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Direct Observation of Photoinduced Charge Separation in Ruthenium Complex/ $\text{Ni}(\text{OH})_2$ Nanoparticle Hybrid

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$\text{Ni}(\text{OH})_2$ have emerged as important functional materials for solar fuel conversion because of their potential as cost-effective bifunctional catalysts for both hydrogen and oxygen evolution reactions. However, their roles as photocatalysts in the photoinduced charge separation (CS) reactions remain unexplored. In this paper, we investigate the CS dynamics of a newly designed hybrid catalyst by integrating a Ru complex with $\text{Ni}(\text{OH})_2$ nanoparticles (NPs). Using time resolved X-ray absorption spectroscopy (XTA), we directly observed the formation of the reduced Ni metal site (~60 ps), unambiguously demonstrating CS process in the hybrid through ultrafast electron transfer from Ru complex to $\text{Ni}(\text{OH})_2$ NPs. Compared to the ultrafast CS process, the charge recombination in the hybrid is ultraslow ($\gg 50$ ns). These results not only suggest the possibility of developing $\text{Ni}(\text{OH})_2$ as solar fuel catalysts, but also represent the first time direct observation of efficient CS in a hybrid catalyst using XTA.

Generation of H_2 from water by solar energy conversion is an attractive strategy to partially address the energy crisis and climate issues¹. The overall process for water splitting includes two half-catalytic reactions, i. e. hydrogen (HER) and oxygen evolution reactions (OER)^{2–4}. Each of the reactions requires a photocatalytic system that can efficiently integrate multiple components: a photosensitizer (PS) for light harvesting, a catalyst that can perform the appropriate half-reaction, and finally a reaction medium to allow charge flow from the excited PS to the catalyst and electron/hole acceptors^{2,3}. Since the initial report of light-driven splitting of water⁵, extensive efforts have been devoted to develop such photocatalytic systems. One class that has been intensively studied uses the platinum group metals or metal oxides^{1,6} as catalysts. While these durable catalytic systems can efficiently catalyze HER or OER, their long term practicality is hampered by the scarcity and high cost of noble metals. As a result, various low-cost electrocatalysts such as MoS_2 ^{7–9} and NiMo catalysts^{10,11} for HER and CoPi^{12,13} and metal oxides^{14,15} for OER, have evolved during the past decade. However, most of these earth abundant catalysts are only active or stable in either acidic or basic electrolytes. Given that the overall water splitting requires that both HER and OER catalysts exist in the same electrolyte, it is crucial to develop bifunctional catalysts that can efficiently catalyze HER and OER in the same electrolyte.

$\text{Ni}(\text{OH})_2$ nanocrystals have emerged as solar fuel catalysts due to their low-cost and bifunctional nature^{16–23}. OER can be efficiently driven using $\text{Ni}(\text{OH})_2$ nanocrystals as catalysts in alkaline electrolyte^{16,17}. $\text{Ni}(\text{OH})_2$ modified metal surfaces are also efficient HER catalysts with higher activities than bare metal catalysts^{19,20}. Recently, water photolysis with 12.3% efficiency was achieved using an Fe incorporated $\text{Ni}(\text{OH})_2$ foam electrode as a catalyst for both HER and OER²¹, evidently demonstrating the potential of $\text{Ni}(\text{OH})_2$ as a bifunctional photocatalyst for water splitting. Despite these studies, a fundamental understanding of the photoinduced charge separation (CS) and charge recombination (CR) dynamics in $\text{Ni}(\text{OH})_2$ based photocatalytic systems, the processes crucial for solar fuel generation, is still lacking.

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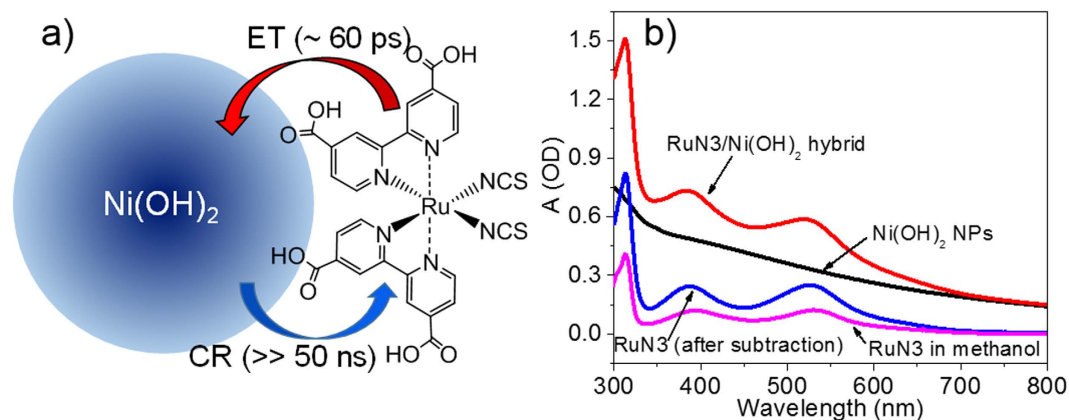


Figure 1. The design of hybrid catalysts. (a) Cartoon of charge separation and recombination dynamics in RuN3/Ni(OH)₂ hybrid. (b) UV-visible absorption spectra of Ni(OH)₂ NPs (black), RuN3/Ni(OH)₂ hybrid (red), RuN3 obtained after subtracting the spectrum of Ni(OH)₂ NPs from the spectrum of the hybrid (blue), and RuN3 in methanol (pink).

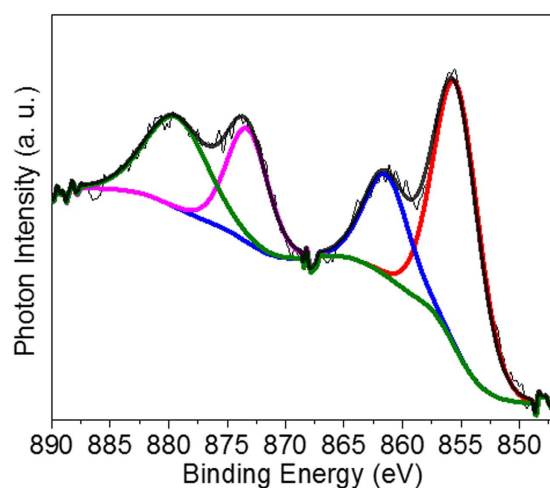


Figure 2. Chemical composition characterization. XPS spectrum of Ni(OH)₂ NPs for binding energies within the range of 840–890 eV.

In this work, we report a new hybrid photocatalytic system employing monodispersed Ni(OH)₂ colloidal nanoparticles (NPs) as catalysts and RuN3 (cis-diisothiocyanato-bis(2,2'-bipyridyl)-4,4'-dicarboxylic acid) ruthenium(II) as a PS (Fig. 1a). We unambiguously demonstrated electron injection from excited RuN3 to Ni(OH)₂ NPs using the combination X-ray (XTA) and optical transient absorption spectroscopy (OTA). RuN3 was chosen as a model PS because it is among the most efficient and robust molecular PS for light harvesting with well-known optical properties, and thus facilitated our photophysical studies. Moreover, RuN3 has –COOH groups which have extensively been used as linkers to bind dye molecules to the surface of metal oxides and can facilitate assembling on the surface of Ni(OH)₂ NPs^{24,25}. The details of the synthesis of Ni(OH)₂ NPs and RuN3/Ni(OH)₂ hybrid are described in supplementary information (SI). The average diameter of Ni(OH)₂ NPs measured by small-angle X-ray scattering (SAXS) is 4.42 ± 0.40 nm (Supplementary Fig. 1), in agreement with our TEM data (Supplementary Fig. 2).

Results and Discussion

Figure 1b shows the UV-visible absorption spectra of Ni(OH)₂ NPs and RuN3/Ni(OH)₂ hybrid in toluene solution. The spectrum of Ni(OH)₂ NPs shows a broad absorption in the whole UV-visible region, which can be attributed to the intra-3d transitions in Ni²⁺ ions^{26,27}. Compared to the spectrum of Ni(OH)₂ NPs, the spectrum of RuN3/Ni(OH)₂ hybrid shows two additional absorption bands at 385 and 535 nm, respectively. These two bands, consistent with those in RuN3 in methanol solution (pink line in Fig. 1b) can be assigned to the metal-to-ligand charge transfer (MLCT) bands of RuN3^{25,28}. Because RuN3 is not soluble in toluene, these additional absorption features in the spectrum of RuN3/Ni(OH)₂ hybrid in toluene solution are due to the directly adsorbed RuN3 on the surface of Ni(OH)₂ NPs.

The chemical composition of the synthesized Ni(OH)₂ NPs were analyzed by X-ray photoelectron spectroscopy (XPS) and steady state X-ray absorption spectroscopy (XAS). Figure 2 shows the high resolution XPS of Ni(OH)₂

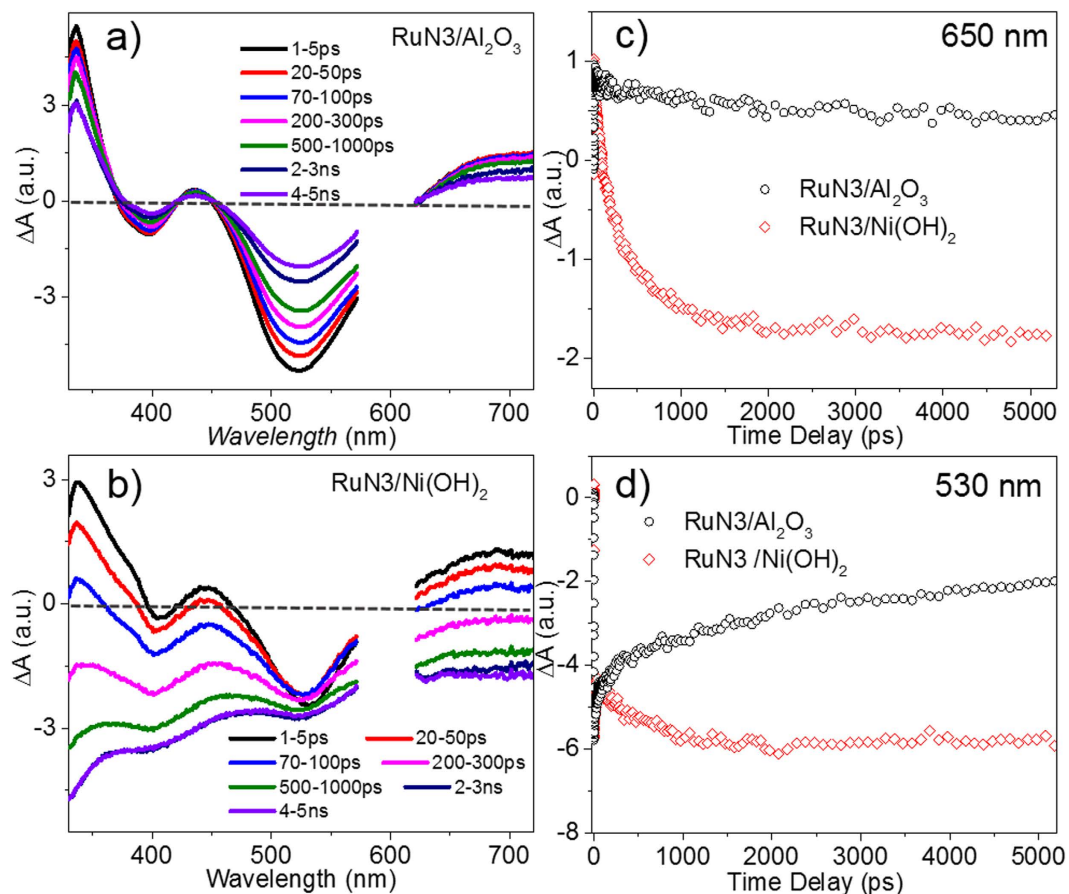


Figure 3. Transient absorption optical spectral features of the hybrid. Femtosecond OTA spectra of RuN3/Al₂O₃ (a) and RuN3/Ni(OH)₂ hybrid (b). Comparison of the femtosecond OTA kinetics of RuN3/Al₂O₃ and RuN3/Ni(OH)₂ hybrid at 650 nm (c) and 530 nm (d).

NPs at Ni 2p region (845–890 eV binding energy). The Ni 2p spectrum consists of two doublet of Ni 2p_{3/2} and Ni 2p_{1/2} transitions: Ni 2p_{3/2} main peak at 855.5 eV and its satellite at 861.5 eV, and Ni 2p_{1/2} main peak at 873.6 eV and its satellite at 879.7 eV. These results are consistent with the literature data of Ni(OH)₂ and thus correspond to the presence of the nickel hydroxide^{29,30}. After subtracting the Shirley-type background, these peaks can be adequately fitted by four curve-fitting bands with Gaussian (50%)-Lorentzian (50%) profile. The five curve-fitting model using either pure Gaussian or Gaussian-Lorentzian function introduced significant noise, together with the fact that the sizes of the NPs (4.42 ± 0.40 nm in diameter) are small enough for XPS to penetrate, suggesting that the NPs are composed of Ni(OH)₂.

The CS dynamics in the RuN3/Ni(OH)₂ hybrid in toluene were investigated by femtosecond OTA (fs-OTA). Figure 3a shows the fs-OTA spectra of RuN3/Al₂O₃ hybrid system, which represents a non-electron injecting model to reveal the intrinsic excited state (ES) dynamics of RuN3 on the surface of NPs. The spectra of RuN3/Al₂O₃ hybrid consist of two negative bands centered at 385 and 535 nm, and two broad positive absorption bands centered at 337 and 680 nm. The two negative bands are consistent with the ground state (GS) MLCT absorption bands and are thus assigned to the GS bleach of RuN3. The two absorption bands centered at 337 and 680 nm, analogous to literature data^{25,28}, can be assigned to the ligand localized $\pi^* \rightarrow \pi^*$ transition and ligand-to-metal charge transfer (LMCT) of RuN3 ES, respectively.

Compared to the fs-OTA spectra of RuN3/Al₂O₃, the spectra of RuN3/Ni(OH)₂ hybrid (Fig. 3b) are drastically different. Although the initial spectra (1–5 ps) of RuN3/Ni(OH)₂ hybrid resemble the spectra of RuN3/Al₂O₃, the broad ES bands of the former decay much faster within 5 ns time window, suggesting enhanced ES decay dynamics in RuN3/Ni(OH)₂ hybrid. Furthermore, the spectra of RuN3/Ni(OH)₂ hybrid evolve to a broad bleach band within the whole spectral region after the ES absorption signal decays substantially (> 100 ps). These features can be further seen by comparing the kinetic traces of RuN3/Ni(OH)₂ hybrid and RuN3/Al₂O₃ at 650 nm (Fig. 3c) and 530 nm (Fig. 3d) which correspond to the ES absorption and GS bleach of RuN3, respectively. Compared to the ES decay kinetics of RuN3/Al₂O₃ due to intrinsic recovery of GS from ES, the kinetic trace of RuN3/Ni(OH)₂ hybrid at 650 nm quickly decays to negative after 100 ps, suggesting that the ES decay of RuN3 in RuN3/Ni(OH)₂ hybrid is accompanied by the formation of negative signal. The formation of negative features were also observed at 530 nm, where the amplitude of the fs-OTA signal grows to more negative in RuN3/Ni(OH)₂ hybrid instead of GS bleach recovery in RuN3/Al₂O₃. We can exclude the contribution of GS bleach of RuN3 to the negative signal at > 650 nm where RuN3 has negligible GS absorption. We can also rule out the possibility that the negative signals are due to

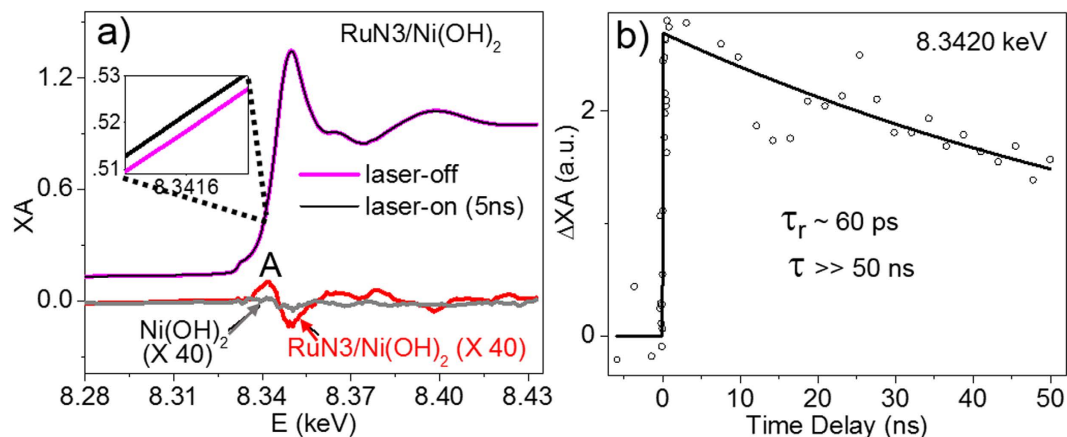


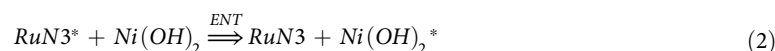
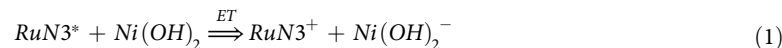
Figure 4. Transient X-ray absorption spectra at Ni K-edge. (a) XANES spectra of RuN3/Ni(OH)₂ hybrid at Ni K edge before (laser-off) and after (laser-on) laser excitation. Inset shows the spectra at 8.3415–8.3417 keV range. The difference spectra obtained from subtracting the laser-off XANES spectrum from laser-on spectrum (5 ns) for RuN3/Ni(OH)₂ hybrid (red solid line) and free Ni(OH)₂ NPs (gray solid line) are presented below the XANES spectra. (b) The time dependence of the transient signal at 8.3420 keV as a function of delay time.

the direct excitation of Ni(OH)₂ NPs, because negligible fs-OTA signals of Ni(OH)₂ NPs were observed under the same experimental conditions (Supplementary Fig. 3). Indeed, this negative signal resembles the GS absorption of RuN3/Ni(OH)₂ hybrid and thus can be assigned to the combination of GS bleach of RuN3 and Ni(OH)₂ NPs.

The formation of GS bleach of Ni(OH)₂ NPs, along with the faster RuN3 ES quenching in RuN3/Ni(OH)₂ hybrid, can arise from two possible pathways: electron transfer (ET) and hole transfer (HT). Although these processes result in different transient products (i.e. oxidized RuN3 and reduced Ni(OH)₂ NPs for ET and reduced RuN3 and oxidized Ni(OH)₂ NPs for HT), it is challenging to identify them using OTA because 1) metal centered states are often optically silent and 2) there are significant spectral overlaps between RuN3 and Ni(OH)₂ NPs. To overcome this problem, element specific XTA, which allows us to selectively probe the specific metal center, was used to track the electron density change at Ni center during the photoinduced process in RuN3/Ni(OH)₂ hybrid. Figure 4a shows XANES spectra of RuN3/Ni(OH)₂ hybrid before (pink line) and 5 ns after 527 nm excitation (black line). The transient signal due to laser excitation was clearly observed in the expanded XANES spectra at 8.3415–8.3417 keV (inset) and the difference XANES spectrum (red solid line) after subtracting the laser-off spectrum from the laser-on spectrum. The positive feature at 8.3420 keV in the difference spectrum indicates that the edge of Ni center shifts to lower energy due to photoexcitation of RuN3 in the RuN3/Ni(OH)₂ hybrid. This feature was not observed in the difference spectrum of Ni(OH)₂ NPs alone (gray solid line), ruling out the possibility that the observed Ni edge shift is due to direct excitation of Ni(OH)₂ NPs. These results unambiguously determine the ET process from excited RuN3 to Ni(OH)₂ NPs.

In order to determine the ET rate, the intensity of the positive feature at 8.3420 keV in the difference spectrum of RuN3/Ni(OH)₂ hybrid as a function of delay time was collected. As shown in Fig. 4b, the kinetics shows an instant rising followed by a slow decay. Fitting the kinetics data with an exponential decay function yields a rising component ($\tau_r \sim 60$ ps, within 80 ps X-ray pulse duration) and a decay component ($\tau \gg 50$ ns), corresponding to ET and CR times, respectively. The much slower CR compared to ET process suggests the potential application of Ni(OH)₂ NPs as solar fuel catalysts.

Although ET and CR processes in RuN3/Ni(OH)₂ hybrid were confirmed by XTA, the possibility of energy transfer (ENT) needs to be considered due to spectral overlap between RuN3 emission and Ni(OH)₂ absorption (Supplementary Fig. 4). In principle, ENT efficiency can be estimated by measuring the emission of Ni(OH)₂ NPs. Unfortunately, due to the extremely low emission quantum yield of Ni(OH)₂ NPs, we are unable to measure the ENT efficiency by measuring the emission of Ni(OH)₂ NPs. As shown in equation (1) and (2),



ENT and ET are two competitive pathways for quenching ES of RuN3. Different from the OTA spectral features of ET process where fast ES quenching is accompanied by long-lived GS, the expected OTA spectral features of ENT is characteristic of simultaneously occurring of fast ES decay and GS recovery. Given the known ultrafast ET rate determined by XTA (~ 60 ps), if ENT is a competitive pathway to ET, an identical ultrafast GS bleach recovery would occur in the spectral range of 500–550 nm in OTA spectra of the hybrid, which however was not observed (Fig. 3b,d). These results suggest that ENT is not the dominating process for RuN3 ES quenching and efficient ET occurs from excited RuN3 to Ni(OH)₂ NPs.

In summary, we have examined the photoinduced CS dynamics in a newly designed RuN3/Ni(OH)₂ hybrid photocatalytic system using the combination of OTA and XTA. Upon photoexcitation of RuN3, ultrafast electron injection from the excited RuN3 to Ni(OH)₂ NPs was observed, which is followed by ultraslow CR. These findings not only suggest the potential application of Ni(OH)₂ NPs as photocatalysts for solar fuel conversion, but also demonstrate the superior capability of XTA in directly capturing the transient charge separated state in hybrid solar fuel catalysts.

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Author Contributions

Y.T. prepared the samples, measured the optical transient absorption data and performed the corresponding data analysis. J.H. initiated the idea, designed the whole experiments and wrote the manuscript. X. Zhang and Q.K.

designed the transient X-ray absorption experiment X. Zhang contributed to manuscript writing and interpretation of the data. A.V.Z. measured the XPS spectrum and X. Zhuo measured the SAXS data. B.P., J.L., A.A. and B.D. measured transient X-ray absorption data with the remaining authors. All authors contributed to the interpretation of the results and preparation of the manuscript.

Additional Information

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