Substrate a developed a simple diagnostic tool for the nature of the steps following the electron transfer. The reduction of imines showed that substrates reacting by a dissociative electron transfer mechanism to normal reductions, which are di-benzyl chloride. The mechanistic understanding gained enabled us also to broaden the scope of photostimulated reactions from reactions. For example, hexamethylphosphoramide (HMPA) enhances the reduction of anthracene while hampering the reaction of ground state. Harnessing the classical knowledge of proton transfer mechanisms to our SmI₂ research enabled us to decipher an old conundrum: why does the combination of water and amine have such an enhancing effect, which depended on the order of the reagents addition. This seminal paper was fundamental to an understanding of how the SmI₂ works and enabled us to later explain various phenomena. For example, it was found that in a given reaction, when MeOH is used as a proton source, a spiro compound is obtained, while a bicyclic product is obtained when t-BuOH is used. Our contribution culminated in formulating guidelines for the rational use of proton donors in SmI₂ reactions.

The need to understand the complexity of the effect of additives on various processes is nicely demonstrated in photoinduced reactions. For example, hexamethylphosphoramide (HMPA) enhances the reduction of anthracene while hampering the reaction of benzyl chloride. The mechanistic understanding gained enabled us also to broaden the scope of photostimulated reactions from substrates reacting by a dissociative electron transfer mechanism to normal reductions, which are difficult to accomplish at the ground state. Harnessing the classical knowledge of proton transfer mechanisms to our SmI₂ research enabled us to decipher an old conundrum: why does the combination of water and amine have such an enhancing effect on the reactivity of SmI₂, which is not typical of these two when used separately. In our studies on the affinity of ligands to SmI₂, we discovered that, in contradistinction to the accepted dogma, SmI₂ is much more azaphilic than it is oxophilic. On the basis of the size difference between Sm³⁺ and Sm²⁺, we developed a simple diagnostic tool for the nature of the steps following the electron transfer. The reduction of imines showed that substrate affinity to SmI₂ plays also a crucial role. In these reactions, new features such as autocatalysis and catalysis by quantum dots were discovered. Several studies of the ligand effect lead to a clear formulation of when an inner sphere or outer sphere electron transfer should be expected. In addition, several reactions where proton-coupled electron transfer (PCET) is the dominant mechanism were identified. Finally, the surprisingly old tool of NMR "shift reagents" was rediscovered and used to directly derive essential information on the binding constants of ligands and substrates to SmI₂.

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## INTRODUCTION

The chemistry of SmI$_2$ displays a rich repertoire of reactions that enable a myriad of synthetic possibilities. In the course of reduction of substrate A$\rightleftharpoons$B, three different intermediates can be obtained (Scheme 1). In principle, each intermediate can lead to a different product. This complexity, which provides great opportunities for the synthetic chemist, also creates a fascinating mechanistic maze.

In retrospect, our first report on SmI$_2$ was seminal to the eventual unraveling of the mechanistic mysteries of this reagent. However, the results were very puzzling. The H/D product isotope effect for the reaction depicted in eq 1 was determined as described in Figure 1. When a THF solution of the substrate containing both MeOH and MeOD (1:1 ratio) was reacted with a THF solution of SmI$_2$, the H/D incorporation ratio into the benzylic carbon was 1.2. However, when the mixing protocol was changed, and methanol was introduced into the SmI$_2$ solution, while MeOD was kept with the substrate solution, the H/D incorporation ratio increased to 7.2. This baffling result, which was confirmed by three generations of postdocs and graduate students, led a referee to suggest that the samarium may insert into the O$\rightarrow$H bond to give a hydride that does not exchange its hydrogen. However, we clearly showed that SmI$_2$ forms reversibly a complex with MeOH, which affects its visible spectrum as shown in Figure 2.

It is reasonable to assume that the isotope effect on MeOH complexation to SmI$_2$ is negligible. Thus, in the first experiment, MeOH and MeOD complex similarly with SmI$_2$. In the second experiment, however, only the MeOH is complexed to SmI$_2$, and protonation occurs within the complex where methanol is coordinated to Sm$^{3+}$, which is ion-paired to the radical anion of the substrate. This unimolecular protonation, which is much faster than a bimolecular protonation by MeOD in the bulk, results in a high H/D isotope effect.

This Account aims to shed light on the main mechanistic channels that characterize the SmI$_2$ reactions. We shall begin with the preassociation effect.

### PREASSOCIATION EFFECT

The association of the substrate radical anion, the Sm$^{3+}$, and the proton donor that is complexed to it, has an immense effect on the chemistry, as will be shown in the following sections. This proximity effect is an entropic factor, which reflects the higher probability for a unimolecular reaction to occur within a reaction cage, as opposed to the much lower probability for a bimolecular reaction. This may result in rate enhancement of up to 13 orders of magnitude.

In a SmI$_2$ reaction that includes protonation, there are three relevant preassociations: (1) proton donor with SmI$_2$, (2) SmI$_2$ with the substrate, and (3) proton donor with the substrate.

**Preassociation of the Proton Donor with SmI$_2$**

As demonstrated above, it is well established that preassociation of the proton donor with SmI$_2$ significantly affects the reaction. Following electron transfer, Sm$^{3+}$I$_2$ complexed with the proton donor is paired to the radical anion of the substrate. The most obvious effect of this preassociation is on the protonation rate. Thus, if protonation of the radical anion is rate

---

**Scheme 1. Mechanistic Options for Reduction by SmI$_2$**

- $A\rightleftharpoons B \xrightarrow{SmI_2} A^- + B^+$
- $H^+ \xrightarrow{HA^{-}} HA^-$
- $SmI_2 \xrightarrow{HA^-} SmI_2HA^-$

**Figure 1. Alternate mixing protocols yield markedly different H/D ratios.**

- H/D = 1.2
- H/D = 7.15

**Figure 2. Effect of MeOH concentration on SmI$_2$ visible spectrum.**

- $Ph = C = CN \xrightarrow{SmI_2, THF, MeOH(MeOD)} Ph = C = CN$ (1)
determining (eq 2), the proximity of the proton donor to the radical anion renders the protonation highly efficient, diminishing the rate of the back electron transfer. If a back electron transfer takes place, the Coulombic attraction, which keeps the ROH–Sm\(^{3+}\) near the radical anion, is lost, and most

---

**Figure 3.** Diode array monitoring of the reaction of naphthalene (0.2 M) and SmI\(_2\) (2.5 mM) in the presence of (a) MeOH (1 M, 200 s) and (b) TFE (1 M, 800 s). Reproduced with permission from ref 13. Copyright 2010 Wiley-VCH.

**Figure 4.** Rate constants as a function of Et\(_3\)N concentration in the reaction of SmI\(_2\) with 3-methyl-2-butanone in the presence of water. Reproduced with permission from ref 2. Copyright 2015 Wiley-VCH.

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**Chart 1. Three Imines Used in the Preassociation Study**
A side issue, which is both very interesting and counter-intuitive, was discovered upon comparison of the kinetic and the product isotope effects using MeOH/MeOD. Most chemists will agree that the charge in the radical anion resides mainly on the carbon $\alpha$ to the cyano group as shown in eq 4. This intuitively suggests that protonation will occur on this carbon. However, four repetitive determinations of the product isotope effect showed that the H/D incorporation isotope effect on the $\alpha$ carbon is $4.5 \pm 0.2$, whereas for the $\beta$ carbon it is $6.8 \pm 0.3$. Yet, the kinetic isotope effect was $6.7 \pm 0.3$, showing that the first protonation, which locks the reaction and prevents its reversal, occurs on the $\beta$ rather than the $\alpha$ carbon. Based on this and other evidence, it was shown that protonation occurs in such a way that the most stable radical is formed.\(^{12}\)

We next used this strategy to broaden the scope of photostimulated SmI$_2$ reactions.\(^{13}\) Irradiation of SmI$_2$ at its $\lambda_{	ext{max}}$ (ca. 600 nm) converts it to a powerful electron donor, which enables an electron transfer even to substrates with a very high LUMO (the excitation energy of SmI$_2$ around 600 nm is equivalent to ca. 2.2 eV, which is much more than the 0.72 V increase in reduction potential effected by 4 equiv of hexamethylphosphoramide (HMPA).\(^{14}\)) The major deficiency of this procedure is that when applied to reduction of resistant substrates, it yields a high-energy radical anion, which will eagerly give back the additional electron to Sm$^{3+}$. Therefore, the photostimulated reductions were successful only in cases of dissociative electron transfer reactions such as shown in eqs 5 and 6. In these reactions, the electron transfer is the rate-determining step, as the cleavage of the leaving group is coupled to the electron transfer. However, the fact that we can bring the proton donor to the reaction center enables the trapping of even very short-lived radical anions such as those of naphthalene and diphenylacetylene. Figure 3 displays diode array kinetics of the reduction of naphthalene in the presence of a MeOH in comparison with the reaction in the presence of TFE, which is a stronger proton donor but unlike MeOH does not complex to SmI$_2$. It should be pointed out that in the dark no reaction takes place.

\[
\text{H-CN} + \text{SmI}_2 \rightarrow \text{H-CN} + \text{I}^- + \text{H} + \text{Sm}^{3+}
\]

Figure 5. Visible spectra of SmI$_2$ as a function of BNO concentration. Reproduced with permission from ref 22. Copyright 2017 Wiley-VCH.

In this system the lifetime of its radical anion is relatively very short. As a result, proton donors that do not complex to SmI$_2$ such as trifluoroethanol (TFE), i-PrOH, or t-BuOH did not lead to any reaction. However, in the presence of water or ethylene glycol, which do coordinate to SmI$_2$, rapid reactions took place ($\tau_{1/2} < 1$ ms). Thus, because of the low electrophilicity of the substrate, the radical anion in this endothermic reaction has a very short lifetime and can only be successfully trapped by a proton donor that resides in its vicinity.

In order to rule out the possibility that the reaction is a hydride transfer as suggested by a referee or that it is a HAT, transfer of a hydrogen atom,\(^{11}\) we performed an experiment with the less stable cis isomer in which it reacted with SmI$_2$ in the absence of a proton donor. As hydride or H transfer reactions are irreversible, the observed conversion of the cis isomer into the trans isomer (eq 4) proves unequivocally that the reaction course is an electron transfer followed by a proton transfer and rules out the two alternative mechanisms.

\[
\begin{align*}
\text{Ph-CN} + \text{SmI}_2 & \rightarrow \text{Ph-CN} + \text{I}^- + \text{H} + \text{Sm}^{3+} \\
\text{Ph-CN} + \text{SmI}_2 & \rightarrow \text{Ph-CN} + \text{I}^- + \text{H} + \text{Sm}^{3+}
\end{align*}
\]
MeOH will protonate the radical anion at a much higher rate, not only because of its higher acidity but also because it changes the protonation mechanism from a bimolecular to a unimolecular one.

We turn to another surprising and important facet of the proximity effect on synthetic feasibility. A paper published in 1995 by Cabri showed that although the addition of water or amine to a reaction mixture does not affect much the reactions of SmI₂, their combination significantly facilitates reaction rates. This discovery lay dormant for seven years until the group of Hilmersson brought it back to the public attention. The water/amine combination received a major push by Procter and Szostak, who, in a long series of papers, got the best out of this “mixture” for synthetic applications. Procter faithfully described the situation in the following words: "the mechanistic details of this process, including the critical role of amine and H₂O additives, remained unclear".

Probing this reaction, we noticed that gradual addition of an amine, such as Et₃N, that does not complex to SmI₂ results in a rate enhancement, which levels off (e.g., 3-methyl-2-butanol, see Figure 4). The kinetic order in the amine before the onset of the plateau is one.

Our working hypothesis was that the role of the amine is to deprotonate a water molecule bound to the samarium cation, with subsequent generation of a hydroxide ion next to it. This would reduce the effective charge on the Sm³⁺ and consequently its ability to accept the electron back from the radical anion. The deprotonation could occur before the electron transfer, in concert with it, or after it. Using the classical tool of physical organic chemistry to distinguish between specific and general base catalysis, we showed that the reaction is a classic case of general base catalysis. Namely, the proton is transferred at the rate-determining step. This and other experiments rule out the first and second aforementioned options and leave us with the third option shown in eq 8 (A = substrate, only one water molecule is shown).

$$\text{A} + \text{SmI}_2 + \text{H}_2\text{O} \rightarrow \text{A}^- + \text{SmI}^3+ + \text{H}_2\text{O}$$

The major conclusion of this section is that the range of substrates that react with SmI₂ can be vastly expanded to include resilient substrates by using proton donors that efficiently complex to SmI₂. We identified two mechanisms by which a proton donor “riding on the back” of the samarium can trap a short-lived radical anion and prevent the back electron transfer to Sm³⁺. The first is by an efficient unimolecular protonation of the radical anion within the ion pair. The second mechanism is by reducing the affinity of the Sm³⁺ to accept back the electron from the radical anion by generating a negatively charged hydroxide ion coordinated to it.

**Preassociation of SmI₂ with the Substrate**

The advantage of uni- over bimolecularity applies here as well, and it is clear that preassociation of SmI₂ with the substrate will significantly enhance the rate of electron transfer. This phenomenon was termed “substrate-directable reactions” in a paper published by Procter, Szostak, and Flowers where the association of SmI₂ with a carbonyl function was enhanced by a neighboring OH group. Similarly, we showed that in the reduction of the imine depicted in Chart 1, the SmI₂ preassociates with the lone pair on the imine nitrogen.

The reactivity order is BPI > BMI > BAI. It does not conform with the electron affinity order of the substrates but rather with the accessibility of the nitrogen lone pair for complexation with SmI₂, demonstrating once again the importance of preassociation with the substrate.

It should be emphasized that this binding may not necessarily be to the reaction center but rather to a remote site in the molecule. This is exemplified in the case of 4-strydrypyridine, where the samarium binds to the lone pair of the pyridine nitrogen while the actual reduction takes place on the double bond (eq 9).

$$\text{I}_2\text{Sm}^- + \text{A} \rightarrow \text{I}_2\text{Sm}^- \text{A}^\text{−}$$

The efficacy of the proximity effect culminates in the reduction of benzene. The reduction of benzene is usually achieved by using strong reducing agents. In order to position the SmI₂ in the vicinity of the benzene ring, we used bidentate ligands (ethanolamine and ethylenediamine) as side arms on the benzene ring. These ligands are known to bind strongly to SmI₂, as demonstrated by visible spectroscopy (Figure 5).

However, due to the resistance of benzene toward reduction, the proximity itself was insufficient to induce a reduction, and we had to use in addition the water/amine magic mixture in order to execute a reduction. After 24 h, 95% conversion into the two classical products of the Birch reduction was observed (eq 10).

$$\text{BNO} + \text{SmI}_2\text{H}_2\text{O/Et}_3\text{N} \rightarrow \text{NH}_2\text{OH} + \text{NH}_2\text{OH}$$

The aforementioned examples show that the association of SmI₂ with the substrate opens new vistas for SmI₂ reactions. Preassociation of the Proton Donor with the Substrate

The benefit of preassociation of the proton donor with the substrate is somewhat less commonly encountered or discernible. If it occurs, its effect can be masked by the rate enhancement due to the protonation step. Moreover, hydrogen bonding between a carbonyl oxygen and the proton donor must be discernible. If it occurs, its efficacy can be masked by the rate enhancement due to the protonation step. Moreover, hydrogen bonding between a carbonyl oxygen and the proton donor must compete with the hydrogen bonding to the oxygen of THF, which is the common solvent for SmI₂ reactions. Therefore, it may be of a low likelihood. Higher probability and visibility of such bonding are with substrates that contain basic nitrogen as shown in the Proton-Coupled Electron Transfer (PCET) section.

**AZA- VERSUS OXOPHILICITY**

It is well established that SmI₂ is an oxophilic reagent. To our surprise, however, we recently discovered that its affinity for nitrogen compounds is much higher than that for their oxygen-based analogs. This was shown using several diagnostic tools. Complexation to SmI₂ is easily detected by visible spectroscopy. In Figure 6, it is shown that 0.1 M pyrrolidine, the nitrogen
analog of THF, is sufficient to fully displace all the solvent THF molecules from the coordination sphere despite the 2 orders of magnitude higher (12.3 M) concentration of THF. Similarly, for ethylene glycol (EG) to reach spectral saturation, concentrations ten times higher than its nitrogen analog ethylenediamine are needed. The higher azaphilicity was also demonstrated by comparing the kinetics of benzophenone and its aza analog. Under the same conditions, benzophenone imine reacts in nearly the dead time of the stopped-flow spectrometer whereas the reduction of benzophenone itself is about 30 times slower despite its much higher reduction potential (Figure 7).25

The higher azaphilicity was demonstrated for mono- and bidentate ligands, cyclic and acyclic molecules, and crown ethers, as well as for both sp³ and sp² hybridized systems. It is more pronounced for Sm³⁺ suggesting that it is governed by the same factors that control basicity. As Sm³⁺ is harder than Sm²⁺, it bears a higher resemblance to a proton, and therefore the more basic nitrogen compounds show a larger affinity to it. This is in fact the origin of the larger enhancing effect of nitrogen ligands on the reduction potential of SmI₂ relative to their oxygen analogs.

### OPTIMAL COORDINATION LEVEL

Figure 8 shows the effect of EG on the reduction potential of SmI₂ and on the rate constants for the benzyl chloride reaction.26 As can be seen, the two graphs coincide. However, with anthracene and cyclohexanone, the graphs are separate, and the one for cyclohexanone is closer to the reduction potential curve than the one for anthracene (Figure 9).

Similar results were found for other ligands and substrates. The variance in behavior is indicative of the nature of the rate-determining step. When the electron transfer is rate determining, as for benzyl chloride, the reduction potential and reaction rate constant curves overlap. However, when protonation is rate determining (anthracene and cyclohexanone), a larger concentration of ligand is needed to reach the leveling off region. The faster is the protonation, the more the curve will resemble that of the reduction potential. In general, protonation on oxygen is much faster than on carbon.18 If this is correct for the radical anions of cyclohexanone and anthracene as well, then it is clear why the curve for cyclohexanone is closer to that of the reduction potential than that of anthracene.

An interesting question is why does the concentration of ligand needed for reaching maximum reduction potential differ from the concentration needed for the maximal protonation rate? The answer is that different tasks need different coordination levels. Ligands increase the reduction potential of SmI₂ as the energy gain by coordination to Sm³⁺ is larger than that obtained by coordination to Sm²⁺. As the surface area of Sm²⁺ is larger by more than 50% than that of Sm³⁺ (Figure 10)27 and as the reduction potential is affected by coordination to
Sm$^{3+}$, it is superfluous to maintain on Sm$^{2+}$ more ligand molecules than Sm$^{3+}$ can accommodate. Therefore, the plateau is achieved early. However, for reactions involving protonation, the more proton donor molecules are in the vicinity of the radical anion, the greater the probability for protonation. Therefore, ligand saturation of the larger Sm$^{2+}$ is beneficial, and the plateau is achieved at higher ligand concentrations.

**PROTON-COUPLED ELECTRON TRANSFER (PCET)**

In this Account, we shall use the term PCET in its original context. Namely, we shall refer only to cases where the proton and electron are transferred simultaneously to and from two different sites.\(^{28}\) In recent years, PCET was also used to denote a hydrogen atom transfer (HAT). It would seem that the HAT mechanism cannot be very common in SmI$_2$ reactions, as most of the reactions are accelerated by the water/amine system.\(^{15}\) Catalysis by this magic mixture is a clear indication of an electron transfer in a pre-equilibrium step, which immediately rules out a HAT mechanism. In addition, HAT cannot operate in the reduction of functional groups for which an inner sphere electron transfer was proven.

The first proven PCET mechanism in the chemistry of SmI$_2$ was found in the water/amine reaction of benzyl chloride.\(^2\) As mentioned before, increasing the concentration of the amine induces rate enhancement, which eventually levels off as the electron transfer becomes rate determining (Figure 4). However, with benzyl chloride no such leveling off is observed, and the rate order in the amine remains one up to the highest amine concentration used (Figure 11). Hence, this is a clear case of the classical PCET mechanism.

In the two examples below, the PCET mechanism is not fully proven but is highly likely. In the aforementioned case of 4-styrylpyridine (eq 9),\(^{21}\) the reaction is catalyzed both by TFE (Figure 12), which does not bind to SmI$_2$, and by MeOH that binds to it. Catalysis is probably due to hydrogen bonding to the pyridine nitrogen lone pair, which increases its electrophilicity. The mechanistic options for this reaction are shown in Figure 13.

Motion along the periphery of the square diagram implies a stepwise mechanism, whereas motion along the diagonal implies a concerted PCET mechanism. The likelihood of this latter mechanism is high due to the hydron bond, which positions the proton in the optimal position for proton transfer. As proton transfer between two heteroatoms is very fast, it is highly likely that the proton transfer from the alcohol to the nitrogen will be synchronous with the electron transfer. It should be noted that the catalysis by proton donors competes with the catalytic effect resulting from coordination of the SmI$_2$ to the nitrogen lone pair. The relative effect depends on the binding equilibrium constants, the concentrations ratios, and the individual rate constants.

A similar case is encountered in the mesolytic cleavage of benzyl halides substituted by groups such as p-CN and p-CO$_2$Me. In these cases, we have shown\(^{30}\) that, in accordance with the Bunnett and Rossi rule,\(^{31}\) these systems will undergo an inverse mesolytic cleavage (eq 12b). The halide thus depar as a radical, while the negative charge remains on the benzylic system. The said activating groups are capable of hydrogen bonding to proton donors. Added methanol was indeed found to catalyze only the reactions of the p-CN and the p-CO$_2$Me derivatives and, in line with the inverse cleavage mechanism, did not affect the rate of the other substituted benzyl halides. Thus, in the first
step, the MeOH forms a hydrogen bond with a sigma lone pair on these activating groups. When the electron is transferred to the \( \pi \) system of the ring, a concomitant tightening of the hydrogen bond in the orthogonal sigma system takes place to form the covalent N–H bond that neutralizes the negative charge on the ring (eq 13 for \( p\)-CN). Departure of a Br radical and aromatization of the \( p \)-quinoidic structure lead eventually to formation of the corresponding toluene.

\[
\text{ArCH}_2\text{X} + \text{Sml}_2 \rightarrow \text{ArCH}_2^\cdot + X^\ominus \quad \text{normal (12a)} \\
\text{ArCH}_2^\Theta + X^\cdot \quad \text{inverse (12b)}
\]

\[
\text{MeO}^\cdot \cdots \text{CN} \begin{array}{c}
\text{CH}_2 \cdots \text{Br} \\
\rightarrow \text{Sml}_2 \\
\rightarrow \text{H} \cdots \text{NC} = \text{CH}_2 \cdots \text{Br}
\end{array} 
\]

\[
= \text{Br}^* \\
\rightarrow \text{H} \cdots \text{NC} = \text{CH}_2 \rightarrow \text{NC} \cdots \text{CH}_2
\]

\( (13) \)

**INNER AND OUTER SPHERE ELECTRON TRANSFER**

The issue of inner versus outer sphere electron transfer in the reactions of \( \text{Sml}_2 \) was the subject of several publications.\(^{13,32,33} \)

Thus, when the charge is localized on an electrophilic atom of the substrate, having the \( \text{Sm}^{3+} \) in its close vicinity is energetically highly beneficial, leading to an inner sphere electron transfer mechanism. However, in cases where the negative charge is stabilized by delocalization, as in arenes, close interaction with the \( \text{Sm}^{3+} \) is counterproductive as it causes charge localization. Therefore, an outer sphere electron transfer mechanism is preferred in these cases. Consequently, an intensive coordination of ligands to \( \text{Sml}_2 \) may prevent the association of the \( \text{Sm}^{3+} \) with an electronegative atom on the radical anion and force an outer sphere electron transfer mechanism in cases where an inner sphere is preferred. This is nicely exemplified in the photocatalyzed reactions of benzyl chloride and anthracene with EG as an additive.\(^{34} \)

Increasing the concentration of the ligand reduces the rate for benzyl chloride but enhances the rate for anthracene (cf. Figures 14 and 15).

In the case of benzyl chloride, a voluminous coordination sphere around the \( \text{Sm}^{3+} \) interferes with its ability to stabilize the negative charge on the departing chloride ion, leading to rate retardation. The reaction of anthracene, however, is enhanced because the stabilization of \( \text{Sm}^{3+} \) by the ligand hampers the electron from hopping back to the \( \text{Sm}^{3+} \), and at the same time provides proton donors for the protonation of the radical anion.

To conclude this section, in photoinduced reactions, when ligand addition causes rate retardation, the reaction proceeds by inner sphere electron transfer. On the other hand, a rate enhancement upon ligand addition indicates with high likelihood an outer sphere electron transfer.

In this respect, it is interesting to analyze ground state reactions in a similar manner.\(^{35} \)

![Figure 14. Pseudo-first-order rate constants for the photoinduced reaction of benzyl chloride as a function of the concentration of ethylene glycol. Reproduced with permission from ref 34. Copyright 2019 Wiley-VCH.](https://dx.doi.org/10.1021/acs.accounts.0c00497)

![Figure 15. Pseudo-first-order rate constants for the photoinduced reaction of anthracene as a function of the concentration of EG. Reproduced with permission from ref 34. Copyright 2019 Wiley-VCH.](https://dx.doi.org/10.1021/acs.accounts.0c00497)

Whether the electron is transferred by an inner or an outer sphere mechanism depends on the mode by which the substrate stabilizes the negative charge of the radical anion. We showed that the Coulombic attraction between the \( \text{Sm}^{3+} \) and a localized negative charge in the radical anions of benzophenones contributes tens of kilocalories per mole to the stability of the system.\(^{50} \)

![Figure 16. Effect of coordination number on factors that affect the reaction rate.](https://dx.doi.org/10.1021/acs.accounts.0c00497)
SmI₂ coordination to the lone pair, thereby significantly reducing the reaction rate.

QUANTITATIVE DETERMINATION OF EQUILIBRIUM CONSTANTS OF SUBSTRATES AND LIGANDS WITH SMII₂

We used two methods to determine the equilibrium constants for the complexation of a ligand to SmI₂. One is based on cyclic voltammetry and the other on NMR. The first method was used to determine the equilibrium constants for tridentate ligands shown in Chart 2, as well as the contribution of each added ligand to the reduction potential. This was enabled as the CV gave several peaks, each corresponding to a different coordination number (Figure 17). The three successive equilibrium constants for ONO, for example, are 209 ± 32, 444 ± 95, and 313 ± 120 M⁻¹, and the corresponding reduction potentials are 1.55, 1.82, and 2.05 V, respectively.

The NMR method, which is much more accurate than the CV, was based on the nearly abandoned concept of lanthanides as shift reagents. We used it for the determination of the equilibrium constants for binding ligands and substrates. Figure 18 shows the effect of various SmI₂ concentrations on the chemical shift of p-chloroethylbenzene.

The equilibrium constant for the ligation of the first molecule of HMPA to SmI₂ was found to be ca. 2500 M⁻¹, compared with 103 M⁻¹ for EG and only 4 M⁻¹ for MeOH.

A unique mechanistic insight was gained by analyzing the equilibrium constants with arenes and haloarenes. The data shows that SmI₂ binds to the benzene ring rather than the chlorine atom despite the availability of lone pairs on the latter. Thus, in the case of benzyl chloride, the data suggests that the reaction takes place in two steps. In the first step, the SmI₂ binds to the aromatic nucleus, and in the second step, it migrates to the chlorine atom to assist in its departure (eq 14).

EPILOGUE

SmI₂ has now entered its fifth decade in organic chemistry. A plot of the publications per year in this field during this period as a function of SmI₂-related words in the Title or Topic (Figure 19) is very telling and informative. Looking at the early years of SmI₂ chemistry, one can see that it took a whole decade until the scientific community realized the huge potential of this reagent. Following this latency period, the annual number of papers exploring the scope and limitations of this reagent, rose within less than a decade by more than 10-fold. In the subsequent decade, the field seems to have attained a measure of maturity. Interestingly, if we follow the annual number of publications in which SmI₂ appears in the “Title”, there is a decline around the year 2000. This presumably reflects the fact that SmI₂ had become pretty much of a conventional reagent, much like NaBH₄. However, its overall use, as evaluated by the “Topic”...
search (Figure 19), seems to remain stable, with a slight increase as more and more people join the “users” group.38 Regarding studies of the reaction mechanism, it is true that “there is always room for more.” Nevertheless, it seems that the basic forces underlying the chemistry of SmI₂ are now largely understood. The contribution of our laboratory to this understanding is firmly based on the foundation laid by those who preceded us in this field. I owe a debt as well, to my contemporaries who supplied a constant flow of novel facts and mechanistic insight, which provided the stimuli and support for our work. We are grateful to them all.

AUTHOR INFORMATION

Corresponding Author

Shmaryahu Hoz — Department of Chemistry and Institute for Nanotechnology & Advanced Materials, Bar-Ilan University, Ramat Gan 5290002, Israel; orcid.org/0000-0003-3097-8264; Email: shoz@biu.ac.il

Complete contact information is available at: https://pubs.acs.org/10.1021/acsa000497

Notes

The author declares no competing financial interest.

Biography

Shmaryahu Hoz was born in Jerusalem. He received a B.Sc. in Chemistry and Physics and a M.Sc. in Chemistry from the Hebrew University and a Ph.D. in Chemistry from Bar Ilan University. After postdoctoral studies at UCSC with Prof. Joseph F. Bunnett, he joined the Chemistry Department at Bar Ilan University in 1975. Professor Hoz held several administrative positions at Bar Ilan University. He served as the Head of the Department of Chemistry and subsequently as Vice President for Research. His research interests are physical organic chemistry, the chemistry of SmI₂, computational nano-technology, and the effects of electric fields on structure and reactivity.

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(37) Based on data gathered from the “Web of Science”, searches for appearance of SmI₂, samarium iodide, samarium diiodide, and Sm(II) in the “Title” or “Topic”.

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