

Identification of Photoexcited Electron Relaxation in a Cobalt Phosphide Modified Carbon Nitride Photocatalyst

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Transition metal phosphides have been recognized as efficient cocatalysts to boost the activity of semiconductor photocatalysts. However, a rigorous and quantitative understanding is still to be developed about how transition metal phosphides influence photoexcited electron dynamics. Here, we present a nanosecond time-resolved transient absorption spectroscopy (TAS) study of the photoexcited electron dynamics in carbon nitrides (g-C₃N₄) before and after Co and/or P modifications. Our spectroscopic study showed that Co or P lowered the initial electron density, whereas they promoted the photoexcited electron relaxation of g-C₃N₄, with their half-life times (t_{50%}) of 2.5 and 1.8 ns, respectively. The formation of a CoP co-catalyst compound promoted the electron relaxation ($t_{50\%}$ =2.8 ns) without significantly lowering the charge separation efficiency. Density functional theory (DFT) calculations were undertaken to explore the underlying fundamental reasons and they further predicted that CoP, compared to Co or P modification, better facilitates photoexcited electron transfer from $g-C_3N_4$ to reactants.

Solar-to-fuel conversion is a promising route to store solar energy in a way that compensates for increasing environmental

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issues caused by fossil fuel consumption.^[1] In principle, the formation of electron-hole pairs in photocatalysts upon illumination and their subsequent involvement in redox reactions are the main processes in photocatalytic conversion. To achieve highly efficient photochemical conversion, the main challenge in photocatalysis is to maximize the extraction of charge carriers while suppressing their recombination.^[2]

Co-catalyst/semiconductor heterostructures have been the focus of many investigations in recent years for photocatalysis and solar energy conversion. Co-catalysts will not only serve as active sites, but also promote the photo-excited charge separation and transfer,[3] thus enhancing the photocatalytic activity. Significant attention has been drawn to transition metal phosphides (e.g., FeP, CoP, and NiP), [4] because of their low cost, earth abundance and high electrochemical activity. CoP in particular has been extensively studied as an effective co-catalyst for photocatalytic water splitting and CO₂ reduction reactions. [5] The reason for its enhanced O₂ evolution activity has been ascribed to the increased life-time of photogenerated holes. [6] However, its contribution to photogenerated electron dynamics remains elusive, although Co and/or P can significantly enhance the catalytic activity towards H2 evolution and CO₂ reduction reactions.^[5b-d]

Carbon nitride (g-C₃N₄), known as a polymeric semiconductor for H₂ evolution since the 19th century, is attracting significant research interest because of its visible light photocatalytic performance, good physicochemical stability and facile synthesis. [7] However, only a handful of studies have focused on its fundamental photophysical processes, especially the photoexcited electron dynamics, which control its photocatalytic activity.[8] It has been reported that CoP-loaded g-C₃N₄ has demonstrated higher photocatalytic activity for the H₂ evolution reaction than pristine or Co- or P-modified g- C_3N_4 , while the photoexcited electron dynamics of Co- and/or P-loaded q-C₃N₄ remain elusive. The current work seeks to uncover Co and/or P effects on the photoexcited electron dynamics in g-C₃N₄ by combining transient absorption spectroscopy (TAS) with density functional theory (DFT) calculations to investigate the structural differences and possible origins for the affected electron relaxation process. This study enables new insights into the fundamental processes that govern electron relaxation dynamics at the transition metal phosphide co-catalyst/semiconductor interface, and therefore provides a unique perspective for the development of low-cost co-catalyst/semiconductor heterostructures for efficient solar-to-fuel conversion.

The experimental investigation was realized by using nanosecond time-resolved pump-probe $\mathsf{TAS}^{\text{[10]}}$ with a temporal

resolution of lower than 1 ns and probe spectral range of 350–900 nm (see SI for details). The charge carrier dynamics were monitored after laser excitation at 355 nm, at which wavelength the steady-state absorption spectra were similar for the studied samples. The $g-C_3N_4$ used in this study was

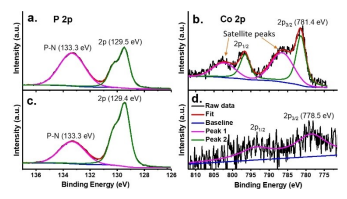


Figure 1. X-ray photoelectron spectroscopy (XPS) data of (a, c) P 2p and (b, d) Co 2p regions in (a) P/g-C₃N₄, (b) Co/g-C₃N₄ and (c, d) CoP/g-C₃N₄, the intensity of (d) was magnified by 1.5 fold to clearly show the peaks. The binding energy in the P 2p region around 133.3 eV corresponds to the interaction of P with N. The featured satellite peaks in the Co 2p region indicates the oxidized states of cobalt in Co/g-C₃N₄, and the oxidation state of CoP/g-C₃N₄ was similar to that of CoP structure.

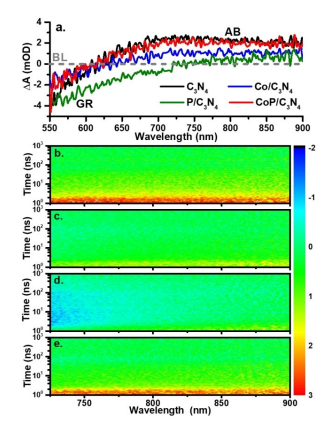


Figure 2. (a) Spectral slice taken from the transient absorption spectroscopy (TAS) data at the pump–probe time delay of 1 ns. The baseline (BL), ground state bleaching (GR) and photoinduced positive absorption (AB) were labelled. Hyperspectral TAS images are shown in (b) $g-C_3N_4$, (c) Co/ $g-C_3N_4$, (d) P/ $g-C_3N_4$ and (e) CoP/ $g-C_3N_4$, after excitation with a pump laser pulse of 355 nm (2 μJ/pulse, 1000 Hz).

synthesized by a typical thermal condensation method using urea as the precursor, [11] followed by pyrolysis and phosphorization reactions to obtain Co- and/or P-modified samples [9] (Co/g-C₃N₄, P/g-C₃N₄ and CoP/g-C₃N₄). Relatively low Co loading of ca. 0.5 wt % was chosen to ensure high optical transparency and minimize the light absorption of the CoP/g-C₃N₄ sample. The structure was confirmed by XRD, UV-vis and TEM (EDX) (Figures S1–S4). X-ray photoelectron spectroscopy (XPS) (Figure 1) demonstrated that P interacted with N in P/g-C₃N₄ and cobalt existed as oxidized states in Co/g-C₃N₄, while a CoP compound was formed in the CoP/g-C₃N₄ sample, which was in accordance with the reported results. [9,12]

Figure 2a shows the TA spectral slice taken at a time delay of 1 ns for the synthesized samples in water. The spectrum of $q-C_3N_4$ was similar to that of Godin et al., [8c] with the ground state bleaching around 550 nm and a positively broad absorption range above 600 nm. The positive absorption range has been ascribed to the photogenerated electrons, [8c,d] and was also confirmed by our TAS study, which showed that the broad absorption range was obviously enhanced when TEOA was added as the hole scavenger (Figure S5), while significant changes were observed in the TA spectra for the same g-C₃N₄ after Co or P modification. Co/g-C₃N₄ demonstrated lower positive absorption intensity, though with similar extended absorption range to that of g-C₃N₄. P/g-C₃N₄ displayed both lower positive absorption intensity and narrower absorption range, with the positive to negative crossover point of the TA spectrum moved to longer wavelength (~750 nm). However, the TA spectrum of CoP/g-C₃N₄ demonstrated obviously positive absorption in the compared range and displayed almost the same spectrum as that of g-C₃N₄ at the compared time slice of 1 ns.

The hyperspectral TAS images after 725 nm are shown in Figures 2b–e, which monitored the electron-induced TA spectra evolution. The broad photoexcited electron-induced hyperspectral image in Figure 2b indicates the long-lived electrons extended to long-timescale (μ s) in g-C₃N₄. The absorption intensity was obviously lowered for Co/g-C₃N₄ (Figure 2c) and P/g-C₃N₄ (Figure 2d). In P/g-C₃N₄ in particular, the positive absorption at short wavelength changed to negative bleaching with prolonged time. CoP/g-C₃N₄ clearly displayed high photoexcited electron absorption intensity, while with prolonged time, the spectra evolution was obviously different from that of g-C₃N₄.

To elucidate the dynamics of photogenerated electrons, we monitored and compared the TA kinetics around 800 nm for all the samples. As shown in Figure 3a, following the same laser excitation intensity, g-C₃N₄ displayed high initial TA intensity, then showed a power law kinetic decay with a life-time longer than our spectral detection time limit (Figure S6a), which is due to the formation of photocatalytically inactive electron trap states,^[8c,d] thus leading to low photoactivity. For Co/g-C₃N₄, the TA intensity decreased, which may be due to the decreased charge separation efficiency,^[13] but it still displayed a life-time of up to microsecond timescale (Figure S6b). Compared with g-C₃N₄, an interesting phenomenon occurred for P/g-C₃N₄, which showed that not only did the TA intensity decrease, but

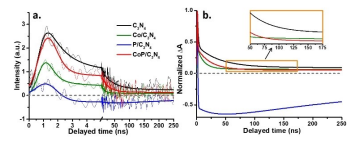


Figure 3. Transient absorption (TA) kinetic decays probed at 800 nm of g-C₃N₄, Co/g-C₃N₄, P/g-C₃N₄ and CoP/g-C₃N₄ in water after laser excitation at 355 nm (2 μ J/pulse, 1000 Hz): (a) before normalization, and (b) normalized fitting results with the inset shows the inverse phenomenon.

it also changed to a negative bleaching around 2 ns, with the bleaching extended outside of our spectral detection limit (Figure S6c), which may be ascribed to the trapping of photogenerated charges over P sites and will be discussed in the following section. After the formation of the CoP structure, CoP/g-C₃N₄ demonstrated completely different features compared to both Co/g-C₃N₄ and P/g-C₃N₄. The highest TA intensity of CoP/g-C₃N₄ only slightly decreased compared with that of pristine g-C₃N₄, while it demonstrated a different decreasing decay time around 3 ns, and with prolonged time the TA intensity was lower than that of pristine g-C₃N₄.

Furthermore, we use t_{50%} (the time at which the TA intensity of the photoexcited electrons has decreased by 50%)[13-14] to quantitatively compare the photogenerated electron dynamics. The $t_{50\%}$ of all the analyzed samples are in the order of $g-C_3N_4 > CoP/g-C_3N_4 > Co/g-C_3N_4 > P/g-C_3N_4$ (4.0, 2.8, 2.5 and 1.8 ns, respectively). Whereas Co and/or P modifications have been reported to exhibit higher photoactivity than pristine g-C₃N₄ for water splitting and CO₂ reduction, [5c-d,9] recent timeresolved spectroscopy studies have also shown that deeply trapped electrons with the life-time of longer than a microsecond were responsible for the limited H2 evolution activity of g-C₃N₄; [8c,d] thus, the photocatalytic activity of g-C₃N₄ would be mainly affected by the photogenerated electron dynamics on the nanosecond timescale. The decreasing $t_{50\%}$ is an indication of promoted relaxation of photoexcited electrons in g-C₃N₄ after Co and/or P modifications. The transfer times of photoexcited electrons from g-C₃N₄ to Co and/or P are estimated to be 6.7, 4.9 and 3.3 ns for CoP/g- C_3N_4 , Co/g- C_3N_4 and P/g- C_3N_4 , respectively (see SI for calculation details). These results indicate that the promoted electron relaxation occurred on the timescale of several nanoseconds, and their capabilities to extract the photoexcited electrons from g-C₃N₄ were in an order of $P/g-C_3N_4 > Co/g-C_3N_4 > CoP/g-C_3N_4$. Furthermore, the weak signals of Co/g-C₃N₄ and P/g-C₃N₄ indicate the ultrafast quenching of the photogenerated electrons, which might be ascribed to the ultrafast bulk recombination of the electronhole pairs, and thus may reduce the possibility of electron transfer to Co or P on picosecond timescales.

The normalized fitting results were compared in Figure 3b, where an inversed phenomenon around 75 ns (inset of Figure 3b) was observed between CoP/g-C₃N₄ and Co/g-C₃N₄. Such inversed phenomenon indicates that the electron trapping is

less strong over CoP/g-C₃N₄, which may contribute to the higher photoactivity of the CoP co-catalyst rather than that of Co or P modification.^[5c,9] This is also verified by the calculated work functions for Co/g-C₃N₄ and CoP/g-C₃N₄ which were 4.68 and 4.37 eV, respectively. With the lower work function for CoP/g-C₃N₄, the photoexcited electrons in CoP/g-C₃N₄ are more easily accessible for reactions, which may correspond to the inversed phenomenon of the spectra fitting results (inset of Figure 3b).

Further insight into the underpinning origins for the different TA kinetics was gained by DFT analysis. Compared with pristine $g-C_3N_4$ (Figure 4a), modification of the electronic band gap of the $g-C_3N_4$ was observed after Co and P doping, as shown in Figure 4b and c. Consistent with our XPS analysis and previous reports, P was preferentially substituted at the C site (Figure 4c), whereas the Co was interstitially doped at the hollow site, bridging two N atoms (Figure 4b). The band gap of the pristine $g-C_3N_4$ was predicted at 2.80 eV, which was found to narrow upon Co (1.84 eV) and P (2.17 eV) doping. The valence band edge of the pristine $g-C_3N_4$ is dominated by N-p orbitals. Compared to P doping, the valence band edge is modified by Co doping, with the Co-d orbitals dominating the valence band edge of Co/ $g-C_3N_4$. The average C-N distance in the pristine $g-C_3N_4$ is calculated at 1.335 Å, compared to a P-N

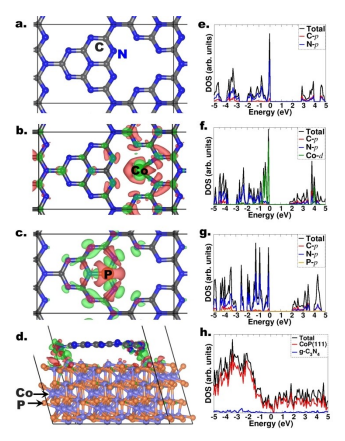


Figure 4. Optimized structures of (a) $g-C_3N_4$, (b) Co/C_3N_4 , (c) P/C_3N_4 and (d) $CoP(111)/C_3N_4$, with their corresponding projected density of states (e)–(h). The green and red isosurface contours from different charge density analysis denote regions of charge density accumulation and depletion, respectively.



distance of 1.501 Å in the doped P/g-C₃N₄, whereas the average bridging Co–N bond distance in Co/g-C₃N₄ is predicted at 2.081 Å. The interface formation between g-C₃N₄ and the CoP (111) surface is found to induce a much larger reduction in the bandgap of g-C₃N₄, to 1.68 eV, which is consistent with the enhanced visible light absorption of the CoP/g-C₃N₄ sample (Figure S3). The significant reduction in the bandgap of g-C₃N₄ in the interface structure can be attributed to the intrinsic metallicity of bulk CoP (Figure S8) and its CoP(111) surface (Figure 4h).

The electron density redistribution and transfer between CoP and g-C₃N₄ was analyzed via differential charge density isosurfaces and Bader charge analyses. In the P/g-C₃N₄ system, the extra electron from the C substitution by a P atom delocalizes to the π -conjugated triazine ring, creating positively charged P⁺ centers, which can promote efficient separation of photogenerated charges. Consistently, the calculated Bader charge of the P site $(+5.00 e^{-})$ in P/g-C₃N₄ is higher than that of the interstitial Co sites (+0.97 e⁻) in Co/g-C₃N₄. The larger positive charge and thus stronger Lewis acid site of the P⁺ centers indicates that the transfer of photogenerated electrons from g-C₃N₄ to P⁺ would be much easier/faster than to the Co⁺ centers, which is in accordance with the electron transfer time of our TA kinetic results (Figure 3). It has been verified that the photoexcited electrons will be produced by the N sites, [7a,9] and the XPS results in Figure 1a shows that there are strong P-N interactions in the P/q-C₃N₄ sample. Thus, the larger positive charge of the P⁺ sites will also be responsible for the trapping of electrons to form catalytically inactive states, which induces the transition from a positive to a negative bleaching with prolonged time (Figure 3). Although P/g-C₃N₄ showed the fastest extraction of the photoexcited electrons in $g\text{-}C_3N_4$, the photoactivity is low due to electron trapping. [5c-d,9] It is evident from Figure 4d that electron density accumulates (green contours) around the centers of the newly formed C-Co bonds at the g-C₃N₄/CoP(111) interface, and Bader charge analysis demonstrates that the g-C₃N₄ loses about 0.67 electrons to the top of the CoP(111) surface, which may correspond to the relatively low capability to extract photogenerated electrons (Figure 3a). However, the work functions for g-C₃N₄, Co/g-C₃N₄, P/g-C₃N₄ and CoP/g-C₃N₄ were calculated at 4.88, 4.68, 4.59 and 4.37 eV, respectively. The lowest work function of CoP/g-C₃N₄ shows that the photoexcited electrons over CoP/g-C₃N₄ are more easily accessible for reactions.

In summary, the influence of Co and P on photoexcited electron dynamics in $g\text{-}C_3N_4$ is directly monitored on the nanosecond timescale, which is the timescale of the photoexcited electron transfer processes taking place in a $g\text{-}C_3N_4$ photocatalytic material. Combined with DFT calculations, further insights are provided for different electron relaxation processes, thus explaining the current reported results, which showed that a CoP co-catalyst displays excellent photocatalytic performance. The origin of this behavior may come from the promotion of photoexcited electron relaxation in $g\text{-}C_3N_4$ without significantly lowering the charge separation efficiency; these photoexcited electrons would be easily available for reactions. Furthermore, a special interface could form between

CoP and $g-C_3N_4$ to enhance the visual light absorption and promote the photoexcited charge separation, which will also benefit the photocatalytic activity.

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Conflict of Interest

The authors declare no conflict of interest.

Keywords: carbon nitride · cobalt phosphide · electron relaxation · photocatalysis · transient absorption spectroscopy

- [1] a) Y. Tachibana, L. Vayssieres, R. J. Durrant, Nat. Photonics 2012, 6, 511–518; b) C. Jiang, S. J. A. Moniz, A. Wang, T. Zhang, J. Tang, Chem. Soc. Rev. 2017, 46, 4645–4660.
- [2] a) R. Marschall, Adv. Funct. Mater. 2014, 24, 2421–2440; b) A. Kafizas, R. Godin, J. R. Durrant, Semicond. Semimetals 2017, 97, 3–46; c) P. Zhang, T. Wang, X. Chang, J. Gong, Acc. Chem. Res. 2016, 49, 911–921.
- [3] a) X. Li, J. Yu, M. Jaroniec, X. Chen, Chem. Rev. 2019, 119, 3962–4179;
 b) J. Yang, D. Wang, H. Han, C. Li, Acc. Chem. Res. 2013, 46,1900–1909;
 c) K. Wu, H. Zhu, T. Lian, Acc. Chem. Res. 2015, 48, 851–859.
- [4] a) P. Jiang, Q. Liu, Y. Liang, J. Tian, A. M. Asiri, X. Sun, Angew. Chem. Int. Ed. 2014, 53, 12855–12859; Angew. Chem. 2014, 126, 13069–13073;
 b) E. J. Popczun, C. G. Read, C. W. Roske, N. S. Lewis, R. E. Schaak, Angew. Chem. Int. Ed. 2014, 53, 5427–5430; Angew. Chem. 2014, 126, 5531–5534;
 c) Z. Huang, Z. Z. Chen, Z. Z. Chen, C. Lv, H. Meng, C. Zhang, ACS Nano 2014, 8, 8121–8129.
- [5] a) D. K. Zhong, M. Cornuz, K. Sivula, M. Grätzel, D. R. Gamelin, Energy Environ. Sci. 2011, 4, 1759; b) P. Huang, J. Huang, S. A. Pantovich, A. D. Carl, T. G. Fenton, C. A. Caputo, R. L. Grimm, A. I. Frenkel, G. Li, J. Am. Chem. Soc. 2018, 140, 16042–16047; c) R. Shi, H.-F. Ye, F. Liang, Z. Wang, K. Li, Y. Weng, Z. Lin, W.-F. Fu, C.-M. Che, Y. Chen, Adv. Mater. 2018, 30, 1705941; d) Y. Zhang, T. Mori, J. Ye, M. Antonietti, J. Am. Chem. Soc. 2010, 132, 6294–6295.
- [6] a) M. Barroso, A. J. Cowan, S. R. Pendlebury, M. Grätzel, D. R. Klug, J. R. Durrant, J. Am. Chem. Soc. 2011, 133, 14868–14871; b) M. Barroso, C. A. Mesa, S. R. Pendlebury, A. J. Cowan, T. Hisatomi, K. Sivula, M. Grätzel, D. R. Klug, J. R. Durrant, Proc. Natl. Acad. Sci. USA 2012, 109, 15640–15645; c) Y. Ma, Andreas Kafizas, S. R. Pendlebury, F. L. Formal, J. R. Durrant, Adv. Funct. Mater. 2016, 26, 4951–4960.
- [7] a) X. Wang, K. Maeda, A. Thomas, K. Takanabe, G. Xin, J. M. Carlsson, K. Domen, M. Antonietti, Nat. Mater. 2009, 8, 76–80; b) J. Liu, Y. Liu, N. Liu, Y. Han, X. Zhang, H. Huang, Y. Lifshitz, S-T. Lee, J. Zhong, Z. Kang, Science 2015, 347, 970–974; c) W.-J. Ong, L.-L. Tan, Y. H. Ng, S.-T. Yong, S.-P. Chai, Chem. Rev. 2016, 116, 7159–7329; d) M. Z. Rahman, Chaillins, Acc. Chem. Res. 2019, 52, 248–257; e) F. Podjaski, J. Kröger, B. V. Lotsch, Adv. Mater. 2018, 30, 1705477; f) V. W. Lau, I. Moudrakovski, T.



- Botari, S. Weinberger, M. B. Mesch, V. Duppel, J. Senker, V. Blum, B. V. Lotsch, *Nat. Commun.* **2016**, *7*, 12165.
- [8] a) C. Merschjann, S. Tschierlei, T. Tyborski, K. Kailasam, S. Orthmann, D. Hollmann, T. Schedel-Niedrig, A. Thomas, S. Lochbrunner, Adv. Mater. 2015, 27, 7993–7999; b) H. Zhang, Y. Chen, R. Lu, R. Li, A. Yu, Phys. Chem. Chem. Phys. 2016, 18, 14904–14910; c) R. Godin, Y. Wang, M. A. Zwijnenburg, J. Tang, J. R. Durrant, J. Am. Chem. Soc. 2017, 139, 5216–5224; d) K. L. Corp, C. W. Schlenker, J. Am. Chem. Soc. 2017, 139, 7904–7912; e) Z. Chen, Q. Zhang, Y. Luo, Angew. Chem. Int. Ed. 2018, 57, 5320–5324; Angew. Chem. 2018, 130, 5418–5422; f) M. A. Khan, P. Maity, M. Al-Oufi, I. K. Al-Howaish, H. Idriss, J. Phys. Chem. C 2018, 122, 16779–16787; g) L. Jing, R. Zhu, D. L. Phillips, J. C. Yu, Adv. Funct. Mater. 2017, 27, 1703484–9.
- [9] W. Liu, L. Cao, W. Cheng, Y. Cao, X. Liu, W. Zhang, X. Mou, L. Jin, X. Zheng, W. Che, Q. Liu, T. Yao, S. Wei, Angew. Chem. Int. Ed. 2017, 56, 9312–9317; Angew. Chem. 2017, 129, 9440–9445.
- [10] G. X. Pei, J. H. J. Wijten, B. M. Weckhuysen, Phys. Chem. Chem. Phys. 2018, 20, 9806–9811.
- [11] D. J. Martin, K. Qiu, S. A. Shevlin, A. D. Handoko, X. Chen, Z. Guo, J. Tang, Angew. Chem. Int. Ed. 2014, 53, 9240–9245; Angew. Chem. 2014, 126, 9394–9399.

- [12] a) H. Yang, J. Ouyang, A. Tang, J. Phys. Chem. B 2007, 111, 8006–8013;
 b) A. P. Grosvenor, S. D. Wik, R. G. Cavell, A. Mar, Inorg. Chem. 2005, 44, 8988–8998;
 c) Y. Zhang, L. Gao, E. J. M. Hensen, J. P. Hofmann, ACS Energy Lett. 2018, 3, 1360–1365.
- [13] S. R. Pendlebury, A. J. Cowan, M. Barroso, K. Sivula, J. Ye, M. Grätzel, D. R. Klug, J. Tang, J. R. Durrant, *Energy Environ. Sci.* 2012, 5, 6304–6312.
- [14] S. A. Haque, Y. Tachibana, D. R. Klug, J. R. Durrant, J. Phys. Chem. B 1998, 102, 1745–1749.
- [15] J. Feng, D. Zhang, H. Zhou, M. Pi, X. Wang, S. Chen, ACS Sustainable Chem. Eng. 2018, 6, 6342–6349.

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