

# Generation of Halomethyl Radicals by Halogen Atom Abstraction and Their Addition Reactions with Alkenes

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Supporting Information

**ABSTRACT:**  $\alpha$ -Aminoradicals undergo halogen atom abstraction to form halomethyl radicals in reactions initiated by the combination of *tert*-butyl hydroperoxide, aliphatic trialkylamine, halocarbon, and copper(I) iodide. The formation of the  $\alpha$ -aminoradical circumvents preferential hydrogen atom transfer in favor of halogen atom transfer, thereby releasing the

halomethyl radical for addition to alkenes. The resulting radical addition products add the *tert*-butylperoxy group to form  $\alpha$ -peroxy- $\beta$ , $\beta$ -dichloropropylbenzene products that are convertible to their corresponding  $\beta$ , $\beta$ -dichloro-alcohols and to novel pyridine derivatives. Computational analysis clearly explains the deviation from traditional HAT of chloroform and also establishes formal oxidative addition/reductive elimination as the lowest energy pathway.

#### ■ INTRODUCTION

Radical addition reactions of alkenes with dissociable molecular species (X—Y) provide convenient access to vicinal disubstituted products with increased molecular and functional complexity (Figure 1a). The initiation of these reactions effects atom transfer that forms radical Y, which adds to the carbon—carbon double bond to form a carbon radical that abstracts X from X—Y to enter the catalytic cycle. In one of the most

**Figure 1.** (a) General approach to functionalize olefins by dissociable molecular species. (b) Kharasch's first examples of halogen atom abstraction from carbon tetrachloride and HAT from chloroform. (c) Selective chlorine atom abstraction from chloroform to form α-peroxy- $\beta$ , $\beta$ -dichloropropylbenzenes.

notable examples of these reactions, Kharasch et al. reported in 1945 the acetyl peroxide-induced Markovnikov addition of Cl and CCl<sub>3</sub> from CCl<sub>4</sub> to alkenes. The Kharasch reaction has been extensively investigated, and numerous examples of atom-transfer radical addition using transition-metal catalysts to initiate CCl<sub>3</sub> radical generation have been reported.<sup>4-</sup> However, when this process was applied with chloroform (Figure 1b), the initiation step generated the CCl<sub>3</sub> radical by H-atom transfer (HAT), rather than forming the dichloromethyl radical by chlorine-atom transfer. In fact, the dichloromethyl radical can be generated by a HAT from CH<sub>2</sub>Cl<sub>2</sub>, albeit indirectly via an aryl radical formed from a boronic acid or an aryldiazonium salt, and a catalytic methodology has been reported to utilize the chloroform analogue CHCl<sub>2</sub>Br in the presence of stoichiometric amounts of tri- and tetra-tertiary amines as an alternative. Transitionmetal compounds, especially those of low valent platinum, 10 are well-known to undergo oxidative addition with halocarbon solvents, but there has not been a report of a catalytic process through which the dichloromethyl radical has been generated from chloroform. We wish to report an effective indirect methodology for preferential chlorine-atom abstraction from chloroform and its application to the regioselective addition of CHCl<sub>2</sub> and the tert-butylperoxy radical to styrenes, resulting in  $\alpha$ -peroxy- $\beta$ , $\beta$ -dichloropropylbenzene products in moderate-tohigh yields (Figure 1c), and their conversion to the corresponding  $\beta$ , $\beta$ -dichloro-alcohols and novel pyridine

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derivatives. Experimental data along with density functional theory (DFT) calculations demonstrate the essential role of the  $\alpha$ -aminoradical that changes the preference of atom extraction from CHCl<sub>3</sub>. An excellent correlation that is found to be valuable for selecting appropriate radical and a general mechanism are also proposed.

#### ■ RESULTS AND DISCUSSION

We envisioned that generation of the dichloromethyl radical from chloroform would be a function of the relative energies for C-Cl versus C-H homolytic bond cleavage from various radical initiators. Those atom-transfer agents that have been reported (\*BuOO, \*BuO, OH, and Ar) favor C-H bond cleavage, 8a,11 but this list is limited. 12 As shown in Scheme 1,

Scheme 1. Different Polar Effects of Atom-Transfer Reaction

$$R^{\cdot} + CHCI_{3}$$

$$\begin{bmatrix} 8^{+} & 8^{-} \\ [R^{---}H^{--}CCI_{3}]^{\dagger} & R^{-}CI + \dot{C}CI_{3} \end{bmatrix}$$

$$\begin{bmatrix} 8^{+} & 8^{-} \\ [R^{---}CI^{--}CHCI_{2}]^{\dagger} & R^{-}H + \dot{C}HCI_{2} \end{bmatrix}$$

we expected that reversing radical polarity would lead to free radical species that could favor C-Cl bond cleavage and thereby provide a viable pathway for the generation of halogenated carbon radicals.

A search for appropriate radicals was initiated with examining adiabatic ionization potentials (IPs) of carbon radicals. 13 The calculated IPs and differences in free energy barriers of hydrogen-atom transfer (HAT) versus chlorineatom transfer (CAT)  $(\Delta G_H^{\neq} - \Delta G_{Cl}^{\neq})$  are listed in Table 1 and plotted in Figure 2.

Indeed, the IPs indicate the potential of releasing an electron and a good correlation between IP and the preference of HAT versus CAT was found. Results in Table 1 and Figure 2 suggested that  $\alpha$ -amino radicals would favor C-Cl homolytic cleavage to generate the dichloromethyl radical. <sup>14</sup> More specifically,  $\alpha$ -amino radicals were indicated to have the highest potential to undergo chlorine-atom abstraction with energy differences greater than those of  $\alpha$ -alkoxy carbon radicals (Table 1, entries 1–6). Alkyl,  $\alpha$ -carbonyl, and phenyl radicals, in contrast, have a higher preference for C-H bond homolytic cleavage (Table 1, entries 7-9).

On this basis, we searched for those reactants that preferentially form an intermediate radical capable of undergoing chlorine-atom abstraction from chloroform to produce the CHCl<sub>2</sub> radical. 15 Tertiary amines were surveyed, but with the caveat that generation of  $\alpha$ -amino radicals is subject to the oxidation potential of the amine. Specifically, electron transfer (ET) is the initial step in the catalytic oxidation of N,Ndialkylanilines to their corresponding iminium ions by tertbutyl hydroperoxide (TBHP), 16 whereas hydrogen atom transfer (HAT) occurs preferentially as the initial step in the similar oxidation of tetrahydroquinolines.<sup>1</sup>

Because of their demonstrated suitability for free radical addition reactions, styrenes were selected as the target substrates. 18 Reactions were performed in various solvents containing excess chloroform, amines capable of HAT, and selected catalysts known to form the tert-butylperoxy radical. Table 2 provides an abbreviated selection of various screening parameters (see Supporting Information for full screening

Table 1. Calculated Gibbs Free Energy Barrier  $(\Delta G_{x}^{\neq})/$ Energy Differences  $(\Delta G_{\text{H}}^{\neq} - \Delta G_{\text{Cl}}^{\neq})$  of Various Radicals

 $\textbf{R'+ H-CX}_3 \xrightarrow{\Delta G_{\textbf{H}}^{\neq}} \textbf{R-H+CX}_3 \ \, \textbf{R'+ X-CHX}_2 \xrightarrow{\Delta G_{\textbf{X}}^{\neq}} \textbf{R-X+CHX}_2$  $\Delta G_{H}^{\neq}$  (kcal/mol) for free energy barriers of hydrogen atom transfer  $\Delta G_{X}^{\neq}$  (kcal/mol) for free energy barriers of halogen atom transfer

Entry	Radical	IP	$\Delta G_{\text{H}^{\#}}$	$\Delta G_{Cl}^{\sharp}$	$\Delta G_{\text{H}}{}^{\!$
1	N	4.7	17.4	9.2	8.2
2	N	4.9	12.9	6.7	6.2
3	N <sub>Ph</sub>	5.3	24.2	17.9	6.3
4	N	5.1	12.9	7.8	5.1
5	N	5.6	14.7	12.8	1.9
6	<b>5</b>	7	14.6	13.6	1.0
7	•	7.1	13.3	15.8	-2.5
8		8.3	17.2	21.9	-4.7
9	<b>5</b>	8.1	5.4	10.7	-5.3

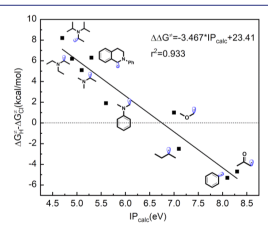


Figure 2. Correlation between IPs and the difference in free energy barrier of HAT vs CAT.

parameters). As expected, trialkylamine bases performed well under the reaction conditions in the presence of Rh<sub>2</sub>(cap)<sub>4</sub> (Table 2, entries 1-3). When 3° benzylamines and anilines were used, no conversion of the styrene reactant to product was observed (Table 2, entries 4 and 5). Rh<sub>2</sub>(OAc)<sub>4</sub> was ineffective at room temperature or 65 °C, and silver-based catalysts showed moderate product formation (Table 2, entries 7 and 8). However, use of CuI gave nearly quantitative formation of the desired product (Table 2, entry 9). No significant drop in yield was observed at ambient temperature,

Table 2. Optimization of Reaction Conditions for Dichloromethyl Radical Addition<sup>a</sup>

	Cat. (1 mol		Me 2a	
Me 1a	TBHP (5.0 eq.), Ami CHCl <sub>3</sub> , Ten			
entry	amine	catalyst	yield <sup>b</sup>	
1	NEt <sub>3</sub>	$Rh_2(cap)_4$	71%	
2	$NCy_2Me$	$Rh_2(cap)_4$	71%	
3	DIPEA	$Rh_2(cap)_4$	87%	
4	$N(Bn)_3$	$Rh_2(cap)_4$	0%	
5	PhNMe <sub>2</sub>	$Rh_2(cap)_4$	0%	
6	DIPEA	Rh <sub>2</sub> OAc <sub>4</sub>	0%	
7	DIPEA	AgOTf	55%	
8	DIPEA	AgOAc	61%	
9	DIPEA	CuI	97%	
$10^c$	DIPEA	CuI	95%	
11 <sup>c</sup>	NEt <sub>3</sub>	CuI	62%	
12 <sup>c</sup>	NCy <sub>2</sub> Me	CuI	67%	
13 <sup>c</sup>	$N(Hex)_3$	CuI	36%	
14 <sup>c</sup>	DIPEA	CuOAc	74%	
15 <sup>c</sup>	DIPEA	CuBr	82%	
16 <sup>c</sup>	DIPEA	Cu(ACN) <sub>4</sub> PF <sub>6</sub>	82%	
$17^c$	DIPEA		0%	
18 <sup>c</sup>		CuI	0%	

<sup>a</sup>Reaction conditions: unless indicated otherwise, the reaction of 1a (1.0 mmol), catalyst (1 mol %), TBHP (5.0 equiv), and amine (5.0 equiv) was carried out in CHCl $_3$  (5.0 mL) at 65 °C.  $^b$ Isolated yield. <sup>c</sup>CHCl<sub>3</sub>:acetone were used in a 1:1 ratio as cosolvents (5.0 mL) at rt.

although the low solubility of CuI in organic solvents led to inconsistent initiation/yields. To overcome this problem, we used acetone as a cosolvent, and this change provided high reproducibility to these reactions (Table 2, entry 10). Furthermore, screening of the amine and the copper catalyst showed that the DIPEA and CuI provide the best result (Table 2, entries 11–16). No product could be detected without both 3° aliphatic amine and CuI (Table 2, entries 17 and 18, and Supporting Information). 18

The optimum catalyst, amine, and reaction conditions were applied to substituted styrenes, and the results obtained showed generality for this transformation (Scheme 2). Styrenes with both electron-rich and electron-deficient substituents at the para-, meta-, and ortho-positions of the phenyl moiety furnished product in moderate-to-high yields (2a-2k). Furthermore, substitution at the  $\alpha$ -position of styrene also gave product with good yields (2l-2m), and 1-vinylnaphthalene and 2-vinylnaphthalene were also compatible with this catalytic process (2n-2o). In reactions with vicinal disubstituted alkenes, however, an additional 3.0 equiv of TBHP were necessary to obtain higher yields (2p-2q). Cumyl peroxide was also compatible under the reaction conditions (2r). However, di-tert-butylperoxide (DTBP) and benzoyl peroxide were unreactive.

Other perhalogenated compounds that could undergo hydrogen-atom abstraction were also found to be selective for halogen-atom abstraction (Table 3, except entries 1, 5, and 6). Several methodologies undergo similar addition of alkyl halides but oftentimes are limited to substrates wherein  $\alpha$ halogens are excluded. 19 Under our standard reaction conditions where mixed halogenated compounds were used, selectivity for I > Br > Cl > F removal was observed (Table 3,

Table 3. Substrate Scope of Alkyl Halide for Addition to Aryl Alkenes<sup>a</sup>

Entry	Alkyl Halide	Product	Yield <sup>b</sup>
1	Br Br	'BuOO Br F F	81%
2	I CF <sub>3</sub>	'BuOO CF <sub>3</sub>	83%
3	Br Cl H Cl	'BuOO CI	53%
4	Br OEt	<sup>1</sup> BuOO CO <sub>2</sub> Et	69%
5	Br F O OEt	'BuOO F F CO <sub>2</sub> Et	71%
6	CICI	'BuOO CI CI CI CI See See See See See See See See See Se	91%
7	Br Br	'BuOO Br Br Me 3f	73%
8	H	<sup>'BuOO</sup> 3g	43%

<sup>a</sup>Reaction conditions: Unless indicated otherwise, the reaction of 1 (1.0 mmol), R-X (5.0 equiv), CuI (1 mol %), TBHP (5.0 equiv), and DIPEA (5.0 equiv) was carried out in acetone (5.0 mL) at rt. <sup>b</sup>Isolated yields.

entries 1-3) in moderate-to-high yields. Carbonyl substrates containing  $\alpha$ -halogenated esters also underwent addition in good yields (Table 3, entries 4 and 5). Finally, while reactivity remained high for chloroform derivatives and bromoform (Table 3, entry 7) and for chlorine-atom transfer from tetrachloromethane (Table 3, entry 6), iodoform (Table 3, entry 8) consistently led to lower conversions, possibly because of subsequent reactions.

Moreover, the dichloromethyl addition reaction could be carried out on a gram scale with high product yield using only 0.025 mol % CuI (eq 1). Considering that the reaction does

not give 2a without the CuI catalyst (Table 2, entry 17), we conclude that CuI is very efficient but also essential. Furthermore, without the  $3^{\circ}$  aliphatic amine the reaction takes an entirely different course.

The synthetic utility of the peroxide products was studied next.<sup>20</sup> Different from "Russell mechanism" where the alcohol is a derived in a 1:1 mixture with their corresponding carbonyl

Scheme 2. Substrate Scope of Alkenes for Dichloromethyl Radical Additiona

<sup>a</sup>Reaction conditions: Unless indicated otherwise, reactions of 1 (1.0 mmol), CuI (1 mol %), TBHP (5.0 equiv), and DIPEA (5.0 equiv) were carried out in CHCl<sub>3</sub>/acetone (1:1, 5.0 mL) at rt. <sup>b</sup>After 30 min, an additional 3.0 equiv of TBHP were added. <sup>c</sup>Cumyl hydroperoxide was used as an alternative to TBHP, and an additional 3.0 equiv of cumyl hydroperoxide were added after 30 min.

products,<sup>21</sup> the peroxide products can be easily converted to their alcohol counterparts by a rarely applied reduction that occurs in acetic acid with zinc dust (Scheme 3).<sup>22</sup> In all cases,

Scheme 3. Transformation of Peroxide Products to Benzyl Alcohols

<sup>a</sup>After 48 h an additional 3.0 equiv of Zn dust was added. <sup>b</sup>After 48 h, an additional 5.0 equiv of Zn dust was added. Product yield includes 10% acetate ester.

once starting material was consumed, a simple basic aqueous workup led to the desired product with no need for additional purification. For the para-methoxyphenyl substrate, the main product was the alcohol 4d, but 10% of acylated 4d' was also isolated.

Another example of the utility of the dichloromethyl addition process is evident in the conversion of the addition products to substituted pyridine derivatives, a recent example of which is a palladium-catalyzed condensation of acetonitrile with homopropargylic alcohols.<sup>23</sup> We were delighted to observe that formation of substituted pyridines occurred in moderate yields in the presence of KO<sup>t</sup>Bu in acetonitrile

(Scheme 4). A similar process, known as the Thorpe-Ziegler reaction, is used in the self-condensation reactions of aliphatic

Scheme 4. Transformation of Peroxide Products into **Substituted Pyridines** 

nitriles that results in the synthesis of enamines that provide ketone products after acid hydrolysis.<sup>24</sup> The X-ray structure of 5b was obtained for full characterization. A full study of this reaction and its implications is ongoing.

The radical nature of the transformation is suggested by the reaction conditions that have been employed, but additional experimentation confirms this inference (Scheme 5). Reaction

Scheme 5. Mechanistic Experiments To Probe the Radical **Pathway** 

of 1a under standard reaction conditions along with TEMPO as an additive provided no conversion of the starting material (Scheme 5, eq 1). Additionally, with use of vinylcyclopropane 6 as the substrate to ascertain the fate of the initially formed reaction intermediate and its lifetime, several products were formed in addition to 7a:ketone 7b and ring-opening products 7c and 7c' (Scheme 5, eq 2). Major product 7a is consistent with the overall addition process from which products 7c and 7c' are formed and whose yields suggest the relative rates for capture by the tert-butylperoxy radical versus ring-opening/ hydrogen-atom transfer. 25 The formation of cleavage product 7b is due to the initial addition of the tert-butylperoxy radical to 6, and its formation in reactions with  $O_2$  or  $H_2O_2$  as the oxidant has been reported.<sup>26</sup>

From a mechanistic standpoint, the most likely initiation of this reaction is by TBHP and CuI to form the tert-butylperoxy radical by hydrogen-atom transfer from TBHP to the initially formed *tert*-butoxy radical<sup>27</sup> (Scheme 6). This process has been extensively studied.<sup>28</sup> The Cu<sup>II</sup> species produced by this oxidative reaction, (HO)CuI, is proposed to be in equilibrium

## Scheme 6. Initiation of the Process Leading to the Dichloromethyl Radical through Formation of α-Aminoradical 9

with ICu-OO<sup>t</sup>Bu (8).<sup>29</sup> A HAT of the *tert*-butylperoxy radical with the trialkylamine generates the  $\alpha$ -aminoradical (9) that is the key intermediate which preferentially undergoes chlorineatom abstraction from chloroform. The CHCl<sub>2</sub> radical generated from chloroform adds regioselectively to the terminal carbon of styrene to generate the benzyl radical.

That the  $\alpha$ -aminoradical is the key intermediate is not only inferred from the calculations reported in Table 1 but also its involvement is strongly supported by the direct observation of iminium ion 11 using <sup>1</sup>H NMR spectral analysis under the standard conditions (Scheme 7). Addition of H<sub>2</sub>O sequestered

## Scheme 7. <sup>1</sup>H NMR Spectral Observation of Iminium Ion 11 and Acetaldehyde 12 in the Standard Reaction

the intermediate quantitatively to acetaldehyde 12. The reaction was also monitored by <sup>1</sup>H NMR under the standard reaction conditions, and acetaldehyde was observed to ultimately form in a 1:1 ratio to the peroxide product (see Supporting Information).

Hydrogen-atom abstraction from tertiary amine bases is key to understanding the formation of the dichloromethyl radical from chloroform, and may also contribute to understanding recently published copper(II) hydroxide-catalyzed generation of the dichloromethyl radical from CHCl2Br whose transformations are dependent on the stoichiometric uses of tertiary amine bases (Scheme 8, eq 1).9 The requirement for amines

# Scheme 8. Copper-Catalyzed Dichloromethylazidation of Alkene Using CHBrCl, or CHCl3 as the Dichloromethylating Reagent

Wang's work: CHBrCl<sub>2</sub> as the dichloromethylating reagent 
$$\begin{array}{c} \text{Cu}(\text{CH})_2 \text{ (10 mol\%)} \\ \text{Me} & \text{Tal} \end{array} + \begin{array}{c} \text{CHBrCl}_2 \\ \text{Me}_6\text{TREN} \text{ (20 mol\%)} \\ \text{Me}_8\text{(1.5 eq.)} \\ \text{(2.0 eq)} \end{array} + \begin{array}{c} \text{Cu}(\text{CH})_2 \text{ (10 mol\%)} \\ \text{Me}_8\text{(1.5 eq.)} \\ \text{Me}_8\text{(2.5 eq.)} \\ \text{Me}_8\text{(2.0 eq.)} \end{array} + \begin{array}{c} \text{CHCl}_2 \\ \text{Me}_8\text{(3.0 eq.)} \\ \text{Me}_8\text{(3.0 eq.)} \\ \text{TMSN}_3 \text{ (3.0 eq.)} \\ \text{Me} \end{array} + \begin{array}{c} \text{CHCl}_2 \\ \text{Me} \\ + \begin{array}{c} \text{CHCl}_2 \\ \text{Me} \\ \text{Me} \end{array} + \begin{array}{c}$$

may also indicate that the amine plays an important role in the generation of the dichloromethyl radical rather than being only a ligand or base. Under our conditions with CuI, TBHP, and DIPEA, reaction with TMS-azide produces the same azide addition product as was reported by Wang and co-workers [in competition with formation of 2a (Scheme 8, eq 2)]; and increasing the stoichiometric amount of TMSN<sub>3</sub> also increases the yield of the azide 13 relative to 2a (see Supporting Information); this outcome suggests that the indirect methodology makes possible a broad spectrum of radical oxidative addition processes.

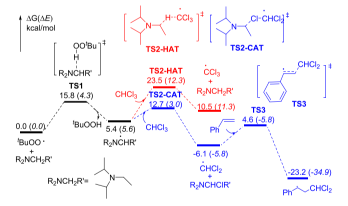
As we were submitting this article for publication, Li and coworkers reported the use of cobalt(II) acetylacetonate as a catalyst and triethylamine as a base for difluoroalkylationperoxidation of arylalkenes (eq 2). 15d In their survey of

$$\begin{array}{c}
R^{2} \\
\nearrow \\
R^{1}
\end{array}
+ ROOH + X-CF_{2}CO_{2}Et \xrightarrow{Co(acac)_{2}, NEt_{3}} \xrightarrow{R^{1}} \begin{array}{c}
R^{2} \\
\nearrow \\
ROO \\
CF_{2}CO_{2}Et
\end{array}$$
(2)

applicable halogenation reagents they reported one example using chloroform in which the dichloromethyl product 2h was formed in 41% yield, and we have confirmed this result. The role of trimethylamine was proposed to be solely that of a base to neutralize acid formation, but the results that we report suggests that the tertiary amine is essential for halogen-atom abstraction.

The selectivity of CAT versus HAT for the generation and oxidative addition of the dichloromethyl radical was subjected to DFT analysis (Scheme 9). Hydrogen-atom abstraction from

Scheme 9. Computed Free Energy Diagram for  $\alpha$ -Amino Radical Mediated Reactions with Chloroform<sup>a</sup>

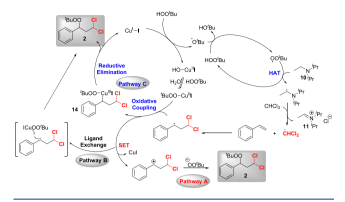


<sup>a</sup>Calculations were at the B3LYP-D3 level.

the  $\alpha$ -position of DIPEA to the *tert*-butylperoxy radical occurs with a relatively low activation barrier of 15.8 kcal/mol. The HAT from chloroform to the DIPEA radical was calculated to have a higher energy barrier (23.5 kcal/mol) than the CAT from chloroform to the DIPEA radical (12.7 kcal/mol). This preference is sufficiently great to direct the reaction exclusively to the dichloromethyl radical.

The formation of peroxide products 2 and 3 in relatively high yields narrows the possible outcomes to three distinct pathways (Scheme 10). In Pathway A a single-electron transfer (SET) by Cu<sup>II</sup> would yield a benzyl carbocation which could be intercepted by TBHP or, in theory, by other suitable nucleophiles, including water in T-HYDRO. However, when the reaction was conducted in the presence of alternative

Scheme 10. Three Possible Mechanistic Pathways for Product Formation



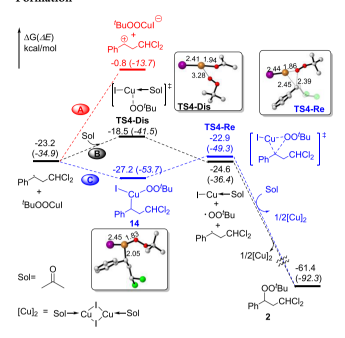
nucleophiles, no deviation in product selectivity for 2 was observed (i.e., with MeOH, NaOH/H<sub>2</sub>O, indole). In Pathway B the formation of 2 occurs through a two-step outer-sphere ligand transfer between ICu-OO'Bu and the benzyl radical. Alternatively, in Pathway C addition of Cu<sup>II</sup> to the benzyl radical, resulting in Cu<sup>III</sup> species 14, followed by reductive elimination regenerates the catalyst and affords the observed product. This pathway has been used to explain other coppercatalyzed addition reactions involving TBHP, 30 but there has not been experimental evidence for this pathway, and asymmetric induction from the use of chiral ligands for copper(I) (and, alternatively, for dirhodium(II) carboxamidates) offered an attractive experiment that might distinguish this pathway from Pathway B. However, only borderline enantioselectivity was observed in the presence of chiral ligands (see Supporting Information), possibly because of ineffective metal-ligand coordination. Because the lack of enantioselectivity does not fully discredit this particular pathway, we turned to DFT calculations to further distill the key reaction intermediates as well as which route was more plausible.

The three pathways for C–O bond formation were probed (Scheme 11). A single-electron transfer to form the benzylic carbocation was calculated to have the highest energy profile (Pathway A) which explains why no product loss due to the presence of other nucleophiles is observed. The direct outer sphere 'BuOO ligand transfer had the next highest energy barrier (TS4-Dis in Pathway B), whereas the formal oxidative addition of copper to styrene followed by reductive elimination (TS4-Re in Pathway C) was calculated to have the lowest activation free energy for the two steps and, so, is considered to be the most favorable pathway. The Cu–C and O–C distances (2.45 and 2.39 Å) in TS4-Re are much longer than the reported enantioselective cases.<sup>31</sup> Such ineffective coordination accounts for the experimental observation of the lack of enantioselectivity (see Supporting Information).

#### CONCLUSIONS

In conclusion, a previously unrealized chlorine-atom transfer from chloroform was achieved by an indirect process through which catalytic TBHP-induced formation of an  $\alpha$ -aminoradical causes chlorine-atom transfer from chloroform, forming the dichloromethyl radical that undergoes regioselective addition to the carbon—carbon double bond of styrenes. This process was optimized to form various substituted  $\alpha$ -peroxy- $\beta$ , $\beta$ -dichloropropylarene products and was also suitable for

Scheme 11. Computed Free Energy Diagram for C-O Bond Formation  $^a$ 



<sup>a</sup>Calculations were at the B3LYP-D3 level and distances are in Å.

halogen-atom abstraction of additional alkyl halides to form diverse  $\alpha$ -peroxy- $\beta$ -substituted ethylbenzene products. An excellent correlation between IPs and the difference in free energy barrier of HAT versus CAT was found. The correlation provides a general method to select appropriate radical for halogen-atom transfer reaction. Furthermore, the mechanism for halogen-atom transfer involving  $\alpha$ -amino radical intermediates unifies disparate reports and establishes a cohesive framework for further advances. A detailed computational investigation clearly demonstrated the deviation from traditional HAT of chloroform and also established a formal oxidative addition/reductive elimination as being the lowest energy pathway. The tert-butylperoxide products formed by this methodology are reducible to their corresponding alcohols under mild conditions and are also convertible to highly substituted pyridine derivatives in acetonitrile.

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.9b05921.

General experimental procedures and detailed optimization and product analyses, experimental procedures for mechanistic analyses, computational details, references, and NMR spectra (PDF)
Crystallographic data for **5b** (CIF)

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The authors declare no competing financial interest.

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# REFERENCES

- (1) Kharasch, M.; Jensen, E.; Urry, W. Addition of Carbon Tetrachloride and Chloroform to Olefins. Science 1945, 102, 128.
- (2) (a) Kharasch, M. S.; Urry, W. H.; Jensen, E. V. Addition of Derivatives of Chlorinated Acetic Acids to Olefins. J. Am. Chem. Soc. 1945, 67, 1626. (b) Kharasch, M. S.; Jensen, E. V.; Urry, W. H. Reactions of Atoms and Free Radicals in Solution. X. The Addition of Polyhalomethanes to Olefins. J. Am. Chem. Soc. 1947, 69, 1100. (c) Kharasch, M. S.; Kuderna, B. M.; Urry, W. Reactions of Atoms and Free Radicals In Solution. XV. The Additions of Bromodichloromethane and Dibromodichloromethane to Olefins. The Preparation of 2-Alkenyls. J. Org. Chem. 1948, 13, 895.
- (3) (a) Kharasch, M. S.; Mayo, F. R. Peroxide Effect in the Addition of Reagents to Unsaturated Compounds. I. The Addition of Hydrogen Bromide to Allyl Bromide. J. Am. Chem. Soc. 1933, 55, 2468. (b) Igbal, J.; Bhatia, B.; Nayyar, N. K. Transition Metal-Promoted Free-Radical Reactions in Organic Synthesis: The Formation of Carbon-Carbon Bonds. Chem. Rev. 1994, 94, 519. (c) Gossage, R. A.; van de Kuil, L. A.; van Koten, G. Diaminoarylnickel(II) "Pincer" Complexes: Mechanistic Considerations in the Kharasch Addition Reaction, Controlled Polymerization, and Dendrimeric Transition Metal Catalysts. Acc. Chem. Res. 1998, 31, 423. (d) Clark, A. J. Atom transfer radical cyclisation reactions mediated by copper complexes. Chem. Soc. Rev. 2002, 31, 1.
- (4) Balili, M. N. C.; Pintauer, T. Persisitent Radical Effect in Action: Kinetic Studies of Copper-Catalyzed Atom Transfer Radical Addition in the Presence of Free-Radical Diazo Initiators as Reducing Agents. Inorg. Chem. 2009, 48, 9018.
- (5) Muñoz-Molina, J. M.; Sameera, W. M.; Álvarez, E.; Maseras, F.; Belderrain, T. R.; Pérez, P. J. Mechanistic and Computational Studies of the Atom Transfer Radical Addition of CCl4 to Styrene Catalyzed by Copper Homoscorpionate Complexes. Inorg. Chem. 2011, 50, 2458.
- (6) Nair, R. P.; Kim, T. H.; Frost, B. J. Atom Transfer Radical Addition Reactions of CCl<sub>4</sub>, CHCl, and p-Tosyl Chloride Catalyzed by Cp'Ru(PPh<sub>3</sub>)(PR<sub>3</sub>)Cl Complexes. Organometallics 2009, 28, 4681.
- (7) Ueda, M.; Doi, N.; Miyagawa, H.; Sugita; Takeda, N.; Shinada, T.; Miyata, O. Reaction of Cyclopropenes with Trichloromethyl Radical: Unprecedented Ring-Opening Reaction of Cyclopropanes with Migration. Chem. Commun. 2015, 51, 4204.
- (8) (a) Lu, M. Z.; Loh, T. P. Iron-Catalyzed Cascade Carbochloromethylation of Activated Alkenes: Highly Efficient Access to Chloro-Containing Oxindoles. Org. Lett. 2014, 16, 4698. (b) Li, X.; Xu, J.; Gao, Y.; Fang, H.; Tang, G.; Zhao, Y. Cascade Arylalkylation of Activated Alkenes: Synthesis of Chloro- and Cyano-Containing Oxindoles. J. Org. Chem. 2015, 80, 2621. (c) Liu, Y.; Zhang, J.-L.; Song, R.-J.; Li, J.-H. 1,2-Alkylarylation of activated alkenes with dual C-H bonds of arenes and alkyl halides toward polyhalo-substituted oxindoles. Org. Chem. Front. 2014, 1, 1289. (d) Sheng, W.; Jin, C.; Shan, S.; Jia, Y.; Gao, J. Metal-Free Polychloromethylation/ Cyclization of N-Aryl Acrylamides. Youji Huaxue 2016, 36, 325. (e) Tian, Y.; Liu, Z.-Q. Metal-free radical cascade dichloromethylation of activated alkenes using CH2Cl2: highly selective activation of the C-H bond. RSC Adv. 2014, 4, 64855.
- (9) (a) Bao, Y.; Wang, G.-Y.; Zhang, Y.-X.; Bian, K.-J.; Wang, X.-S. Copper-catalyzed formylation of alkenyl C-H bonds using BrCHCl<sub>2</sub> as a stoichiometric formylating reagent. Chem. Sci. 2018, 9, 2986. (b) Zhang, Y.-X.; Jin, R.-X.; Yin, H.; Li, Y.; Wang, X.-S. Copper-

- Catalyzed Dichloromethylazidation of Alkenes Using BrCCl<sub>2</sub>H as a Stoichiometric Dichloromethylating Reagent. Org. Lett. 2018, 20,
- (10) Abo-Amer, A.; McCready, M. S.; Zhang, F.; Puddephatt, R. J. The Role of Solvent in Organometallic Chemistry- Oxidative Addtion with Dichloromethane or Chloroform. Can. J. Chem. 2012, 90, 46.
- (11) (a) Ueda, M.; Doi, N.; Miyagawa, H.; Sugita, S.; Takeda, N.; Shinada, T.; Miyata, O. Reaction of cyclopropenes with a trichloromethyl radical: unprecedented ring-opening reaction of cyclopropanes with migration. Chem. Commun. 2015, 51, 4204. (b) Liu, Y.; Zhang, J.-L.; Song, R.-J.; Li, J.-H. 1,2-Alkylarylation of activated alkenes with dual C-H bonds of arenes and alkyl halides toward polyhalo-substituted oxindoles. Org. Chem. Front. 2014, 1, 1289. (c) Pan, C.; Gao, D.; Yang, Z.; Wu, C.; Yu, J. T. Metal-free radical cascade chloromethylation of unactivated alkenes: synthesis of polychloro-substituted indolines. Org. Biomol. Chem. 2018, 16, 5752. (d) Liu, Y.; Zhang, J.- L.; Song, R.-J.; Li, J.-H. Visible-Light-Facilitated 5-exo-trig Cyclization of 1,6-Dienes with Alkyl Chlorides: Selective Scission of the C(sp<sup>3</sup>)-H Bond in Alkyl Chlorides. Eur. J. Org. Chem. 2014, 2014, 1177. (e) Nair, R. P.; Kim, T. H.; Frost, B. J. Atom Transfer Radical Addition Reactions of CCl<sub>4</sub>, CHCl<sub>3</sub>, and p-Tosyl Chloride Catalyzed by Cp'Ru(PPh3)(PR3)Cl Complexes. Organometallics 2009, 28, 4681. (f) Li, S.; Fan, W. Y. FTIR and computational studies of gas-phase hydrogen atom abstraction kinetics by t-butoxy radical. Chem. Phys. Lett. 2006, 427, 276. (g) Sekušak, S.; Sabljić, A. The role of complexes in hydrogen abstraction from haloethanes by the hydroxyl radical. A case of guided reactions. Chem. Phys. Lett. 1997, 272, 353.
- (12) De Vleeschouwer, F.; Van Speybroeck, V.; Waroquier, M.; Geerlings, P.; De Proft, F. Electrophilicity and Nu-cleophilicity Index for Radicals. Org. Lett. 2007, 9, 2721.
- (13) Fu, Y.; Liu, L.; Yu, H.-Z.; Wang, Y.-M.; Guo, Q.-X. Quantum-Chemical Predictions of Absolute Standard Redox Potentials of Diverse Organic Molecules and Free Radicals in Acetonitrile. J. Am. Chem. Soc. 2005, 127, 7227.
- (14) (a) Lalevée, J.; Allonas, X.; Fouassier, J. P. Halogen Abstraction Reaction between Aminoalkyl Radicals and Alkyl Halides: Unusual High Rate Constants. Chem. Phys. Lett. 2008, 454, 415. (b) Lalevée, J.; Fouassier, J. P.; Blanchard, N.; Ingold, K. U. Reaction between aminoalkyl radicals and akyl halides: Dehalogenation by electron transfer? Chem. Phys. Lett. 2011, 511, 156.
- (15) (a) Munoz, Z.; Cohen, A. S.; Nguyen, L. M.; McIntosh, T. A.; Hoggard, P. E. Photocatalysis by tetraphenylporphyrin of the decomposition of chloroform. Photochem. Photobiol. Sci. 2008, 7, 337. (b) Thommes, K.; Icli, B.; Scopelliti, R.; Severin, K. Atom-Transfer Radical Addition (ATRA) and Cyclization (ATRC) Reactions Catalyzed by a Mixture of [RuCl<sub>2</sub>Cp\*(PPh<sub>3</sub>)] and Magnesium. Chem. - Eur. J. 2007, 13, 6899. (c) Asscher, M.; Vofsi, D. Chlorine-activation by redox-transfer. Part III. The "abnormal" addition of chloroform to olefins. J. Chem. Soc. 1963, 0, 3921. (d) Chen, Y.; Li, L.; Ma, Y.; Li, Z. Cobalt-Catalyzed Three-Component Difluoroalkylation-Peroxidation of Alkenes. J. Org. Chem. 2019, 84, 5328.
- (16) Catino, A. J.; Nichols, J. M.; Nettles, B. J.; Doyle, M. P. The Oxidative Mannich Reaction Catalyzed by Dirhodium Caprolactamate. J. Am. Chem. Soc. 2006, 128, 5648.
- (17) (a) Boess, E.; Schmitz, C.; Klussmann, M. A Comparative Mechanistic Study of Cu-Catalyzed Oxidative Coupling Reactions with N-Phenyltetrahydroisoquinoline. J. Am. Chem. Soc. 2012, 134, 5317. (b) Boess, E.; Wolf, L. M.; Malakar, S.; Salamone, M.; Bietti, M.; Thiel, W.; Klussmann, M. Competitive Hydrogen Atom Transfer to Oxyl- and Peroxyl Radicals in the Cu-Catalyzed Oxidative Coupling of N-Aryl Tetrahydroisoquinolines Using tert-butyl Hydroperoxide. ACS Catal. 2016, 6, 3253.
- (18) During our investigations a report appeared which claimed the dichloromethyl radical from chloroform was formed in reactions with TBHP and styrene without a catalyst or amine, but our attempts to repeat these reactions were unsuccessful: Chen, C.; Tan, H.; Liu, B.; Yue, C.-C.; Liu, W.-B. ATRA-like alkylation-peroxidation of alkenes

with trichloromethyl derivatives by the combination of *t*-BuOOH and NEt<sub>3</sub>. Org. Chem. Front. **2018**, 5, 3143.

- (19) (a) Luo, Q.; Liu, C.; Tong, J.; Shao, Y.; Shan, W.; Wang, H.; Zheng, H.; Cheng, J.; Wan, X. Cu-Catalyzed Multicomponent Reaction of Styrenes, Perflouroalkyl Halide, Alcohol, and *tert*-Butyl Hydroperoxide: One-Pot Synthesis of (Z)- $\beta$ -Alkoxyperfluoroalkenone. *J. Org. Chem.* **2016**, *81*, 3103. (b) Li, Y.; Liu, J.; Zhao, S.; Du, X.; Guo, M.; Zhao, W.; Tang, X.; Wang, G. Copper-Catalyzed Fluoroolefination of Silyl Enol Ethers and Ketones toward the Synthesis of  $\beta$ -Fluoronones. *Org. Lett.* **2018**, *20*, 917.
- (20) Kropf, H. Houben-Weyl Methoden der Organishen Chemie, Peroxo-Verbindungen; Kropf, H., Ed.; Thieme: Stuttgart, 1988; p 1102. (21) (a) Howard, J. A.; Ingold, K. U. The Self-Reaction of sec-Butylperoxy Radicals. Confirmation of the Russell Mechanism. J. Am. Chem. Soc. 1968, 90, 1056. (b) Di Mascio, P. D.; Martinez, G. R.; Miyamoto, S.; Ronsein, G. E.; Medeiros, M. H. G.; Cadet, J. Singlet Molecular Oxygen Reactions with Nucleic Acids, Lipids, and Proteins. Chem. Rev. 2019, 119, 2043.
- (22) Cheng, J.-K.; Loh, T.-P. Copper- and Cobalt-Catalyzed Direct Coupling of sp<sup>3</sup>  $\alpha$ -Carbon of Alcohols with Alkenes and Hydroperoxides. *J. Am. Chem. Soc.* **2015**, *137*, 42.
- (23) Sarkar, D.; Rout, N.; Ghosh, M. K.; Giri, S.; Neue, K.; Reuter, H. Atom-Economical Palladium Carbon-Catalyzed de Novo Synthesis of Trisubstituted Nicotinonitriles. *J. Org. Chem.* **2017**, *82*, 9012.
- (24) Condensation of ethyl cyanoacetate with its sodium: Baron, H.; Remfry, F. G. P.; Thorpe, J. F. *J. Chem. Soc., Trans.* **1904**, *85*, 1726. (25) (a) Zhang, H.; Pu, W.; Xiong, T.; Li, Y.; Zhou, X.; Sun, K.; Liu, Q.; Zhang, Q. Copper-Catalyzed Intermolecular Aminocyanation and Diamination of Alkenes. *Angew. Chem., Int. Ed.* **2013**, *52*, 2529. (b) Zhang, H.; Song, Y.; Zhao, J.; Zhang, J.; Zhang, Q. Regioselective Radical Aminofluorination of Styrenes. *Angew. Chem., Int. Ed.* **2014**, *53*, 11079.
- (26) (a) Gonzalez-de-Castro, A.; Xiao, J. Green and Efficient: Iron-Catalyzed Selective Oxidation of Olefins to Carbonyls with O<sub>2</sub>. *J. Am. Chem. Soc.* **2015**, *137*, 8206. (b) Wang, T.; Jing, X.; Chen, C.; Yu, L. Organoselenium-Catalyzed Oxidative C=C Bond Cleavage: A Relatively Green Oxidation of Alkenes into Carbonyl Compounds with Hydrogen Peroxide. *J. Org. Chem.* **2017**, *82*, 9342.
- (27) (a) Minisci, F.; Fontana, F.; Araneo, S.; Recupero, F.; Banfi, S.; Quici, S. Kharasch and Metalloporphyrin Catalysis in the Functionalization of Alkanes, Alkenes, and Alkylbenzenes by t-BuOOH. Free Radical Mechanisms, Solvent Effect, and Relationship with the Gif Reaction. J. Am. Chem. Soc. 1995, 117, 226. (b) Avila, D. V.; Ingold, K. U.; Lusztyk, J.; Green, W. H.; Procopio, D. R. Dramatic Solvent Effects on the Absolute Rate Constants for Abstraction of the Hydroxylic Hydrogen Atom from ferf-Butyl Hydroperoxide and Phenol by the Cumyloxyl Radical. The Role of Hydrogen Bonding. J. Am. Chem. Soc. 1995, 117, 2929.
- (28) (a) Boess, E.; Wolf, L. M.; Malakar, S.; Salamone, M.; Bietti, M.; Thiel, W.; Klussmann, M. Competitive Hydrogen Atom Transfer to Oxyl- and Peroxyl Radicals in the Cu-Catalyzed Oxidative Coupling of N-Aryl Tetrahydroisoquinolines Using tert-Butyl Hydroperoxide. ACS Catal. 2016, 6, 3253. (b) Boess, E.; Schmitz, C.; Klussmann, M. A Comparative Mechanistic Study of Cu-Catalyzed Oxidative Coupling Reactions with N-Phenyltetrahydroisoquinoline. J. Am. Chem. Soc. 2012, 134, 5317. (c) Li, Z.; Li, C.-J. CuBr-Catalyzed Efficient Alkynylation of sp<sup>3</sup> C-H Bonds Adjacent to a Nitrogen Atom. J. Am. Chem. Soc. 2004, 126, 11810. (d) Li, Z.; Li, C.-J. Highly Efficient Copper-Catalyzed Nitro-Mannich Type Reaction: Cross-Dehydrogenative-Coupling between sp<sup>3</sup> C-H Bond and sp<sup>3</sup> C-H Bond. J. Am. Chem. Soc. 2005, 127, 3672. (e) Zhang, Y.; Fu, H.; Jiang, Y.; Zhao, Y. Copper-Catalyzed Amidation of sp<sup>3</sup> C-H Bonds Adjacent to a Nitrogen Atom. Org. Lett. 2007, 9, 3813.
- (29) DFT determination gives this conversion a reaction energy of 16.6 kcal/mol that provides an equilibrium constant of  $1.4 \times 10^{12}$ ..
- (30) Ren, Z.-H.; Zhao, M.-N.; Guan, Z.-H. CuI-catalyzed oxidative cross coupling of oximes with tetrahydrofuran: a direct access to Otetrahydrofuran-2-yl oxime ethers. *RSC Adv.* **2016**, *6*, 16516.

(31) (a) Zhu, X.; Deng, W.; Chiou, M.-F.; Ye, C.; Jian, W.; Zeng, Y.; Jiao, Y.; Ge, L.; Li, Y.; Zhang, X.; Bao, H. Copper-Catalyzed Radical 1,4-Difunctionalization of 1,3-Enynes with Alkyl Diacyl Peroxides and N-Fluorobenzenesulfonimide. *J. Am. Chem. Soc.* 2019, 141, 548. (b) Zhang, W.; Wang, F.; McCann, S. D.; Wang, D.; Chen, P.; Stahl, S. S.; Liu, G. Enantioselective cyanation of benzylic C–H bonds via copper-catalyzed radical relay. *Science* 2016, 353, 1014.