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Nicholas Dolan, Ryan J Scamp, Tzuhsiung Yang, John F. Berry, and Jennifer M Schomaker

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Catalyst-controlled and tunable, chemoselective silver-catalyzed intermolecular nitrene transfer: Experimental and computational studies

Nicholas S. Dolan,[†] Ryan J. Scamp,[†] Tzuhsiung Yang,[†] John F. Berry* and Jennifer M. Schomaker*

Department of Chemistry, University of Wisconsin, Madison, Wisconsin 53706, United States

ABSTRACT: The development of new catalysts for selective nitrene transfer is a continuing area of interest. In particular, the ability to control the chemoselectivity of intermolecular reactions in the presence of multiple reactive sites has been a long-standing challenge in the field. In this paper, we demonstrate examples of silver-catalyzed, non-directed, intermolecular nitrene transfer reactions that are both chemoselective and flexible for aziridination or C–H insertion, depending on the choice of ligand. Experimental probes present a puzzling picture of the mechanistic details of the pathways mediated by $[(^{t}Bu_{3}tpy)AgOTf]_{2}$ and (tpa)AgOTf. Computational studies elucidate these subtleties and provide guidance for the future development of new catalysts exhibiting improved tunability in group transfer reactions.

Introduction

Amines are important functional groups occurring in a diverse array of molecules with significant biological and therapeutic value. One effective strategy for streamlining the synthesis of amines involves the selective introduction of a new C-N bond at a specific site in a hydrocarbon precursor. Diverse transition metals¹⁻⁸ are known to catalyze the insertion of metal nitrenes into either a C-H or C=C bond to yield amines or aziridines, respectively. Substrate control has traditionally been the primary factor dictating the chemoselectivity of the amination event; for example, intramolecular aminations of homoallylic sulfamates tend to yield the aziridination product using popular dinuclear Rh(II) tetracarboxylate catalysts.² One notable exception is the dinuclear, chiral carboxamidate complex Rh₂(S-nap)₄, which can favor intramolecular C-H bond insertion over aziridination.9 More recently, it has been shown that the chemoselectivity of nitrene transfer can be altered to favor C-H insertion by changing the metal identity to Ru, Co, Mn or Fe, which results in a change from a concerted to a step-wise nitrene transfer event.^{3,4b,5,}

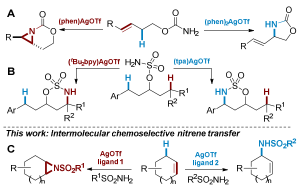
Achieving catalyst control over the chemoselectivity of intermolecular nitrene transfer using a single metal is even more challenging, particularly when compared to metal-catalyzed carbene transfer reactions.¹⁰ A rare example of a Rh-based system selective for intermolecular nitrene insertion into allylic C-H bonds employs $Rh_2((S)-nta)_4$ (nta = (S)-N-1,8-naphthoylalanine) in the presence of chiral sulfonimidamides;¹¹ however, this behavior relies on a strong match between the chiral Rh catalyst and the optically pure nitrene precursor. In general, most dinuclear Rh catalysts that are successful for intramolecular nitrene transfer perform poorly in intermolecular reactions.¹² Two major factors impeding the development of more modular and less expensive catalysts for nitrene transfer include a poor understanding of how reactivity differs in intra- vs. intermolecular processes, as well as how the mechanism and selectivity of the nitrene transfer responds to modifications to the supporting ligands for a particular metal. A better understanding of these issues is necessary to spur further progress towards our ultimate goal of developing new, inexpensive catalysts for predictable, tunable and intermolecular

site-selective nitrene transfer reactions, a long-standing goal in the field.

We have demonstrated that silver(I) catalysis offers unique opportunities to tune the chemo- and site-selectivity of intramolecular nitrene transfer (Scheme 1A-B).^{13,14,15} Changes to the ligand, the AgOTff:ligand ratio, and the nitrene precursor all influence the coordination geometry and reactivity of the putative nitrene. For example, preference for aziridination or C–H bond insertion in a homoallylic carbamate can be controlled by simply changing the ratio of 1,10-phenanthroline (phen) to AgOTf (Scheme 1A).¹³ Enforcing different coordination geometries at Ag(I) using diverse ligands, including 4,4'-di-*tert*-butyl-2,2'-bipyridine ('Bubpy) and tris(2-pyridyl-methyl)amine (tpa), enables site-selective amination of chemically distinct C—H bonds (Scheme 1B).^{14,15} Expanding Ag(I) catalysis to tunable, intermolecular nitrene transfer reactions has been challenging; in this work, we describe initial experimental and computational studies to better understand these reactions (Scheme 1C).

Scheme 1. Ag-catalyzed chemo- and site-selective aminations.

Previous work: Intramolecular chemo- and site-selectivity



Results and Discussion

Initial experimental studies. The reaction of cyclohexene (1), 2,2',2"-trichloroethylsulfamate (TcesNH₂, **2**) and iodosobenzene (PhIO) with Ag catalysts supported by a variety of readily available.

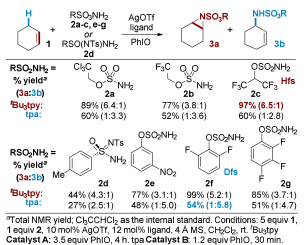
ble N-donor ligands (see the Supporting Information for further details) was carried out (Table 1). We found that AgOTf ligated by *tert*-butylterpyridine **4** (entry 1) gave good selectivity for production of the aziridine **3a** over the allylic amine **3b**.^{7d-f} X-ray crystallographic and DOSY NMR studies showed that the silver catalyst supported by **4** exists in both the solid and solution states as the dimer [($^{t}Bu_{3}tpy$)AgOTf]₂ (hereafter called catalyst **A**), consisting of two three-coordinate Ag centers.^{7e,13} Interestingly, employing the tpa ligand **5** (entry 2) resulted in a preference for allylic C–H amination to **3b** over aziridination. A crystal structure of a silver(I) complex bearing a ligand similar to **5** has been reported, suggesting the formula (tpa)Ag(OTf) for catalyst **B**.¹⁶

Table 1. Impact of the ligand on silver-catalyzed nitrene transfer.

1 AgOTf NHTces NTces ^tBu ligand for ligand + Catalyst A PhIO Cl₃C 0 Q 2 'Ru ^tBu `S Ό `NH₂ 3b 3a Ń <u>entry</u>^a PhIO (ea) 3a : 3b 4 ^tBu₃tpy h vield 4 89% 6.4:1 1 3.5 4 ligand for Catalyst B 5 1.2 60% 1:3.3 2 1 ^aNMR yield; Cl₃CCH₂Cl internal standard Conditions: 5 equiv 1, 1 equiv 2, 10 mol% AgOTf, 12 mol% ligand, PhIO, 4 Å MS, CH₂Cl₂. 5 tpa

Having demonstrated the ability to control the chemoselectivity of Ag-catalyzed amination by changing the nature of the ligand, the identity of the nitrene precursor was varied in an attempt to further improve selectivity (Table 2). While the 3a:3b selectivity for aziridination catalyzed by A was not improved with alkylsulfamate 2c as compared to 2a, the yield did increase to 97%. Other nitrene precursors resulted in either lower yields or decreased selectivity for the aziridine **3a**. When **B** was employed, the C-H amination product was favored to a greater extent employing arylsulfamates 2e-2g, with 2f displaying the highest **3b:3a** ratio of 5.8:1. It is worth noting that in the majority of examples, the overall preference of A for aziridination and that of B for C-H amination is independent of the identity of the nitrene precursor, supporting catalyst-control of nitrene transfer. The one exception is the chiral sulfonimidamide 2d previously employed by Dauban in Rh-catalyzed nitrene transfer; in this case, aziridination was preferred using both A and B.¹¹

 Table 2. Influence of the nitrene precursor on chemoselectivity.



The scope of tunable nitrene transfer was briefly explored with selected substrates containing both C=C and C-H functionalities

(Table 3). Five- and six-membered rings (entries 1-4) give chemodivergent amination. However, as the size of the ring increases (entries 5-6), chemoselectivity for the aziridine product also increases with both catalysts A and B. The lesser selectivity for C-H amination with **B** in this case may be attributed to poor stabilization of an allylic radical via conjugation with the double bond in the 7-membered ring due to competing transannular interactions, providing indirect evidence that **B** may operate *via* a step-wise nitrene-transfer pathway. Reaction of *trans*-hex-2-ene 8 (entries 7-8) gave no isomerization in either aziridination or C-H amination reactions, but did show a rearranged product using **B** (entry 8), also indicative of a potential stepwise pathway. A is not overly sensitive to steric effects, giving excellent selectivity for aziridination of both tri- and tetrasubstituted alkenes 9 and 10 (entries 9, 11). However, achieving selectivity for C-H insertion in acyclic substrates employing **B** is challenging, as aziridination becomes competitive (entries 10, 12). In order to rationalize these observations and achieve a better understanding of the reasons for the lack of chemocontrol, extensive computational studies were carried out as described later in this report.

Table 3. Tunable, catalyst-controlled amination.

R^{1} R^{2} R^{2} R^{2} R^{3} R^{3		AgOTf ligand PhIO 3a,b	NSO ₃ R R ² R ³ 6-10a,b	R ¹ R ³ 3c,d 6-1	IHSO₃R
substrate	entry	ligand	A ^a	l ^a	A : I
6	1	^t Bu₃tpy	61% 6a	11% 6c	5.5 : 1
	2	tpa	13% 6b	35% 6d	1 : 2.7
_н з	3	^t Bu₃tpy	84% 3a	13% 3c	6.5 :1
	4	tpa	8% 3b	46% 3d	1: 5.8
Н 7	5	^t Bu₃tpy	59% 7a	2% 7c	30 :1
	6	tpa	20% 7b	24% 7d	1: 1.2
H 8	7	^t Bu₃tpy	58% ^b 8a	4% ^b 8c	15 :1
	8	tpa	8% 8b	25% ^c 8d	1: 3.1
9	9	^t Bu₃tpy	72% ^b 9a	2% ^b 9c	36 :1
	10	tpa	29% 9b	25% ^d 9d	1.2 :1
10	11	^t Bu₃tpy	97% 10a	0% 10c	> 19 :1
	12	tpa	25% 10b	0% 10d	> 19 :1

A: HfsNH₂ (0.25 mmol), 10 mol% AgOTf, 12 mol% ¹Bu₃tpy, 3.5 equiv PhIO, CH₂Cl₂, 4 A MS, 4 h, rt. I: DfsNH₂ (0.25 mmol), 10 mol% AgOTf, 12 mol% tpa, 1.2 equiv PhIO, CH₂Cl₂, 4 A MS, 1 h, rt. ^aIsolated yield. ^bNMR yield, Cl₃CCH₂Cl internal standard. ^c*trans* only, 6% rearranged. ^d3% rearranged.

A series of substituted cyclohexenes of varying complexity were investigated further in tunable silver-catalyzed amination (Table 4). Addition of a Me group to the alkene of 11 increases selectivity for aziridination with A (entry 1), and improves the preference for C–H amination with **B**; amination of the less hindered allylic C-H_a bond is favored over C-H_b by a ratio of 3.0:1 (entry 2). Aryl-substituted alkenes 12 and 14 (entries 3 and 7) resulted in good selectivity using A, but competing epoxidation occurs with electron-rich 13 (entry 5). Excellent selectivity for C-H amination is observed for 12-14 using **B** (entries 4, 6, 8). Catalyst **B** shows sensitivity to steric effects, as amination of the less congested H_a in 12-14 is preferred over the more hindered H_b by a ratio of approximately 6:1 in all three cases. Conjugation of an alkyne with the cyclohexene of 15 gives good chemoselectivity with both catalysts (entries 9-10); however, the aziridine is not stable to chromatography, undergoing rearrangement and hydrolysis to aldehyde 15a.¹⁷ Good site-selectivity for the amination of H_a over $H_{\rm b}$ was also noted using 15 (entry 10). Moving the Me group from the vinyl carbon of **11** to the allylic position in **16** (entries 11-12) decreases selectivity for aziridination with A. In contrast, the selectivity of C-H amination was improved with B; however, allylic transposition occurs to give 11d and 16d. The addition of an allylic Ph group in 17 further decreases selectivity in aziridination

 Table 4. Tunable amination of C=C and C-H bonds.

R ¹	11-20	+	0 0 RO ^{´S} 2c or	NH ₂	OTf and nIO	NSO ₃ R	11a,b-20a	ı,b ⁺	R ¹	11c,d-20c,c ISO₃R	I
substrate	entry	ligand	A ^a	l (H _a + H _b) ^a	A : I	substrate	entry	ligand	A ^a	l (H _a + H _b) ^a	A : I
H _b Me	1	^t Bu₃tpy	73% 11a	7% 11c	10 : 1	Me	11	^t Bu₃tpy	40% 16a 1:4.8 <i>syn:anti</i>	16% ^e 11c	2.5 :1
11	2	tpa	7% 11b	61% 11d 3.0:1 H _a :H _b	1 : 8.7		12	tpa	3% 16b	39% ^e 11d	1 : 13
Ĥ _a H _b				16% 12c		Ph 17	13	^t Bu₃tpy	22% 17a 1:19 <i>syn:anti</i>	20% ^e 12c	1.1 : 1
C ₆ H ₅	3	^t Bu ₃ tpy	73% ^b 12a	3.4:1 H _a :H _b	4.6 : 1		14	tpa	2% 17b	32% ^e 12d	1 : 16
H _a 12	4	tpa	20% 12b	60% 12d 5.6:1 H _a :H _b	1 : 3.2	18 AcO	15	^t Bu₃tpy	65% 17a 1:11 syn:anti	10% 18c	6.5 :1
H _b 4-MeOC ₆ H	I ₄ 5	^t Bu₃tpy	0% ^c 13a	0% 13c		Т Т _н	16 OAc	tpa	12% 18b	49% 18d 4.4:1 H _a :H _b	1 : 4.1
H _a H _b 13	6	tpa	2% 13b	65% 13d 6.4:1 H _a :H _b	1:33	19	\downarrow	^t Bu₃tpy	55% 19a (<i>dr</i> 2.4:1) (a:b >20:1)	19% 19c (<i>dr</i> 3.9:1)	2.9 : 1
4-CIC ₆ H ₄	7	^t Bu₃tpy	57% ^b 14a	6% 14c 5.0:1 H _a :H _b	9.5 : 1	Haa	, 18	tpa	3% 19b	46% 19d H_a (2.2:1 <i>anti:syn</i>)	1 : 15.3
H _a H _b ⁿ Pent	8	tpa	3% 14b	51% 14d 5.9:1 H _a :H _b	1 : 17		19	^t Bu₃tpy	72% 20a (a:b 2.4:1)	0% 20c	>19 : 1
	9	^t Bu₃tpy	51% 15a ^d	17% 15c 9.0:1 H _a :H _b	3.0 : 1	20 H _b		tpa	29% 20b (a:b 6.3:1)	18% 20d >19:1 H _a :H _b	1.6 : 1
H _a 15	10	tpa	8% 15b	62% 15d 11:1 H _a :H _b	1 : 7.8	Me ⁻ OMe	a		(4.2 0.0.1)		

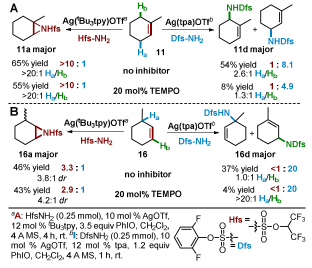
A: 10 mol% AgOTf, 12 mol% ¹Bu₃tpy, **2c**, 3.5 equiv PhIO, CH₂Cl₂, 4 A MS, 4 h, rt. **I**: 10 mol% AgOTf, 12 mol% tpa, **2f**, 1.2 equiv PhIO, CH₂Cl₂, 4 A MS, 1 h, rt. ^a Isolated yields. ^b Aziridine was opened with MeOH. ^c Formation of the epoxide as the major product. ^d 56% NMR yield of aziridine, isolated as the aldehyde in 51% yield. ^e Transposition of the allylic bond was noted to give the trisubstituted alkene as the product. The ratio of expected to allylic product was approximately 1:1 while transposition was highly dominant for **17**.

(entry 13), but showed increased preference for insertion to **12d** (entry 14). These experiments provide insights into the mechanistic details of Ag-catalyzed intermolecular nitrene transfer that will be further discussed in the context of our computational studies (*vide infra*).

A series of natural product-derived substrates 18-20, containing multiple alkene and allylic C-H functionalities, were explored. Alkene 18 gives good selectivity for aziridination and a high 11:1 anti:syn ratio (entry 15). The sensitivity of B to steric effects is illustrated in entry 16, where activation of H_a is now preferred over H_b by a ratio of 4.4:1 (compare to entry 2). The α -ionone derivative **19** shows poorer selectivity for aziridination (entry 17), perhaps due to the increased steric demand of the more electronrich alkene a. Both chemo- and site-selectivity for C-H insertion (entry 18) are excellent. Finally, only aziridination of bisabolyl methyl ether 20 is observed with A (entry 19), with the exocyclic alkene preferred over the endocyclic alkene by a ratio of 2.4:1. The presence of two electron-rich alkenes in 20 results in poor selectivity for C-H amination (entry 20); however, site-selectivity for the less hindered H_a is excellent, favoring H_a over H_b by a ratio of > 19:1.

Mechanistic studies. Further experimental studies were carried out to better understand the mechanistic details of amination pathways promoted by **A** and **B** with both cyclic and acyclic substrates. In Scheme 2A, subjecting **11** to aziridination conditions in the absence and presence of TEMPO as a radical inhibitor showed a modest 10% decrease in yield, which is not a definitive indication that radical intermediates are involved. In contrast, subjecting **11** to C–H amination conditions did result in a substantial decrease in yield in the presence of TEMPO, indicating the possibility of a radical intermediate. These results suggest that catalysts Aand B operate under mechanistically distinct pathways. However, we note that the C–H amination products of 16 and 17 employing either catalyst A or B contain a transposed double bond (Table 4,

Scheme 2. Influence of TEMPO on the amination of 11 and 16.

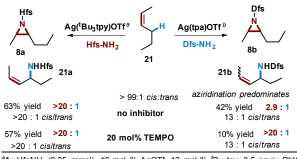


entries 11-14). This suggests *both* silver catalysts are capable of promoting step-wise nitrene transfer pathways involving radical intermediates. Surprisingly, TEMPO inhibition studies with **16** (Scheme 2B) show decreased yields of the insertion product, but

have little effect on the aziridination yield. Explaining these puzzling results is important for developing better Ag(I) catalysts and were addressed using computational studies (*vide infra*).

Further mechanistic studies with acyclic alkenes (Schemes 3-4) reveal that catalyst **A** generates **21a** exclusively as the *cis* isomer (Scheme 3), while **B** gives **21b** as a 13:1 mixture of *cis:trans* isomers. Both catalysts favor aziridination, although the selectivity with **B** (2.9:1) is much lower than that observed with **A** (>20:1). Slight isomerization of the alkene from >20:1 *cis:trans* to 13:1 *cis:trans* in the C–H amination product **21b** occurs in both the absence and presence of TEMPO.

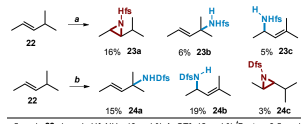
Scheme 3. Influence of the radical inhibitor TEMPO on the amination of acyclic alkene 21.



^aA: HfsNH₂ (0.25 mmol), 10 mol % AgOTf, 12 mol % ^tBu₃tpy, 3.5 equiv PhIO, CH₂Cl₂, 4 A MS, 4 h, rt. ^bI: DfsNH₂ (0.25 mmol), 10 mol % AgOTf, 12 mol % tpa, 1.2 equiv PhIO, CH₂Cl₂, 4 A MS, 1 h, rt.

While the addition of TEMPO using catalyst **B** supported the presence of radical intermediates in the nitrene transfer pathway, the minimal isomerization of **21** (Scheme 3) with both **A** and **B** was unexpected. To shed more insight into this question, a sterically congested alkene **22** was subjected to both aziridination and C–H amination conditions (Scheme 4). These experiments suggest the presence of radical intermediates in both pathways, as transposition of the alkene double was noted using both catalysts. The question was why catalyst **A** appeared in many cases to give concerted nitrene transfer, as opposed to the clear-cut evidence that **B** proceeded through a stepwise nitrene transfer pathway.

Scheme 4. Allylic isomerization studies.



a: 5 equiv **22**, 1 equiv HfsNH₂, 10 mol % AgOTf, 12 mol % ^fBu₃tpy, 3.5 equiv PhIO, 0.05 M CH₂Cl₂, 4 A MS, 4 h, rt. *b*: 5 equiv **22**, 1 equiv DfsNH₂, 10 mol % AgOTf, 12 mol % tpa, 1.2 equiv PhIO, 0.05 M CH₂Cl₂, 4 A MS, 1 h, rt.

Computational studies

The only previously reported, extensive computational study on Ag-catalyzed nitrene transfer explores aziridination using a trispyrazolyl borate complex (catalyst C).^{7a} Aziridination was found to proceed through an initial C–N bond formation step, but a minimum energy crossing point (MECP) between the triplet and closed-shell singlet surface induces direct formation of the aziridine from the C–N bond formation transition state, resulting in retention of the stereochemistry of the alkene. The reactivity observed with our ligands on Ag(I) led to an initial hypothesis that

catalyst A follows a single-step mechanism similar to that described by Pérez and coworkers, and catalyst **B** follows a stepwise pathway.^{7a} However, our experimental probes of the mechanism (Schemes 2-4) yield puzzling results. Experiments in Scheme 2 on cyclic alkenes appear to support our hypothesis, as the use of TEMPO as a radical inhibitor did not impact the aziridination pathways involving catalyst A. The same is true for the acyclic alkene 21 in Scheme 3; however, the data in Scheme 4 suggest that even catalyst A is capable of promoting step-wise nitrene transfer, as allylic C=C double bond transposition is observed. To find a mechanistic explanation for these seemingly contradictory observations, as well as to investigate the reasons for the chemoselectivity differences between A and B, we decided to employ quantum chemical methods to study the reactivity of these two catalysts at the level of density functional theory. Here, B3LYP-D3 is used for the structures of critical points along the reaction coordinates and we employ complete active space selfconsistent field (CASSCF) for the electronic structure of the nitrene complexes. The validity of our choice of B3LYP-D3 is demonstrated by benchmarking various functionals against the crystal structure of catalyst A and B3LYP-D3 was found to be one of the most robust functional in reproducing the observed molecular geometry (see Table S1-4 for quantitative assessments of the functionals).

Structural features of nitrene intermediates. To begin our computational studies, the nitrene complex of **B**, $(tpa)Ag(OTf)(NSO_3R)$ B(N), was optimized with the nitrene ligand bound in the equatorial site (cis to the tertiary amine) and the OTf⁻ ion bound in the axial site. This nitrene complex of catalyst B was chosen for computational studies of its reactivity owing to the fact that the chemoselectivity of **B** is affected by the identity of the Ag salt employed supporting the hypothesis that the anionic OTf ligand remains bound to B(N) during catalysis (See the Supporting Information for the discussion of other potential structure of **B(N)**. Catalyst A has been shown to exist as a dimer with one of the two OTf ions dissociated in its crystal structure, leaving two inequivalent Ag sites.^{7e} Ag_N has a coordination sphere composed entirely of ^tBu₃tpy N atoms, while Ag₀ also has a coordinated OTf ion. To improve computational efficiency, the tert-butyl groups on the ^tBu₃tpy ligand were replaced by H atoms ($[(tpy)AgOTf]_2^+$, called A'). The nitrene complex of A' was optimized with the nitrene ligand bound to the Ag_N, denoted as A'(N). This coordination isomer was chosen due to the fact that catalyst A displays minimal counteranion effects. The effect of the truncation on the structures of A and A(N) is also assessed in Figures S1-3 and S1-4 in the SI (see Tables S1-9, S1-20 and S1-22 for a quantitative comparison and A'(N) and A'(N₀), nitrene binding to the AgO atom of catalyst A, between A and A', and A(N) and A'(N), respectively). The optimized structures of **B(N)** and **A'N** are shown in Figure 1.

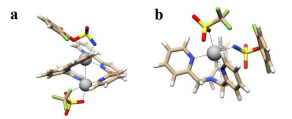


Figure 1. The optimized structures of the Ag-nitrene intermediates investigated. **B(N)** and **A'(N)** are shown on the left (a) and right (b), respectively.

Electronic features of nitrene intermediates. The electronic structures of B(N) and A'(N) have been investigated both at the density

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functional (DFT) and complete active space self-consistent field (CASSCF) levels of theory. For both B(N) and A'(N), the lowest energy state was found by DFT to be a triplet state, in which a Ag(II) ion is ferromagnetically coupled to a doublet nitrene anion radical $(Ag(II)\uparrow \uparrow \{NSO_3R\}^-)$. The ferromagnetic alignment of spins in this complex is due to the orthogonality of the Ag–N σ^* and N p_{π} magnetic orbitals. Analysis of the Kohn-Sham orbitals shows that the Ag–N bond is composed of a 3-electron σ component and a 3-electron π component, with an overall calculated Mayer bond order of 0.56 and 0.57 for the (antiferromagnetically coupled) open-shell (OS) singlet and triplet B(N), respectively, and 0.63 and 0.55 for OS singlet and triplet A'(N), respectively. The σ^* orbital in both complexes is strongly covalent with ~20% polarization toward Ag, whereas the π^* orbital is strongly polarized towards the nitrene N atom by 85-90%. The OS singlet excited state is low-lying, at 4.13 and 8.27 kcal mol⁻¹ above the triplet ground state for B(N) and A'(N), respectively. The frontier orbitals are shown in Figure S1-5. Analysis of the CASSCF wave function indicates that there is only one configuration that contributes strongly to the electronic structure of both complexes in their triplet state. This configuration can be described as $(\sigma)^2 (\pi^*)^2 (\pi^{**})^1 (\sigma^*)^1$. On the other hand, the OS singlet has multiconfigurational character with a major configuration (~70%) of $(\sigma)^2(\pi'^*)^2(\pi''^*)^2(\sigma^*)^0$ and a minor configuration (~30%) of $(\sigma)^2(\pi'^*)^2(\pi''^*)^0(\sigma^*)^2$. A second closed-shell (CS) singlet excited state, discussed previously in the literature,^{7a} was located 7.23 and 12.1 kcal mol⁻¹ above the triplet ground state for B(N) and A'(N), respectively. This particular excited state, which could be described as either a Ag(I) ion bound to a singlet nitrene or a Ag(III)-imido complex, is not discussed due to its high energy.

We recently discussed the necessity for a metal-nitrene intermediate to have a pair of empty α and β molecular orbitals with N character in order to be able to undergo concerted nitrene transfer.¹⁸ Given this constraint, we would expect nitrene transfer from these silver complexes with triplet ground states (i.e., having two empty β molecular orbitals with N character) to occur via a stepwise mechanism. However, Pérez and coworkers found in their computational study that it is possible for intersystem crossing to occur directly following an initial single-electron transition state in such a way as to avoid radical-containing intermediate species.^{7a} Thus, an examination of the nitrene transfer potential energy surface (PES) for both **A'(N)** and **B(N)** is warranted.

Reactivity landscape for nitrene transfer. PESs for nitrene transfer from **B**(N) to substrate 1 modeling experimental chemoselectivity. Two distinct transition states (TSs) have been located as substrate 1 approaches B(N). The first involves partial formation of an N-H bond (C-H bond amination (I), TS_{tpa,I,1}, the numerical subscript denotes the substrate identity) on either the triplet (³TS) or OS singlet (^{OS1}TS) potential energy surface (PES). The second TS involves partial formation of a C-N bond (aziridination (A), $TS_{tpa,A,1}$ on the triplet (³TS) PES. Both $^{OS1}TS_{tpa,I,1}$ and $^{3}TS_{tpa,I,1}$ were found to be lower in energy than the $^{3}TS_{tpa,A,1}$ (10.1 and 10.2 vs 10.7 kcal·mol⁻¹, respectively), consistent with the experimentally observed chemoselectivity. After HAT is complete, intermediates (Int) were found on both the OS singlet $(^{OS1}Int_{tpa,I,1})$ and triplet (³Int_{tpa,I,1}) PES (-19.2 and -19.7 kcal mol⁻¹, respectively), with a slight favoring of the triplet state. Radical rebound occurs without an energetic barrier after the complete formation of $OSI_{Int_{tpa,I,1}}$ on the OS singlet surface to give the aminated product and regenerate catalyst **B** (-71.4 kcal mol⁻¹). The complete reaction coordinate for both reactions catalyzed by B(N) is shown in Figure 2. The presence of an intermediate after hydrogen-atom abstraction on both the OS singlet and triplet PESs and the fact that the two intermediates are close in energy suggest that the ³Int_{tpa.L1} could be long-lived enough to be trapped by radical inhibitors. This finding is consistent with our experimental observations that a radical quencher lowers the yield of the C–H inserted product of **B(N)** in the reactions of **11** and **21** (Schemes 2-3). In contrast, substrate aziridination by **B(N)** occurs on the triplet PES *via* ³**TS**_{tpa,A,1}; no ^{OS1}**TS**_{tpa,A,1} was located. Complete formation of the C–N bond gives ³**Int**_{tpa,A,1}; a reorganization of the electronic structure is presumed to occur to give ^{OS1}**Int**_{tpa,A,1} (kcal·mol⁻¹ in terms of vertical excitation energy), which then leads to the formation of the aziridine product, ¹**PC**_{A,tpa,1}. Similar to the case for C–H insertion, the presence of ³**Int**_{tpa,A,1}, along with the absence of a fast recombination ^{OS1}**TS**_{tpa,A,1}, suggest that trapping by radical inhibitors is possible (*vide infra*), a hypothesis that is consistent with our experimental observations.

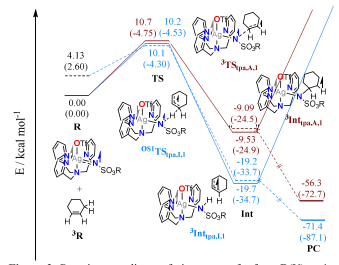


Figure 2. Reaction coordinate of nitrene transfer from B(N) to 1 to give either the C-H amination (blue) or aziridine product (brown). **R** = reactant, **TS** = transition state, **Int** = intermediate, and PC = product complex. Subscript I = C-H insertion and A =aziridination: superscript OS1 and 3 denote open-shell singlet and triplet spin states, respectively. Solid lines denote the triplet potential energy surface (PES) and dashed lines denote the openshell singlet PES. Arrows and the size of the arrows represent α or β spin populations for the electrons on Ag (silver), blue for N (blue), and black (organic substrate). Black arrows inside the ring indicate that electron density is delocalized over the allylic radical and arrows outside the ring represent electron density localized on the carbon proximal to the arrow. Numbers shown are Gibbs free energies; numbers enclosed in parentheses are enthalpies, both taking into account the solvent (CH₂Cl₂). Note: the energy for the blue dashed line enclosed by quotation marks was obtained from converging the single point energy at the OS singlet state based on the geometry of the solid blue line below it.

PESs for nitrene transfer from A'(N) to substrate 1 modeling experimental chemoselectivity. Four distinct TSs are located as 1 approaches A'(N). Partial formation of the N–H bond (TS_{tpy,I,1}) or C–N bond (TS_{tpy,A,1}) occurs on either the triplet or OS singlet PES. ${}^{3}TS_{tpy,I,1}$ and ${}^{3}TS_{tpy,A,1}$ are lower in energy than ${}^{OS1}TS_{tpy,I,1}$ and ${}^{OS1}TS_{tpy,A,1}$ by 5.60 and 7.62 kcal·mol⁻¹, respectively. ${}^{3}TS_{tpy,I,1}$ and ${}^{3}TS_{tpy,A,1}$ lead to the formation of triplet state intermediates, ${}^{3}Int_{tpy,I,1}$ and ${}^{3}TS_{tpy,A,1}$ lead to the formation of triplet state intermediates, ${}^{3}Int_{tpy,I,1}$ and ${}^{3}Int_{tpy,A,1}$ (-16.0 and -8.32 kcal·mol⁻¹, respectively). However, these intermediates do not evolve to form products on the triplet surface. Instead, it was found that ${}^{OS1}TS_{tpy,I,1}$ and ${}^{1}PC_{tpy,A,1}$ lead to the direct formation of product complexes, ${}^{1}PC_{tpy,I,1}$ and ${}^{1}PC_{tpy,A,1}$, respectively (-81.7 and -62.5 kcal·mol⁻¹, respectively) without accessing an intermediate. The

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complete reaction coordinate for both reactions by A'(N) is shown in Figure 3. The experimental observation that A is resistant to inhibition by TEMPO (Schemes 2 and 3) suggests that direct access of the OS PES after ³Int_{tpy} (the crosses in Figure 3) to yield products is the major reaction pathway (*vide infra*).

The shape of the PES and the experimentally observed tolerance of the aziridination pathway towards radical inhibition are strongly reminiscent of the system described by Pérez and coworkers for Ag complexes supported by tripodal ligands.^{7a} These resemblances prompted us to perform a comparison between **B**(**N**), **A'**(**N**), and the Pérez (Tp^{Me2})Ag(NTs) (**C**(**N**); Tp^{Me2} = hydro(3,5dimethyl-1-pyrazolylborate)) system in order to achieve a deeper understanding of reactivity in our silver-catalyzed intermolecular nitrene transfer reactions.

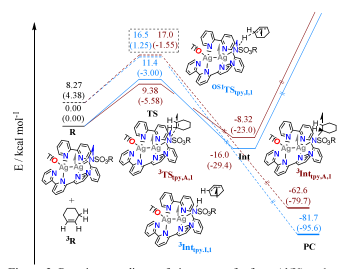
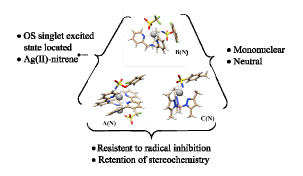


Figure 3. Reaction coordinate of nitrene transfer from A'(N) to 1 to give either C–H insertion (blue) or aziridine products (brown). **R** = reactant, **TS** = transition state, **Int** = intermediate, and **PC** = product complex. Subscript **I** = C–H insertion and **A** = aziridination and superscript OS1 and 3 denote open-shell singlet and triplet spin states, respectively. Solid lines denote the triplet potential energy surface (PES) and dashed lines denote the open-shell singlet PES. Arrows and the size of the arrows represent α or β spin populations for the electrons on Ag (silver), N (blue), and organic substrate (black). Black arrows inside the ring represent electron density delocalized over the allylic radical and arrows outside the ring represent electron density localized on the carbon proximal to the arrow. Numbers shown are Gibbs free energies and numbers enclosed in parentheses are enthalpies, both taking into account the solvent (CH₂Cl₂).

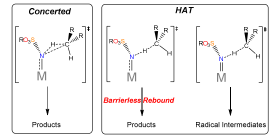
Differential mechanisms for nitrene transfer from B(N), A'(N), and C(N) to organic substrates. Results for the reaction coordinates of nitrene transfer from B(N) and A'(N) to substrate 1 (vide supra) contrast with those recently reported for Ag-nitrene intermediates supported by tripodal Tp ligands.^{7a,19} In previously reported examples, only a triplet and CS singlet state were located, with the triplet favored in energy by 7.6 kcal·mol^{-1,7a} The large spin population of 1.32 on the N atom, along with the inability to locate an OS singlet state, led to the description of C(N) as a Ag(I) complex with a coordinated triplet nitrene ligand. In contrast, we located OS singlet states for B(N) and A'(N) with spin populations (-0.78 on N and 0.78 on (tpa)Ag for B(N) and -0.74 on N and 0.74 on (tpy)₂Ag₂ for A'(N)), which prompt us to describe these silver nitrene complexes as Ag(II) ions coupled to nitrene anion radicals, despite the fact that spin densities on the triplet states are similar to that of C(N) (1.26 on N and 0.74 on (tpa)Ag for B(N) and 1.27 on N and 0.64 on (tpy)₂Ag₂ for A'(N)).

Unlike either A'(N) or C(N), B(N) performs nitrene transfer via a mechanism that generates Int_{tpa} on both the triplet and OS singlet PESs. This allows radical inhibition of the reaction, in contrast to the tolerance towards TEMPO exhibited by A'(N) and C(N) (Scheme 5). Importantly, the divergence in mechanism between these three catalysts occurs after the initial transition state. Therefore, the differences in the mechanism for nitrene transfer from B(N), A'(N), and C(N) are not likely due to their different ground state electronic structures. Instead, the nature of the transition states needs to be examined along with the thermodynamic driving forces *en route* to product formation.





Nature of the C–H amination transition states from A(N), B(N), and C(N). It is useful to examine the general geometrical features of nitrene transfer transition states. For simplicity, we will analyze in detail only the C-H amination transition states. In concerted insertions of nitrenes into C-H bonds, as is seen in Rh2-catalyzed reactions, the transition state involves simultaneous (though asynchronous) C-N and N-H bond formation and C-H cleavage.¹⁸ These three interactions take place within a triangular, threemembered ring transition state (left, Scheme 6). Stepwise reactions proceed first through a hydrogen atom transfer (HAT) transition state leading to radical intermediates. The transition state involves a linear N···H···C geometry (right, Scheme 6). These geometries are electronically dictated: the three-membered ring transition state requires a stabilizing pair of electrons and therefore an empty N orbital (or a pair of α/β holes with N characters to accept the electrons of the C-H bond. The linear N···H···C structures result from three-center/three-electron or three-center/fourelectron bonds and are



Scheme 6. Diagrams of concerted and HAT transition state geometries for C–H amination, including the features of stepwise transition states vs those leading to barrierless rebound.

seen for the more electron-rich systems. All silver-nitrene complexes are electron rich and will therefore use HAT transition states for C–H amination (and corresponding single-electron transition states for aziridination). The major question of this work,

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59 60 however, is to explain when the HAT transition state leads to the formation of radical intermediates vs when it is followed by a *barrierless* rebound.

Geometric details of the C-H amination transition states of catalysts A', B, and C are given in Table 5 (a more complete set of data for catalyst A' and B is given in Tables S1-23, S1-24 and Tables S1-27, S1-28, respectively, in the SI). Since catalyst A performs C-H amination via a barrierless recombination pathway and **B** via a stepwise pathway, it is useful to compare their geometric differences. Both catalysts feature typical HAT transition states with N···H···C angles very close to linearity. However, the lowest-energy transition state (triplet) for catalyst A' also clearly shows signs of Ag-N bond cleavage, since its Ag-N bond length is increased by 0.05 Å from the geometry of A'(N). The Ag-N bond length for **B** is unchanged (0.01 Å). The observation that transition states leading to barrierless recombination involve Ag-N bond cleavage is also true of aziridination transition states. Catalysts A and C undergo aziridination via barrierless rebound, and their Ag–N bond distances are elongated by 0.06 and 0.04 Å, respectively, in their transition state geometries.

Table 5. Geometrical features of optimized transition states.

	C-C Aminatio	n Transition States				
Catalyst	Spin State ^a	∠N…H…C, °	$\Delta d(Ag-N)^{b}$			
			Å			
A'	Singlet	171	0.03			
	Triplet	172	0.05			
B	Singlet	173	- 0.03			
	Triplet	176	0.01			
Aziridination Transition States						
Catalyst	Spin State ^a	$\Delta d(N - C)^{c}$,	$\Delta d(\text{Ag} - \text{N})^{\text{b}}$			
		Å	Å			
A'	Singlet	0.627	0.01			
	Triplet	0.736	0.06			
В	Triplet	0.626	0.03			
С	Singlet	0.335	- 0.06			
	Triplet	0.625	0.04			

^a The singlet spin states are all optimized open-shell singlets.

^b $\Delta d(Ag-N)$ is defined as the difference in Ag-N(nitrene) bond distance between the transition state geometry and the reactant complex geometry.

^c $\Delta d(N - C)$ is defined as the difference between the two N···C distances in the transition state.

This geometrical observation is also supported by an energetic analysis. First, we compare ΔH^{\ddagger} of the singlet and triplet transition states. In contrast to B(N), which has a similar enthalpic driving force at both ^{OS1}TS_{tpa,I,1} and ³TS_{tpa,I,1} ($\Delta\Delta H_{CH_2Cl_2}^{\pm,0S1-3} = 0.23$ kcal·mol⁻¹), ^{OS1}TS_{tpy,1} for A'(N) is much less exothermic than ³TS_{tpy,1} ($\Delta\Delta H_{CH_2Cl_2}^{\pm,0S1-3} = 3.08$ kcal·mol⁻¹ for TS_{tpy,4,1} and ³TS_{tpy,4,1} and $\Delta\Delta H_{CH_2Cl_2}^{+10S1-3} = 3.34 \text{ kcal} \cdot \text{mol}^{-1} \text{ for } \mathbf{TS}_{tpy,I,I}). \text{ Since the } \Delta H_{CH_2Cl_2}^{+10S1-3} \text{ for } \mathbf{TS}_{tpy,I,I}).$ the formation of the first C-N bond in aziridination and the N-H bond in C-H insertion should be very similar on both PESs, the difference in $\Delta H_{CH_2Cl_2}^{\ddagger}$ between $^3TS_{tpy}$ and $^{OS1}TS_{tpy}$ must result mainly from the breaking of the Ag–N bond during $^{OS1}TS_{tpy}$. Comparing the Ag-N bond lengths in B(N) and A'(N), they differ by ~ 0.07 Å, a difference that may be attributed to partial chelating character of the nitrene ligand in A'(N), which contains a weak Ag...O interaction of 2.7 Å, reminiscent of the quasibidentate nitrene coordination mode in C(N). This implies that the bond between Ag and N is stronger in the OS singlet A'(N) than in the triplet A'(N); thus, it takes more energy to break the Ag-N bond during $^{OSi}TS_{tpy}$ to form the bond with the incoming atom

from the substrate ($\Delta(Ag-N)_{TS-R} = +0.062$ Å for ^{OS1}TS_{tpy,A,1} and $\Delta(Ag-N)_{TS-R} = +0.077$ Å for ^{OS1}TS_{tpy,L1} vs $\Delta(Ag-N)_{TS-R} = -0.01$ Å for ^{OS1}TS_{tpa,I}). Once past ^{OS1}TS_{tpy}, the Ag–N bond has elongated enough to form ¹PC_{tpy} in a barrierless radical recombination step. The transient process leading to the formation of ¹PC_{tpy} means that the A'(N)-substrate complex must cross from ³TS_{tpy} to ¹PC_{tpy} through an MECP (where dashed and solid lines cross in Figures 2 and 3). Through this analysis, we observe that nitrene complexes with a semi-chelating coordination mode to Ag lead to nitrene transfer transition states having significant Ag–N bond cleavage character, which results in barrierless recombination and an absence of radical intermediates.

Interplay of enthalpy and entropy of activation on the differential chemoselectivity of nitrene transfer from B(N) and A'(N) to substrate 1. The differential chemoselectivity of nitrene transfer from B(N) and A'(N) to substrate 1 is satisfactorily reproduced by DFT studies of the reaction coordinates on both the triplet and OS singlet PESs $((P_l: P_A)_{1N_{eq}}^{calc} \approx (k_l: k_A)_{1N_{eq}}^{calc} = 2.60:1 \text{ using } {}^{OS1}TS_{tpa,1,I}$ or $(k_l: k_A)_{1N_{eq}}^{calc} = 2.3:1 \text{ using } {}^{3}TS_{tpa,1,I} \text{ vs } (P_l: P_A)_{1N_{eq}}^{exp} = 5.8:1;$ $(P_{I}: P_{A})_{2}^{\text{calc}} \approx (k_{I}: k_{A})_{2}^{\text{calc}} = 1:30.7 \text{ vs } (P_{I}: P_{A})_{2}^{even} = 1:5.2 ;$ see below for the origin of the approximately equal signs). The agreement with experiment justifies a further analysis of thermodynamic parameters, namely enthalpy, entropy, and Gibbs free energy of activation/formation. Activation of the C=C bond on the triplet PES is the most exothermic step among all TSs for both B(N) and A'(N) (Table 6). This is sensible, as breaking the C=C π -bond in an alkene is less energetically costly than breaking the C-H σ-bond at an allylic position.²⁰ However, when the entropy of activation is taken into account, B(N) suffers more than A'(N) for both C-N bond formation and HAT. The fact that B(N) is more negatively impacted by ΔS_A^{\ddagger} is due to steric hindrance around the nitrene site, which restricts the flexibility of the incoming substrate, as can be seen from space-filling models of **B(N)** and A'(N) (Figure 4). This larger negative entropic contribution of ${}^{3}TS_{tpa,1}$ in general, and of ${}^{3}TS_{tpa,A,1}$ compared to ${}^{3}TS_{tpa,I,1}$, is the major thermodynamic factor in the differential chemoselectivity of B(N) and A'(N). Hence, the chemoselectivity of A'(N) towards aziridination of 1 is enthalpy-driven while that of B(N) towards amination of **1** is entropy-driven.

Table 6. Enthalpy (kcal·mol⁻¹), entropy (cal·mol⁻¹·K⁻¹), and Gibbs free energy (kcal·mol⁻¹) of activation for nitrene transfer from **B(N)** to 1 and 21 on either the triplet or OS singlet surface.

D (1) to T and ZT on ender the unplet of OS singlet surface.							
TS	substrate	$\Delta H^{\ddagger}_{CH_2Cl_2}$	ΔS^{\ddagger}	$\Delta G^{\dagger}_{CH_2Cl_2}$			
OS1TS _{tpa,I}	1	-4.30	-48.3	10.1			
³ TS _{tpa,I}	1	-4.53	-49.5	10.2			
³ TS _{tpa,A}	1	-4.75	-51.7	10.7			
³ TS _{tpv,I}	1	-3.00	-48.3	11.0			
³ TS _{tnv} , A	1	-5.58	-50.2	9.38			
USI TS _{tpa,I}	21	-4.89	-57.0	12.1			
³ TS _{tpa,I}	21	-0.51	-47.5	13.7			
³ TS _{tpa,A}	21	-4.69	-53.4	11.2			

Fine-tuning the chemoselectivity of nitrene transfer from B(N) to cyclic vs acyclic alkenes. To further examine our hypothesis that the interplay of ΔH^{\ddagger} and ΔS^{\ddagger} fine-tunes the chemoselectivity of Ag-nitrene complexes, the reaction coordinate for nitrene-transfer from B(N) to acyclic substrate 21 was studied. Replacing cyclic alkenes by acyclic alkenes (i.e., 1 by 21) changes the experimental chemoselectivity of B(N). Correspondingly, we found that the lowest energy TS is the aziridination transition state ${}^{3}TS_{tpa,A,21}$, which is 0.89 kcal·mol⁻¹ more stable than the lowest energy C – H amination transition states $TS_{tpa,I,21}$, ${}^{OS1}TS_{tpa,I,21}$. This gives

 $k_A/k_I = 4.52 \approx P_A/P_I$, assuming that the product-determining steps are the rate-determining steps. Note that there are only two internal secondary allylic C–H bonds in **21**, compared to four accessible C=C sites comprised of C2-*re*, C2-*si*, C3-*re* and C3-*si*. This could potentially increase the P_A/P_I to 9.04. The calculated

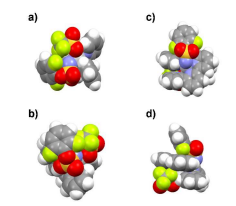


Figure 4. Space-filling models of B(N) a) viewed from the top of the nitrene and b) viewed from the side. Space-filling models of A'(N) c) viewed from the top of the nitrene and d) viewed from the side. Color code: silver:grey, sulfur:yellow, fluorine:green, oxygen:red, nitrogen:blue, carbon:black, and hydrogen:white.

product ratios match the experimental product ratio of 2.9:1 well and, more importantly, the observed chemoselectivity. To understand the influence of ΔH^{\ddagger} and ΔS^{\ddagger} on the change of chemoselectivity from 1 to 21, their contributions to the TS_{tpa} are summarized in Table 6. $\Delta H^{\ddagger}_{CH_2Cl_2}$ is more favorable for HAT than C–N bond formation for 21 as compared to 1, because the migrating allylic C–H bond can adopt a nearly coplanar alignment with the π orbitals of the C=C bond to allow some degree of resonance stabilization as the C-H bond starts to break. This stabilization is less prominent in 1 due to the constraint imposed by the cyclic structure (Figure 5). However, the staggered conformation of 21 forces the propyl substituent to impose steric hindrance on **B(N)**, a feature that is absent in 1. The reason for this is the constraint imposed by the cyclic structure of 1 that forces C4 to stay in a pseudo-gauche conformation with respect to the C=C bond (Figure 6). This steric effect reflects heavily on ΔS^{\ddagger} , hence making ${}^{3}TS_{tna,A,21}$ more energetically accessible than $^{OS1}TS_{tpa,I,21}$.

The C=C cis/trans isomerization of acyclic alkene 21 during nitrene transfer from B(N). Radical intermediates are found on both triplet and OS singlet surfaces for **B**, with ³Int_{tpa,I} lower in energy by only 0.6 kcal·mol⁻¹. However, ${}^{3}Int_{tpa,I}$ cannot lead to the formation of products without first accessing ${}^{OS1}Int_{tpa,I}$ via a minimum energy crossing point (MECP: where crosses are located in Figure 3) to allow the conservation of spin symmetry. The presence of an MECP after the initial formation of ³Int_{tpa,I} suggest that this species will be relatively long-lived. In turn, this increased lifetime allows cis/trans isomerization of the allylic C=C bond via C-C rotation of the diradical ³Int_{tpa,I} species to occur. This also explains why TEMPO inhibition is observed (Scheme 3). Similar arguments apply to $Int_{tpa,A}$. To further support this hypothesis, the *trans*-isomer of $Int_{tpa,21}$ ($Int_{tpa,21}^{iso}$) was considered. Indeed, ${}^{3}Int^{iso}_{tpa,A,21}$ and ${}^{3}Int^{iso}_{tpa,I,21}$ were located and found to be more stable than their *cis*-isomers $(-7.02 \text{ kcal mol}^{-1})$ for ${}^{3}Int_{tpa,I,21}^{iso}$ and -3.55 kcal·mol⁻¹ for ${}^{3}Int_{tpa,A,21}^{iso}$). This result implies that a pre-equilibrium between the two isomers occurs before radical recombination to yield the trans isomer as the major product. However, the distribution of different isomers also

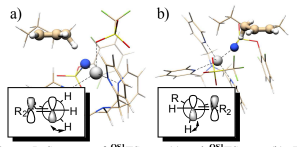


Figure 5. Structure of $^{OS1}TS_{tpa,I,1}$ (a) and $^{OS1}TS_{tpa,I,21}$ (b). The allylic group, nitrene N atom, and the Ag atom are shown in spheres for emphasis. Insets: Neumann projections of substrate looking down along C2–C3 bond (a) or C3–C4 bond (b).

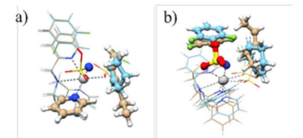


Figure 6. a) Overlay of 3 Int_{tpa,A,21} (mostly in beige) and 3 Int_{tpa,A,1} (mostly in blue). Ag, the nitrene N, the pyridyl group closest to substrates, and substrates are shown in spheres for clarity. b) Overlay of 3 Int_{tpa,I,21} (mostly in beige) and 3 Int_{tpa,I,1} (mostly in blue). Ag, the aryl sulfamate, and substrates are shown in spheres for clarity.

depends on the kinetic barrier of the isomerization, which, in turn, depends on the specific steric constraints of the substrate and the degree of C=C bond character in $Int_{tpa,21}$, i.e. the C=C bond is completely broken in $Int_{tpa,A,1}$ to give a carbon radical at C3 while the C=C bond is half broken in $Int_{tpa,I,1}$ to stabilize the carbon radical at C4 after HAT through resonance (Figure 7). This can be seen structurally in Figure 7a where the methyl group on C2 and the propyl group on C3 are no longer coplanar in ³Int_{tpa,A,21}.

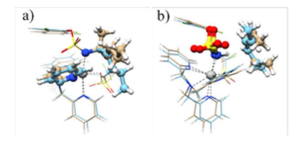


Figure 7. a) Overlay of ³ Int_{tpa,21,A} (mostly in beige) and ³Int_{tpa,21,A} (mostly in blue). The rest of the molecule except Ag, nitrene N, the pyridyl group closed to 21, and substrate 21 is omitted for clarity. b) Overlay of ³Int_{tpa,1,21} (mostly in beige) and ³Int_{tpa,21,21} (mostly in blue), the rest of the molecule except Ag, the sulfamate, and substrate 21 is omitted for clarity.

Timing of intersystem crossing and C=C bond transposition. MECPs are required to be present after the transition states in the reaction trajectories from both A'(N) and B(N). Yet, the two Agnitrene complexes display dissimilar reactivity. As discussed

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above, for catalyst **B**, the MECP must occur *after* intermediate formation and this feature "delays" the formation of products allowing a by-product shunt to occur through bimolecular collision of Int_{tna} with TEMPO. On the other hand, for catalyst A, the MECP must occur before the formation of intermediates, because no radical byproducts are observed, with the exception of C=C bond transposition products seen with substrate 16. Thus, the potential energy surface for this substrate was examined more closely. For this substrate, as the allylic C-H bond on 16 breaks in the (triplet) transition state, electronic reorganization occurs almost instantaneously, as compared to the nuclear movement to form the C-N bond giving products. The HAT nature of the transition state yields an allylic radical with spin population on both C1 and C3. Bond formation between C1 and N occurs immediately on the OS singlet PES to give 16d. The driving force favoring C1 over C3, which is mostly steric, is due to ΔS^{\ddagger} (vide infra), hence giving the thermodynamic product. Indeed, 16d is formed directly (with no intermediates) as the product after $^{OS1}TS_{tpy}$, consistent with experimental observations (Table 1 and Scheme 2). Thus, although C=C double bond transposition in this case was initially suggestive of a stepwise mechanism with radical intermediates, we find computationally that this transposition can occur via a barrierless recombination and does not necessitate a long-lived diradical intermediate. This reactivity is therefore consistent with an HAT mechanism with barrierless rebound from A'(N).

Conclusions

In conclusion, we have reported the first examples of simple silver catalysts capable of tunable, non-directed and intermolecular chemoselective amination. We have found catalyst systems that demonstrate complementary chemoselectivities in alkenes containing sites for both aziridination and C-H amination in nitrene transfer, largely independent of substrate identities. Computational studies showed that the tunable chemoselectivity between tpa and ^tbu₃tpy ligands is a result of the steric profile around the Ag-nitrene intermediates. The means to tune the steric bulk around the nitrene site without changing the electronic structure of the silver-nitrene intermediates, such as the size of the bound counteranion, could potentially afford an additional tunability within the same ligand system. Furthermore, two distinct nitrene-transfer mechanisms are observed in the computational results, the first involving a very late transition-state from catalyst A, followed by a barrierless recombination step that preserves stereochemistry and the second having a very early transition state from catalyst B to yield radical intermediates. The major distinction between these two mechanisms appears to be the extent to which the Ag-N bond breaks during the HAT transition state. Future studies are focused on the development of increasingly selective amination catalysts using a broader array of supporting ligands and counteranions. The ability to correlate the coordination geometry of the catalyst with both reaction mechanism pathways and selectivity are goals currently under investigation.

Supporting Information

Experimental procedures and characterization for all new compounds is included in the Supporting Information. This material is available free of charge via the Internet at http://pubs.acs.org.

Corresponding Authors

schomakerj@chem.wisc.edu, berry@chem.wisc.edu

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