Absolute Rate Expressions for Hydrogen Atom Abstraction from Molybdenum Hydrides by Carbon-Centered Radicals

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Abstract: A new family of basis rate expressions for hydrogen atom abstraction by primary, secondary, and tertiary alkyl radicals in dodecane and benzyl radical in benzene from the molybdenum hydride Cp*Mo-(CO)3H and for reactions of a primary alkyl radical with CpMo(CO)3H in dodecane are reported (Cp* = η5-pentamethylcyclopentadienyl, Cp = η5-cyclopentadienyl). Rate expressions for reaction of primary, secondary, and tertiary radical clocks with Cp*Mo(CO)3H were as follow: for hex-5-enyl, log(k/M^−1 s^−1) = (9.27 ± 0.13) − (1.36 ± 0.22)/θ, θ = 2.303RT kcal/mol; for hept-6-en-2-yl, log(k/M^−1 s^−1) = (9.12 ± 0.42) − (1.91 ± 0.74)/θ; and for 2-methylhept-6-en-2-yl, log(k/M^−1 s^−1) = (9.36 ± 0.18) − (3.19 ± 0.30)/θ (errors are 2σ). Hydrogen atom abstraction from CpMo(CO)3H by hex-5-enyl is described by log(k/M^−1 s^−1) = (9.53 ± 0.34) − (1.24 ± 0.62)/θ. Relative rate constants for 1°, 2°, 3° alkyl radicals were found to be 26:7:1 at 298 K. Benzyl radical was found to react 1.4 times faster than tertiary alkyl radical. The much higher selectivities for Cp*Mo-(CO)3H than those observed for main group hydrides (Bu3SnH, PhSeH, PhSH) with alkyl radicals, together with the very fast benzyl hydrogen-transfer rate, suggest the relative unimportance of simple enthalpic effects and the dominance of steric effects for the early transition-state hydrogen transfers. Hydrogen abstraction from Cp*Mo(CO)3H by benzyl radicals is described by log(k/M^−1 s^−1) = (8.89 ± 0.22) − (2.31 ± 0.33)/θ.

Introduction

Transition metal hydrides are particularly important chemical species because of their role in catalytic and stoichiometric hydrogenation reactions.1−4 Metal−hydrogen bond strengths,5,6 hydrogen atom7,11 and hydride-transfer reactions,12−14 and radical-mediated reduction of olefins via transition metal hydrides have been the subject of many studies.15−19 Although individual rate constants for reaction of alkyl radicals with several transition metal hydrides have been reported,7,8,11 Arrhenius expressions have been reported only for abstraction reactions of the trityl radical with organometallic hydrides.9,10 Whereas main group tin and silicon hydrides have been widely used in kinetic measurements and organic synthesis,20−22 transition metal hydrides have received relatively little attention as reagents for kinetic measurements and in free-radical-based synthetic transformations. Rate expressions for hydrogen abstraction from the metal hydride by organic free radicals and for the corresponding reactions of the resulting metal-centered radicals with halides and other groups provide basis rates for design and control of carbon−carbon and carbon−heteroatom bond formation in radical-based synthetic reactions.23,24 Thus, we present rate expressions for reactions of primary, secondary, and tertiary radical clocks and benzyl radical with molybdenum hydrides. We show that, unlike the less exothermic, but completely unselective, reactions of thiophenol and phenylselenol with primary, secondary, and tertiary alkyl radicals,25,26

1 Pacific Northwest National Laboratory.
2 Norfolk State University.
19 Connolly, J. W. Organometallics 1984, 3, 1333−1337.
21 Ungvary, F.; Marko, L. Organometallics 1982, 1, 1120−1125.

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Cp*Mo(CO)₃H exhibits appreciable selectivity in abstraction reactions, reflecting a trend, primary > secondary > benzyl > tertiary, that is predominately steric, rather than enthalpic, in origin. The present kinetic results offer a new family of basis rate expressions for hydrogen-transfer reactions of alkyl and benzyl radicals for use in competition kinetics and the design of radical-based synthetic strategies.

Results

Alkyl Radical Clock Kinetics. The rates of hydrogen atom abstraction from Cp*Mo(CO)₃H and CpMo(CO)₃H by alkyl radicals were determined using the appropriate alkyl radical clocks (Scheme 1). For primary and tertiary clock radicals, an integrated rate expression which accounts for changing hydride concentration was employed to determine the relative rate constants for radical clock rearrangement vs abstraction, \( k_{rel} \), where \( k_{rel} = k_{abs}k_{abs} \). The abstraction rate constants, \( k_{abs} \), were determined from ratios of concentrations of unrearranged, reduced hydrocarbon to rearranged hydrocarbon using values of \( k_{rel} \) from Arrhenius rate expressions for the clock reactions available in the literature. Both alkyl bromides and tert-butyl peresters of alkenoic acids were used to generate the desired primary and tertiary radicals in reactions with Cp*Mo(CO)₃H (Scheme 2). The ring-unsubstituted hydride CpMo(CO)₃H was found to react rapidly at room temperature with adventitious hydrogen donor and \( k_{abs} \) is an adventitious hydrogen donor and \( k_{abs} \) is the associated rate constant for abstraction. A near-zero intercept \( k' \) indicates the lack of significant hydrogen sources in the solvent, or other alternate sources of the unreacted hydrocarbon not proportional to the primary donor concentration.

The Arrhenius plots of data for the reaction of Cp*Mo(CO)₃H with primary, secondary, and tertiary alkyl radicals are presented in Figure 2. These plots include results from both alkyl bromide and perester radical precursors for the primary and tertiary radicals, while the secondary radical rate was determined using the bromide clock precursor. The Arrhenius and Eyring parameters for hydrogen abstraction from Cp*Mo(CO)₃H by alkyl radicals are presented in Table 1.

Both perester and bromide precursors were used over the entire temperature range and gave identical results within statistical error limits. Unlike the rapid iodine atom-exchange reactions between radicals and organic iodides, which may...
Table 1. Arrhenius* and Eyring Parameters for the Reaction of Primary, Secondary, and Tertiary Radical Clocks and Benzyl Radicals with Cp*Mo(CO)_3H and Primary Radicals with CpMo(CO)_3H in Dodecane

<table>
<thead>
<tr>
<th>radical</th>
<th>hydride</th>
<th>log A (M^(-1) s^-1)</th>
<th>(\Delta H^0) (kcal/mol)</th>
<th>k (M^(-1) s^-1)</th>
<th>temp range (K)</th>
<th>mean temp (T_{av}, K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>hex-5-enyl</td>
<td>Cp*Mo(CO)_3H</td>
<td>9.27 ± 0.13</td>
<td>-19.8 ± 1.7</td>
<td>1.36 ± 0.22</td>
<td>9.18 ± 10^6</td>
<td>309 ± 400</td>
</tr>
<tr>
<td>hept-6-en-2-yl</td>
<td>Cp*Mo(CO)_3H</td>
<td>9.12 ± 0.42</td>
<td>-19.3 ± 2.3</td>
<td>1.91 ± 0.74</td>
<td>5.2 ± 10^7</td>
<td>318 ± 458</td>
</tr>
<tr>
<td>2-methyl-hept-6-en-2-yl</td>
<td>Cp*Mo(CO)_3H</td>
<td>9.36 ± 0.18</td>
<td>-18.2 ± 1.7</td>
<td>3.19 ± 0.30</td>
<td>1.1 ± 10^7</td>
<td>343 ± 448</td>
</tr>
<tr>
<td>hex-5-enyl</td>
<td>CpMo(CO)_3H</td>
<td>9.53 ± 0.34</td>
<td>-17.5 ± 1.2</td>
<td>1.24 ± 0.62</td>
<td>4.2 ± 10^8</td>
<td>318 ± 458</td>
</tr>
<tr>
<td>benzyl</td>
<td>Cp*Mo(CO)_3H</td>
<td>8.89 ± 0.22</td>
<td>-20.0 ± 2.0</td>
<td>2.31 ± 0.33</td>
<td>1.5 ± 10^7</td>
<td>297 ± 373</td>
</tr>
</tbody>
</table>

* Eyring and Arrhenius parameters are related by \(\Delta H^0 = -R(\beta \ln(k_{av}/T)/\theta(1/2)) = E_r - RT_{av}\), where \(T_{av}\) is the mean temperature of the kinetics, and \(\Delta \Delta S = R(\ln(k_{av}/T_{av}) + AT_{av} = R \ln(b/b_0)T_{av}\), where \(k_r\) is Boltzmann’s constant, \(b\) is Planck’s constant, \(k_{av}\) is the rate constant from the Arrhenius expression at mean temperature, and \(e\) is the base of natural logarithms. * The radical cyclization rates used in this work are (ref 29): for hex-5-enyl, \(\log(k_{abs}(s^{-1})) = 10.42 - 6.85\theta/\theta(1/2)\); for hept-6-en-2-yl, cyclization to cis-2-methylcyclopentylmethyl is given by \(\log(k_{abs}(s^{-1})) = 9.79 - 6.50\theta\); for 2-methyl-hept-6-en-2-yl, \(\log(k_{abs}(s^{-1})) = 10.0 - 6.1\theta/\theta(1/2)\). * The reaction of benzyl radical with Cp*Mo(CO)_3H was carried out in benzene. Errors are 2σ.

compromise unrearranged/rearranged hydrocarbon ratios and affect their use in synthetic transformations, 31 bromine atom exchange between the alkylbenzyl bromides and the cyclized radical is unimportant, reflected in the present results, that show close agreement between perester and bromide precursors.

The Arrhenius plot for hydrogen abstraction from CpMo(CO)_3H by hex-5-enyl is also shown in Figure 2, and the rate expression for abstraction of hydrogen by the primary clock from CpMo(CO)_3H is presented in Table 1.

Benzyl Radical Kinetics. Rate constants for hydrogen abstraction by the benzyl radical from Cp*Mo(CO)_3H were determined by a competition of abstraction to form toluene with benzyl radical self-termination to produce bibenzyl during photolysis of dibenzyl ketone (DBK) in benzene solutions. The benzyl radical was produced by broadband visible UV irradiation (>330 nm) of optically dilute solutions of the ketone and hydride, as shown in Scheme 3. Figure 3 shows the time-dependent evolution of products during photolysis of a 0.02 M solution of DBK in benzene at 373 K.

Under conditions of constant rate of photolysis of dibenzyl ketone, the absence of photolysis of starting hydride, and no significant absorption of light by photoproducts, the relationship between toluene, bibenzyl (BB), and the initial hydride concentration, [MoH]_0, that accounts for consumption of MoH by given by eq 1a. At low conversion of hydride, the exponent term, \(k_{abs}(BB)^{(1/2)}k_{1/2}\) in eq 1a is small, and simplifies to eqs 1b and 1c, where [MoH]_0 represents the average concentration of the molybdenum hydride over a period of photolysis time, \(\Delta t\):

\[
\text{[toluene]} = [\text{MoH}]_0 \left( 1 - e^{-k_{abs}(BB)^{(1/2)}k_{1/2}} \right) \tag{1a}
\]

\[
\text{[toluene]} = [\text{MoH}]_0 \frac{k_{abs}(BB)^{(1/2)}k_{1/2}}{k_{1/2}} \tag{1b}
\]

\[
k_{abs} = \frac{[\text{toluene}]^{1/2}}{[\text{BB}]^{1/2}[\text{MoH}]_0 \Delta t^{1/2}} \tag{1c}
\]

In practice, conversion of hydride up to 35% leads to less than about 10% error in rate constant \(k_{abs}\) using eq 1c. In the present experiments, the yield of toluene was used to calculate an average hydride concentration at each photolysis time. Combining experimental values of bibenzyl and toluene at 0.5, 1.0, and 1.5 s photolysis time with an independent value of the rate constant for self-termination of benzyl, \(k_t\) (see below), provided values of \(k_{abs}\) at each temperature.

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Figure 3. Product evolution of toluene (■) and bibenzyl (♦) at 373 K at initial concentrations of [Cp*Mo(CO)_3H], 0.000105 M, and [dibenzy1 ketone], 0.02 M. Rate constants are determined using data at 0.5, 1.0, and 1.5 s and eq 1c. The solid lines are the predicted concentrations of toluene and bibenzyl from minimizing the sum of the squares of residuals for toluene and bibenzyl in a numerical integration of Scheme 3 using a rate constant for photolysis of DBK, 8.7 × 10^-6 M/s, \(k_t = 5.6k_{abs} = 6.2 × 10^9 M^{-1} s^{-1}\), giving \(k_{abs}\) = 34.4 × 10^7 M^-1 s^-1. The agreement of the two approaches is due to the near linearity of the product curves through about 50% conversion of hydride. All of the points of Figure 3 can be alternatively fit to eq 1a, allowing \(k_{abs}\) and the initial hydride concentration (MoH) to vary in a nonlinear least-squares optimization. This results in a predicted initial concentration of hydride of 9.13 × 10^-5 M and a rate constant \(k_{abs}\) = 3.46 × 10^7 M^-1 s^-1. Alternatively, the first three points (0.5, 1.0, and 1.5 s) can be fit to eq 1a. This results in values of \(k_{abs}\) identical to those resulting from an average of 0.5, 1.0, and 1.5 s points using eq 1c.

Scheme 3

\[
\text{PhCH}_2\text{C}=\text{CH}_2\text{Ph} \xrightarrow{k_{abs}} \text{PhCH}_2\text{C}=\text{CH}_2\text{Ph} + 2 \text{PhCH}_2 + \text{CO}
\]

\[
\text{PhCH}_2\text{•} + \text{Cp}^*\text{Mo(CO)}_3\text{H} \xrightarrow{k_{abs}} \text{PhCH}_3
\]

\[
\text{PhCH}_2\text{•} + \text{PhCH}_2\text{•} \xrightarrow{2k_c} \text{PhCH}_2\text{CH}_2\text{Ph}
\]

\[
2 \text{Cp}^*\text{Mo(CO)}_3\text{•} \xrightarrow{2k_c} \text{Cp}^*\text{Mo(CO)}_2\text{PhCH}_2\text{Ph}
\]

\[
\text{Cp}^*\text{Mo(CO)}_3\text{H} \xrightarrow{k_{abs}} \text{Products}
\]

The rate of photolysis of DBK employed in the kinetic measurements was in the range [DBK] × (7–9) × 10^-6 s^-1, corresponding to a steady-state concentration of benzyl radical in the range (2–5) × 10^-8 M. The rate of formation of toluene and bibenzyl was found to be approximately linear during the first 2 s of photolysis. The evolution of toluene and bibenzyl with time was examined by numerical integration of the...
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$\text{CpMo(CO)}_2 H$. 32 Photolysis of the dimer $[\text{CpMo(CO)}_3]_2$ involves CO loss and subsequent reactions of the 16-electron $\text{CpMo(CO)}_3 H$. 32 Photolysis of the dimer $[\text{CpMo(CO)}_3]_2$ results in CO loss, giving $\text{CpMo}_2(\text{CO})_4$ or metal-metal bond homolysis to form the radical $\text{CpMo} \cdot \text{(CO)}_2$. 33 In the benzylic radical experiments, a 1-kW high-pressure (Pyrex-filtered) xenon arc lamp was used. Those conditions led to a small extent of photolysis ($<10\%$) of the hydride during the kinetic experiments, whereas in the experiments utilizing halide radical clock precursors, the Pyrex-filtered, diffuse light of a 100-W long-wavelength sunlamp ($\lambda_{\text{max}} = 365 \text{ nm}$) was employed, which led to significant photolysis of the hydride only after prolonged photolysis. (In the halide radical clock precursor experiments, radical initiation most probably occurred via photolysis of small amounts of the dimer $[\text{Cp*Mo(CO)}_3]_2$ present in the hydride $\text{Cp*Mo(CO)}_3 H$.) Kinetic modeling of case 1, by reducing the initial hydride concentration, or of case 2, by adding an additional decay pathway in the numerical integration of the equations of Scheme 3 ($k_h$, in Scheme 3), produced an overall better fit of the data through complete consumption of hydride but poorer agreement with the experimental concentrations of hydride used and poorer agreement with initial rates of reaction calculated using eq 1c. Careful analysis of material balances revealed no more than $6-7\%$ decrease in the rate of photolysis of DBK over 5 s; thus, case 3 was found insignificant. Because of the small discrepancy at longer photolysis times, we preferred to use eq 1c for data obtained through typically 1.5 s, or less than one half-life of hydride conversion, to avoid ambiguities of modeling the system at high conversion. For times between 0.5 and 1.5 s, values of $k_{\text{dbn}}$ found by numerical integration (Scheme 3, neglecting $k_h$) were found to be in excellent agreement with the rates calculated with eq 1c from initial product ratios ($[\text{PhCH}_2]\text{H}/[\text{PhCH}_2\text{CHCH}_2\text{Ph}]^{1/2}$) when the kinetic simulation employs $k_{\text{dbn}}$ as the only variable and values of $k_h$, rate of photolysis of DBK, and hydride concentration are experimental constants. The Arrhenius plot for reaction of benzylic radical with $\text{Cp*Mo(CO)}_3 H$ is shown in Figure 4, and the rate expression is presented in Table 1.

Discussion

Alkyl and Benzylic Radical Selectivities in Hydrogen Atom Abstractions of Molybdenum Hydrides. The rate constant for hydrogen atom abstraction by a primary radical from $\text{Cp*Mo(CO)}_3 H$ at 298 K ($1.9 \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$) is similar in magnitude to rates of abstraction of alkyl radicals from thiophenol: e.g., for the reaction of n-butyl radical with thiophenol, $k_{\text{dbn}} = 1.4 \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$ and is about an order of magnitude less than that for phenylselenol. 25,26 Thiophenol and phenylselenol have the useful properties (as kinetic radical trapping agents) of exhibiting, within experimental error, no selectivity between tertiary, secondary, and primary alkyl radicals. Relative rates for tertiary, secondary, and primary and benzylic radicals are shown in Table 2 for thiophenol, 25 phenylselenol, 26 and tributylstannane. The $\text{S}^\cdot \text{H}$ and $\text{Se}^\cdot \text{H}$ bond strengths of thiophenol and phenylselenol are 80 $\pm$ 2 kcal/mol 34-36 and 78 $\pm$ 2 kcal/mol, 37 respectively. The $\text{M}^\cdot \text{H}$ bond strengths of $\text{Cp*Mo(CO)}_3 H$ and $\text{CpMo(CO)}_3 H$ are 69 and 70 kcal/mol, respectively. 5,6,38 Table 2 illustrates that the relative magnitudes of rate constants and activation barriers for highly exothermic hydrogen abstraction reactions are not related by simple enthalpy considerations. For main group hydrides, stabilized radicals react more slowly than simple alkyl radicals. Thus, benzyl radical abstracts hydrogen atom about 2–300 times more slowly from PhSH and Bu3SnH than alkyl radicals, reflecting the trend in $\text{C}^\cdot \text{H}$ bond dissociation energies for primary (1-propyl $\text{C}^\cdot \text{H}$), secondary (2-propyl $\text{C}^\cdot \text{H}$), tertiary ($\text{tert}$-butyl $\text{C}^\cdot \text{H}$), and benzyl (toluene $\text{C}^\cdot \text{H}$) radicals: 101, 99, 95, and 89 kcal/mol, respectively. 39-41 Thus, the selectivity of alkyl radicals in the much more highly exothermic reactions of $\text{Cp*Mo(CO)}_3 H$ with alkyl radicals is quite remarkable compared to the selectivity in reactions of primary, secondary, and tertiary alkyl radicals with PhSH, Bu3SnH, and PhSeH. The relative reactivity of benzyl radical with $\text{Cp*Mo(CO)}_3 H$ is also surprising by comparison with the main-group hydrides, exhibiting a rate constant higher than that of tertiary alkyl radical. Since the benzylic radical is slower than the secondary alkyl radical, enthalpic effects are almost, but not completely, removed. The results of Table 2 suggest that the selectivity of the hydrogen abstraction reactions of $\text{Cp*Mo(CO)}_3 H$ can be attributed in significant extent to metal ligand/radical steric effects. In comparing the results of thiophenol and $\text{Cp*Mo(CO)}_3 H$ in Table 2, it will be noted that the reaction of benzyl radical with $\text{Cp*Mo(CO)}_3 H (\Delta H^\circ \approx -20 \text{ kcal/mol})$ is similar in exothermicity to the reaction of secondary (\Delta H^\circ \approx -21 \text{ kcal/mol}) or tertiary radical (\Delta H^\circ \approx -19 \text{ kcal/mol}) with PhSH.

Several examples of the steric effects of pentamethyl substitution of the cyclopentadiene group in homolytic displacement and atom-transfer chemistry can be cited. In the present work, (34) Colussi, A. J.; Benson, S. W. Int. J. Chem. Kinet. 1977, 9, 925.
CpMo(CO)_3H reacts an additional factor of 1.9 faster than Cp*Mo(CO)_3H with primary radicals. Pentamethyl substitution on the peresters used in this study, tert-butyl perhept-6-enoate, were found to undergo reaction with CpMo(CO)_3H decreased with temperature (°C), the rate of unimolecular decomposition of the tertiary and secondary peresters is sufficiently fast to provide ample radical production compared to the induced decomposition pathway of reduction by Cp*Mo(CO)_3H. Note that the site of the reaction of molybdenum radicals, carbonyl or peroxodic oxygen, is not specified in Figure 4. Although CpMo(CO)$_2$ radical is known to react at the carbonyl groups of quinones, reactions of alkyl and other radicals with peresters have been established to occur at the peroxidic oxygen. Whether attack occurs at carbonyl or peroxodic oxygen, the efficiency of the induced decomposition pathway suggests that pyridinethiocarbonylic acid esters, employed as convenient radical precursors, will also be useful as radical precursors in conjunction with the molybdenum hydrides.

### Conclusions

A new family of rate expressions using transition metal hydrides has been developed for use in competition kinetics and radical-based synthetic transformations, and for estimation of the efficiency of the induced decomposition pathway of reduction by Cp*Mo(CO)$_2$H. Note that the site of the reaction of molybdenum radicals, carbonyl or peroxodic oxygen, is not specified in Figure 4. Although CpMo(CO)$_2$ radical is known to react at the carbonyl groups of quinones, reactions of alkyl and other radicals with peresters have been established to occur at the peroxidic oxygen. Whether attack occurs at carbonyl or peroxodic oxygen, the efficiency of the induced decomposition pathway suggests that pyridinethiocarbonylic acid esters, employed as convenient radical precursors, will also be useful as radical precursors in conjunction with the molybdenum hydrides.

### Scheme 4

![Scheme 4](image-url)

![Scheme 4](image-url)
of fundamental steps in hydrogen transfer involving the molybdenum hydride group in the context of catalysis. The rate constants for hydrogen abstraction from Cp*Mo(CO)₃H and CpMo(CO)₃H by alkyl radicals, and the efficiency of induced decomposition reactions of the corresponding molybdenum radicals Cp*Mo(CO)₃ and CpMo(CO)₃⁺ in reactions with peresters, have been shown to be influenced by the steric bulk of the metal ligands and the α-alkyl substituents on the radical center, or by the effects of α-alkyl substitution of the perester. The influence of steric interactions on the rates of highly exothermic reactions of the hydrides with carbon-centered radicals is generally recognized.¹⁰

**Experimental Section**

CpMo(CO)₃H.²² CP*Mo(CO)₃H.²² 6-bromohexene.⁵¹ 2,2-dimethylpent-6-enoic acid in CH₂Cl₂, to give the acid chloride, were similarly prepared as clear liquids in 25% yield. tert-Butyl 2,2-dimethylperhept-6-enoate. The acid chloride was produced by refluxing 2,2-dimethylhept-6-enoic acid with excess SOCl₂ in CH₂Cl₂, to give the acid chloride, which was converted to the perester by reaction with tert-butyl hydroperoxide in CH₂Cl₂ at 0°C. Chromatography of the crude perester on silica gel using 2.5% methyl tert-butyl ether in pentane gave a fraction containing pure liquid perester, 0.48 g (17% yield).

**Kinetic Procedure for Alkyl Radical Clock Kinetics.** Solutions of halide or perester radical precursor, hydride, and internal GC standard were freeze-dried, sealed, and heated through four half-lives of the peresters or heated and photoinitiated (halide precursors) with a tungsten filament lamp. The usual integrated rate expression, with the Spernol correction factor for consumption of hydride from the induced decomposition (eqs 5a,b, etc.), was calculated using eq 6, which was in near-perfect agreement with experimental values (refs 61, 62).

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\[ \text{RCO₂Bu-t + 2CPMo(CO)₃H} \rightarrow \]
\[ \text{R-H} + \text{t-BuOH} + \text{CO₂} + [\text{CPMo(CO)₃}]_2 \] (4)

\[ \text{RCO₂Bu-t + CP*Mo(CO)₃O₂CR + r-BuO} \] (5a)

\[ \text{r-BuO} + \text{CP*Mo(CO)₃H} \rightarrow \text{r-BuOH} + \text{CP*Mo(CO)} \] (5b)

Product concentrations are corrected for temperature-dependent density changes.⁵⁹

**Rate Constants for Reaction of Benzyl Radical with CPMo(CO)₃H**. Pyrex reaction tubes containing solutions of 0.02 M DBK, 10⁻⁴ M CP*Mo(CO)₃H, and 10⁻³ M tert-butybenzene were freeze-dried, sealed, and photolyzed with a water-filtered 1-kW high-pressure xenon arc lamp for controlled periods of time (0.50–5.00 ± 0.005 s) using a Uniblitz model 22S0AT0T522952 computer-controlled electronic shutter. Reagent concentrations were corrected for solvent vapor pressure and density.⁶⁰ Rate constants for hydrogen abstraction were calculated using eq 1 at 0.5–1.5 s photolysis times, using the measured product concentrations, the calculated average molybdenum hydride donor concentration, and the benzyl self-radical termination rate, kd, calculated from the expression ln[2k₂M⁻¹s⁻¹] = 27.23 – 2952k₂RT. This expression was calculated using the von Smoluchowski equation modified, eq 6, using the Spernol–Wirtz modification of the Debye–Einstein equation, eq 7, where f in eq 7 is the SW microfriction factor.⁶¹

\[ 2k_d = \frac{(8\pi/1000)\sigma D_{AB}N}{(6)} \]
\[ D_{AB} = kT/6\pi\eta r_f \]

Fischer and co-workers have measured self-termination rates for benzyl radical in cyclohexane, ln(2k₂M⁻¹s⁻¹) = 26.2 – 2497.6RT, and in toluene, ln(2k₂M⁻¹s⁻¹) = 27.05 – 2797.4RT. For self-termination of benzyl in hexane, we developed the expression ln(2k₂M⁻¹s⁻¹) = 26.0 – 1803.6RT.₆₅ Parameters for estimation of the rate expression employed in these kinetics are presented.⁶⁶ The pathway of the peresters forming tert-butyl alcohol. For halide radical precursors, f = 1. For peresters, 2 equiv of hydride is consumed for each event of unimolecular perester decomposition (eq 4), or 1 equiv of hydride per chain-reaction-induced decomposition (eqs 5a,b, etc.).
absolute error in estimation of self-termination rate constants by this method is estimated to be less than ca. 25%\textsuperscript{61}.

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**Supporting Information Available:** Tabulations of kinetic data for the entries in Table 1 (PDF). This material is available free of charge via the Internet at http://pubs.acs.org.