

Figure 2. 3-21G geometries of transition structures for reactions of hydride with ethylene, methyl anion with ethylene, and hydride with acetylene.



Figure 3. Possible mechanism for the addition of an alkyllithium tetramer to ethylene.

with some lateral displacement and a significant decrease of the $C^{\delta\bar{-}} - Li^+$ distance.^17

The angle of attack of H⁻ or Me⁻ on ethylene or acetylene is significantly larger than tetrahedral,¹⁸ even though little CC bonding has developed in the transition structures. The presence of the Li⁺ counterion decreases this angle slightly, as compared to isolated anion additions.

Activation energies and transition structures for additions to formaldehyde are not influenced significantly by dimerization of LiH or MeLi,⁷ perhaps because the aggregation of organolithium compounds arises primarily from electrostatic effects.¹⁹ Α conceivable mechanism of addition of a solvated lithium tetramer to ethylene is shown in Figure 3. Displacement of solvent and coordination of ethylene should be slightly exothermic, since initial solvation energies of alkyllithium tetramers with THF are typically 7-8 kcal/mol, and 2-6 kcal/mol with ether.²⁰ Since the lone pair of the methyl anion is pointed toward the center of the lithium tetrahedron, two of the methyl-lithium bonds must be lengthened in order to allow rotation of the lone-pair to interact with the ethylene terminus. The activating effect of Lewis basic solvents upon the addition reaction^{2a} may be due to greater ease of lengthening of these bonds when lithium is additionally coordinated in the transition state.

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Registry No. LiH, 7580-67-8; CH₃Li, 917-54-4; ethylene, 74-85-1; acetylene, 74-86-2.

Supplementary Material Available: Listings of geometries and energies (4 pages). Ordering information is given on any current masthead page.

Multiple Substitutions in Radical-Chain Chlorinations. A New Cage Effect

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It is widely agreed that radical-chain Cl₂ chlorinations of aliphatic systems in noncomplexing solvents or in gas phase are slower when electronegative substituents are present:1a For example, alkyl chlorides react less rapidly than the corresponding alkanes. Thus, the behavior we report here appears to be anomalous: In CCl₄ or CFCl₃ solvents, photochlorinations with low conversions of cyclohexane, isobutane, neopentane, or 2,3-dimethylbutane, with careful exclusion of O_2 , result in formation of unexpectedly large proportions of multiply chlorinated products, resulting from further chlorination of the monochloro products. Although this is an effect which gives the appearance that the chloro derivatives are more reactive than the alkane, the dependence on alkane concentration leads us to a novel proposal, a type of elementary process which appears to have been unrecognized heretofore.

For example 1000 μ mol of purified cyclohexane (purity 99.99%; 0.10 M) with 309 μ mol of Cl₂ in CCl₄ (10 mL) shows no product formation in the dark, but on brief exposure to a tungsten lamp results in formation of 74 μ mol of cyclohexyl chloride, 67 μ mol of dichlorides, and 39 μ mol of a mixture of trichlorides; *these are produced in the presence of more than 800 \mumol of unreacted cyclohexane*. The composition of these multiply halogenated products is readily recognizable with GC-mass spectrum analysis. Thus, cyclohexyl chloride appears to be far more reactive than cyclohexane. The more usual behavior, cyclohexane more reactive than cyclohexyl chloride, is observed in chlorination of undiluted cyclohexane (18.5 mmol; no added solvent) with 483 μ mol of Cl₂, resulting in formation of 460 μ mol of cyclohexyl chloride and 15 μ mol of dichlorides (no trichlorides).

With low conversion conditions we report here that the amount of polychlorinated product relative to the total amount of chlorinated product increases with decreasing concentration of cyclohexane. This can be seen in Table I. In the first four reactions, each starting with 10 mol % Cl₂ with respect to cyclohexane, the percentage of polychlorinated product increases from 6% in the photochlorination of neat cyclohexane to 56% in the reaction carried out with 0.030 M cyclohexane in CCl₄. The same increases in the percentage of polychlorinated product were observed in the reactions that initially contained 30 mol % Cl₂. Despite the 3-fold greater conversion of the cyclohexane, there is only a slightly increased percentage of polychlorinated product. This effect is independent of the Cl₂ concentration and is attributable to the change of cyclohexane concentration and to the percentage of its conversion. For example, a reaction of 0.020 M cyclohexane in CCl_4 in the presence of 0.031 M Cl_2 , stopped early by quenching with corn oil, resulted in $\sim 20\%$ loss of the original 200 μ mol of cyclohexane and production of 12.4 μ mol of cyclohexyl chloride, 16.0 μ mol of dichlorides, and 11.0 μ mol of a mixture of trichlorides, 68% polychlorinated product. A reaction identical except for 0.006 M Cl₂ resulted in an identical product composition. Thus, the percentage of polychlorination is independent of $[Cl_2]$ and dependent on $[C_6H_{12}]$.

An analogous result is obtained with low conversion photochlorinations of 2,3-dimethylbutane (DMB). In pure DMB the two isomeric C_6H_{13} Cl's are produced in ~100% yield; with 0.10 M DMB in CFCl₃ (10 mL), 102 μ mol of Cl₂ produces 41 μ mol of monochlorides and 24 μ mol of dichlorides. This result is independent of either the presence of HCl produced in the reaction, or its presence initially at 0.5 M, or of scavenging of HCl by anhydrous K₂CO₃.

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Table I. Photochlorinations of Cyclohexane at 20 °C

[cyclohexane], M	solvent	100[Cl ₂]/ [C ₆ H ₁₂]	polychlorinated product, % ^a
9.2	none	10	6
1.0	CCl₄	9.7	29
0.10	CCl4	10	53
0.030	CCl ₄	11	56
3.0	CCl₄	32	20
1.0	CCl₄	30	33, 33
0.30	CCl₄	32	51
0.10	CCl4	31	57, 59
0.050	CCl4	30	63, 64
0.020	CCl4	33	63, 64, 68
1.0	CFCl ₃	34	33
0.10	CFCl	31	58
0.020	CFCl ₃	33	63, 68, 70

^a Polychlorides \times 100 \div \sum monochlorides + polychlorides.

Low-conversion photochlorinations of 0.10 M neopentane or 0.10 M isobutane in inert solvents similarly result in formation of unexpectedly large amounts of the dichlorides.

But, if benzene is used as the solvent, in place of CCl₄ or CFCl₃, the anomalous behavior is not observed: Chlorination of 0.10 M cyclohexane or 0.10 M DMB results in nearly exclusive formation of monochlorides in excellent yields, based on Cl₂.

In contrast, "normal" behavior is observed with methyl chloride, methylene chloride, ethyl chloride, and 1,1-dichloroethane under all conditions. For example, chlorination of 1000 μ mol of methyl chloride (0.10 M) in CFCl₃ solvent (10-mL solution) with 103 μ mol of Cl₂ (0.01 M) produces 78 μ mol of CH₂Cl₂. Similar experiments (1) with methylene chloride in CFCl₃ produce 83 μ mol of CHCl₃ and (2) with ethyl chloride in CFCl₃ produce 90 μ mol of C₂H₄Cl₂'s. With these compounds similar results are obtained at high concentrations of substrate. There is no concentration effect with these substrates such as that observed with alkanes.

These results are understandable if one recognizes a consequence of attributing an encounter-controlled rate constant to the reaction of Cl. with an alkane.^{1b} The alkyl radicals react with Cl₂ to produce the alkyl chloride and a chlorine atom. These geminate reaction products are in the same solvent cage and react by transfer of a hydrogen atom from the alkyl chloride to the Cl, thus leading ultimately to the dihalides etc.

$$c-C_6H_{12} + Cl \rightarrow HCl + c-C_6H_{11}.$$
 (1)

$$c - C_{\varepsilon} H_{11} + C l_{2} \rightarrow [c - C_{\varepsilon} H_{11} C l_{1} + C l_{1}]_{\varepsilon}$$

$$(2)$$

$$[c-C_6H_{11}Cl + Cl\cdot]_c \rightarrow HCl + c-\dot{C}_6H_{10}Cl \qquad (3)$$

$$[c-C_6H_{11}Cl + Cl\cdot]_c \xrightarrow{\text{diff}} c-C_6H_{11}Cl + Cl\cdot$$
(4)

Reaction of the cage partners accounts for the multiple chlorinations; escape of Cl. from the cage leads to monochlorides. This cage effect should be most notable at low concentrations of alkane in relatively inert solvents. With the alkane as the solvent, cage "walls" would also consist of alkane molecules which react faster than the alkyl chloride with the chlorine atom, thus giving a normal product distribution. The same result is obtained with any solvent for which the rate of reaction with Cl. is encounter controlled, as, for example, benzene.²

With low reactivity substrates, for which the rate constants for reaction with Cl- are substantially below the cage-escape rate, multiple consecutive cage chlorinations should not be important pathways. Substrates such as CH₂Cl₂ and CHCl₃ react with chlorine atoms at rates 1-2 orders of magnitude less than those of primary C-H's of alkanes,^{1a,3} thus explaining the failure to

observe the anomalous behavior with the substrates CH₃Cl, CH_2Cl_2 , etc.

One can predict this anomalous effect, resulting from reactions of the geminate pair, will be observed when (1) the solvent is relatively inert, (2) the substrate concentration is low enough to make improbable that substrate and chlorination product molecules will be found in the same cage, and (3) the rate of reaction of the geminate pair is equal to or greater than the rate of cage escape of the smallest member of the pair.

We are seeking other instances of this "anomaly".

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Registry No. DMB, 79-29-8; cyclohexane, 110-82-7; methyl chloride, 74-87-3; methylene chloride, 75-09-2; ethyl chloride, 75-00-3.

Electrochemistry of Polymer Films Not Immersed in Solution: Electron Transfer on an Ion Budget

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This laboratory recently described¹ steady-state electron conduction through submicron films of electroactive polymeric transition-metal complexes sandwiched between two electrodes. Complexes like $[Os(bpy)_2(vpy)_2](ClO_4)_2$ were electropolymerized^{2,3} onto polished Pt and then overlaid with a porous film of evaporated Au, which was contacted by an electrolyte solution containing reference and auxillary electrodes. In this paper, we show that voltammograms with large limiting currents can be obtained for similarly prepared¹ $Pt/poly[Os(bpy)_2(vpy)_2]$ - $(ClO_4)_2$ /Au sandwiches in the absence of an electrolyte solution, bathed only in acetonitrile vapor or dry N_2 gas.

The essential features of the previous¹ sandwich voltammetry in electrolyte solution are summarized in Figure 1A,B. Figure 1A is a cyclic voltammogram where only E_{Pt} is controlled (vs. SSCE) and shows waves for the Os(III/II), Os(II/I), and Os(I/0)(formal) couples. In Figure 1B, where both E_{Pt} and E_{Au} are controlled (vs. SSCE), E_{Au} at 0 V and E_{Pt} being varied, a steady-state current-potential wave appears when $E_{\rm Pt}$ passes each region of film electroactivity. In the wave at positive E_{Pt} , for instance, the limiting current $(i_{111/11})$ means that all of the polymer next to the Pt electrode is Os(III) and all that next t o the Au electrode is Os(II), with linear concentration gradients of Os(III) and Os(II) states in the interior of the film as in Figure 1B inset. This $i_{\rm HI}/_{\rm H}$ limiting current is controlled by the rate of electron

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an ionic impurity the limiting currents were less reproducible upon successive potential scans.

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