the 91^+ base peak was about twice as large for **2** as for toluene while the parent M^+ was smaller.

It has been postulated that the relative amounts of benzyl and tropylium cations in the $C_7H_7^+$ population can be determined on the basis of their reactivity. Deuterium labeling studies have shown that those $C_7H_7^+$ ions that are reactive with a wide variety of neutral compounds are probably of the benzyl structure,^{4,5} while tropylium ion is known to be unreactive with most compounds.⁴ We find that at 30-eV electron energy more than 90% of the 91^+ ions from **2** react with the neutral to give 105^+ ions but that the corresponding reaction in **1** proceeds for only ca. 40% of the originally formed $C_7H_7^+$. These data are compared in Figure 1

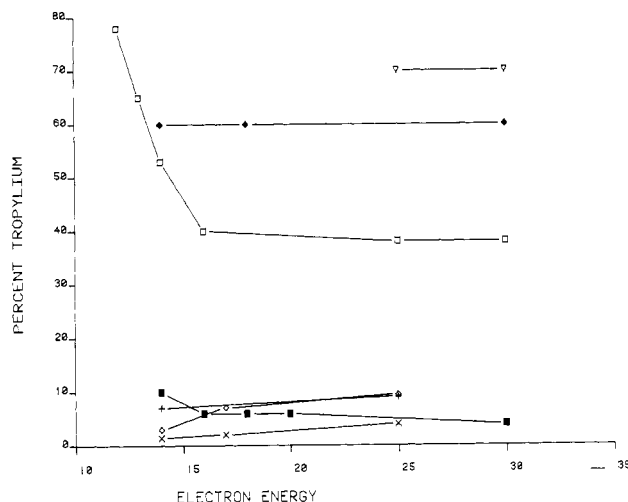


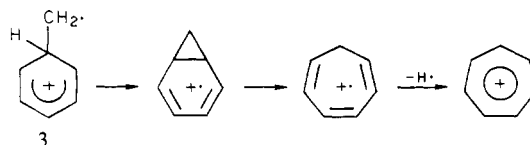
Figure 1. Percent of tropylium in the $C_7H_7^+$ population as a function of electron-impact energy on various precursors. (\blacklozenge) **1**; (\blacksquare) **2**; this work. (\square) toluene; (∇) norbornadiene; (\diamond) benzyl chloride; (\times) *o*-chlorotoluene; ($+$) *m*-chlorotoluene, ref 5.

Table I. Rate Constants and Efficiencies for Cationic Reactions of the Toluene Tautomers

tautomer	$92^+ \rightarrow 93^+$		$91^+ \rightarrow 105^+$	
	k^a	E^b	k^a	E^b
1	2.2	0.18	0.6	0.05
2	1.7	0.14	2.7	0.22
toluene	0.0	0.0 ^c	1.6	0.13

^a Units of $10^{10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, $\pm 20\%$. ^b Efficiency, $k_{\text{obsd}}/k_{\text{ADO}}$. ^c No 93^+ ion observed.

with data for other precursors. It has been stated that the ratio of the two $C_7H_7^+$ forms is not set by direct interconversion but by preequilibration of the parent M^+ ions and subsequent hydrogen loss.⁵ We believe it significant that the tropylium population is appreciable only for those precursors that can give **3**, a logical intermediate to tropylium as shown, by a simple bond cleavage, as with norbornadiene, or by a 1,2-hydrogen shift, as with **1** or toluene.¹¹



The other ion-molecule chemistry of **1** and **2** also differs significantly from that of toluene. The parent M^+ ions of **1** and **2**, unlike toluene, both react with their neutrals to give $M + 1$ ions. We assume protonation occurs on the exo-methylene group in **1** and **2** to give a ring-protonated toluene plus benzyl radical. That ion is the most stable of the possible products and should be identical for **1** and **2**, due to scrambling. Using literature values for ΔH_f° for these species, plus the thermochemistry obtained in this work for the reactants, the 92^+ to 93^+ reaction should be 37 ± 6 kcal/mol exothermic for **1** and **2** but 4 ± 3 kcal/mol endothermic for toluene. The rate constants in Table I for this proton-transfer reaction are comparable for **1** and **2**. In contrast, the alkylation rate for the benzyl ion reacting with the neutral to give 105^+ ion is appreciably faster for **2** than for **1**, as with the corresponding neutral chemistry.⁶ The rates cannot be strictly compared with the reaction in toluene, since there is the possibility of different 105^+ structures in the reactions, such as 1-phenethyl vs. xylyl ions. Both **1** and **2** give many ionic products in the

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106–250 amu range, while toluene yields only a small amount of 128⁺ and 129⁺ above mass 105.

Acknowledgment. We thank J. J. Gajewski for the samples of **1** and **2** and for helpful discussions and the National Science Foundation for financial support.

Registry No. 1, 20679-59-8; **2,** 3217-87-6; toluene, 108-88-3.

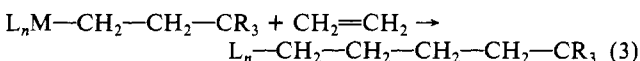
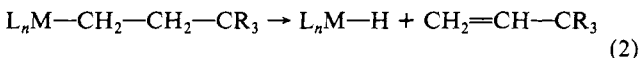
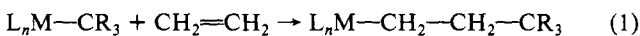
Ziegler–Natta Polymerization: The Lanthanide Model¹

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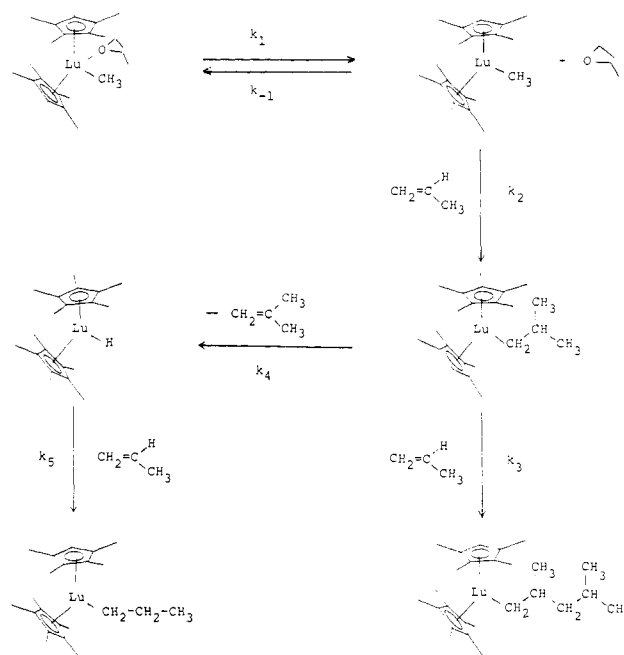
Received September 3, 1981

The formal insertion of an olefin into a metal–carbon (M–C) or metal–hydrogen (M–H) bond is of fundamental significance in a large number of catalytic reactions. While complexes arising from insertion of olefins into the M–H bond have been observed directly^{2–4} primary insertion products from the analogous reaction with M–C bonds (reaction 1) have proved experimentally elusive.^{5–8} The existence of the latter type of insertion product has often been inferred from the isolation of secondary products which derive from the initial insertion product via such processes as β -hydrogen elimination^{9,10} (reaction 2) or further olefin insertion¹¹ (reaction 3).



The direct insertion shown in reaction 3 has been postulated¹² to be a major reaction pathway for olefin polymerization using Ziegler–Natta catalysts^{13,14} (group 6 transition metal with aluminum alkyl). One important difference between this insertion mechanism¹⁵ and the alternative McKinney¹⁶ or Green¹⁷ mech-

Scheme I



anisms for olefin polymerization is that the latter two schemes require oxidative additions in order to generate the active metallacyclic or carbenoid intermediates. While these routes may be reasonable for transition metals having at least two d electrons, they are not so reasonable for the lanthanide elements where reversible two-electron couples are generally precluded by the lack of accessible oxidation states.

During our investigations of the reactivity of lanthanide–carbon bonds, we observe insertion of a variety of olefins and acetylenes into the ytterbium- and lutetium–methyl bonds of $M(\eta^5-C_5Me_5)_2CH_3$ -ether ($M = Yb, Lu$). In these reactions the primary insertion products are observable by spectroscopic and chemical techniques. This preliminary communication describes the synthesis of $M(\eta^5-C_5Me_5)_2CH_3$ -ether (**1**, $M = Lu$) and the reaction of **1** with propene to give the isobutyl complex $M(\eta^5-C_5Me_5)_2-CH_2-CH(CH_3)_2$ (**2**, $M = Lu$). The chemistry of $Lu(\eta^5-C_5Me_5)_2CH_3$ -ether and $Yb(\eta^5-C_5Me_5)_2CH_3$ -ether is qualitatively similar with respect to the olefin insertion reaction.

The dimethyl complexes $[M(\eta^5-C_5Me_5)_2(CH_3)_2]Li(THF)_3$ ($M = Yb, Lu$) are prepared as described¹⁸ earlier for $M = Yb$. Conversion of these complexes to the neutral monomethyl species is achieved in several steps. The THF-free materials, $[M(\eta^5-C_5Me_5)_2(CH_3)_2]Li$ (obtained from the solvated precursors by heating under vacuum at 75 °C), react with $Al(CH_3)_3$ in pentane to give the soluble adducts $M(\eta^5-C_5Me_5)_2Al(CH_3)_4$.¹⁹ Cleavage of these adducts by dissolution in diethyl ether yields $M(\eta^5-C_5Me_5)_2CH_3$ -ether ($M = Yb, Lu$)^{20,21} which can be crystallized

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