Inorganic Chemistry

Tunable Electrochemical and Catalytic Features of BIAN- and BIAO-Derived Ruthenium Complexes

Arijit Singha Hazari,[†] Ankita Das,[†] Ritwika Ray,[†] Hemlata Agarwala,[†] Somnath Maji,[†] Shaikh M. Mobin,[‡] and Goutam Kumar Lahiri^{*,†}

[†]Department of Chemistry, Indian Institute of Technology Bombay, Powai, Mumbai 400076, India

[‡]Discipline of Chemistry, School of Basic Sciences, Indian Institute of Technology Indore, Indore 452017, India

S Supporting Information

ABSTRACT: This article deals with a class of ruthenium–BIAN-derived complexes, $[Ru^{II}(tpm)(R-BIAN)CI]CIO_4$ (tpm = tris(1-pyrazolyl)methane, R-BIAN = bis(arylimino)acenaphthene, R = 4-OMe ([1a]CIO_4), 4-F ([1b]-CIO_4), 4-CI ([1c]CIO_4), 4-NO_2 ([1d]CIO_4)) and $[Ru^{II}(tpm)(OMe-BIAN)-H_2O]^{2+}$ ([3a](CIO_4)₂). The R-BIAN framework with R = H, however, leads to the selective formation of partially hydrolyzed BIAO ([*N*-(phenyl)imino]-acenaphtenone)-derived complex $[Ru^{II}(tpm)(BIAO)CI]CIO_4$ ([2]CIO_4). The redox-sensitive bond parameters involving -N=C-C=N- or -N=C-C=O of BIAN or BIAO in the crystals of representative [1a]CIO₄, [3a](PF₆)₂, or [2]CIO₄ establish its unreduced form. The chloro derivatives $1a^+-1d^+$ and 2^+ exhibit one oxidation and successive reduction processes in CH₃CN within the potential limit of ± 2.0 V versus SCE, and the redox potentials follow the order $1a^+ < 1b^+ < 1c^+ < 1d^+ \approx 2^+$. The electronic



potentials follow the order $1a^+ < 1b^+ < 1c^+ < 1d^+ \approx 2^+$. The electronic structural aspects of $1a^n-1d^n$ and 2^n (n = +2, +1, 0, -1, -2, -3) have been assessed by UV-vis and EPR spectroelectrochemistry, DFT-calculated MO compositions, and Mulliken spin density distributions in paramagnetic intermediate states which reveal metal-based ($Ru^{II} \rightarrow Ru^{III}$) oxidation and primarily BIAN- or BIAO-based successive reduction processes. The aqua complex $3a^{2+}$ undergoes two proton-coupled redox processes at 0.56 and 0.85 V versus SCE in phosphate buffer (pH 7) corresponding to { $Ru^{II}-H_2O$ }/{ $Ru^{III}-OH$ } and { $Ru^{III}-OH$ }/{ $Ru^{IV}=O$ }, respectively. The chloro ($1a^+-1d^+$) and aqua ($3a^{2+}$) derivatives are found to be equally active in functioning as efficient precatalysts toward the epoxidation of a wide variety of alkenes in the presence of PhI(OAc)₂ as oxidant in CH₂Cl₂ at 298 K, though the analogous 2^+ remains virtually inactive. The detailed experimental analysis with the representative precatalyst $1a^+$ suggests the involvement of the active { $Ru^{IV}=$ O} species in the catalytic cycle, and the reaction proceeds through the radical mechanism, as also supported by the DFT calculations.

INTRODUCTION

The ruthenium complexes of α -diimine (—N=C-C=N—)derived ligands have drawn significant attention over the decades primarily due to their diverse applications in dyesensitized solar cells, molecular electronics, sensor technology, homogeneous catalysis, biomedical research, and supramolecular chemistry.¹ In this regard, metal complexes (main group as well as transition elements) of the robust framework of bis(arylimino)acenaphthene (BIAN) involving α -diimine fragment conjugated with the naphthalene ring (Chart 1) have





recently been employed in catalysis and to understand other fundamental issues.² The corresponding ruthenium complexes of BIAN are however limited to a few recent reports,³ dealing primarily with the structural and electronic aspects as well as their potential applications in catalysis.

The low-lying unoccupied molecular orbitals of BIAN, $\pi^*(\alpha$ diimine) (5b2) and $\pi^*(naphthalene)$ (4a2), facilitate the successive electron uptake processes particularly on metalation, which in turn provides a multielectron reservoir platform suitable for selective application in catalysis.^{3b-d,2m} The tetraanionic state of 1,2-bis[(2,6-diisopropylphenyl)imino]acenaphthene has also been reported,²ⁿ involving the stepwise two-electron reductions of the α -diimine group followed by the addition of two more electrons over the multiple cation (Na⁺) coordinated naphthalene π system.

The structural rigidity, stereoelectronic tunability based on the "R" groups (Chart 1), and facile successive electron-



Received: March 18, 2015

Chart 2. Representation of Complexes



[3a](CIO₄)₂

H₂O

H₃CO

Article

accepting feature of BIAN collectively make it an attractive ligand to design metal complexes suitable for electron-transfer processes and catalytic application.

R= F

R=CI

R=NO₂

R= OCH3 : [1a]CIO4

:[1b]ClO₄

:[1c]CIO4

: [1d]CIO4

¢CIO

[2]CIO₄

In view of the aforestated unique features of BIAN, the present article is centered around the designing of ruthenium–BIAN-based complexes (Chart 2) of $[Ru^{II}(tpm)(R-BIAN)Cl]^+$ (tpm = tris(1-pyrazolyl)methane, R-BIAN = bis(arylimino)-acenaphthene, R = 4-OMe (1a⁺), 4-F (1b⁺), 4-Cl (1c⁺), 4-NO₂ (1d⁺)), $[Ru^{II}(tpm)$ (OMe-BIAN)H₂O]²⁺ (3a²⁺), as well as partially in-situ-hydrolyzed BIAO for R = H (BIAO: [*N*-(phenyl)imino] acenapthenone) derived $[Ru^{II}(tpm)(BIAO)-Cl]^+$ (2⁺) with the broader perspectives of understanding their electronic structural aspects and catalytic potential.

Herein, we report the synthesis and characterization of $[1a]ClO_4-[1d]ClO_4$, $[2]ClO_4$, and $[3a](ClO_4)_2$, molecular structures of representative $[1a]ClO_4$, $[2]ClO_4$, and $[3a]-(PF_6)_2$, and assessment of electronic structures of the chloro derivatives $1a^n-1d^n$ and 2^n (n = +2, +1, 0, -1, -2, -3) in accessible redox states via structure, UV-vis, and EPR spectroelectrochemistry in conjunction with DFT/TD-DFT calculations. The significant impact of "R" groups in the BIAN frameworks in $1a^+-1d^+$ and BIAO in 2^+ on the redox processes has also been highlighted.

Further, the potential of $1a^+-1d^+/2^+$ or $3a^{2+}$ involving the labile Ru^{II}-Cl or Ru^{II}-OH₂ bond, respectively, as precatalyst for the epoxidation of a wide variety of alkenes has been explored with special references to (i) "R" groups in BIAN, (ii) {Ru^{II}-Cl} versus {Ru^{II}-H₂O}, and (iii) BIAN versus BIAO. The limitation of the traditional method for epoxidation of alkenes by simple organic peracids involving the narrow substrate scope and inconvenience in product separation⁴ has led to introduction of the metal complex-catalyzed efficient epoxidation process in the presence of suitable oxidants.⁵ In this context, the initial report of Nishiyama et al. of ruthenium complex/TBHP (*tert*-butylhydroperoxide) catalyzed epoxidation of olefins in 1997⁶ has spurred the subsequent utilization of varying ruthenium complexes in combination with suitable oxidants for the epoxidation reactions.⁷

RESULTS AND DISCUSSION

Synthesis, General Characterization, and Structure. The 1:1 electrolytic and diamagnetic complexes $[Ru^{II}(tpm)(R-BIAN)CI]CIO_4$ (red powder) (tpm = tris(1-pyrazolyl)methane and BIAN = bis(arylimino)acenaphthene, R = 4-OMe [1a]CIO_4, 4-F [1b]CIO_4, 4-CI [1c]CIO_4, 4-NO_2 [1d]CIO_4) have been prepared from the metal precursor $Ru^{III}(tpm)Cl_3$ in combination with respective R-BIAN in refluxing methanol under a dinitrogen atmosphere (Charts 1 and 2). The isolated perchlorate salts are purified by column chromatography on a neutral alumina column using 4:1 CH₂Cl₂–CH₃CN mixture as eluant. To our surprise, under identical reaction conditions the use of R-BIAN with R = H (Charts 1 and 2) selectively results in [2]ClO₄ (blue powder) encompassing [*N*-(phenyl)imino]-acenapthenone (BIAO) via the hydrolysis of one of the imine bonds ($-C=N-Ph \rightarrow -C=O$)⁸ of BIAN. Complex 2⁺ represents the first example of a ruthenium-coordinated BIAO ligand; the corresponding mercury,⁹ cobalt,¹⁰copper,¹⁰ and zinc¹¹ complexes were however reported recently. The representative yellow-colored aqua derivative [Ru^{II}(tpm)(4-OMe-BIAN)(H₂O)](ClO₄)₂ [**3a**](ClO₄)₂ (Chart 2) has been prepared from the corresponding chloro derivative ([**1a**]ClO₄) in the presence of excess AgClO₄ in refluxing 1:3 acetone–water via the Cl/H₂O exchange process.^{12,13}

The complexes exhibit satisfactory microanalytical data and molar conductivities in CH₃CN (Experimental Section). The ESI mass spectral data of $[1a]ClO_4$ - $[1d]ClO_4$, $[2]ClO_4$, and $[3a](ClO_4)_2$ in CH₃CN and CH₃OH, respectively, match with their calculated values (Figure S1, Supporting Information and Experimental Section).

The molecular identities of the representative complexes $[1a]ClO_4$, $[2]ClO_4$, and $[3a](PF_6)_2$ are authenticated by their single-crystal X-ray structures (Figures 1–3; Tables 1, 2, and S1, Supporting Information). We failed to generate suitable single crystal for $[3a](ClO_4)_2$; however, the corresponding $[3a](PF_6)_2$ results in a single crystal suitable for its structure



Figure 1. ORTEP¹⁴ diagram of the cationic part of $[1a]ClO_4$: 2CH₂Cl₂. Ellipsoids are drawn at the 50% probability level. ClO_4^- anion, hydrogen atoms, and solvent molecules are omitted for clarity.



Figure 2. ORTEP¹⁴ diagram of the cationic part of [2]ClO₄. Ellipsoids are drawn at the 40% probability level. ClO₄⁻ anion and hydrogen atoms are omitted for clarity.



Figure 3. ORTEP¹⁴ diagram of the dicationic part of $[3a](PF_6)_2$. C₃H₆O. Ellipsoids are drawn at the 40% probability level. PF₆⁻ anions, hydrogen atoms, and solvent molecule are omitted for clarity.

determination. The tridentate tpm ligand is bonded to the metal ion in $[1a]ClO_4$ or $[2]ClO_4$ or $[3a](PF_6)_2$ through its pyrazolyl nitrogen donors in the usual facial mode, resulting in two shared six-membered chelates.¹³ The N,N or N,O donors of BIAN or BIAO in $[1a]ClO_4/[3a](PF_6)_2$ or $[2]ClO_4$ respectively, form a five-membered chelate with the metal ion. The bite angles of the six-membered chelates involving the tpm ligand, N3-Ru1-N5 (86.37(10)°)/N5-Ru1-N8 $(83.87(9)^{\circ})$, N2-Ru1-N4 $(85.16(9)^{\circ})/N4$ -Ru1-N7 (85.73(9)°), and N3-Ru1-N5 (85.86 (15)°)/N5-Ru1-N8 $(83.49(17)^{\circ})$, are appreciably larger than those of the corresponding five-membered chelates involving BIAN, N1-Ru1-N2 $(79.08(10)^{\circ})$ in [1a]ClO₄, N1-Ru1-N2 $(78.59(16)^{\circ})$ in $[3a](PF_6)_2$ or BIAO, and N1-Ru1-O1 $(80.14(8)^{\circ})$ in [2]ClO₄ (Table S1, Supporting Information). This in conjunction with the smaller average trans angles, 176.60°, 176.91°, and 176.71° in [1a]ClO₄, [2]ClO₄, and $[3a](PF_6)_2$, respectively (Table S1, Supporting Information),

suggest distorted octahedral geometries. The Ru^{II} -N(tpm),¹³ Ru^{II} -N(BIAN),^{3a-c} Ru^{II} -Cl,^{13a,b,15} and Ru^{II} -O(H₂O)^{13a,b,16} bond lengths in [1a]ClO₄, [2]ClO₄, and $[3a](PF_6)_2$ match fairly well with those of the reported analogous systems.

The feasibility of accepting successive two electrons by the low-lying π^* orbitals of the conjugated $-N=C-C=N-(\alpha$ diimine) or $-N=C-C=O(\alpha$ -ketoimine) fragment of BIAN or BIAO, respectively, leads to the three redox states, fully oxidized neutral form (A) to intermediate one-electron reduced radical state (\mathbf{B}) to doubly reduced dianionic form (\mathbf{C}) (Scheme 1) with distinctive C-N (1.30, 1.35, 1.38 Å), C-O (1.25, 1.28, 1.31 Å), and C-C (≥1.47, ~1.43, ~1.38 Å) bond distances, respectively.^{3a-c} The experimentally observed singlebond C-C length $[C8-C18 = 1.469(4) ([1a]ClO_4)]$ and 1.476(7) Å ($[3a](PF_6)_2$), C7–C17 = 1.474(4) Å ($[2]ClO_4$)] and double bond C-N length [C8-N1/C18-N2 = 1.298(4)/1.307(3) ($[1a]ClO_4$) and 1.291(6)/1.299(6) Å ($[3a](PF_6)_2$) and C7-N1/C17-O1 1.303(3)/1.249(3) Å ([2]ClO₄)], however, unequivocally establish that the unreduced form of BIAN or BIAO links to the ruthenium(II) ion in the complexes.

The calculated bond parameters of $1a^+-1d^+$, 2^+ , and $3a^{2+}$ based on their DFT-optimized structures (B3LYP-LANL2DZ/ SDD basis set) in the singlet ground state (S = 0) (Table 2 and Tables S2–S11, Figure S2, Supporting Information) match fairly well with the X-ray data of representative complexes.

The IR spectra (KBr pellets) of $[1a]ClO_4$ - $[1d]ClO_4$, $[2]ClO_4$, and $[3a](ClO_4)_2$ display characteristic strong ClO_4^- vibrations at around 1100 and 625 cm^{-1,3a} the C=O vibration of $[2]ClO_4$ at 1690 cm^{-1,17} and OH vibration of $[3a](ClO_4)_2$ at 3438 cm^{-1 13a} (Experimental Section).

¹H NMR spectra of $1a^+-1d^+$ and 2^+ in CDCl₃ (Figures 4 and S3, Supporting Information and Experimental Section) and $3a^{2+}$ in D₂O (Figure 4) display partially overlapping calculated number of aromatic proton resonances (tpm = 9, BIAN = 14, BIAO = 11) within the chemical shift range δ 6–9 ppm. The slightly acidic CH proton resonance of tpm for $1a^+-1d^+$ or 2^+ appears at a sharp and moderately strong singlet at $\delta \sim 10$ ppm;¹³ however, the same for $3a^{2+}$ in D₂O resolves as a weak but distinct peak at δ 9.7 ppm. The proton resonances associated with the six-membered rings of BIAN or BIAO in $1a^+ - 1d^+/3a^{2+}$ or 2^+ , respectively, can easily be distinguished from those of the five-membered rings of tpm, primarily based on their difference in spin-spin coupling constant (J/Hz) values (7–8 for BIAN or BIAO and 2–3 for tpm).^{3a,13} The OMe signal for $1a^+$ or $3a^{2+}$ appears at δ /ppm 3.80 or 3.79, respectively. The variation of chemical shift values based on the substituents at the BIAN framework in $1a^+-1d^+$ is rather insignificant.

Electrochemistry, MO Composition, and Spin Density Distribution. The chloro derivatives $1a^+-1c^+$ and 2^+ exhibit similar electrochemical features with one oxidation (O1) and two successive reductions (R1 and R2), while complex $1d^+$ incorporating *para*-NO₂-substituted BIAN (Chart 2) possesses one oxidation (O1) and four reductions (R1–R4) in CH₃CN within the potential window of ±2.0 V versus SCE (Figure 5, Table 3).

The redox potentials of $1a^+-1d^+$ vary systematically depending on the electron-donating or -withdrawing effect of the substituents in the BIAN framework, and it follows the order $1a^+ < 1b^+ < 1c^+ < 1d^+$. The impact of the -N=C-C=O chromophore of BIAO in 2^+ with regard to the -N=C-C=C=N- fragment of BIAN in analogous $1a^+-1d^+$ has been reflected in its relatively higher oxidation potential (0.93 V), even comparable to that of the *para*-NO₂-substituted BIAN in $1d^+$ (0.94 V).

Table 1. Selected Crystallographic Data for $[1a]ClO_4$, $[2]ClO_4$, and $[3a](PF_6)_2$

	$[1a]ClO_4 \cdot 2CH_2Cl_2$	[2]ClO ₄	$[\mathbf{3a}](\mathrm{PF}_6)_2 \cdot \mathrm{C}_3\mathrm{H}_6\mathrm{O}$
empirical formula	$\mathrm{C}_{38}\mathrm{H}_{34}\mathrm{Cl}_{6}\mathrm{N}_{8}\mathrm{O}_{6}\mathrm{Ru}$	$\mathrm{C}_{28}\mathrm{H}_{21}\mathrm{Cl}_{2}\mathrm{N}_{7}\mathrm{O}_{5}\mathrm{Ru}$	$C_{39}H_{38}N_8O_4F_{12}P_2Ru$
fw	1012.50	707.49	1073.78
cryst syst	monoclinic	monoclinic	triclinic
space group	$P2_1/n$	$P2_1/c$	P-1
a (Å)	11.7233 (3)	14.48414 (14)	11.650 (2)
b (Å)	17.2500 (4)	12.64471 (11)	14.944 (3)
c (Å)	20.1702 (5)	15.76167 (15)	16.267 (3)
α (deg)	90	90	71.09 (3)
β (deg)	94.8260 (10)	92.8914 (9)	69.90 (3)
γ (deg)	90	90	72.02 (3)
V (Å ³)	4064.50 (17)	2883.04 (5)	2453.3 (9)
Z	4	13	2
$\mu \ (\mathrm{mm}^{-1})$	0.838	6.554	0.476
T (K)	100 (2)	150 (2)	100 (2)
$D_{\text{calcd}} (\text{g cm}^{-3})$	1.655	1.630	1.454
F(000)	2048	1424	1084
θ range (deg)	1.55-29.78	3.05-72.16	1.48-24.29
data/restraints/params	11 546/0/534	5627/0/388	7908/1/607
R1, wR2 $[I > 2\sigma(I)]$	0.0520, 0.1223	0.0357, 0.1031	0.0584, 0.1396
R1, wR2(all data)	0.0776, 0.1368	0.0376, 0.1058	0.0864, 0.1508
GOF	1.048	1.057	1.001
largest diff. peak/hole (e $Å^{-3}$)	1.60/-1.06	0.56/-0.51	1.18/-0.71

Table 2. Experimental and DFT-Calculated Selected Bond Lengths (Angstroms) for [1a]ClO₄, [2]ClO₄, and [3a](PF₆)₂

	[1a]ClO ₄		[2]ClC	[2]ClO ₄		[3a](PF ₆) ₂	
	X-ray	DFT	X-ray	DFT	X-ray	DFT	
Ru1–N1	2.021(2)	2.088	2.020(2)	2.077	2.038(4)	2.104	
Ru1–N2	2.033(2)	2.087	2.057(2)	2.112	2.036(4)	2.116	
Ru1–N3	2.047(3)	2.118			2.023(4)	2.070	
Ru1–N4			2.068(2)	2.095			
Ru1–N5	2.070(2)	2.127			2.075(4)	2.147	
Ru1–N7			2.042(2)	2.071			
Ru1–N8	2.083(2)	2.129			2.060(5)	2.133	
Ru1-O1			2.0818(19)	2.108	2.126(4)	2.218	
Ru1-Cl1	2.3965(8)	2.430	2.3690(7)	2.391			
C8-N1	1.298(4)	1.302			1.291(6)	1.303	
C18-N2	1.307(3)	1.302			1.299(6)	1.306	
C7-C17			1.474(4)	1.490			
C8-C18	1.469(4)	1.483			1.476(7)	1.495	
C17-O1			1.249(3)	1.252			
C6-N1			1.443(3)	1.428			
C7-N1	1.445(4)	1.428	1.303(3)	1.307	1.446(6)	1.430	
C20-N2	1.430(4)	1.428			1.419(6)	1.431	

Scheme 1. Varying Redox States of BIAN and BIAO



The origin of the redox processes of $1a^+-1d^+$ and 2^+ (Figure 5, Table 3) has been ascertained via the DFT-calculated MO compositions (Tables 4 and S12–S34, Supporting Information) and Mulliken spin density distributions (Table 5 and Figure 6) in paramagnetic intermediate states. The metal-based HOMO of $1a^+-1d^+/2^+$ in the S = 0 state (~60%) and β -

LUMO of $1a^{2+}-1d^{2+}/2^{2+}$ in the S = 1/2 state (>60%) as well as metal-dominated spin (>0.75) collectively suggest the correspondences of O1 to a Ru^{II} \rightarrow Ru^{III} oxidation process. In consequence, the redox-sensitive C-C, C-N, and C-O bond distances of BIAN and BIAO remain virtually invariant on the oxidation process (O1) (Scheme 1, Tables 6 and S2, S4, S6, S8, and S10, Supporting Information). The metal-based oxidation process has been further evidenced by the highly anisotropic EPR spectrum of coulometrically oxidized representative $1a^{2+}$ with $g_1 = 2.674$, $g_2 = 2.212$, $g_3 = 1.769$, $\Delta g = 0.905$, $\langle g \rangle = 2.248$ ($\Delta g = g_1 - g_3$ and $\langle g \rangle = \{1/3(g_1^2 + g_2^2 + g_3^2)\}^{1/2}$) (Figure 7a).²²

It should be noted that an appreciable contribution of Cl in the HOMO of $1a^+-1d^+/2^+$ (>20%) and β -LUMO of $1a^{2+}-1d^{2+}/2^{2+}$ (9–14%) has also been predicted due to the effect of Ru–Cl covalency. A similar sizable Cl contribution in the DFT-

Article



Figure 4. ¹H NMR spectra of (i) $[1a]ClO_4$, (ii) $[2]ClO_4$ in $CDCl_3$, and (iii) $[3a](ClO_4)_2$ in D_2O (t = tpm, b = BIAN or BIAO).

calculated MOs has also been pointed out earlier in analogous ruthenium complexes. $^{\rm 23}$

The presence of two strong π -acceptor ligands, BIAN³ and tpm,¹³ in the complexes raises the pertinent question of their specific involvements in the successive reduction processes (Figure 5). The greater than 80% BIAN/BIAO contribution in the LUMO of $1a^+ - 1d^+/2^+$ (*S* = 0) and SOMO of 1a - 1d/2 (*S* = 1/2) as well as -N=C-C=N-(BIAN)/-N=C-C=O (BIAO) dominated spin (>90%) (Table 4), however, supports the BIAN/BIAO-based first reduction process (R1 in Figure 5). This has been further supported by the free radical EPR with g = 2.001 (line width = 25 G) for the electrogenerated representative 2 (Figure 7b).²² On the other hand, the mixed MO compositions of β -LUMO (BIAN/tpm, 50-60%/30-40%; BIAO/tpm, 69/25%) in **1a-1c** or **2** (S = 1/ 2) and HOMO (BIAN/tpm, 61-83%/10-37%; BIAO/tpm, 81/12% in $1a^--1c^-$ or 2^- (S = 0) suggest the partial involvement of tpm along with the BIAN- or BIAO-based orbitals in the second reduction process (R2 in Figure 5). On the contrary, the β -LUMO of 1d (S = 1/2, BIAN, 96%) and HOMO of $1d^{-}$ (S = 0, BIAN 89%) imply solely a BIAN (-N=C-C=N- function) based second reduction (R2 in Figure 5). The involvement of BIAN (-N=C-C=N-) or BIAO (-N=C-C=O) in successive electron uptake processes (R1 and R2) finds further justification via the sequential shortening and lengthening of the DFT-calculated redox-sensitive C-C and C-N/C-O bond distances, respectively (Scheme 1, Table 6 and S2, S4, S6, S8, and S10, Supporting Information). The subsequent two reductions R3 and R4 of $1d^+$ in Figure 5 essentially involve the NO₂ group of BIAN (~95% BIAN contribution in LUMO of $1d^{-}$ (S = 0) and



Figure 5. Cyclic voltammograms of (i) $[1a]ClO_4$, (ii) $[1b]ClO_4$, (iii) $[1c]ClO_4$, (iv) $[1d]ClO_4$, and (v) $[2]ClO_4$ in CH₃CN. (Inset) Cyclic voltammograms (black) and differential pulse voltammograms (green) of reversible oxidation (O1) and first reduction (R1) couples.

SOMO of $1d^{2-}$ (S = 1/2) as well as 98% BIAN-dominated spin in $1d^{2-}$) and a mixed tpm—BIAN-based situation, respectively (α -LUMO of $1d^{2-}$ (S = 1/2) = 88% tpm, SOMO of $1d^{3-}$ (S =1) = 68% tpm/28% BIAN, and spin density of $1d^{3-}$ = BIAN(0.998)/tpm(0.951)).

Remarkably, the appreciable positive shift of the redox potential on selective incorporation of the electron-withdrawing NO₂ group in the BIAN framework in 1d⁺ (Table 3, Figure 5) facilitates detecting two additional reductions (R3 and R4) covering the NO₂ function/naphthalene ring of BIAN as well as the reduction of tpm as has been revealed by MO compositions and spin density plots.

The representative aqua derivative $3a^{2+}$ displays two successive one-electron oxidation processes, E°_{298} ,V (ΔE_{p} , mV) at 0.56(60) and 0.85(70) in phosphate buffer versus SCE at pH 7.0, which undergo anodic shifting with lowering pH (Figure S4, Table S35, Supporting Information), attributed to the simultaneous one-electron/one-proton transfer processes, leading to the {Ru^{IV}=O} state: Ru^{II}-OH₂/Ru^{III}-OH and Ru^{III}-OH/Ru^{IV}=O.^{13a,24} Unfortunately, we failed to check the shift in potential of $3a^{2+}$ with the further change in pH primarily due to the associated precipitation problem.

The comparison of the oxidation potentials of the chloro $(1a^+-1d^+, 2^+)$ and aqua $(3a^{2+})$ derivatives with the reported analogous {Ru^{II}(tpm)(L)Cl} and {Ru^{II}(tpm)(L)H₂O} com-

				$E^{\circ}_{298}/V (\Delta E_{\rm p}/{\rm mV})^{\ell}$,		
complex	01	O2	R1	R2	R3	R4	ref
			chloro derivative	s			
[1a]ClO ₄ ^c	0.77(60)		-1.12(60)	-1.57(120)			this work
[1b]ClO ₄ ^{<i>c</i>}	0.82(60)		-1.01(70)	-1.54(70)			this work
[1c]ClO ₄ ^c	0.87(70)		-0.94(70)	-1.43(220)			this work
[1d]ClO ₄ ^c	0.94(60)		-0.77(60)	-0.85(60)	-1.31(180)	-1.71^{d}	this work
$[2]ClO_4^c$	0.93(90)		-0.62(70)	-1.32(230)			this work
[Ru ^{II} tpm(L ¹)Cl]ClO ₄ ^c	0.82						13a
$[Ru^{II}tpm(L^2)Cl]Cl^c$	0.68						18
[Ru ^{II} tpm(L ³)Cl]Cl ^e	0.65						19
[Ru ^{II} tpm(L ⁴)Cl]BF ₄ ^e	0.83						20
$[Ru^{II}tpm(L^5)Cl]^e$	0.24						20
[Ru ^{II} tpm(L ⁶)Cl]BF ₄ ^e	0.79						13b
$[Ru^{II}tpm(L^7)Cl]^e$	0.22						13b
[Ru ^{II} tpm(L ⁸)Cl]ClO ₄ ^c	1.02						21
aqua derivatives							
$[3a](ClO_4)_2^f$	0.56(60)	0.85(70)					this work
$[\operatorname{Ru}^{II}\operatorname{tpm}(\operatorname{L}^{1})\operatorname{OH}_{2}](\operatorname{ClO}_{4})_{2}{}^{g}$	0.44	0.64					13a
$[\operatorname{Ru}^{II}\operatorname{tpm}(\operatorname{L}^2)\operatorname{OH}_2](\operatorname{ClO}_4)_2^h$	0.40	0.71					18
$[\operatorname{Ru}^{II}\operatorname{tpm}(\mathrm{L}^{3})\operatorname{OH}_{2}](\operatorname{PF}_{6})_{2}^{I}$	1.00						19
[RuIItpm(L4)OH2](BF4)2j	0.38						20
[Ru ^{II} tpm(L ⁵)OH ₂]BF ₄ ^j	0.17	0.37					20
$[\operatorname{Ru}^{II}\operatorname{tpm}(\operatorname{L}^6)\operatorname{OH}_2](\operatorname{PF}_6)_2^{j}$	0.38						13b
$[Ru^{II}tpm(L^7)OH_2]PF_6^{j}$	0.16						13b

^{*a*}L¹: *o*-benzoquinonediimine. L²: 2,2'-bipyridine. L³: 4-methyl-4'-(N-phenothiazylmethyl)-2,2'-bipyridine. L⁴: 4,4'-dibenzyl-4,4',5,5'-tetrahydro-2,2'-bioxazole. L⁵: 2-[((1'S)-1'-(hydroxymethyl)-2'-phenyl)ethylcarboxamide]-(4S)-4-benzyl-4,5-dihydrooxazole. L⁶: ((4S,4'S)-(-)-4,4',5,5'-tetrahydro-4,4'-bis(1-methylethyl)-2,2'-bioxazole. L⁷: N-(1-hydroxy-3-methylbutan-(2S)-(-)-2-yl)-(4S)-(-)-4-isopropyl-4,5-dihydrooxazole-2-carbimidate. L⁸: 2-phenylazopyridine. ^bPotential in volts versus SCE; peak potential difference $\Delta E_p/mV$ (in parentheses). ^cIn CH₃CN/0.1 M Et₄NClO₄ at 100 mV s⁻¹. ^dIrreversible. ^eIn CH₃CN/0.1 M Bu₄NPF₆ at 100 mV s⁻¹. ^fIn phosphate buffer, pH 7 at 100 mV s⁻¹. ^gIn H₂O/0.1 M Kcl, pH 7 at 100 mV s⁻¹.

plexes, respectively, integrating bidentate ligands (L) with different electronic properties (Table 3) reveals the significant variation in potential based on the π -accepting or σ -donating strength of L.

Spectroelectrochemistry and TD-DFT Calculations. The complexes $(1a^+-1d^+ and 2^+)$ exhibit a similar spectral profile in the UV-vis region in CH₃CN, and the transitions are assigned based on the TD-DFT calculations (Figure 8, Table 7). The moderately intense three visible bands in the range 700–400 nm are originated via BIAN/BIAO-targeted $(d\pi)$ Ru $\rightarrow (\pi^*)$ BIAN/BIAO MLCT (metal-to-ligand charge transfer)²⁵ and $(d\pi)Ru/(\pi)BIAN/BIAO \rightarrow (\pi^*)BIAN/BIAO$ MLLCT (metal/ligand-to-ligand charge transfer)²⁵ transitions. The reversible one-electron oxidized species $1a^{2+}-1d^{2+}$ and 2^{2+} (O1, Figure 5) display two weak bands in the range 800-600 nm followed by an intense band near 400 nm, which are predicted to be originated via (π) BIAN/BIAO $\rightarrow (d\pi)$ Ru LMCT (ligand-to-metal charge transfer) and (π) BIAN/BIAO/ $(d\pi)Ru \rightarrow (d\pi)Ru$ LMLCT (ligand/metal-to-metal charge transfer) transitions. One-electron reversible reduction of $1a^+$ - $1d^+$ to 1a-1d (R1, Figure 5) causes the marginal change in band position as well as intensity of the BIAN-targeted MLCT or MLLCT band, though an appreciable change is noticed on moving from 2^+ to 2. The higher energy UV region bands in all cases are assigned to ligand-based $((\pi)BIAN/BIAO \rightarrow$ (π^*) BIAN/BIAO) transitions.

Catalytic Epoxidation. The accessibility of high-valent ruthenium-oxo $\{Ru^{IV}=O\}$ species in the selective molecular frameworks has led to the exploration of their potential

application in the catalytic epoxidation process.^{7,13a,24a} In this context, the newly designed molecular sets $1a^+-1d^+$, 2^+ , and $3a^{2+}$ have been tested as possible precatalysts toward the epoxidation of a wide variety of alkenes. The initial standardization process has been monitored with the precatalyst $1a^+$ (chloro derivative) and styrene as the model substrate in combination with H₂O₂, TBHP, IDA, and *m*-CPBA (H₂O₂ = hydrogen peroxide, TBHP = *tert*-butyl hydrogen peroxide, IDA = iodobenzene diacetate, *m*-CPBA = *m*-chloroperbenzoic acid) as possible oxidants in CH₂Cl₂, CH₃CN, and C₂H₅OH solvents at 298 K (Table 8). The optimization study reveals that IDA and CH₂Cl₂ are the best oxidant and solvent, respectively, under the catalyst:substrate:oxidant molar ratio of 1:100:200, which gives maximum conversion of 88% (entry 4, Table 8). The above optimal condition (eq 1) with styrene as the model

substrate has therefore been applied for the precatalysts $1a^+$ - $1d^+$, which shows a marginal difference based on the electrondonating (OMe) or -withdrawing (NO₂, Cl, F) group at the para position of the arylimino moieties of BIAN. The representative aqua derivative $3a^{2+}$ extends comparable activity to that of $1a^+$ with selected tested alkenes (Table 9); therefore, the epoxidation reaction of a wide variety of alkenes has been performed selectively with $1a^+$. It should be noted that under the above reaction conditions (eq 1) no epoxide formation takes place in the absence of catalyst.

Table 4. DFT-Calculated Selected MO Compositions of $1a^n-1d^n$ and 2^n

complex	МО	fragments	% contribution
$\mathbf{1a}^+ (S=0)$	НОМО	Ru/Cl/BIAN/tpm	56/23/11/10
	LUMO	BIAN/Ru/tpm/Cl	82/11/06/01
$1a^{2+}(S = 1/2)$	β -LUMO	Ru/BIAN/tpm/Cl	61/20/09/09
1a (S = 1/2)	SOMO	BIAN/Ru/tpm/Cl	85/09/05/01
	β -LUMO	BIAN/tpm/Ru/Cl	55/40/04/01
$\mathbf{1a}^{-}(S=0)$	НОМО	BIAN/tpm/Ru	61/37/02
$\mathbf{1b}^+ (S=0)$	НОМО	Ru/Cl/BIAN/tpm	59/23/09/09
	LUMO	BIAN/Ru/tpm/Cl	84/10/04/01
$1b^{2+}(S = 1/2)$	β -LUMO	Ru/Cl/BIAN/tpm	67/13/10/10
1b $(S = 1/2)$	SOMO	BIAN/Ru/tpm/Cl	85/09/05/01
	β -LUMO	BIAN/tpm/Ru/Cl	54/41/04/01
$\mathbf{1b}^{-}(S=0)$	НОМО	BIAN/tpm/Ru/Cl	81/12/06/01
$\mathbf{1c}^+ (S=0)$	НОМО	Ru/Cl/tpm/BIAN	60/23/09/09
	LUMO	BIAN/Ru/tpm/Cl	84/10/04/01
$1c^{2+}(S=1/2)$	β -LUMO	Ru/Cl/BIAN/tpm	67/13/10/10
1c $(S = 1/2)$	SOMO	BIAN/Ru/tpm/Cl	85/09/05/01
	β -LUMO	BIAN/tpm/Ru/Cl	61/34/04/01
$\mathbf{1c}^{-}(S=0)$	НОМО	BIAN/tpm/Ru/Cl	83/10/06/01
$\mathbf{1d}^+ \ (S=0)$	НОМО	Ru/Cl/tpm/BIAN	59/24/09/09
	LUMO	BIAN/Ru/tpm/Cl	86/09/03/01
$1d^{2+}(S = 1/2)$	β -LUMO	Ru/Cl/tpm/BIAN	67/13/10/09
1d (S = 1/2)	SOMO	BIAN/Ru/tpm/Cl	86/09/04/01
	β -LUMO	BIAN/Ru/tpm/Cl	96/02/01/00
$\mathbf{1d}^{-}(S=0)$	НОМО	BIAN/tpm/Ru/Cl	89/06/04/01
	LUMO	BIAN/Ru/tpm	95/03/02
$1d^{2-}(S=1/2)$	SOMO	BIAN/Ru/tpm	94/03/02
	α-LUMO	tpm/Ru/BIAN	88/06/06
$1d^{3-}(S=1)$	SOMO1	tpm/BIAN/Ru	68/28/04
$2^+ (S=0)$	НОМО	Ru/Cl/tpm/BIAO	60/24/11/05
	LUMO	BIAO/Ru/tpm/Cl	80/13/05/02
$2^{2+}(S = 1/2)$	β -LUMO	Ru/Cl/tpm/BIAO	68/14/11/06
2 $(S = 1/2)$	SOMO	BIAO/Ru/tpm/Cl	83/11/05/02
	β -LUMO	BIAO/tpm/Ru/Cl	69/25/05/01
$2^{-}(S=0)$	НОМО	BIAO/tpm/Ru/Cl	81/12/06/01

Table 5. Mulliken Spin Densities for Paramagnetic Forms of $1a^n-1d^n$ and 2^n

complex	Ru	BIAN/BIAO	tpm	Cl
$1a^{2+}(S=1/2)$	0.755	0.098	0.028	0.118
1a $(S = 1/2)$	0.047	0.941	0.010	0.004
$\mathbf{1b}^{2+}(S = 1/2)$	0.782	0.032	0.062	0.147
1b $(S = 1/2)$	0.048	0.943	0.005	0.004
$1c^{2+}(S = 1/2)$	0.781	0.037	0.043	0.147
1c $(S = 1/2)$	0.049	1.001	0.010	0.005
$1d^{2+}(S = 1/2)$	0.777	0.035	0.036	0.152
1d (S = 1/2)	0.051	0.937	0.006	0.007
$1d^{2-}(S=1/2)$	0.024	0.98	-0.001	-0.001
$1d^{3-}(S=1)$	0.053	0.998	0.951	-0.002
$2^{2+}(S = 1/2)$	0.805	0.006	0.039	0.155
2 $(S = 1/2)$	0.040	0.946	0.008	0.005

Surprisingly, under identical reaction conditions (eq 1) the precatalyst 2^+ (chloro derivative) involving partially hydrolyzed BIAN ligand (i.e., BIAO) has failed to deliver any reasonable catalytic activities with any of the tested alkenes (Table 9). The observed inertness of 2^+ could possibly be attributed to its inability to form the necessary {Ru^{IV}==O} intermediate species (see below), as revealed by the independent reaction of 2^+ in

 CH_2Cl_2 or CH_3CN or C_2H_5OH with the oxidant (IDA or TBHP or H_2O_2 or *m*-CPBA).

Though the metal-catalyzed epoxidation process can also facilitate oxidative cleavage of the olefinic double bond resulting in aldehyde or ketone or alcohol,²⁶ the precatalyst $1a^+$ or $3a^{2+}$ is found to be chemoselective toward epoxide formation without any trace of carbonyl derivative as ascertained by GC-MS or ¹H NMR.

The efficiency of the catalyst $1a^+$ has also been further tested by varying the catalyst loading with respect to the substrate vinyl cyclopentane, which gives an appreciably high TON of 2100 even with 0.001 mol % of the catalyst (Table S36, Supporting Information).

The individual role of the metal ion, the free ligand (tpm or BIAN), and the metal ion plus ligand toward the epoxidation process has been tested with vinyl cyclopentane under the reaction conditions in eq 1 (Table S37, Supporting Information), which divulges that Ru as RuCl₃ or free ligand (1:1 tpm-BIAN) or Ru as RuCl₃ + tpm (0.5:1 mol ratio) fails to facilitate the epoxidation process at all. However, Ru as RuCl₃ in combination with BIAN (0.5:1 mol ratio) yields slight conversion (18%) with moderate epoxide formation (30% selectivity). On the contrary, the infusion of free ligand (tpm + BIAN) and RuCl₃ in a 1:1 molar ratio (i.e., the in-situgenerated catalyst) enhances the yield up to 66% with 80% chemoselectivity, which is however less than that achieved by using the corresponding preformed catalyst $1a^+$ (88% conversion, 100% selectivity, Table 9).

In general, complex $1a^+$ is found to be an effective precatalyst for the oxidation of a wide variety of alkenes to respective epoxides under the standardized reaction conditions in eq 1, which gives excellent to good selectivity. It also reveals that unactivated olefins such as vinyl cyclohexane, 1-phenyl cyclohexene, vinyl cyclopentane, cyclooctene, cyclooctadiene, 1-octene, and 1-decene (entries 10-16, Table 9) exhibit comparable to better activity in terms of both % conversion and selectivity to that of aromatic alkenes. Though cis-stilbene and trans-stilbene (entries 7 and 8, Table 9) show chemoselective epoxidation to cis-stilbene oxide and trans-stilbene oxide, respectively, without any side product, α -methylstyrene (entry 2, Table 9) yields overoxidized acetophenone along with the desired epoxide as the primary product. The substrate cyclooctadiene yields the double epoxide as the main product (76%, entry 14, Table 9) with the partially oxidized monoepoxide minor product (8%). The controlled experiment comprising of 1:1 mixture of 1-octene and trans-5-decene as substrate reveals the preferential formation of terminal oxide (1,2-epoxyoctane, conversion 80%, selectivity 100%) over the internal oxide (1,2-epoxydecane, conversion 20%) (entry 17, Table 9).

As proposed earlier,²⁴ the ruthenium-catalyzed epoxidation reaction of alkene proceeds through the in situ formation of $\{Ru^{III}-OH\}$ or $\{Ru^{IV}=O\}$ active intermediate. The use of hydroxyl or radical scavenger specific for each intermediate, however, can distinguish between the nature of the active species.²⁷ The extent of variation in % conversion of the representative substrate 1-octene to its corresponding oxide in the presence of specific hydroxyl (*N*,*N'*-dimethylthiourea, DMTU) or radical (TEMPO, 2,2,6,6-tetramethylpiperidinyloxy radical) scavenger has thus been tested under similar reaction conditions (eq 1). Though the % conversion of 1-octene to its corresponding epoxide remains virtually unaltered in the presence of DMTU (Table S38, Supporting Information), the



Figure 6. DFT-calculated Mulliken spin density plots of $1a^n - 1d^n$ and 2^n .

Table 6. DFT(B3LYP/LANL2DZ/SDD)-0	Calculated	Selected
Bond Distances of Coordinated BIAN or	BIAO in	$1a^n - 1d^n$
or 2^n		

8	18		$\langle \rangle$			\rightarrow	
Ar—N ₁	"N 1a"	2 Ar A	r — N ₁ 1	"N ₂ b"-1d"	Ar Pi	η—Ν ₁	'O ₁ 2"
complex	C8- C18	C7– C17	N1- C8	N1– C7	N2- C18	N2- C17	01– C17
1a ²⁺	1.496		1.301		1.301		
$1a^+$	1.483		1.302		1.302		
1a	1.431		1.339		1.340		
1a ⁻	1.396		1.381		1.381		
1b ²⁺		1.496		1.304		1.304	
1b ⁺		1.484		1.301		1.301	
1b		1.431		1.340		1.340	
1b-		1.395		1.382		1.382	
1c ²⁺		1.496		1.304		1.304	
$1c^+$		1.484		1.301		1.301	
1c		1.431		1.340		1.341	
1c ⁻		1.395		1.382		1.382	
1d ²⁺		1.499		1.301		1.301	
$1d^+$		1.485		1.301		1.301	
1d		1.434		1.340		1.339	
1d-		1.402		1.374		1.374	
1d ²⁻		1.420		1.370		1.372	
1d ³⁻		1.420		1.383		1.383	
2 ²⁺		1.508		1.303			1.257
2 ⁺		1.490		1.307			1.252
2		1.433		1.341			1.287
2-		1.395		1.396			1.314

complete inertness of alkene toward the epoxide formation on addition of TEMPO suggests the involvement of $\{Ru^{IV}=O\}$ as an active intermediate (Scheme 2).

The formation of active $\{Ru^{IV}=O\}$ intermediate has also been justified by the following controlled steps.



Figure 7. EPR spectra of electrogenerated (a) $1a^{2+}$ and (b) 2 in CH₃CN/0.1 M Et₄NClO₄ at 77 K.

(1) Addition of IDA to a solution of precatalyst $1a^+$ in CH₂Cl₂ immediately changes the color of the solution from red-brown to yellow due to the formation of $\{Ru^{IV}=O\}$ species which causes the disappearance of the visible transitions of $1a^+$ at 565 and 688 nm with the concomitant growth of two higher energy new bands at 482 and 430 nm (Figures S5 and S6, Supporting Information).^{13a,24a,b} This has also been supported by the mass spectrometry which shows the peaks at m/z = 823.06 and 838.06 corresponding to $\{Ru^{IV}(tpm)(BIAN)-O]ClO_4\}^+$ (calcd 823.20) (4^{2+}) and $\{Ru^{II}(tpm)(BIAN)-OCH_3\}^+$ (calcd 838.23), respectively (Figure S7, Supporting Information). The identification of the



Figure 8. UV–vis spectroelectrochemical plots in $CH_3CN/0.1$ M NEt_4ClO_4 .

solvated species ({ $Ru^{II}(tpm)(BIAN)OCH_3$ }⁺) implies the reactive feature of the intermediate { $Ru^{IV}=O$ } species (4²⁺).

- (2) The yellow solution of the intermediate 4^{2+} reverts back to its original red-brown color $(1a^+)$ upon addition of Me₂S, which is attributed to the oxidation of dimethyl sulfide to dimethyl sulfoxide (GC) with the concomitant reduction of Ru(IV) to Ru(II) (Figure S5, Supporting Information).
- (3) Addition of PPh₃ to the freshly prepared 4^{2+} in CDCl₃ in a 1:1 ratio results in Ph₃P=O as evidenced by its characteristic ³¹P NMR resonance at δ 30 ppm (Figure S8, Supporting Information).^{24a}

The olefin approaches the metal—oxo species either via sideon fashion or end-on fashion.²⁸ However, the observed appreciably higher % conversion in case of *trans*-stilbene (76%) as compared to *cis*-stilbene (66%) (entries 7 and 8, Table 9) suggests that the less hindered side-on approach is preferred. The probable five intermediates involving the transfer of oxygen of {M=O} to the olefinic double bond are (a) concerted transition state, (b) radical, (c) carbocation, (d) π -radical cation, and (e) metallaoxetene^{13a,24a,29} (Scheme 3).

The DFT calculations on the optimized adduct of ruthenium-oxo species $(4^{2+}, \Delta E(S = 0)-(S = 1) = 7506 \text{ cm}^{-1})$ with styrene as a model substrate i.e., [5] (Scheme 4 and

Figure S9, Supporting Information), however, reveals the involvement of a radical intermediate step (**b** in Scheme 3) as it results in the lengthening of the $C_{\alpha}-C_{\beta}$ bond of styrene from 1.32 to1.50 Å in [**5**] with a simultaneous stretching of the Ru–O bond of 1.77 Å in 4²⁺ to 1.93 Å in [**5**]. The formation of a C_{β} -O single bond (1.43 Å) in [**5**] leads to a radical center at C_{α} as has been corroborated by the Mulliken spin density of 0.76 on C_{α} . Moreover, the almost identical calculated Mulliken charge of 1.00 and 0.92 on Ru in optimized 4²⁺ and [**5**], respectively, favors the radical mechanism (Schemes 2–4). The DFT-supported radical pathway can also be substantiated by the fact of no epoxide formation from 1-octene as a model substrate in the presence of radical scavenger TEMPO under the experimental conditions stated in eq 1.

CONCLUSION

In conclusion, the salient features of the article as well as the outlook are stated below.

- The newer classes of Ru–BIAN-derived chloro $[Ru^{II}(tpm)(R-BIAN)(CI)]^+$ (R = 4-OMe, 4-F, 4-Cl, 4-NO₂, $1a^+-1d^+$) and aqua $[Ru^{II}(tpm)(OMe-BIAN)-(H_2O)]^{2+}$ ($3a^{2+}$) derivatives are reported. The BIAN framework with R = H, however, results in a partially hydrolyzed BIAO ([*N*-(phenyl)imino]acenapthenone)-derived first ruthenium complex $[Ru^{II}(tpm)(BIAO)-(CI)]^+$ (2^+) (Chart 2).
- Appreciable variation in the redox potential based on the electronic nature of the "R" group of BIAN in $1a^+-1d^+$ or BIAO in 2^+ has been evident which in effect leads to the experimental observation of four successive reductions particularly in $1d^+$ encompassing $R = NO_2$ group.
- The sensitivity of the -N=C-C=N- or -N=C-C=O bond parameters of BIAN or BIAO in accessible redox states of 1aⁿ-1dⁿ or 2ⁿ (n = +2, +1, 0, -1, -2, -3), respectively, has been reflected in experimental and DFT-optimized structures.
- The formation of the {Ru^{IV}=O} state ([Ru^{IV}(tpm)(R-BIAN)(O)]²⁺, 4²⁺) for the BIAN-derived precursor chloro (1a⁺-1d⁺) or aqua (3a²⁺) derivative makes it an efficient precatalyst for the epoxidation of a wide variety of alkenes. The inaccessibility of the {Ru^{IV}=O} state of the analogous BIAO-derived 2⁺, however, makes it virtually ineffective for the epoxidation process.

The present article thus demonstrates enormous potential of the suitably designed BIAN-derived newer molecular frameworks toward the tunable electrochemical and catalytic applications.

EXPERIMENTAL SECTION

Materials. The ligands tris(1-pyrazolyl)methane $(tpm)^{30}$ and substituted bis(arylimino)acenaphthene (BIAN)^{31} and the precursor complex Ru(tpm)Cl₃·1.5H₂O³² were prepared as reported previously in the literature. The starting material pyrazole was purchased from Sigma-Aldrich. Other solvents and reagents were of reagent grade and used as received. For spectroscopic and electrochemical studies, HPLC-grade solvents were employed.

Physical Measurements. The electrical conductivities of the complexes in CH₃CN were checked by using a Systronic 305 conductivity bridge. ¹H NMR and ³¹P NMR spectra were recorded using Bruker Avance III 400 MHz. FT-IR spectra were recorded on a Nicolet spectrophotometer with samples prepared as KBr pellets. Cyclic voltammetry measurements were performed on a PAR model 273A electrochemistry system. Glassy carbon working electrode,

Table 7. TD-DFT(B3LYP/CPCM/CH₃CN)-Calculated Electronic Transitions for $1a^n - 1d^n$ and 2^n

λ /nm expt (DFT)	$\varepsilon/M^{-1} \text{ cm}^{-1}(f)$	transitions	character
		$1a^{2+}(S=1/2)$	
822(817)	810(0.039)	$HOMO-2 \rightarrow LUMO(98)$	$BIAN(\pi) \rightarrow Ru(d\pi)$
660(614)	1170(0.004)	$HOMO-4(\beta) \rightarrow LUMO+1(\beta)(95)$	$BIAN(\pi) \rightarrow Ru(d\pi)$
435(452)	17 150(0.052)	$HOMO-10(\beta) \rightarrow LUMO(\beta)$ (91)	$BIAN(\pi) \rightarrow Ru(d\pi)$
330(329)	11 890(0.073)	$HOMO-2(\beta) \rightarrow LUMO+2(\beta)$ (64)	$BIAN(\pi)/Ru(d\pi) \rightarrow BIAN(\pi^*)$
277(284)	30 670(0.004)	$HOMO-6(\beta) \rightarrow LUMO+2(\beta)(64)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
2//(201)		$1a^{+}(S=0)$	
658(597)	2710(0.016)	$HOMO \rightarrow LUMO(61)$	$\operatorname{Bu}(d\pi) \to \operatorname{BIAN}(\pi^*)$
552(474)	9200(0.139)	$HOMO = 3 \rightarrow LUMO(69)$	$\operatorname{Ru}(d\pi)/\operatorname{BIAN}(\pi) \to \operatorname{BIAN}(\pi^*)$
478(448)	15 600(0.041)	$HOMO-3 \rightarrow LUMO(51)$	$\operatorname{Ru}(d\pi)/\operatorname{BIAN}(\pi) \rightarrow \operatorname{BIAN}(\pi^*)$
270(270)	7360(0.022)	$HOMO-3 \rightarrow LUMO(31)$	$\operatorname{Ru}(\mathfrak{u}_{n})/\operatorname{Budv}(\mathfrak{u}) \to \operatorname{Budv}(\mathfrak{u})$ $\operatorname{Budv}(\mathfrak{u}_{n})/\operatorname{Budv}(\mathfrak{u}_{n}) \to \operatorname{Budv}(\mathfrak{u}^{*})$
3/9(3/0)	13250(0.022)	$HOMO=5 \rightarrow LUMO+1(65)$	$\operatorname{BIAN}(\pi) \to \operatorname{BIAN}(\pi^*)$
328(318)	13330(0.231)	$HOMO = 4 \rightarrow LUMO + 4(61)$	$\operatorname{DIAN}(n) \to \operatorname{DIAN}(n^*)$ $\operatorname{DIAN}(\pi) \to \operatorname{DIAN}(\pi^*)$
288(201)	211/0(0.018)	$HOMO=4 \rightarrow LOMO+4(01)$	$\operatorname{BIAN}(n) \to \operatorname{BIAN}(n^{+})$
(50((40))	740(0.004)	Ia (S = 1/2)	$\mathbf{D}_{-}(1_{-}) \rightarrow \mathbf{D}(\mathbf{A}\mathbf{N}(-*))$
058(048)	/40(0.004)	$HOMO(p) \rightarrow LOMO(p)(98)$	$\operatorname{Ku}(d\pi) \to \operatorname{BIAN}(\pi^*)$
5/5(562)	6840(0.003)	$HOMO-I(\alpha) \rightarrow LUMO(\alpha)(58)$	$\operatorname{Ru}(\mathrm{d}\pi) \to \operatorname{BIAN}(\pi^*)$
485(462)	15 680(0.223)	$HOMO-1(\alpha) \rightarrow LUMO(\alpha)(69)$	$\operatorname{Ru}(d\pi)/\operatorname{BIAN}(\pi) \to \operatorname{BIAN}(\pi^*)$
379(378)	7470(0.040)	$HOMO-4(\beta) \rightarrow LUMO(\beta)(61)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
332(329)	13 360(0.002)	$HOMO-4(\beta) \rightarrow LUMO+1(\beta)(75)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
288(289)	20 820(0.013)	$HOMO-3(\beta) \rightarrow LUMO+3(\beta)(83)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)/tpm(\pi^*)$
		$1b^{2+} (S = 1/2)$	
793(743)	950(0.017)	$HOMO-2(\beta) \rightarrow LUMO(\beta)(97)$	$BIAN(\pi) \rightarrow Ru(d\pi)$
517(484)	3190(0.014)	$HOMO-3(\beta) \rightarrow LUMO+1(\beta)(62)$	$\operatorname{Ru}(\mathrm{d}\pi) \to \operatorname{BIAN}(\pi^*)$
442(439)	11 210(0.050)	$HOMO-2(\beta) \rightarrow LUMO+1(\beta)(74)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
328(319)	11 560(0.004)	$HOMO(\alpha) \rightarrow LUMO+2(\alpha)(64)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
283(281)	20 070(0.013)	$HOMO-6(\beta) \rightarrow LUMO+2(\beta)(84)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
		$\mathbf{1b}^+ (S=0)$	
662(590)	2140(0.022)	$HOMO-1 \rightarrow LUMO(55)$	$\operatorname{Ru}(d\pi) \to \operatorname{BIAN}(\pi^*)$
556(503)	7670(0.079)	$HOMO-2 \rightarrow LUMO(51)$	$\operatorname{Ru}(d\pi) \to \operatorname{BIAN}(\pi^*)$
476(441)	11 920(0.252)	HOMO \rightarrow LUMO+1(62)	$\operatorname{Ru}(d\pi) \to \operatorname{BIAN}(\pi^*)$
328(311)	13 660(0.006)	$HOMO-4 \rightarrow LUMO+1(52)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
314(307)	15 100(0.031)	$HOMO-4 \rightarrow LUMO+1(56)$	$BIAN(\pi)/Cl(\pi) \rightarrow BIAN(\pi^*)$
289(273)	16 120(0.005)	$HOMO-6 \rightarrow LUMO+1(69)$	$BIAN(\pi)/Cl(\pi) \rightarrow BIAN(\pi^*)$
		1b $(S = 1/2)$	
590(559)	6730(0.007)	$HOMO-1(\alpha) \rightarrow LUMO(\alpha)(52)$	$\operatorname{Ru}(d\pi) \to \operatorname{BIAN}(\pi^*)$
558(543)	9370(0.020)	$HOMO-1(\beta) \rightarrow LUMO(\beta)(82)$	$\operatorname{Ru}(d\pi) \to \operatorname{BIAN}(\pi^*)$
498(493)	13360(0.004)	$HOMO - 2(\beta) \rightarrow LUMO(\beta)(85)$	$\operatorname{Bu}(d\pi) \to \operatorname{BIAN}(\pi^*)$
331(352)	13300(0.001) 11.720(0.038)	$HOMO - 5(\beta) \rightarrow LUMO(\beta)(82)$	$\operatorname{BIAN}(\pi) \to \operatorname{BIAN}(\pi^*)$
316(306)	13 840(0.006)	$HOMO - 4(\beta) \rightarrow LUMO + 1(\beta)(68)$	$\operatorname{BIAN}(\pi) \to \operatorname{BIAN}(\pi^*)$
201(204)	15 400(0.010)	$HOMO = 6(\beta) \rightarrow LUMO(\beta)(68)$	$\operatorname{BIAN}(\pi) \to \operatorname{BIAN}(\pi^*)$
291(294)	13 400(0.010)	$1c^{2+}(S-1/2)$	$\operatorname{DEU}(n) \to \operatorname{DEU}(n)$
706(741)	1050(0.014)	$HOMO-2(\beta) \rightarrow LLIMO(\beta)(98)$	$\operatorname{BIAN}(\pi) \rightarrow \operatorname{Bu}(4\pi)$
790(741) 576(562)	1030(0.014)	$HOMO = 2(p) \rightarrow LOMO(p)(98)$	$\operatorname{DIAN}(n) \to \operatorname{Ru}(dn)$ $\operatorname{PLAN}(\pi) \to \operatorname{Pu}(d\pi)$
570(505)	1390(0.004)	$HOMO = O(p) \rightarrow LOMO(p)(82)$	$BIAN(n) \rightarrow Ru(dn)$
521(515)	3840(0.022)	$HOMO = 8(\beta) \rightarrow LOMO(\beta)(95)$	$BIAN(\pi) \to Ru(d\pi)$
445(438)	12 /00(0.063)	$HOMO-2(\beta) \rightarrow LOMO+1(\beta)(74)$	$BIAN(\pi) / Ru(d\pi) \rightarrow BIAN(\pi^{*})$
329(323)	13 130(0.006)	$HOMO-I(\beta) \rightarrow LUMO+I(\beta)(91)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
286(266)	23 110(0.009)	$HOMO-7(\beta) \rightarrow LUMO+2(\beta)(87)$	$BIAN(\pi)/Cl(\pi) \rightarrow BIAN(\pi^*)$
	<i>(</i>)	$\mathbf{1c}^+ (S=0)$	
664(593)	2250(0.024)	$HOMO-1 \rightarrow LUMO(55)$	$\operatorname{Ru}(\mathrm{d}\pi)/\operatorname{BIAN}(\pi) \to \operatorname{BIAN}(\pi^*)$
561(506)	8810(0.085)	$HOMO-2 \rightarrow LUMO(51)$	$\operatorname{Ru}(\mathrm{d}\pi) \to \operatorname{BIAN}(\pi^*)$
477(441)	13 570(0.241)	$HOMO \rightarrow LUMO+1(51)$	$\operatorname{Ru}(\mathrm{d}\pi) \to \operatorname{BIAN}(\pi^*)$
329(318)	16 340(0.039)	$HOMO-9 \rightarrow LUMO(67)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
315(307)	17 680(0.009)	$HOMO-5 \rightarrow LUMO+1(63)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
290(264)	19 270(0.001)	$HOMO-7 \rightarrow LUMO+1 (59)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
		1c (S = 1/2)	
590(567)	9440(0.049)	$HOMO-1(\beta) \rightarrow LUMO(\beta)(72)$	$\operatorname{Ru}(d\pi) \to \operatorname{BIAN}(\pi^*)$
564(504)	11 290(0.007)	$HOMO-1(\beta) \rightarrow LUMO(\beta)(79)$	$\operatorname{Ru}(d\pi) \to \operatorname{BIAN}(\pi^*)$
497(461)	14 410(0.051)	$HOMO-1(\alpha) \rightarrow LUMO(\alpha)(79)$	$\operatorname{Ru}(d\pi)/\operatorname{Cl}(\pi) \to \operatorname{BIAN}(\pi^*)$
330(330)	13 990(0.008)	$HOMO(\beta) \rightarrow LUMO(\beta)(81)$	$\operatorname{Ru}(d\pi)/\operatorname{BIAN}(\pi) \to \operatorname{BIAN}(\pi^*)$
317(311)	15 200(0.006)	$HOMO-5(\alpha) \rightarrow LUMO(\alpha)(61)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
291(279)	15 400(0.001)	HOMO-4(α) \rightarrow LUMO+2(α)(79)	$BIAN(\pi) \rightarrow BIAN(\pi^*)/tpm(\pi^*)$

Table 7. continued

λ /nm expt (DFT)	$\varepsilon/M^{-1} \text{ cm}^{-1} (f)$	transitions	character
		$1d^{2+}(S=1/2)$	
827(879)	810(0.046)	$HOMO(\beta) \rightarrow LUMO(\beta)$ (98)	$BIAN(\pi) \rightarrow Ru(d\pi)$
657(594)	1140(0.001)	$HOMO-6(\beta) \rightarrow LUMO(\beta)(74)$	$BIAN(\pi) \rightarrow Ru(d\pi)$
434(417)	17 200(0.030)	HOMO-3(β) \rightarrow LUMO+1 (β)(74)	$\operatorname{Ru}(d\pi) \to \operatorname{BIAN}(\pi^*)$
330(331)	11 950(0.002)	HOMO-9(β) \rightarrow LUMO+1 (β)(51)	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
277(274)	30 680(0.150)	HOMO-6(α) \rightarrow LUMO+3 (α)(61)	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
		$\mathbf{1d}^+ \ (S=0)$	
652(595)	2780(0.028)	$HOMO-1 \rightarrow LUMO(50)$	$\operatorname{Ru}(d\pi) \to \operatorname{BIAN}(\pi^*)$
552(507)	9230(0.049)	HOMO \rightarrow LUMO+1(54)	$\operatorname{Ru}(d\pi) \to \operatorname{BIAN}(\pi^*)$
478(454)	15 570(0.028)	$HOMO-1 \rightarrow LUMO+1(66)$	$\operatorname{Ru}(d\pi) \to \operatorname{BIAN}(\pi^*)$
330(301)	13 230(0.099)	$HOMO-4 \rightarrow LUMO+1(52)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
287(281)	21 180(0.021)	HOMO-6 \rightarrow LUMO+2(53)	$BIAN(\pi)/Cl(\pi) \rightarrow BIAN(\pi^*)$
		1d (S = 1/2)	
801(836)	1310(0.010)	$HOMO(\beta) \rightarrow LUMO(\beta)$ (96)	$\operatorname{Ru}(\mathrm{d}\pi) \to \operatorname{BIAN}(\pi^*)$
578(578)	10 260(0.002)	HOMO \rightarrow 1(α) \rightarrow LUMO+1(α)(77)	$\operatorname{Ru}(\mathrm{d}\pi) \to \operatorname{BIAN}(\pi^*)$
488(455)	16 010(0.009)	HOMO $\rightarrow 1(\alpha) \rightarrow LUMO + 2(\alpha)(59)$	$\operatorname{Ru}(d\pi) \to \operatorname{BIAN}(\pi^*)$
316(314)	30 580(0.005)	HOMO \rightarrow 7(α) \rightarrow LUMO(α)(58)	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
283(295)	36 740(0.009)	$HOMO-6(\beta) \rightarrow LUMO(\beta)(62)$	$BIAN(\pi) \rightarrow BIAN(\pi^*)$
		$2^{2+}(S=1/2)$	
397(442)	6120(0.067)	HOMO $-7(\beta) \rightarrow$ LUMO (β)(88)	$BIAO(\pi^*) \rightarrow Ru(d\pi)$
354(347)	11 270(0.048)	$HOMO(\alpha) \rightarrow LUMO+1(\alpha)(57)$	$BIAO(\pi) \rightarrow BIAO(\pi^*)$
255(326)	17 860(0.001)	$HOMO-2(\alpha) \rightarrow LUMO+1(\alpha)(56)$	$BIAO(\pi) \rightarrow BIAO(\pi^*)$
		$2^+ (S=0)$	
638(566)	10 550(0.170)	$HOMO-2 \rightarrow LUMO(47)$	$\operatorname{Ru}(\mathrm{d}\pi) \to \operatorname{BIAO}(\pi^*)$
493(439)	6620(0.016)	$HOMO-1 \rightarrow LUMO+1(67)$	$\operatorname{Ru}(\mathrm{d}\pi) \to \operatorname{BIAO}(\pi^*)$
378(372)	3630(0.006)	$HOMO-5 \rightarrow LUMO(68)$	$\operatorname{Cl}(\pi)/\operatorname{BIAO}(\pi) \to \operatorname{BIAO}(\pi^*)$
314(337)	16 090(0.013)	$HOMO \rightarrow LUMO+2(48)$	$\operatorname{Ru}(\mathrm{d}\pi) \to \operatorname{BIAO}(\pi^*)$
301(305)	16 700(0.022)	$HOMO-4 \rightarrow LUMO+1(66)$	$BIAO(\pi)/Cl(\pi) \to BIAO(\pi^*)$
<i>(</i>)	<i>(</i>)	2(S = 1/2)	
483(459)	13 700(0.001)	$HOMO-2(\alpha) \rightarrow LUMO(\alpha)(79)$	$\operatorname{Ru}(d\pi) \to \operatorname{BIAO}(\pi^*)$
362(365)	16 480(0.008)	$HOMO-4(\beta) \rightarrow LUMO(\beta)(62)$	$BIAO(\pi) \rightarrow BIAO(\pi^*)$
337(326)	17 000(0.060)	$HOMO-4(\alpha) \rightarrow LUMO(\alpha)(54)$	$BIAO(\pi) \rightarrow BIAO(\pi^*)$
249(317)	21 740(0.019)	$HOMO-5(\alpha) \rightarrow LUMO(\alpha)(38)$	$BIAO(\pi)/Cl(\pi) \to BIAO(\pi^*)$
		$HOMO-9(\beta) \rightarrow LUMO(\beta)(37)$	$Cl(\pi)/BIAO(\pi) \rightarrow BIAO(\pi^*)$

Table 8. Optimization of the Critical Reaction Parameters^a

entry	solvent	catalyst 1a ⁺ (mmol %)	oxidant	% conversion ^b
1	CH_2Cl_2	1	H_2O_2	22
2	CH_2Cl_2	1	TBHP	39
3	CH_2Cl_2	1	m-CPBA	27
4	CH_2Cl_2	1	IDA	88
5	CH_2Cl_2	0.5	IDA	75
6	CH_2Cl_2	2	IDA	88
7	C ₂ H ₅ OH	1	H_2O_2	27
8	C ₂ H ₅ OH	1	TBHP	32
9	C ₂ H ₅ OH	1	m-CPBA	19
10	C ₂ H ₅ OH	1	IDA	62
11	CH ₃ CN	1	H_2O_2	23
12	CH ₃ CN	1	TBHP	28
13	CH ₃ CN	1	m-CPBA	24
14	CH ₃ CN	1	IDA	68

^{*a*}1 mmol of styrene, solvent = 3 mL, oxidant = 2 equiv, catalyst:substrate:oxidant = 1:100:200, 298 K, 6 h under aerobic conditions. ^{*b*}Percent (%) conversions were calculated by GC with *n*-decane as an internal standard as well as by GC-MS. TBHP: *tert*-butyl hydrogen peroxide. IDA: iodobenzene diacetate. *m*-CPBA: *m*-chloroperbenzoic acid. H_2O_2 : hydrogen peroxide.

platinum wire auxiliary electrode, and a saturated calomel reference electrode (SCE) were used in a standard three-electrode configuration

cell. A platinum wire-gauze working electrode was used for the constant potential coulometry experiment. UV-vis spectral studies were performed on a PerkinElmer Lambda 950 spectrophotometer. UV-vis spectroelectrochemical studies were performed on a BAS SEC2000 spectrometer system. The supporting electrolyte was Et₄NClO₄, and the solute concentration was $\sim 10^{-3}$ M. All electrochemical experiments were carried out under a dinitrogen atmosphere at 298 K. The half-wave potential E° was set equal to $0.5(E_{pa} + E_{pc})$, where E_{pa} and E_{pc} are anodic and cathodic cyclic voltammetry peak potentials, respectively. The elemental analyses were carried out on a Thermoquest (EA 1112) microanalyzer. Electrospray mass spectra (ESI-MS) were recorded on a Bruker's Maxis Impact (282001.00081). The EPR measurements were made with a JEOL model FA200 electron spin resonance spectrometer. GC analysis was done by a Shimadzu GC-2014 gas chromatograph with a FID detector using a capillary column (112-2562 CYCLODEXB, from J&W Scientific, length 60 m, inner diameter 0.25 mm, film 0.25 μ m), and GC-MS analysis was performed on an Agilent Technologies 7890A GC system coupled with a 5975C inert XL EI/CI Mass Selective Detector (GCMSD) with its triple-axis detector.

Crystallography. Single crystals of $[1a]ClO_4$ and $[2]ClO_4$ were grown by slow evaporation of their 1:2 dichloromethane-*n*-hexane solutions, while those of $[3a](PF_6)_2$ were obtained from 1:1 acetone-*n*-hexane solution. X-ray crystal data for $[1a]ClO_4$, $[2]ClO_4$, and $[3a](PF_6)_2$ were collected on RIGAKU SATURN-724+, Agilent Technologies (Oxford Diffraction) SUPER NOVA, and Bruker SMART APEX-II, CCD diffractometers, respectively. The data collection for $[1a]ClO_4$ and $[2]ClO_4$ were evaluated using the

Table 9. Catalytic Epoxidation of Olefins^a

entry	substrate	product	%conversion (selective		vity) ^b
			1a ⁺	2+	3a ²⁺
1	\bigcirc	C	88(80)	10(50)	89(82)
2			84(58+26)	0	NC
3	$\bigcirc \frown$		82(80)	0	NC
4	$\bigcirc \sim$		84(80)	0	NC
5	F	F C C	82(100)	0	NC
6	NC	NC	84(100)	18(10)	NC
7	$\bigcirc \bigcirc \bigcirc$		66(100)	0	70(80)
8			76(100)	12(10)	NC
9	ССНО	CHO O CHO	84(80)	18	NC
10	$\bigcirc \frown$	$\bigcirc \checkmark $	82(100)	0	NC
11			84(75)	0	NC
12	$\bigcirc \frown$	$\bigcirc \checkmark \rega $	88(100)	0	NC
13	\bigcirc		66(70)	0	NC
14	\bigcirc	\$	8/76(70)	12(0)	NC
15			90(100)	25	92(100)
16	~~~~		67(100)	0	NC
17	$\overset{+}{\overset{+}{\overset{-}{\overset{-}{\overset{-}{\overset{+}}}}}$		80(100) + 20	0	NC

^{*a*}1 mmol of styrene, solvent = 3 mL, oxidant = 2 equiv, catalyst:substrate:oxidant = 1:100:200, 298 K, 6 h under aerobic conditions. Products are analyzed by GC with *n*-decane as an internal standard as well as by GC-MS. ^{*b*}Selectivity in terms of epoxide formation. NC = Not determined.

CrysAlisPro CCD and CrystalClear-SM Expert software, respectively. The data for $[1a]ClO_4$ were collected by the standard $\phi-\omega$ scan techniques, scaled and reduced using CrysAlisPro RED software, and by the ω scan technique for $[2]ClO_4$. The cell refinement and data reduction in $[3a](PF_6)_2$ were performed with SAINT V7.68A. The structures were solved by direct method using SHELXS-97 and refined by full matrix least-squares with SHELXL-97, refining on $F^{2,33}$ All non-hydrogen atoms were refined anisotropically. The hydrogen atoms were placed in geometrically constrained positions and refined with isotropic temperature factors, generally 1.2Ueq of their parent atoms.

Except H1A and H1B (protons of aqua molecule in $[3a](PF_6)_2$), the other hydrogen atoms were added in the refinement process as per the riding model. The disordered solvent molecules in $[1a]ClO_4$ and $[3a](PF_6)_2$ were SQUEEZED by the PLATON³⁴ program. CCDC 1052730 ($[1a]ClO_4$), 1052731 ($[2]ClO_4$), and 1052732 ($[3a](PF_6)_2$) contain supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Center via www.ccdc.cam.ac.uk/data_request/cis.

Computational Details. Full geometry optimizations were carried out by using the density functional theory method at the (R)B3LYP Scheme 2. Proposed Mechanistic Pathway



Scheme 3. Probable Intermediates in Metal-Catalyzed Oxygen Transfer Reaction from Metal–Oxo to Olefins



Scheme 4. Proposed Radical Intermediate During Epoxidation



level for $1a^+\!-\!1d^+\!,\,1a^-\!-\!1d^-\!,\,2^+\!,\,2^-\!,$ and $3a^{2+}$ and (U)B3LYP level^{35} for $1a^{2+}-1d^{2+}$, 1a-1d, $1d^{2-}$, $1d^{3-}$, and 2^{2+} , and 2 used the Gaussian 09 program package.³⁶ Except ruthenium all other elements were assigned the 6-31G* basis set. The SDD (in 2^{2+} , 2^+ , $2, 2^-$) and LANL2DZ (in $1a^{2+}-1d^{2+}$, $1a^+-1d^+$, 1a-1d, $1a^--1d^-$, $3a^{2+}$) basis sets with effective core potential were employed for the ruthenium atom.³⁷ The vibrational frequency calculations were performed to ensure that the optimized geometries represent the local minima and there are only positive eigenvalues. All calculations were performed with the Gaussian09 program package. Vertical electronic excitations based on (R)B3LYP/(U)B3LYP-optimized geometries were computed for $1a^{n}-1d^{n}$ (n = +2, +1, 0) and 2^{n} (n = +2, +1, 0) using the timedependent density functional theory (TD-DFT) formalism³⁸ in acetonitrile using a conductor-like polarizable continuum model (CPCM).³⁹ Chemissian 1.7⁴⁰ was used to calculate the fractional contributions of various groups to each molecular orbital. All calculated structures were visualized with ChemCraft.⁴

General Procedure for Catalytic Epoxidation Studies. The catalyst [1a]ClO₄ (4.13 mg, 0.0049 mmol, 1 mol %) in 3 mL of CH₂Cl₂ was taken in a vial and stirred for ~15 min at 298 K. The olefin (0.50 mmol) was then added to it with simultaneous stirring followed by the addition of iodobenzene diacetate (2 equivalents with respect to olefin) as the oxidant. The mixture was stirred at 298 K for 6 h. The reaction mixture was then diluted with water and extracted with ethyl acetate (10 mL) three times. The combined organic extracts were dried over Na₂SO₄ and concentrated under reduced pressure, and the yields were determined by gas chromatography with respect to the internal standard, *n*-decane (1 mmol). A 2.0 μ L aliquot was subjected for analysis by GC-FID in each case.

Preparation of the Complexes. The complexes $[1a]ClO_4$ – $[1d]ClO_4$ and $[2]ClO_4$ were synthesized by following a general procedure using respective R-BIAN. For $[2]ClO_4$ R-BIAN with R = H was used. The details are given below for $[1a]ClO_4$.

Synthesis of [Ru(tpm)(OMe-BIAN)CI]CIO₄, [1a]CIO₄. Ru(tpm)-Cl₃·1.5 H₂O (101 mg, 0.24 mmol) and the ligand R-BIAN (R = OCH₃, Chart 2) (106 mg, 0.27 mmol) were taken in 40 mL of methanol. The reaction mixture was heated at reflux with stirring under a dinitrogen atmosphere for 8 h, and the color of the solution turned to reddish brown. The solvent was then evaporated under reduced pressure. The solid mass thus obtained was dissolved in 2 mL of acetonitrile followed by addition of a saturated aqueous solution of NaClO₄. The solution was then refrigerated for 6 h, and the resulting solid was filtered through a Buchner funnel and dried under vacuum. It was then purified on a neutral alumina column by using 4:1 CH₂Cl₂– CH₃CN as eluant. Evaporation of the solvent under reduced pressure yielded the solid product of [1a]ClO₄.

[1a]ClO₄: Yield 114 mg (57%). ¹H NMR in CDCl₃ [δ/ppm (J/Hz)]: 9.82 (s, 1H), 8.77 (d, 2.6, 1H), 8.36 (d, 2.4, 2H), 8.03 (d, 8.2, 2H), 7.70 (d, 7.2, 2H), 7.49 (t, 7.4, 2H), 7.24 (m,4H), 7.14 (d, 7.2, 2H), 6.99 (d, 2.1, 2H), 6.71 (d, 7.5, 2H), 6.47 (d, 2.6, 1H), 6.19 (t, 2.5, 3H), 3.80 (s, 6H). MS (ESI+, CH₃CN): m/z [1a⁺] calcd 743.122; found 743.128. IR (KBr): ν (ClO₄⁻⁷, cm⁻¹): 1093, 622. Molar conductivity ($\Lambda_{\rm M}$ (Ω^{-1} cm² M⁻¹), CH₃CN): 118. Anal. Calcd for C₃₆H₃₀N₈Cl₂O₆Ru: C, 51.31; H, 3.59; N, 13.30. Found: C, 51.31; H, 3.50; N, 13.18.

 $\begin{array}{l} [1b] \text{ClO}_4: \text{ Yield 112 mg (58\%).} \ ^{1}\text{H NMR in CDCl}_3 \ [\delta/\text{ppm }(J/\text{Hz})]: 9.89 \ (s, 1H), 8.80 \ (d, 2.6, 1H), 8.38 \ (d, 2.6, 2H), 8.07 \ (d, 8.2, 2H), 7.78 \ (m, 2H), 7.52 \ (d, 7.8, 2H), 7.36 \ (m, 2H), 7.24 \ (m, 1H), 7.17 \ (d, 7.2, 2H), 7.00 \ (d, 1.6, 2H), 6.94 \ (m, 2H), 6.50 \ (t, 2.4, 1H), 6.30 \ (m, 2H), 6.23 \ (t, 2.4, 2H). \text{MS (ESI+, CH}_3\text{CN}): m/z \ \{1b^+\} \text{ calcd 719.082; found 719.104. IR \ (KBr): $\nu(\text{ClO}_4^-, \text{cm}^{-1}): 1089, 622. \text{ Molar conductivity } (\Lambda_{\text{M}} \ (\Omega^{-1} \ \text{cm}^2 \ \text{M}^{-1}), \ \text{CH}_3\text{CN}): 114. \text{ Anal. Calcd for C}_{34}\text{H}_{24}\text{N}_8\text{Cl}_2\text{F}_2\text{O}_4\text{Ru}: \text{C}, 49.89; \text{H}, 2.96; \text{N}, 13.69. \text{Found: C}, 49.74; \text{H}, 2.82; \text{N}, 13.23. \end{array}$

 $\begin{array}{l} [1c] ClO_4: \mbox{ Yield 105 mg (52\%)}. \ ^{1}\mbox{H NMR in CDCl}_3 \ [\delta/\mbox{pm }(J/\mbox{Hz})]: 9.87 (s, 1H), 8.79 (d, 2.7, 1H), 8.37 (d, 2.6, 2H), 8.08 (d, 8.2, 2H), 7.74 (d, 6.6, 2H), 7.64 (d, 7.1, 2H), 7.52 (t, 8.1, 2H), 7.20 (m, 5H), 7.01 (d, 2.2, 2H), 6.51 (t, 2.6, 1H), 6.26 (d, 8.9, 1H), 6.24 (m, 3H). \mbox{MS (ESI+, CH}_3 CN): $m/z \ \{1c^+\}$ calcd 751.023; found 751.060. \label{eq:KBr} (ClO_4^-, cm^{-1}): 1086, 622. \mbox{Molar conductivity } (\Lambda_{\rm M} \ (\Omega^{-1} \ cm^2 \ M^{-1}), \ CH_3 CN): 122. \mbox{ Anal. Calcd for $C_{34}{\rm H}_{24}{\rm N}_8{\rm Cl}_4{\rm O}_4{\rm Ru}: C, 47.96; \ H, 2.84; \ N, 13.16. \ Found: C, 48.12; \ H, 2.87; \ N, 13.04. \end{array}$

47.96; H, 2.84; N, 13.16. Found: C, 48.12; H, 2.87; N, 13.04. **[1d]**ClO₄: Yield 118 mg (57%). ¹H NMR in CDCl₃ [δ /ppm (J/Hz)]: 9.98 (s, 1H), 8.86 (d, 2.8, 1H), 8.57 (d, 8.3, 2H), 8.42 (d, 2.7, 2H), 8.15 (d, 8.2, 4H), 8.01 (d, 7.5, 2H), 7.56 (t, 7.8, 2H), 7.28 (d, 3.9, 1H), 7.12 (d, 7.2, 2H), 7.00 (d, 2.1, 2H), 6.57 (m, 1H), 6.50 (m, 2H), 6.23 (t, 2.6, 2H). MS (ESI+, CH₃CN): m/z {1d⁺} calcd 773.071; found 773.080. IR (KBr): ν (ClO₄⁻⁷, cm⁻¹): 1091, 622. Molar conductivity ($\Lambda_{\rm M}$ (Ω^{-1} cm² Λ^{-1}), CH₃CN): 108. Anal. Calcd for C₃₄H₂₄N₁₀Cl₂O₈Ru: C, 46.80; H, 2.77; N, 16.05. Found: C, 46.95; H, 2.80; N, 15.92.

[2]ClO₄: Yield 74 mg (44%). ¹H NMR in CDCl₃ [δ/ppm (J/Hz)]: 9.96 (s, 1H), 8.72 (d, 2.7, 1H), 8.56 (d, 2.1, 1H), 8.53 (d, 2.7, 1H), 8.42 (m, 1H), 8.41 (d, 1.8, 1H), 8.36 (d, 8.1, 1H), 8.18 (d, 8.2, 1H), 7.87 (t, 7.2, 2H), 7.53 (m, 6H), 6.97 (d, 2.1, 1H), 6.82 (d, 2.2, 1H), 6.68 (t, 2.5, 1H), 6.44 (t, 2.6, 1H), 6.14 (t, 2.6, 1H). MS (ESI+, CH₃CN): m/z [2⁺] calcd 608.053; found 608.037. IR (KBr): ν (ClO₄⁻, cm⁻¹): 1090, 623; ν (C=O, cm⁻¹): 1690. Molar conductivity ($\Lambda_{\rm M}$ (Ω^{-1} cm² M⁻¹), CH₃CN): 92. Anal. Calcd for C₂₈H₂₁Cl₂N₇O₅Ru: C, 47.54; H, 2.99; N, 13.86. Found: C, 47.68; H, 2.80; N, 13.90.

Preparation of $[Ru(tpm)(OMe-BIAN)(H_2O)](CIO_4)_2$ ([3a]-(CIO_4)_2). The mixture of $[1a]CIO_4$ (100 mg, 0.12 mmol) and AgCIO_4 (100 mg, 0.48 mmol) in 1:3 acetone-water (25 mL) was heated at reflux under stirring for 6 h. The initial reddish brown solution turned to yellow with the precipitation of AgCl. It was then cooled to room temperature and filtered through a sintered glass crucible (G4). The filtrate was concentrated to 5 mL under reduced pressure, and excess NaClO₄ was added to it. The solid $[3a](CIO_4)_2$

thus obtained was filtered off and washed with a few drops of chilled Milli-Q water followed by drying in vacuo over P_4O_{10} .

[3a](ClO₄)₂: Yield 80 mg (73%). ¹H NMR in D₂O [δ/ppm (J/Hz)]: 9.70 (s, 1H), 8.56 (d, 8.0, 1H), 8.46 (d, 3.0, 1H), 8.35 (d, 2.7, 1H), 8.19 (d, 6.0, 1H), 8.13 (d, 7.0, 1H), 7.89 (d, 2.1, 1H), 7.84 (d, 2.2, 1H), 7.53 (m, 2H), 7.16 (m, 8H), 6.70 (m, 1H), 6.61 (d, 3.0, 1H), 6.34 (m, 4H) 3.79 (s, 6H). MS (ESI+, CH₃OH): m/z [3a(ClO₄)₂-ClO₄-H₂O]⁺ calcd 807.102; found 807.092. IR (KBr): ν (OH, cm⁻¹): 3438; ν (ClO₄⁻, cm⁻¹): 1092, 623. Molar conductivity ($\Lambda_{\rm M}$ (Ω^{-1} cm² M⁻¹), H₂O): 195. Anal. Calcd for C₃₆H₃₂N₈Cl₂O₁₁Ru: C, 46.76; H, 3.49; N, 12.12. Found: C, 46.54; H, 3.21; N, 12.32.

Caution! Perchlorate salts of metal complexes with organic ligands are potentially explosive. Heating of dried samples must be avoided; handling of even a small amount has to proceed with great caution.

ASSOCIATED CONTENT

S Supporting Information

X-ray crystallographic files in CIF format for $[1a]ClO_4$, $[2]ClO_4$, and $[3a](PF_6)_2$, mass spectra, cyclic and differential pulse voltammograms of $[3a](PF_6)_2$, electronic spectra, ¹H NMR, ³¹P NMR spectra, DFT-optimized structures, bond parameters, MO compositions, and energies of DFT optimized states. The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/ acs.inorgchem.5b00615.

AUTHOR INFORMATION

Corresponding Author

*E-mail: lahiri@chem.iitb.ac.in.

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

Financial support received from the Department of Science and Technology, University Grants Commission (fellowship to A.H. and R.R.), Council of Scientific and Industrial Research (fellowship to A.D., H.A. and S.M.), New Delhi (India) is gratefully acknowledged.

REFERENCES

(1) (a) Argazzi, R.; Iha, N. Y. M.; Zabri, H.; Odobel, F.; Bignozzi, C. A. Coord. Chem. Rev. 2004, 248, 1299-1316. (b) Sun, L.; Hammarström, L.; Åkermark, B.; Styring, S. Chem. Soc. Rev. 2001, 30, 36-49. (c) Concepcion, J. J.; Jurss, J. W.; Brennaman, M. K.; Hoertz, P. G.; Patrocinio, A. O. T.; Iha, N. Y. M.; Templeton, J. L.; Meyer, T. J. Acc. Chem. Res. 2009, 42, 1954-1965. (d) Balzani, V.; Credi, A.; Venturi, M. Chem. Soc. Rev. 2009, 38, 1542-1550. (e) Balzani, V.; Credi, A.; Silvi, S.; Venturi, M. Chem. Soc. Rev. 2006, 35, 1135-1149. (f) Wolfbeis, O. S.; Klimant, I.; Werner, T.; Huber, C.; Kosch, U.; Krause, C.; Neurauter, G.; Dürkop, A. Sens. Actuators, B 1998, 51, 17-24. (g) Hurst, J. K. Coord. Chem. Rev. 2005, 249, 313-328. (h) Zhang, S.; Baldino, S.; Baratta, W. Organometallics 2013, 32, 5299-5304. (i) Ma, G.; McDonald, R.; Ferguson, M.; Cavell, R. G.; Patrick, B. O.; James, B. R.; Hu, T. Q. Organometallics 2013, 26, 846-854. (j) Barton, J. K.; Olmon, E. D.; Sontz, P. A. Coord. Chem. Rev. 2011, 255, 619-634. (k) Lim, M. H.; Song, H.; Olmon, E. D.; Dervan, E. E.; Barton, J. K. Inorg. Chem. 2009, 48, 5392-5397. (1) Puckett, C. A.; Barton, J. K. Biochemistry 2008, 47, 11711-11716. (m) Schulz, M.; Karnahl, M.; Schwalbe, M.; Vos, J. G. Coord. Chem. Rev. 2012, 256, 1682-1705. (n) Hoogenboom, R.; Schubert, U. S. Chem. Soc. Rev. 2006, 35, 622-629.

(2) (a) O'Brien, C.; Wong, M. Y.; Cordes, D. B.; Slawin, A. M. Z.; Zysman-Colman, E. Organometallics **2015**, *34*, 13–22. (b) Nunes, C. D.; Vaz, P. D.; Félix, V.; Veiros, L. F.; Moniz, T.; Rangel, M.; Realista, S.; Mourato, A. C.; Calhorda, M. J. Dalton Trans. **2015**, *44*, 5125– 5138. (c) Schmitz, M.; Seibel, M.; Kelm, H.; Demeshko, S.; Meyer, F.;

Krüger, H.-J. Angew. Chem., Int. Ed. 2014, 53, 5988-5992. (d) Tao, W.-J.; Li, J.-F.; Peng, A.-Q.; Sun, X. -L; Yang, X.-H.; Tang, Y. Chem.-Eur. J. 2013, 19, 13956-13961. (e) Fedushkin, I. L.; Sokolov, V. G.; Piskunov, A. V.: Makarov, V. M.: Baranov, E. V.: Abakumav, G. A. Chem. Commun. 2014, 50, 10108-10111. (f) Wen, C.; Yuan, S.; Shi, Q.; Yue, E.; Liu, D.; Sun, W.-H. Organometallics 2014, 33, 7223-7231. (g) Bridges, C. R.; McCormick, T. M.; Gibson, G. L.; Hollinger, J.; Seferos, D. S. J. Am. Chem. Soc. 2013, 135, 13212-13219. (h) Fedushkin, I. L.; Markina, O. V.; Lukoyanov, A. N.; Morozov, A. G.; Baranov, E. V.; Maslov, M. O.; Ketkov, S. Y. Dalton Trans. 2013, 42, 7952-7961. (i) Papanikolaou, P. A.; Gdaniec, M.; Wicher, B.; Akrivos, P. D.; Tkachenko, N. V. Eur. J. Inorg. Chem. 2013, 5196-5205. (j) Fedushkin, I. L.; Maslova, O. V.; Morozov, A. G.; Dechert, S.; Demeshko, S.; Meyer, F. Angew. Chem., Int. Ed. 2012, 51, 10584-10587. (k) Li, L.; Lopes, P. S.; Rosa, V.; Figueira, C. A.; Lemos, M. A. N. D. A.; Duarte, M. T.; Avilés, T.; Gomes, P. T. Dalton Trans. 2012, 41, 5144-5154. (1) Clark, K. M.; Bendix, J.; Heyduk, A. F.; Ziller, J. W. Inorg. Chem. 2012, 51, 7457-7459. (m) Hill, N. J.; Vargas-Baca, I.; Cowley, A. H. Dalton Trans. 2009, 240-253. (n) Fedushkin, I. L.; Skatova, A. A.; Chudakova, V. A.; Fukin, G. K. Angew. Chem., Int. Ed. 2003, 42, 3294-3298. (o) van Asselt, R.; Elsevier, C. J.; Smeets, W. J. J.; Spek, A. L. Inorg. Chem. 1994, 33, 1521-1531.

(3) (a) Mondal, P.; Agarwala, H.; Jana, R. D.; Plebst, S.; Grupp, A.; Ehret, F.; Mobin, S. M.; Kaim, W.; Lahiri, G. K. *Inorg. Chem.* **2014**, *S3*, 7389–7403. (b) Ragaini, F.; Cenini, S.; Borsani, E.; Dompé, M.; Gallo, E. *Organometallics* **2001**, *20*, 3390–3398. (c) Singh, S. K.; Dubey, S. K.; Pandey, R.; Mishra, L.; Zou, R.-Q; Xu, Q.; Pandey, D. K. *Polyhedron* **2008**, *27*, 2877–2882. (d) Viganó, M.; Ragaini, F.; Buonomenna, M. G.; Lariccia, R.; Caselli, A.; Gallo, E.; Cenini, S.; Jansen, J. C.; Drioli, E. *ChemCatChem* **2010**, *2*, 1150–1164.

(4) (a) Crawford, K.; Rautenstrauch, V.; Uijttewaal, A. Synlett 2001, 1127–1128. (b) Wahren, U.; Sprung, I.; Schulze, K.; Findeisen, M.; Buchbauer, G. Tetrahedron Lett. 1999, 40, 5991–5992. (c) Kelly, D. R.; Nally, J. Tetrahedron Lett. 1999, 40, 3251–3254.

(5) (a) Grigoropoulou, G.; Clark, J. H.; Elings, J. A. Green Chem.
2003, 5, 1-7. (b) Lane, B. S.; Burgess, K. Chem. Rev. 2003, 103, 2457-2473. (c) Beller, M. Adv. Synth. Catal. 2004, 346, 107-108. (d) Legros, J.; Bolm, C. Chem.—Eur. J. 2005, 11, 1086-1092. (e) Bregeault, J. M. Dalton Trans. 2003, 3289-3302. (f) Herrmann, W. A.; Fischer, R. W.; Marz, D. W. Angew. Chem., Int. Ed. 1991, 30, 1638-1641. (g) Rudolf, J.; Reddy, K. L.; Chiang, J. P.; Sharpless, K. B. J. Am. Chem. Soc. 1997, 119, 6189-6190. (h) Hermann, W. A.; Kratzer, R. M.; Ding, H.; Thiel, W. R.; Glas, H. J. Organomet. Chem. 1998, 555, 293-295. (i) Shi, Y. Acc. Chem. Res. 2004, 37, 488-496. (j) Gerlach, A.; Geller, T. Adv. Synth. Catal. 2004, 346, 1247-1249. (k) Yang, D. Acc. Chem. Res. 2004, 37, 497-505.

(6) Nishiyama, H.; Shimada, T.; Itoh, H.; Sugiyama, H.; Motoyama, Y. Chem. Commun. **1997**, 1863–1864.

(7) (a) Dakkach, M.; López, M. I.; Romero, I.; Rodríguez, M.; Atlamsani, A.; Parella, T.; Fontrodona, X.; Llobet, A. Inorg. Chem. **2010**, 49, 7072–7079. (b) Chatterjee, D. Inorg. Chim. Acta **2008**, 361, 2177–2182. (c) Chatterjee, D.; Sengupta, A.; Mitra, A. Polyhedron **2007**, 26, 178–183. (d) Benet-Buchholz, J.; Comba, P.; Llobet, A.; Roeser, S.; Vadivelu, P.; Wadepohl, H.; Wiesner, S. Dalton Trans. **2009**, 5910–5923. (e) Tse, M. K.; Döbler, C.; Bhor, S.; Klawonn, M.; Mägerlein, W.; Hugl, H.; Beller, M. Angew. Chem., Int. Ed. **2004**, 43, 5255–5260. (f) Tse, M. K.; Bhor, S.; Klawonn, M.; Anilkumar, G.; Jiao, H.; Spannenberg, A.; Döbler, C.; Mägerlein, W.; Hugl, H.; Beller, M. Chem.—Eur. J. **2006**, 12, 1875–1888. (g) Tse, M. K.; Klawonn, M.; Bhor, S.; Döbler, C.; Anilkumar, G.; Hugl, H.; Mägerlein, W.; Beller, M. Org. Lett. **2005**, 7, 987–990. (h) Schröder, K.; Enthaler, S.; Join, B.; Junge, K.; Beller, M. Adv. Synth. Catal. **2010**, 352, 1771–1778.

(8) (a) Mariappan, K.; Alaparthi, M.; Caple, G.; Balasubramanian, V.; Hoffman, M. M.; Hudspeth, M.; Sykes, A. G. *Inorg. Chem.* **2014**, *53*, 2953–2962. (b) Marcazzan, P.; Abu-Gnim, C.; Seneviratne, K. N.; James, B. R. *Inorg. Chem.* **2004**, *43*, 4820–4824. (c) Chanda, N.; Mondal, B.; Puranik, V. G.; Lahiri, G. K. *Polyhedron* **2002**, *21*, 2033– 2043. (9) Visentin, L. C.; Ferreira, L. C.; Bordinhão, J.; Filgueiras, C. A. L. J. Braz. Chem. Soc. 2010, 21, 1187–1194.

(10) Anga, S.; Paul, M.; Naktode, K.; Kottalanka, R. K.; Panda, T. K. Z. Anorg. Allg. Chem. **2012**, 638, 1311–1315.

(11) Anga, S.; Rej, S.; Naktode, K.; Pal, T.; Panda, T. K. J. Chem. Sci. 2015, 127, 103–113.

(12) (a) Liu, Y.; Ng, S.-M.; Yiu, S.-M.; Lam, W. W. Y.; Wei, X.-G.; Lau, K.-C.; Lau, T.-C. Angew. Chem., Int. Ed. 2014, 53, 14468–14471.
(b) Rilak, A.; Bratsos, I.; Zangrando, E.; Kljun, J.; Turel, I.; Bugarčić, Ž. D.; Alessio, E. Inorg. Chem. 2014, 53, 6113–6126.

(13) (a) Agarwala, H.; Ehret, F.; Chowdhury, A. D.; Maji, S.; Mobin, S. M.; Kaim, W.; Lahiri, G. K. Dalton Trans. 2013, 42, 3721–3734.
(b) Serrano, I.; López, M. I.; Ferrer, Í.; Poater, A.; Parella, T.; Fontrodona, X.; Solà, M.; Llobet, A.; Rodríguez, M.; Romero, I. Inorg. Chem. 2011, 50, 6044–6054. (c) Yoshida, M.; Masaoka, S.; Abe, J.; Sakai, K. Chem.—Asian J. 2010, 5, 2369–2378.

(14) Farrugia, L. J. J. Appl. Crystallogr. 2012, 45, 849-854.

(15) (a) Du, W.; Wang, Q.; Wang, L.; Yu, Z. Organometallics 2014, 33, 974–982. (b) Das, A.; Scherer, T. M.; Chowdhury, A. D.; Mobin, S. M.; Kaim, W.; Lahiri, G. K. Inorg. Chem. 2012, 51, 1675–1684.

(16) (a) Sala, X.; Romero, I.; Rodríguez, M.; Escriche, L.; Llobet, A. *Angew. Chem., Int. Ed.* **2009**, *48*, 2842–2852. (b) Tseng, H.-W.; Zong, R.; Muckerman, J. T.; Thummel, R. *Inorg. Chem.* **2008**, *47*, 11763–11773.

(17) Bag, N.; Lahiri, G. K.; Bhattacharya, S.; Falvello, L. R.; Chakravorty, A. Inorg. Chem. 1988, 27, 4396–4402.

(18) Takeuchi, K. J.; Thompson, M. S.; Pipes, D. W.; Meyer, T. J. Inorg. Chem. 1988, 23, 1845–1851.

(19) Jones, W. E., Jr.; Bignozzi, C. A.; Chen, P.; Meyer, T. J. Inorg. Chem. 1993, 32, 1167–1178.

(20) Serrano, I.; Sala, X.; Plantalech, E.; Rodríguez, M.; Romero, I.; Jansat, S.; Gómez, M.; Parella, T.; Stoeckli-Evans, H.; Solans, X.; Font-Bardia, M.; Vidjayacoumar, B.; Llobet, A. *Inorg. Chem.* **2007**, *46*, 5381–5389.

(21) De, P.; Mondal, T. K.; Mobin, S. M.; Lahiri, G. K. Inorg. Chim. Acta 2011, 372, 250–258.

(22) (a) Patra, S.; Sarkar, B.; Mobin, S. M.; Kaim, W.; Lahiri, G. K. Inorg. Chem. 2003, 42, 6469–6473. (b) Kundu, T.; Mobin, S. M.; Lahiri, G. K. Dalton Trans. 2010, 39, 4232–4242. (c) Mandal, A.; Kundu, T.; Ehret, F.; Bubrin, M.; Mobin, S. M.; Kaim, W.; Lahiri, G. K. Dalton Trans. 2014, 43, 2473–2487. (d) Ghosh, P.; Mondal, P.; Ray, R.; Das, A.; Bag, S.; Mobin, S. M.; Lahiri, G. K. Inorg. Chem. 2014, 53, 6094–6106.

(23) Patra, S.; Sarkar, B.; Ghumaan, S.; Patil, M. P.; Mobin, S. M.; Sunoj, R. B.; Kaim, W.; Lahiri, G. K. Dalton Trans. 2005, 1188-1194. (24) (a) Chowdhury, A. D.; Das, A.; Irshad, K.; Mobin, S. M.; Lahiri, G. K. Inorg. Chem. 2011, 50, 1775-1785. (b) Chanda, N.; Mobin, S. M.; Puranik, V. G.; Niemeyer, M.; Datta, A.; Lahiri, G. K. Inorg. Chem. 2004, 43, 1056-1064. (c) Maji, S.; López, I.; Bozoglian, F.; Benet-Buchholz, J.; Llobet, A. Inorg. Chem. 2013, 52, 3591-3593. (d) Badiei, Y. M.; Polyansky, D. E.; Muckerman, J. T.; Szalda, D. J.; Haberdar, R.; Zong, R.; Thummel, R. P.; Fujita, E. Inorg. Chem. 2013, 52, 8845-8850. (e) Neudeck, S.; Maji, S.; López, I.; Meyer, S.; Meyer, F.; Llobet, A. J. Am. Chem. Soc. 2014, 136, 24-27. (f) Vaguer, L.; Miró, P.; Sala, X.; Bozoglian, F.; Masllorens, E.; Benet-Buchholz, J.; Fontrodona, X.; Parella, T.; Romero, I.; Roglans, A.; Rodríguez, M.; Bo, C.; Llobet, A. ChemPlusChem 2013, 78, 235-243. (g) Ohzu, S.; Ishizuka, T.; Hirai, Y.; Jiang, H.; Sakaguchi, M.; Ogura, T.; Fukuzumi, S.; Kojima, T. Chem. Sci. 2012, 3, 3421-3431.

(25) Juris, A.; Balzani, V.; Barigelletti, F.; Campagna, S.; Belser, P.; von Zelewsky, A. Coord. Chem. Rev. 1988, 84, 85–277.

(26) (a) Pappo, R.; Allen, D. S., Jr.; Lemieux, R. U.; Johnson, W. S. J. Org. Chem. 1956, 21, 478–479. (b) Carlsen, P. H. J.; Katsuki, T.; Martin, V. S.; Sharpless, K. B. J. Org. Chem. 1981, 46, 3936–3938.
(c) Das, A.; Ghosh, T. K.; Chowdhury, A. D.; Mobin, S. M.; Lahiri, G. K. Polyhedron 2013, 52, 1130–1137.

(27) (a) Paull, D. E.; Keagy, B. A.; Kron, E. J.; Wilcox, B. R. J. Surg. Res. **1989**, 46, 333–338. (b) Bunda, S.; Kaviani, N.; Hinek, A. J. Biol. Chem. **2005**, 280, 2341–2351.

(28) (a) Acquaye, J. H.; Muller, J. G.; Takeuchi, K. J. Inorg. Chem. 1993, 32, 160–165. (b) Lai, S.; Lepage, C. J.; Lee, D. G. Inorg. Chem. 2002, 41, 1954–1957. (c) Hamelin, O.; Ménage, S.; Charnay, F.; Chavarot, M.; Pierre, J.-L.; Pécaut, J.; Fontecave, M. Inorg. Chem. 2008, 47, 6413–6420. (d) Chavarot, M.; Ménage, S.; Hamelin, O.; Charnay, F.; Pécaut, J.; Fontecave, M. Inorg. Chem. 2003, 42, 4810–4816.

(29) (a) Groves, J. T.; Kruper, W. J. J. Am. Chem. Soc. 1979, 101, 7613–7615. (b) Groves, J. T.; Nemo, T. E. J. Am. Chem. Soc. 1983, 105, 5786–5791. (c) Jørgensen, K. A. Chem. Rev. 1989, 89, 431–458. (d) Meunier, B. Chem. Rev. 1992, 92, 1411–1456. (e) Jørgensen, K. A.; Schiøtt, B. Chem. Rev. 1990, 90, 1483–1506. (f) Ostovic, D.; Bruice, T. C. Acc. Chem. Res. 1992, 25, 314–320. (g) Fung, W.-H.; Yu, W.-Y.; Che, C.-M. J. Org. Chem. 1998, 63, 7715–7726.

(30) Reger, D. L.; Grattan, T. C.; Brown, K. J.; Little, C. A.; Lamba, J. J. S.; Rheingold, A. L.; Sommer, R. D. *J. Organomet. Chem.* **2000**, *607*, 120–128.

(31) (a) Mak, C. S. K.; Wong, H. L.; Leung, Q. Y.; Tam, W. Y.; Chan, W. K.; Djurišic, A. B. J. Organomet. Chem. 2009, 694, 2770.
(b) Gasperini, M.; Ragaini, F.; Cenini, S. Organometallics 2002, 21, 2950–2957.

(32) Llobet, A.; Doppelt, P.; Meyer, T. J. Inorg. Chem. 1988, 27, 514-520.

(33) (a) Sheldrick, G. M. Acta Crystallogr., Sect. A 2008, A64, 112– 122. (b) Program for Crystal Structure Solution and Refinement; University of Göttingen: Göttingen, Germany, 1997.

(34) Shuis, P. V. D.; Spek, A. L. Acta Crystallogr., Sect. A 1990, 46, 194-201.

(35) Lee, C.; Yang, W.; Parr, R. G. Phys. Rev. B 1998, 37, 785-789. (36) Frisch, M. J.; Trucks, G. W.; Schlegel, H. B.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Scalmani, G.; Barone, V.; Mennucci, B.; Petersson, G. A.; Nakatsuji, H.; Caricato, M.; Li, X.; Hratchian, H. P.; Izmaylov, A. F.; Bloino, J.; Zheng, G.; Sonnenberg, J. L.; Hada, M.; Ehara, M.; Toyota, K.; Fukuda, R.; Hasegawa, J.; Ishida, M.; Nakajima, T.; Honda, Y.; Kitao, O.; Nakai, H.; Vreven, T.; Montgomery, J. A.; Peralta, Jr. J. E.; Ogliaro, F.; Bearpark, M.; Heyd, J. J.; Brothers, E.; Kudin, K. N.; Staroverov, V. N.; Kobayashi, R.; Normand, J.; Raghavachari, K.; Rendell, A.; Burant, J. C.; Iyengar, S. S.; Tomasi, J.; Cossi, M.; Rega, N.; Millam, J. M.; Klene, M.; Knox, J. E.; Cross, J. B.; Bakken, V.; Adamo, C.; Jaramillo, J.; Gomperts, R.; Stratmann, R. E.; Yazyev, O.; Austin, A. J.; Cammi, R.; Pomelli, C.; Ochterski, J. W.; Martin, R. L.; Morokuma, K.; Zakrzewski, V. G.; Voth, G. A.; Salvador, P.; Dannenberg, J. J.; Dapprich, S.; Daniels, A. D.; Farkas, O.; Foresman, J. B.; Ortiz, J. V.; Cioslowski, J.; Fox, D. J. Gaussian 09, Revision A.02: Gaussian, Inc.: Wallingford CT, 2009.

(37) (a) Andrae, D.; Haeussermann, U.; Dolg, M.; Stoll, H.; Preuss, H. *Theor. Chim. Acta* **1990**, 77, 123–141. (b) Fuentealba, P.; Preuss, H.; Stoll, H.; Szentpaly, L. V. *Chem. Phys. Lett.* **1989**, 89, 418–422.

(38) (a) Bauernschmitt, R.; Ahlrichs, R. Chem. Phys. Lett. **1996**, 256, 454–464. (b) Stratmann, R. E.; Scuseria, G. E.; Frisch, M. J. J. Chem. Phys. **1998**, 109, 8218–8225. (c) Casida, M. E.; Jamorski, C.; Casida, K. C.; Salahub, D. R. J. Chem. Phys. **1998**, 108, 4439–4450.

(39) (a) Barone, V.; Cossi, M. J. Phys. Chem. A 1998, 102, 1995–2001. (b) Cossi, M.; Barone, V. J. Chem. Phys. 2001, 115, 4708–4718.
(c) Cossi, M.; Rega, N.; Scalmani, G.; Barone, V. J. Comput. Chem. 2003, 24, 669–681.

(40) O'Boyle, N. M.; Tenderholt, A. L.; Langner, K. M. J. Comput. Chem. 2008, 29, 839–845.

(41) Zhurko, D. A.; Zhurko, G. A. *ChemCraft 1.5*; Plimus: San Diego, CA. Available at http://www.chemcraftprog.com.