Communication

On the Possible Coordination on a $^3$MC State Itself? Mechanistic Investigation Using DFT-Based Methods

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Abstract: Understanding light-induced ligand exchange processes is key to the design of efficient light-releasing prodrugs or photochemically driven functional molecules. Previous mechanistic investigations had highlighted the pivotal role of metal-centered (MC) excited states in the initial ligand loss step. The question remains whether they are equally important in the subsequent ligand capture step. This article reports the mechanistic study of direct acetonitrile coordination onto a $^3$MC state of [Ru(bpy)$_2$]$_2^+$, leading to [Ru(bpy)$_2$(2,2'-bpy)(NCMe)]$^2^+$ in a $^3$MLCT (metal-to-ligand charge transfer) state. Coordination of MeCN is indeed accompanied by the decoordination of one pyridine ring of a bpy ligand. As estimated from Nudged Elastic Band calculations, the energy barrier along the minimum energy path is 20 kcal/mol. Interestingly, the orbital analysis conducted along the reaction path has shown that creation of the metallic vacancy can be achieved by reverting the energetic ordering of key $\sigma^*$ and bpy-based $\pi^*$ orbitals, resulting in the change of electronic configuration from $^3$MC to $^3$MLCT. The approach of the NCMe lone pair contributes to destabilizing the $\sigma^*$ orbital by electrostatic repulsion.

Keywords: ruthenium polypyridine complex; photochemistry; photosolvolysis mechanism; metal-centered excited states; triplet state reactivity; nudged elastic band; DFT; molecular orbitals

1. Introduction

The photophysics of ruthenium polypyridine compounds is governed by the subtle balance between the population of two types of triplet excited states of similar energies: metal-to-ligand charge transfer states (MLCT) and metal-centered states (MC) [1]. $^3$MLCT states are photoluminescent, contrary to $^3$MC states, which quench the luminescence and may lead to ligand loss [2]. Forty years of spectroscopic studies on this family of compounds have provided a wealth of robust experimental data, but MC states are spectroscopically “dark” and therefore their theoretical characterization is essential in the rationalization and the anticipation of photophysical properties. For more than a decade, we have been able to optimize $^3$MC excited states [3] and to explore the topology of the lowest triplet potential energy surface using DFT-based methods [4–9]. Spectacular bond elongations and angular distortions have been characterized in numerous Ru(II) $^3$MC states [10–28] and for other metals, e.g., Ir(III) [29–31]. Our contribution to the field covers photoisomerization mechanisms [32–34] and photoluminescence quenching mechanisms [35,36], as well as exploratory ruthenium(II) [37,38] and iron(II) [39–42] photophysics, in a constant dialogue with experimental chemists.

In the course of our theoretical investigations of photosubstitution mechanisms, we have identified some key $^3$MC states along the pathways for ligand photorelease, particularly
pentacoordinate or pseudopentacoordinate species [7,8]. In that work, these \( ^3 \)MC states have been considered to be involved in such mechanisms via intersystem crossing (ISC) through a neighboring minimum energy crossing point (MECP), allowing the system to populate an electrophilic, coordinatively unsaturated, and closed-shell species. The final coordination of a molecule of incoming ligand is thought to be an efficient process [43–47] and yields a \( \kappa^2 \)-bound intermediate product that requires the absorption of a second photon to fully release the departing bidentate ligand [48,49]. However, an alternative pathway can also be envisaged, overall requiring only one photon, namely the direct reaction between the incoming ligand and the complex in its distorted \( ^3 \)MC state, to form a new complex with triplet spin multiplicity according to Wigner rules. The aim of this work was therefore to investigate this type of reactivity through DFT-based methods, computing minimum energy paths using Nudged Elastic Band calculations and undertaking a thorough orbital analysis along this path. The model reaction we chose is the approach of an acetonitrile molecule on the previously reported \( ^3 \)MC\(_{\text{cis}} \) state of Ru(bpy)\(_2^2^+ \) [50]. This state, repelling the two pyridine fragments of a single bpy ligand, is thought to be prone to bpy loss, or at least more prone than the classical \( ^3 \)MC\(_{\text{trans}} \) state that repels two pyridines from two different ligands (Figure 1).

![Figure 1. Structures of \( ^3 \)MC\(_{\text{trans}} \) and \( ^3 \)MC\(_{\text{cis}} \) in Ru(bpy)\(_2^2^+ \) (H atoms not shown).](image)

Acetonitrile was selected for its charge neutrality, for being a reactant that bears only one lone pair, and for being an aprotic and weakly self-associating solvent, that allows us to consider the approach of a single NCMe molecule as a reasonable model. The aim of this work was to envisage direct NCMe coordination on the \( ^3 \)MC\(_{\text{cis}} \) state of Ru(bpy)\(_2^2^+ \) in order to form an intermediate triplet state bearing one monodentate bpy and one bound acetonitrile ligand (Scheme 1). This process was found to have an energy barrier of 20 kcal/mol. Analysis of the coordination process from an orbital perspective has shown that MeCN approach and change in electronic configuration from \( ^3 \)MC to \( ^3 \)MLCT were intimately related.

![Scheme 1. Coordination of one acetonitrile molecule onto the \( ^3 \)MC\(_{\text{cis}} \) state of Ru(bpy)\(_2^2^+ \).](image)

2. Results and Discussion

Working on the photoinduced loss of bidentate ligands, Elliott et al. reported the crystallographic characterization of intermediate photoproducts of the type
trans-[Ru(bpy)(κ²-btz)(κ¹-btz)(NCMe)]⁺ bearing a singly-bound bitiazolyl (btz) ligand [48,49]. By analogy we have built and optimized trans-[Ru(bpy):κ¹-bpy)(NCMe)]²⁺ in a singlet state, from which an excited state with triplet spin multiplicity was subsequently optimized by unrestricted DFT. This state is unambiguously of MLCT electronic nature and was used as endpoint for a minimum energy path calculation. This state contains a bpy ligand that is a monodentate, κ¹-bpy ligand. The exploration of the envisaged chemical reaction was undertaken starting from the 3MC₆s state having an acetonitrile molecule 5 Å away with its lone pair oriented towards the metal, in the most open quadrant of the complex. Note that NCMe is not interacting with the [Ru(bpy)]⁺ moiety in this structure, as estimated from its total energy that is equal to the sum of the fragments’ energies.

At the starting point of the reaction path the complex is in the 3MC₆s state. The rationale for this choice lies in its presumed higher propensity for ligand loss than the 3MC₃trans state. This is due to its peculiar geometry repelling a single bpy ligand and opening a quadrant, allowing a potentially entering ligand to approach the metal [50]. In the 3MC₆s state, two major elongations are found towards the same bpy ligand (Ru–N5 and Ru–N6, Table 1), the opposite quadrant opens up to 121° (Figure 1), and the Mulliken spin population on ruthenium is 1.7. At the end point, the complex contains one monodentate bpy ligand (the pyridine ring containing N5 is rotated out of plane and Ru–N5 = 3.59 Å); the Ru–N6 distance is standard and the two bonds towards the formally anionic bpy ligand are the shortest (Ru–N3 and Ru–N4). The 3MLCT nature of this state is illustrated by its Mulliken spin population on ruthenium, which is 0.9 (the other spin residing on a bpy ligand).

Table 1. Atom numbering and Ru–N distances (Å) at the start (3MC + MeCN) and end (3MLCT) points of the computed reaction path.

| Ru–N     | 3MC + MeCN | 3MLCT |
|----------|------------|-------|
| Ru–N1    | 2.135      | 2.118 |
| Ru–N2    | 2.080      | 2.110 |
| Ru–N3    | 2.078      | 2.049 |
| Ru–N4    | 2.177      | 2.057 |
| Ru–N5    | 2.530      | 3.593 |
| Ru–N6    | 2.384      | 2.132 |
| Ru–NCMe  | 5.000      | 2.055 |

Minimum energy paths (MEPs) can be efficiently computed using the Nudged Elastic Band method [51,52], which discretizes the reaction path into a series of points, called beads. The initial path consists in a series of single point energy calculations along a geometry interpolation, which we perform using the image dependent pair potential (IDPP) method [53]. Subsequently each intermediate bead is minimized using path gradient and tangent information, until convergence to the MEP. The energy gap between the two endpoints is 17 kcal/mol, and a first estimate of the energy barrier for the reaction is 18 kcal/mol, as shown on Figure 2. The energy profile for this reaction appears as having a very moderate slope up to bead 15 (Ru–NCMe distance of 2.77 Å), after which the energy suddenly rises by about 14 kcal/mol at bead 18.
Figure 2. Minimum energy path (blue) and Mulliken spin population on Ru (red) for the coordination of acetonitrile on the \(^3\)MC state from Nudged Elastic Band calculations.

A sudden drop in the Mulliken spin population on ruthenium is apparent between bead 17 and bead 18, signifying a change between MC and MLCT character. Thus, a second nudged elastic band (NEB) calculation was performed to refine this region. For simplicity the beads in this second calculation have been numbered with intermediate values 17.25, 17.50, and 17.75 (Figure 3).

Figure 3. Minimum energy path (blue) and Mulliken spin population on Ru (red) between beads 17 and 18 of the reaction path.

The switch between MC and MLCT states here occurs between beads 17.50 and 17.75, providing a refined energy barrier of 20 kcal/mol. Wiberg bond indices (WBIs) have been computed at all geometries along the reaction path, and have been used to quantify the Ru–N interactions. In the starting \(^3\)MC structure, two Ru–N bonds are significantly elongated as a result of the population of an antibonding \(\sigma^*\) orbital, namely Ru–N5 (2.53 Å) and Ru–N6 (2.38 Å). The corresponding WBIs are consequently the lowest, 0.19 and 0.26 respectively (Figure 4). Bonds \textit{trans} to these, Ru–N4 and Ru–N1, are slightly elongated with WBIs of 0.36 and 0.41. The two remaining bonds, Ru–N2 and Ru–N3, display WBIs that are similar to the ones found in the ground state, i.e., 0.44. Notably the gradual drop in the Ru–N5 WBI along the reaction path is paralleled by a gradual increase in the Ru–NCMe WBI, until bead 17.50 where this latter WBI doubles to reach 0.45 in the \(^3\)MLCT state. At the endpoint, the highest two WBIs are the ones to the nitrogen atoms of the formally anionic bpy ligand, N3 (0.51) and N4 (0.54), Ru–N3, and Ru–N4 being the shortest bonds (Table 1). The intermediate zone (between beads 7–14), where WBIs towards departing and incoming ligands are
both low, illustrates the fact that the complex gets pentacoordinated and is stabilized by two successive weak Ru–N interactions along this path.

In order to allow the Ru–NCMe bond to form, a low-lying vacancy should be identified on the metal with appropriate symmetry and orientation to enable significant overlap with the incoming nitrogen lone pair. The crucial orbitals in this process are thus: (i) the metallic $d\sigma^*$ orbitals, (ii) the NCMe lone pair that eventually evolves to a Ru–NCMe dative bond, and (iii) the bpy-based $\pi^*$ orbital that will be singly occupied in the final $^3$MLCT state. The eigenvalues of these four molecular orbitals (MOs) are plotted against bead number on Figure 5. All lines are broken between beads 17.50 and 17.75 to signify the rupture in the correlation diagram, e.g., when the NCMe lone pair is replaced by a bonding Ru–NCMe interaction. At this very point along the path, we propose that the electrostatic repulsion between the NCMe lone pair and the singly occupied $d\sigma^1$ orbital (as illustrated by the gradual destabilization of $d\sigma^1$ from bead 12 onwards) is such that the $d\sigma^1$ is destabilized to a point where its energy is higher than that of the bpy $\pi^*$ orbital. As a result, the electron that was in $d\sigma^1$ in the MC state is transferred to the bpy $\pi^*$ orbital, thus producing an MLCT state. This change in electronic configuration is what allows the total energy of the system to decrease towards the endpoint of the path. In addition, the quasi invariance in the energy of $d\sigma^2$ confirms that it is always nonbonding towards the incoming ligand.

![Figure 4](image1.png)

**Figure 4.** Evolution of WBIs along the reaction path. The MC–MLCT transition region is highlighted in yellow.

![Figure 5](image2.png)

**Figure 5.** Selected orbital eigenvalues along the reaction path. The MC–MLCT transition region is highlighted in yellow.
A close inspection of these crucial orbitals, at beads 17.50 and 17.75 (Figure 6), reveals that the formation of the Ru–NCMe bond is accompanied by a repolarization of both do* orbitals, as seen for instance in the disappearance of the Ru–N6 antibonding interaction in do*2'. The interaction of do*1 with the NCMe lone pair produces two new orbitals: the Ru–NCMe bond and the do*1' antibond (Figure S1). In this system, coordination of the incoming ligand is accompanied by the change of electronic state nature, from 3MC to 3MLCT, which has a vacant do*1' (as stated in the introduction, an alternative to create a metallic vacancy is spin crossing to a closed-shell pentacoordinate singlet state [8,54], which would also vacate the required do* orbital).

Figure 6. Partial MO correlation diagram between beads 17.50 and 17.75 (Kohn–Sham orbitals), showing NCMe coordination at bead 17.75 (a more complete MO diagram is given as Figure S2).

3. Computational Details and Methods

Geometry optimizations were performed without symmetry with Orca [55] using the B3LYP functional [56,57], a relativistic small core pseudopotential on Ru (SD28) [58], the def2-TZVP(-f) basis set [59], and the empirical D3 dispersion correction [60,61]. Solvent effects were modelled as the SMD polarizable continuum [62]. The restricted Kohn–Sham formalism was used for ground states, while its unrestricted analogue was used for triplet states. SCF convergence was achieved using the DIIS algorithm followed by a semi-quadratic SOSCF converger. Frequency calculations were run at the same level of theory and the absence of imaginary frequencies ascertained the nature of these points as minima. Molecular orbitals were viewed using Gabedit [63]. Mulliken spin densities on Ru were used as a straightforward descriptor of the electronic nature of the triplet excited state (~0.9 for a 3MLCT state, ~1.8 for a 3MC state). Orbital analysis was systematically undertaken to view the localization of the unpaired electrons.

The 3MLCT–3MC minimum energy paths were optimized with the nudged elastic band (NEB) method [51,52] using a python module developed in the Clancy group [64] that is interfaced with
Orca. The convergence criterion was set to 0.03 eV/Å. A 20-frame initial path was prepared by interpolating start and end geometries using the IDPP method [53]. The geometries were previously processed using lab-developed programs to minimize the discrepancy between start and end geometries. These calculations were performed at the same level of theory as all the geometry optimizations. Convergence of the MEP was achieved using a combination of FIRE and BFGS algorithms.

4. Conclusions

In this work, we have envisaged the direct addition of an acetonitrile molecule on the \(^3\)MC\(_{\text{ox}}\) state of [Ru(bpy)\(_3\)]\(^2+\), a state that we had proposed to consider as potentially photoreactive [50]. Nudged elastic band calculations have provided an energy barrier of 20 kcal/mol for this model reaction, significantly higher than the energy barrier involved in the spin crossing process towards a pentacoordinate ground state species [54] (note that the former pathway requires a single photon excitation, whereas the latter requires a second photon to fully release the departing bidentate ligand). The orbital analysis we have conducted along the reaction path has enabled us to describe the chemical reaction and the MC–MLCT transition. The change in electronic nature, from \(^3\)MC to \(^3\)MLCT, is concomitant to the approach of the NCMe lone pair and triggers the interaction between the newly formed metallic vacancy and the NCMe lone pair. In this view, the \(^3\)MC state itself can be seen as unable to bind MeCN. It is noteworthy that the approach of an incoming nucleophile is able to perturbate the metal complex so as to modify the electronic nature of its excited state. These results offer an interesting glimpse of processes that could be involved in excited state reaction mechanisms under specific experimental conditions such as pulsed irradiation. We are currently extending this work to the mechanistic study of the subsequent steps, i.e., \(\kappa^1\)-bpy ligand loss.

Supplementary Materials: The following are available online at www.mdpi.com/xxx/s1, Figure S1: MO diagram schematizing the interaction between the metallic fragment in the \(^3\)MC state and the approaching acetonitrile ligand; Figure S2: MO diagram at selected points along the reaction path; Cartesian coordinates of the 20 beads along the minimum energy path.

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