

Communication

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Characterization of a Reactive Rh2 Nitrenoid by Crystalline Matrix Isolation

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Supporting Information Placeholder

ABSTRACT: The fleeting lifetimes of reactive intermediates in C–H functionalization chemistry often prevent their direct characterization. For example, the critical nitrenoid intermediates that mediate Rh₂-catalyzed C–H amination have eluded characterization for more than 40 years. In the absence of structural characterization of these species, methodological development is often computationally guided. Here we report the first X-ray crystal structure of a reactive Rh₂ nitrenoid, enabled by N₂ elimination from an organic azide ligand within a single-crystal matrix. The resulting high-resolution structure displays metrical parameters consistent with a triplet nitrene complex of Rh₂. The demonstration of facile access to reactive metal nitrenoids within a crystalline matrix provides a platform for structural characterization.

Unstabilized nitrenes are reactive, high-energy species that feature a hextet electronic configuration at nitrogen (i.e. N-R species).¹ Nitrenes participate in a diverse reaction manifold, including C-H insertion, addition to C-C multiple bonds, and various unimolecular rearrangements, that render these species challenging to utilize as intermediates in selective synthetic chemistry.² Synthetic chemists³ and biologists⁴ have advanced selective nitrene-transfer chemistry that is predicated on leveraging the reactivity of transition metal-stabilized nitrenoid intermediates for C-H functionalization and olefin aziridination. In particular, Rh₂-catalysis has emerged as a broadly useful platform in nitrene-transfer catalysis (Figure 1).5 Critical Rh₂ nitrenoid intermediates have been detected by mass spectrometry⁶ and time-resolved spectroscopic methods,⁷ but due to their fleeting lifetimes, structural data of these transient species has not been available. Significant questions, such as the preferred electronic configuration of Rh₂ nitrenoids, have not been resolved despite interest.⁸

Unambiguous molecular structure determination can be achieved by X-ray diffraction, but requires chemical samples

that are sufficiently kinetically stable to be crystallized. The inherently transient nature of reactive intermediates typically precludes application of X-ray diffraction to the characterization of these species. Two methods have been advanced to gain structural information about reactive intermediates in the condensed phase: 1) structural characterization of synthetic derivatives designed to attenuate the reactivity of the intermediate of interest, for example via introduction of sterically encumbering ligands,⁹ and 2) spectroscopic characterization of photogenerated reactive intermediates by cryogenic matrix isolation.¹⁰ We envisioned that photogeneration of reactive intermediates within a crystalline matrix would combine classical matrix isolation with X-ray diffraction and enable structural characterization of these species without synthetic derivatization.¹¹ Here, we demonstrate the successful application of this strategy to the characterization of a Rh₂ nitrenoid generated by N2 elimination from a Rh2 alkylazide complex within a crystalline matrix.

We targeted characterization of a nitrenoid supported by $Rh_2(esp)_2(1)$, in which the Rh_2 core is supported by two chelating *bis*-carboxylate ligands, ^{5c,12} because complex 1 has emerged as a particularly effective, and widely utilized,



Figure 1. Rh₂-catalyzed nitrene transfer chemistry has emerged as a leading method for introducing nitrogen content in organic molecules. These reactions are proposed to proceed via transient Rh_2 nitrenoids, which have thus far eluded structural characterization.



Figure 2. Synthesis and steady-state photochemistry of organoazide complexes of Rh₂(esp)₂ (1). Treatment of Rh₂(esp)₂ (1) with AdN₃ results in sequential formation of Rh₂(esp)₂(AdN₃) (2a) and Rh₂(esp)₂(AdN₃)₂ (2b). (a) Thermal ellipsoid plot of 2b drawn at 50% probability with H-atoms and solvent removed for clarity; Rh(1)–Rh(1): 2.3968(8) Å, Rh(1)–N(1): 2.335(3) Å, N(1)–C(1): 1.509(5) Å, N(1)–N(2): 1.254(5) Å, N(2)–N(3): 1.135(5) Å, N(1)–N(2)–N(3): 176.1(4)°. (b) UV-vis spectra collected during the photolysis of Rh₂ complex 2a in CH₂Cl₂ (335 nm < λ < 610 nm). Isosbestic points are observed at 411, 455, and 618 nm, which indicate the lack of a steady-state intermediate in the conversion of 2a to 5a. (c) Thermal ellipsoid plot of 5a drawn at 50% probability with H-atoms and solvent removed for clarity. Selected metrical parameters: Rh(1)–Rh(2): 2.3959(5) Å; Rh(1)–N(1): 2.303(4) Å. (d) UV-vis spectra collected during the photolysis of Rh₂ complex 2b in CH₂Cl₂ (335 nm < λ < 610 nm). An isosbestic point is observed at 603 nm, which indicates the lack of a steady-state intermediate in the conversion of 2b to 5b.

nitrene-transfer catalyst.^{5c,e-h} We initiated our studies by exploring the synthesis of Rh2 complexes with organic azide ligands based on the hypothesis that facile N2 extrusion would provide access to Rh_2 nitrenoids. Exposure of $Rh_2(esp)_2$ to CH₂Cl₂ solutions of AdN₃ resulted in sequential formation of two Rh_2 azide adducts: $Rh_2(esp)_2(AdN_3)$ (2a) and $Rh_2(esp)_2(AdN_3)_2$ (2b). Concentration-dependent UV-vis spectra display isosbestic points connecting $Rh_2(esp)_2$ (1) and $Rh_2(esp)_2(AdN_3)(2a)$ at 0–12 mM [AdN₃] and isosbestic points connecting **2a** and $Rh_2(esp)_2(AdN_3)_2$ (**2b**) at 12– 85 mM [AdN₃] implying the absence of steady-state intermediates in these reactions (Figure S1). Further spectral evolution was not observed upon further addition of AdN₃. Jobs analysis confirms that **2a** is a 1:1 adduct of $Rh_2(esp)_2$ and AdN_3 (Figure S2 and Table S1). Rapid exchange of free and bound AdN₃ is evident in the room temperature ¹H NMR

spectra of **2a** and **2b**. Low-temperature ¹H NMR spectroscopy and electrospray ionization-mass spectrometry (ESI-MS) support the formulation of **2a** and **2b** as *mono-* and *bis*azide adducts, respectively (Figures S3 and S4). The AdN₃ ligands are weakly bound; titration of a tetrahydrofuran solution of Rh₂(esp)₂ with AdN₃ results in no spectral changes, which suggests AdN₃ does not displace bound THF ligands at the apical sites of Rh₂(esp)₂.

Single crystals of **2b** were obtained from cooling a CH_2Cl_2 solution of $Rh_2(esp)_2$ and AdN_3 . X-ray diffraction analysis revealed the structure depicted in Figure 2a in which two symmetry-equivalent AdN_3 ligands are bound to the Rh_2 core via $N(\alpha)$. Efforts to crystallize **2a** by lowering the AdN_3 loading consistently provided *bis*-azide adduct **2b** as the exclusive

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crystallization product, which suggests the preferential crystallization of **2b** over *mono*-azide adduct **2a**. The Rh–Rh distance in **2b** (2.3968(8) Å) is similar to that previously reported for Rh₂(esp)₂S₂ complexes (S = solvent-derived ligands, see Table S2). The N₃ fragment of the AdN₃ ligand is nearly linear (N(1)–N(2)–N(3) = 176.2(4)°), the N(1)– N(2) and N(2)–N(3) distances are similar to those in free AdN₃, and the infrared (IR) spectrum of **2b** displays v_{N3} at 2120 and 2093 cm⁻¹ (Figure S5). These metrics are consistent with AdN₃ binding as a σ -donor with insignificant π -backbonding.¹³

Rh₂ complexes 2a and 2b are photoprecursors to Rh₂ nitrenoids. Photolysis of a CH₂Cl₂ solution of **2a** (335 nm $< \lambda$ < 610 nm) resulted in new spectral features that are accessed via well-anchored isosbestic points at 411, 455, and 618 nm (Figure 2b). Crystallization of the photolysis reaction mixture afforded a single crystal of **5a** (Figure 2c). Compound **5a** can be envisioned as arising from C-to-N migration within an adamantyl nitrene ligand to generate a transient 2-azahomoadamant-3-ene ligand and subsequent trapping with adventitious water to give rise to the observed hemiaminal ligand (see Figure S6). The observed structure is consistent both with the known low-temperature rearrangement of adamantyl nitrene and with the electrophilicity of highly strained anti-Bredt imines.¹⁴ The UV-vis spectrum of a CH₂Cl₂ solution of 5a is well-matched to the UV-vis spectrum obtained following photolysis of 2a (Figure S7). Similarly, photolysis of complex **2b** (335 nm $< \lambda < 610$ nm) proceeds via a well-anchored isosbestic point at 603 nm (Figure 2d). The product of photolysis was assigned as 5b by comparison of the final UV-vis spectrum obtained from photolysis of 2b with the spectrum generated by addition of AdN_3 to **5a** (Figure S8).

Matrix-assisted laser desorption-mass spectrometry (MALDI-MS) data provided additional evidence for facile N₂ elimination from **2** (Figure 3). Ablation of a sample of **2a** produces an ion at m/z = 907.5, which is well matched to the expected mass of Rh₂(esp)₂(AdN)⁺ (calc = 907.2), and displays the expected isotopic distribution. Ablation of a sample of



Figure 3. Mass spectrometry evidence for N₂ loss from Rh₂ complexes. MALDI-MS data acquired for ablation of samples of 2a and [¹⁵N]-2a, which indicate the facile loss of N₂ from these complexes to generate Rh₂(esp)₂(AdN) fragments.

[¹⁵N]-**2a**, prepared from monolabeled [¹⁵N]-AdN₃, provided the expected +1 m/z (m/z = 908.4 (expt); 908.2 (calc)) and displays the isotope pattern expected for a 1 : 1 mixture of **2a** and [¹⁵N]-**2a**, which results from incorporation of 50% ¹⁵N at each of N(α) and N(γ) in [¹⁵N]-AdN₃. *In situ* IR analysis of both photolyzed or thermolyzed KBr pellets of **2b** further indicate facile N₂ loss from **2b** (Figures S9 and S10).

We hypothesized that low-temperature N₂ extrusion from 2b within a single crystal habit would enable direct structural characterization of nitrenoid 3b. To this end, we examined the in situ structural evolution of a single-crystal of **2b** by X-ray diffraction during irradiation with a 365 nm light source. Data was collected at 100 K with 50 keV synchrotron radiation.¹⁵ Solid-state reaction progress was monitored by free refinement of the nitrogen occupancies, which indicate that while the occupancy of $N(\alpha)$ was unchanged with time, the occupancy of N(β) decreased. Concurrently, the space group was observed to transition from monoclinic $P2_1/c$ to $P2_1/n$.¹⁶ Refinement of the resulting data indicated elimination of a molecule of N_2 to afford Rh_2 nitrenoid **3b** $\cdot N_2$ (Figure 4). Upon N_2 extrusion, Rh(1)-N(1B) (*i.e.* the nitrenoid linkage) contracts from 2.335(3) Å (**2b**) to 2.12(1) Å (**3b**). Concurrent with N_2 extrusion and Rh(1)-N(1B) contraction, significant contraction of N(1B)-C(1) (*i.e.* the N-C bond in the adamantyl nitrene fragment) is also observed from 1.509(5) Å (2b) to 1.41(2) Å (**3b**). The conversion of **2b** to **3b** is also accompanied by a significant expansion of the Rh(1)-N(B)-C(1) angle from 129.1° to 147.2°. No substantial changes in the C-C



	30 expl.	"[30] comp.	-[3b] comp.
Rh(1)–N(1B) / Å	2.12(2)	2.094	1.924
Rh(1)–N(1A) / Å	2.346(4)	2.408	2.534
Rh(1)–Rh(1) / Å	2.3903(4)	2.438	2.465
Rh(1)–N(1B)–C(1) / °	147.2(9)	136.9	126.8

Figure 4. Solid-state structure of reactive Rh₂ nitrenoid 3b. Thermal ellipsoid plot of **3b**·N₂ generated by solid-state N₂ elimination from **2b**. Ellipsoids are drawn at 50% probability. H-atoms and solvent are removed for clarity. The structure illustrated here results from refinement of a data set collected at 46% conversion; higher conversions can be achieved but at the expense of crystal-linity (see Supporting Information). Comparison of the bond metrics derived from the X-ray structure with those computed for ³[**3b**] and ¹[**3b**] indicate excellent agreement with the triplet electronic configuration.

distances of the adamantyl fragment were observed, and both Rh(1)–Rh(1) (2.3968(8) Å (**2b**); 2.3903(4) Å (**3b**)) and Rh(1)–N(1A) (*i.e.* the Rh–N(Ad)N₂ linkage; 2.335(3) (**2b**); 2.346(4) Å (**3b**)) are essentially unchanged. The Rh centers in **3b**·N₂ are symmetry equivalent, and thus following loss of N₂ the AdN₃ and AdN ligands of **3b** are compositionally disordered (*i.e.* 50% occupancy of each AdN and AdN₃ on each of the Rh centers). Solid-state conversion of up to 70% are well accommodated within the single crystal. Attempts to achieve higher conversions or to promote loss of a second equivalent of N₂ to generate a *bis*-nitrenoid by prolonged irradiation were unsuccessful due to loss of crystallinity (see Supporting Information for details).

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Density Functional Theory (DFT) optimization of the geometry of 3b has been pursued both as a singlet and as a triplet electronic configuration (*i.e.* ¹[**3b**] and ³[**3b**]; M06 functional, LANL2DZ basis set for Rh, 6-31G** for other atoms).¹⁶ The calculated Rh(1)-Rh(1), Rh(1)-N(1A), Rh(1)-N(1B), and N(1B)-C(1) distances and Rh(1)-N(1B)-C(1) angle for ³[**3b**] are in excellent agreement with the experimentally defined parameters (Figure 4). In contrast, the optimized structure of [3b] substantially underestimates both the Rh(1)–N(1B) distance (1.924 Å (comp.); 2.12(2) Å (expt.)) and Rh(1)–N(1B)–C(1) angle (126.8° (comp.); 147.2° (expt.)) and substantially overestimates the Rh(1)-N(1A) distance (2.534 Å (comp.); 2.346(4) Å (expt.)). The observation of ${}^{3}[\mathbf{3b}]$ is consistent with the relative stabilities computed for the ${}^{1}[\mathbf{3b}]$ and ${}^{3}[\mathbf{3b}]$; the singlet structure is calculated to be 5.5 kcal/mol above the triplet.¹⁷ One might expect that a singlet nitrene would be best stabilized in a bent geometry that maximizes back-bonding into a vacant p-orbital. We speculate that the observed linearization of the nitrenoid fragment in 3b enables the triplet nitrene to be stabilized by two half-order π -bonds generated by overlap of filled Rh–Rh π^* orbitals with partially occupied N-centered orbitals.16

In closing, the characterization of nitrenoid 3b reported here establishes the electronic and three-dimensional structures of this critical intermediate. Typically, structural characterization of the transient intermediates involved in the intimate bond-forming and -breaking processes during catalysis is not possible and thus investigations of reaction mechanisms often rely on computational characterization of reactive intermediates using methods optimized for isolated catalyst intermediates. We anticipate that direct characterization of reactive nitrenoid intermediates will inform rational development of C-H amination chemistry. Further, the demonstration of crystalline matrix confinement as a platform for the structural characterization of reactive intermediates raises the tantalizing possibility that proper design of photoactive molecular precursors may represent a general strategy to directly characterize reactive species generated by elimination of small molecules within crystalline samples.

ASSOCIATED CONTENT

Supporting Information. Experimental procedures, spectroscopic and crystallographic data, coordinates of optimized structures (PDF, cif). The Supporting Information is available free of charge on the ACS Publications website.

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(15) Loss of N₂ from **2b** to generate **3b** could be stimulated both by irradiation with 365 nm light and also by prolonged exposure to synchrotron radiation. In addition to the 50 keV radiation used to acquire the data presented above, we have examined the structure of **2b** as a function of time with both 50 and 37.5 keV radiation without 365 nm irradiation and in both cases observed N2 loss to generate 3b.N2 loss was stimulated. The structure generated by photochemically promoted N2 loss is identical to that promoted by X-ray stimulated N2 loss. The X-ray stimulated loss of N2 is similar to observations made in protein crystallography regarding the cleavage of weak bonds upon extended irradiation as a mechanism to dissipate incipient X-ray energy. See, Garman, E. Radiation damage in macromolecular crystallography: what is it and why should we care? Acta Cryst. D. 2010, 66, 339-351. (15) The formation of $3b \cdot N_2$ results in the structure evolution from $P2_1/c$ to $P2_1/n$, with only minor changes in the unit cell parameters [2b: a = 10.8913(10), b = 14.274(1), c = 18.732(2), $\beta = 105.847(1)$, V = 2801.7(4)to **3b**·N₂: $a = 11.030(2) b = 14.251(2), c = 18.763(3), \beta = 107.025(2), V =$ 2820.2(8)]. This is not a transformation of the monoclinic standard spacegroup P2₁/c to the nonstandard space group P2₁/n [matrix $(1 \ 0 \ 1/0 \ 1 \ 0/ \ -1$] 00), new cell : 18.924, 14.275, 10.891, 107.77].

(16) For these calculations, adamantyl groups were truncated as t-butyl groups. The described computational method has previously been utilized to evaluate the structures of Rh2 nitrenoids, see: Valeria-Álvarez, A.; Haines, B. E.; Musaev, D. G. Key mechanistic insights into the intramolecular C-H bond amination and double bond aziridination in sulfamate esters catalyzed by dirhodium tetracarboxylate complexes. J. Organomet. Chem. 2018, 867, 183-192. Computations pursued using the BP86 functional in combination with the TZVP basis set for Rh and 6-31G** for other atoms (ref. 8b) are detailed in the Supporting Information.

(17) Similarly, ³[3a] is computed to be 6.9 kcal/mol lower in energy than ¹[**3a**]. See Supporting Information for details and for discussion of the impact of the apical ligand on the structures of Rh₂ nitrenoids.

