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Cite this: Phys. Chem. Chem. Phys., 2021. 23. 25629

Understanding the mechanism of plasmon-driven water splitting: hot electron injection and a near field enhancement effect†

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Utilizing plasmon-generated hot carriers to drive chemical reactions has currently become an active area of research in solar photocatalysis at the nanoscale. However, the mechanism underlying exact transfer and the generation dynamics of hot carriers, and the strategies used to further improve the quantum efficiency of the photocatalytic reaction still deserve further investigation. In this work, we perform a nonadiabatic excited-state dynamics study to depict the correlation between the reaction rate of plasmon-driven water splitting (PDWS) and the sizes of gold particles, the incident light frequency and intensity, and the near-field spatial distribution. Four model systems, H₂O and Au₂₀@H₂O separately interacting with the laser field and the near field generated by the Au nanoparticle (NP) with a few nanometers in size, have been investigated. Our simulated results clearly unveil the mechanism of PDWS and hot-electron injection in a Schottky-free junction: the electrons populated on the antibonding orbitals of H₂O are mandatory to drive the OH bond breaking and the strong orbital hybridization between Au₂₀ and H₂O creates the conditions for direct electron injection. We further find that the linear dependence of the reaction rate and the field amplitude only holds at a relatively weak field and it breaks down when the second OH bond begins to dissociate and field-induced water fragmentation occurs at a very intensive field, and that with the guarantee of electron injection, the water splitting rate increases with an increase in the NP size. This study will be helpful for further improving the efficiency of photochemical reactions involving plasmon-generated hot carriers and expanding the applications of hot carriers in a variety of chemical reactions.

Received 30th July 2021, Accepted 22nd October 2021

DOI: 10.1039/d1cp03509f

rsc.li/pccp

1 Introduction

Plasmonic metal (typically Au, Ag, and Cu) nanoparticles (MNPs) have extremely important applications in many fields due to their characteristic optical properties¹⁻³ such as localized surface plasmon resonance (LSPR). When the incident light induces the collective oscillation of metal free electrons, LSPR occurs in those MNPs.4,5 However, this coherent electron oscillation can dephase quickly by nonradiative decay,6-9 generating electron-hole pairs with higher energy in a nonequilibrium state (*i.e.*, hot carriers) at the timescale of 1–100 fs.¹⁰ During dephasing, the collective dipole moment decreases along with the near-field interaction and scattering probabilities, with the incident energy converted into a hot carrier distribution. 1,11

When molecules are adsorbed on the metal nanostructure surface, the hot electrons can be transferred to the empty states of adsorbates before thermalization and create a transient negative ion (TNI). 12-15 With the response of molecular geometry, the TNI moves on the excited-state potential energy surface, and therefore their reactive capability is mediated. The plasmoninduced photochemical reactions have different properties from thermal activation, and have the potential to efficiently convert solar energy into chemical energy.^{2,8,16}

In general, there exist two pathways for hot electron transfer, the conventional indirect transfer and the recently proposed direct transfer. 14,17-19 In the indirect electron transfer process,20 hot electrons are first generated in the plasmonic metal nanoparticle, and are then injected into the adsorbate. For direct electron transfer, 18,21 in the presence of empty

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[†] Electronic supplementary information (ESI) available: The DOS of an isolated H₂O molecule, the evolution of an OH bond when an isolated H₂O molecule interacts with the laser field, the time evolution of OH bonds in an Au₂₀@H₂O system under the laser field with $E_{\rm max}$ = 1.60 V Å⁻¹ and $E_{\rm max}$ = 1.70 V Å⁻¹, the absorption spectra of Au_{20} @ H_2O vary with the intensity of the incident field, the absorption spectra of Au tetrahedrons with L = 2.88 nm, 4.32 nm, and 5.76 nm calculated by FDTD, the contour plot of near fields, and the time evolution of the forces acting on the H₂O molecule in the XZ plane under a uniform field and the real near field, respectively. See DOI: 10.1039/d1cp03509f

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hybridized orbitals due to the strong metal-adsorbate coupling, chemical interface damping^{22,23} can induce the plasmon dephasing directly and generate nonthermalized electrons in the empty hybridized orbitals. The plasmon-induced hot carrier transfer has been utilized to foster various photochemistry processes, such as the dissociation of H_2 , $^{24-26}$ N_2 , 27 and O_2 , 28,29 CO_2 reduction, 15,30 water splitting $^{31-33}$ and organic transformations.³⁴

Solar photocatalytic water splitting is a promising method for hydrogen production. This process usually takes place in metal-semiconductor heterojunctions where enhance photoconversion in the semiconductor via three mechanisms, including light trapping, hot electron/hole transfer, and plasmon-induced resonance energy transfer. 16,35 The solar water splitting in a Schottky-free junction has been recently found by Robatjazi et al., 36 who observed large photocurrents as a result of direct injection of hot electrons from plasmonic Au NPs to molecules. Many theoretical studies started to explore the mechanism by investigating the effect of laser intensity and frequency on photocatalytic activity. For example, Meng's group³¹ simulated the H₂O splitting dynamics on Au nanospheres using a TDDFT/Ehrenfest dynamics scheme and found a strong correlation between laser intensity, hot electron transfer, and reaction rates. Zhang et al.25 focused on the H2 dissociation induced by Au spheres and found that the dissociation rate of H₂ was closely related to the molecular initial positions. In the above two studies, the Au particles were identified by the jellium spheres. The jellium model is the simplest physical model for the valence electrons, it replaces the real geometry of the metal ionic core by a featureless positively charged background in a finite volume and treats only delocalized electrons explicitly in the mean-field approximation, such as the Hartree-Fock or the local density approximations. The jellium model neglects the lattice structures of metal spheres, and hardly describes the interaction between the metal and adsorbates, and the effect of lattice vibration. Furthermore, the LSPRs of MNPs are largely dependent on the MNP shapes and sizes. It is essential to involve those effects in describing plasmon-mediated chemical reactions.

In this work, we investigate the dynamic process of water splitting driven by an Au cluster and NPs with a few nanometers in size under a femtosecond laser pulse. We explore the related mechanisms of H₂O splitting and hot electron injection, and reveal the relationships between the reaction rate and the incident light intensity and frequency, and NP sizes. The realspace real-time TDDFT/Ehrenfest dynamics scheme is adopted. Four model systems, H₂O and Au₂₀@H₂O interacting with the laser field and the near field generated by different sized NPs, have been investigated. The initial geometry of the water molecule is set to be adjacent to the one of the vertexes of Au_{20} . Au_{20} has a tetrahedral structure and is highly stable and chemically inert. 37,38 The tetrahedral Au₂₀ is a fragment of the face-centered cubic lattice of bulk gold with a small structural relaxation, a unique molecule with atomic packing similar to that of bulk gold.³⁸ For H₂O and Au₂₀@H₂O systems, their electronic degrees of freedom were treated quantum mechanically within the real-space real-time implementation of time-dependent

density functional theory (TDDFT), while their nuclei were handled classically.

Currently, it is still a great challenge to describe the excitedstate dynamics of the hybrid system of molecule-MNPs with large sizes without using the jellium approximation to the MNPs. Here, we thus divide the nanosized tetrahedron into two parts: the tetrahedral cluster Au₂₀ and the residue, where Au₂₀ is described by TDDFT and the interaction of the residue to Au₂₀@H₂O is identified by the near field to which Au₂₀@H₂O is subjected. The near field is obtained by the numerical solution of Maxwell's equations using the classical computational electrodynamics finite-difference time-domain (FDTD) technique. With this regard, the interaction of metal-molecule is described at the atomic level and the strong near-field enhancement effect of NPs can also get involved.

2 Theoretical and computational details

The real-space real-time TDDFT calculations reported in this work were carried out using the open-source OCTOPUS code (version 9.2). 39,40,41 The simulation grid is localized spherically around each atom with a radius of 8.00 Å, and the grid point is set to be 0.20 Å which is considered as the balance of simulation accuracy and time consumption. The core electrons are approximate by the Troullier-Martins pseudopotentials⁴² with 11, 6 and 1 valence electrons for Au, O and H atoms. Therefore, the Au₂₀@H₂O system has 228 valence electrons and 114 occupied orbitals, and 30 unoccupied states are included in all the calculations. The general gradient approximation exchangecorrelation functional Perdew-Burke-Ernzerhof (PBE)43 with Average-density self-interaction correction⁴⁴ is adopted in all DFT/TDDFT calculations. The time-dependent evolution is performed using the approximated enforced time-reversal symmetry algorithm⁴⁵ with a time step of 1.5 as. To obtain the optical absorption spectrum, a δ function electric field is used.

Nonadiabatic dynamics of Au20@H2O is treated by the TDDFT/Ehrenfest dynamics scheme 46 with temperature fluctuating at around 300 K. The set of equations to be solved for the electron and ion dynamics of Au20@H2O are

$$i\frac{\partial}{\partial t}\phi_i(\mathbf{r},t) = \left[\hat{h}(n(\mathbf{r},t)) + v_{\text{field}}(\mathbf{r},t)\right],$$
 (1)

$$m_{\alpha} \frac{\mathrm{d}^2 R_{\alpha}}{\mathrm{d}^2 t} = F_{\alpha}(\mathbf{R}, t), \tag{2}$$

where $v_{\text{field}}(\mathbf{r}, t)$ describes the time-dependent electric field acting on the system, $\{\phi_i\}(i = 1, 2,...,N)$ are the occupied Kohn-Sham orbitals, \hat{h} is the Fock operator and n is the one-electron density. R_{α} stands for the coordinate of the nucleus labeled α , m_{α} for its mass, and F_{α} for the force exerted on it.

The laser field is assumed to polarize along the +z direction with

the function form:
$$\vec{E}_0(t) = \vec{E}_{\max}^0 \exp\left[-\frac{(t-t_0)^2}{2\tau^2}\right] \cos[\omega_0(t-t_0)],$$

where $|E_{\rm max}^0|$ denotes the amplitude of the external electric field, ω_0 is the excitation frequency, $t_0 = 6.60$ fs is the center of **PCCP** Paper

the Gauss-type laser and $\tau = 1.60$ fs is the width of the laser pulse. When the water molecule is in the proximity of the nanosized Au tetrahedron, the field E(r, t), interacting with $Au_{20}@H_2O$, includes the incident laser field $\vec{E}_0(t)$ and the scattered field $\vec{E}^{\rm sca}(r, t)$ generated by the truncated tetrahedron (Au₂₀ is cut off from this nanosized tetrahedron). This inhomogeneous near field can be expressed as $E(r, t) = E_s(r)E_t(t)$, where $E_s(r)$ is the spatial function that represents the non-uniform distribution of fields and $E_t(t)$ is the temporal function which matches the incident laser pulse. In order to describe the non-uniformity of the enhanced near-field, we need to obtain the expression of the spatial function $E_s(r)$. In this work, we applied a fifth-order polynomial to fit the spatial function $E_s(r)$ via a geodesic Levenberg-Marquardt (LM) algorithm, 47,48 namely,

 $E_{\rm s}(r) \approx a_0 + \sum_{i=1}^{3} a_i r^i$, where *r* represents the coordinate of the space grid and a_i is the fitting parameters.

The classical electromagnetic simulations to the Au NPs are performed using the FDTD++ package. 49,50 The Maxwell's equations are solved using Yee's algorithm. 51,52 Considering that Au₂₀ is a fragment of the face-centered cube lattice of bulk gold, or a part of the whole large-sized tetrahedron, we chose the shape of the Au NP to be tetrahedral. The other reason for our choice is that we know that a weak laser field can induce an intensive near field near the apex of an Au tetrahedron, even when the size of the NP is small. The dielectric function of an Au tetrahedron is described by the Drude-Lorentz model⁵³ in the form of

$$\varepsilon(\omega) = \varepsilon(\infty) - \frac{\omega_{\rm D}^2}{\omega(\omega + i\gamma_{\rm D})} - \sum_{n=1}^2 \frac{\Delta \varepsilon_{\rm Ln} \omega_{\rm Ln}^2}{\omega(\omega + 2i\delta_{\rm Ln}) - \omega_{\rm Ln}^2}.$$
 (3)

The parameters of the dielectric function are shown in Table 1.54 The cubic simulation box with a side length of 40,00 nm and a grid size of 0.20 nm is adopted in all FDTD calculations. We use the geodesic Levenberg-Marquardt (LM) algorithm^{47,48} to fit the FDTD field into a polynomial function which can be read in every TDDFT time step. We note that the current FDTD method is a classic electromagnetic model, the quantum size effect in the Au NPs is ignored, which can be accounted for by the nonlocal hydrodynamic model. 55,56 Neglecting the quantum effect may bring some errors to the field enhancement. For a metal tetrahedron with a side length longer than 3 nm, however, its quantum size effect seems not so obvious.⁵⁷

The excited-state population is computed by using orbital projections. The occupation at time t of the static orbital $\phi_{\rm m}(r)$ is given by projecting all occupied time-dependent orbitals onto the static unoccupied ones⁵⁸

$$p_{\rm m}(t) = 2\sum_{j}^{\rm occ} |\langle \phi_{\rm m}(0)|\phi_{j}(t)\rangle|^{2}, \tag{4}$$

Table 1 The parameters of dielectric function

$oldsymbol{arepsilon}_{\infty}$	Drude		Lorentz		
	ω_{D}	γъ	$\omega_{ m Ln}$	$\Delta arepsilon_{ m Ln}$	δ_{Ln}
5.513	9.013	0.000	3.658 2.833	1.857 1.004	0.714 0.399

and the normalization condition is $\sum\limits_{i}^{
m occ} |\langle \phi_i(0)|\phi_j(t)
angle|^2 +$ $\sum_{j=0}^{\mathrm{dinocc}} |\langle \phi_m(0) | \phi_j(t)
angle|^2 = 1$.

3 Results and discussion

3.1 Electronic structure and optical properties of Au₂₀@H₂O

Before investigating the dynamic processes of PDWS, we calculate the electronic and geometric structures of an isolated H₂O molecule and the Au₂₀(a)H₂O system. The energy gap between the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) for the isolated H₂O molecule is 8.74 eV, much larger than the energy of the visible light photon, indicating that it is difficult to excite H2O to drive water splitting.

During the geometry optimization, H₂O and Au₂₀ are initially optimized separately, then they are combined to form the hybrid system of Au₂₀@H₂O, which is re-optimized with all Au atoms frozen. The optimized distance between O and the apex of Au₂₀ is 3.03 Å as shown in Fig. 1a. The calculated density of state (DOS) shown in Fig. 1b indicates that the energy gap between the AB state and the HOMO of the hybrid system is 2.95 eV, which is much smaller than the HOMO-LUMO gap of H₂O, opening the possibility of excitation-induced electron transfer from the Au cluster to the AB state. In addition, we plot the wavefunction of the AB state of the Au₂₀@H₂O system and the optical absorption spectrum of Au₂₀@H₂O. For comparison, the absorption spectrum of isolated Au₂₀ is also plotted. The major absorption peak in the low energy band of Au₂₀@H₂O lies at 3.20 eV, and this peak is considered to be the plasmon resonance excitation. The spectral difference between Au₂₀ and Au₂₀@H₂O manifests

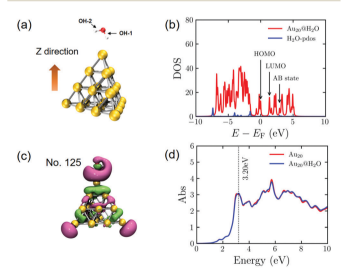


Fig. 1 (a) The optimized $Au_{20}@H_2O$ geometry. (b) The DOS of $Au_{20}@H_2O$ (Fermi-level is set to zero) and PDOS of the ${\rm H_2O}$ molecule (in blue), where we mark the HOMO (No. 114) and LUMO (No. 115, 1.35 eV) of the hybrid system, and the orbital with antibonding (AB) characteristics, namely, AB state (No. 125, 2.95 eV). (c) The MO of No. 125. (d) The absorption spectra of Au₂₀@H₂O and Au₂₀.

the strong orbital coupling between the two components, which can be thought to be caused by the charge transfer excitations and the interface metal/adsorbate hybrid states. 25,29,59

3.2 Electronic/nuclear dynamics of Au₂₀(a)H₂O

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In order to explore the hot electrons transfer mechanism of water splitting reaction, we show the dynamic process of H2O splitting in atomic sight by TDDFT/Ehrenfest dynamics in Fig. 2, where we set $\omega_0 = 3.07$ eV and $E_{\text{max}}^0 = 1.80$ V Å^{-1} (the corresponding laser intensity is $4.30 \times 10^{13} \text{ W cm}^{-2}$).

Fig. 2b shows the time-dependent evolution of the OH bond length (d_{OH}) of the hybrid system under an intensive external laser field. The initial lengths of two OH bonds are 1.03 Å, and during the active time window of the field, one of OH bonds denoted by d_{OH-1} , which is the one with an H atom pointing to the Au₂₀ apex, increases from 1.03 Å to 1.41 Å, and finally d_{OH-1} keeps increasing until it reaches 2.49 Å after 30 fs, and the corresponding OH bond splits. However, the other OH bond doesn't break, d_{OH-2} increases at first and reaches the maximum value of 1.35 Å in t = 17.16 fs, then it decreases and oscillates until t = 30 fs. In other words, the water splitting reaction is asymmetric. The OH bond near the apex of Au₂₀ is prone to breaking firstly, attributed to the initial structural arrangement. As Fig. 1 shows, we set one of the H atoms of H₂O to bind with an Au atom in the apex of Au₂₀ and the other to be away from the apex. With this structural arrangement, the symmetry of H₂O breaks down. As the MO of No. 125 in Fig. 1(c) shows, the wavefunction overlap between two OH bonds of H₂O and Au₂₀ will be different, indicating that their ability to accept electrons will be different. With this regard, the asymmetric dissociation appears in the PDWS process.

To obtain the information of products of water splitting, we compute the Hirshfeld charge⁶⁰ of H and O atoms in real time. The amount of Hirshfeld charge may be underestimated because of the electron loss when they reach the absorbing

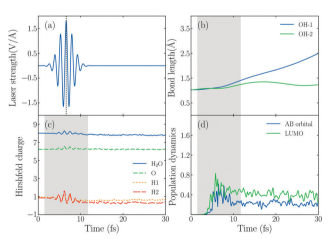


Fig. 2 (a) The laser pulse with $\omega_0 = 3.07$ eV and $E_{\rm max}^0 = 1.80$ V Å $^{-1}$ polarized in the Z direction. (b) Time evolution of O-H bonds in the Au₂₀@H₂O system interacting with the laser field. (c) Time evolution of the Hirshfeld charge of an H₂O molecule. (d) The population dynamics of the LUMO (No. 115) and AB state (No. 125) of Au₂₀@H₂O. The laser illumination in the time window indicated by the gray shaded areas.

boundary.61 The results in Fig. 2c show that the Hirshfeld charge of an H2O molecule oscillates with the change in the external field, and the total charge number of H and O atoms fluctuates around 1 and 8 after the laser is switched off. This result indicates that the H2O molecule in the Au20@H2O system splits into a hydrogen (H) and hydroxyl group (OH) under this laser field.

In order to understand the reaction mechanism, we calculate the time-evolved population of the AB state of the H₂O molecule and LUMO of the hybrid system in Fig. 2d. We observe that the photo-induced electron transfer takes place from the metal to the AB states of the H₂O molecule, indicating that the water splitting occurs when the AB states are populated by the electrons.

To further confirm this, we set a new system of an isolated H₂O molecule interacting with the same laser field. From the result shown in Fig. S2 (ESI†), we observe that the first OH bond oscillates with a period of 8.62 fs and the maximum bond length reaches 0.94 Å, with no signs of bond breaking. The population dynamics on the HOMO and LUMO of an isolated H₂O molecule also indicate that there are rarely electron populations in H2O's AB orbital, namely, this laser doesn't excite the electrons to the unoccupied orbitals of H₂O, so the isolated H₂O molecule doesn't split. These results suggest that the OH bond breaking in Au₂₀@H₂O is attributed to the photon-induced intermolecular electronic transfer from the metal to the H₂O.

Next we examine the dependence of the dissociation rate on the laser intensity and frequency. The dissociation rate of an H₂O molecule is defined as the inverse of the time required for the first OH bond length to reach 2.00 Å. As shown in Fig. 3a, initially, the splitting rate increases linearly as E_{max}^0 increases from 1.80 V \mathring{A}^{-1} to 2.20 V \mathring{A}^{-1} . A maximum rate appears at 2.20 V Å⁻¹. When the field intensity is further increased, the rate doesn't change obviously. To gain insight into this phenomenon, we plot the time-evolution of the OH bond length at $E_{\text{max}}^0 = 2.80 \text{ V Å}^{-1}$ in Fig. 3c, and find that both OH bonds are dissociated in this case. The intensive laser field can drive two OH bonds to break one after another though the reaction rate doesn't change obviously. As the light field increases from 1.80 V Å⁻¹ to 2.80 V Å⁻¹, there is a transition from water splitting to water fragmenting, and the dissociation rate of the OH bond even decreases.

To unveil the possibility of water fragmentation, we perform a calculation of the absorption spectra of Au20@H2O with varied laser intensity. Fig. S4 in the ESI† shows the calculated results. We observe that as the field intensity continuously increases, the intensities of the high-energy absorption bands with excitation energies ≥ 2.9 eV decrease and their peak locations blue-shift while the intensities of low-energy absorption bands continually increase. These phenomena manifest that the intensive laser fields couple with the system Au₂₀@H₂O nonlinearly.⁶⁴ The decrease of the high-energy absorption and the integrated energy shift are attributed to the absorption saturation and the field dressing during the excitation. The appearing and continually increasing intensity of low-energy

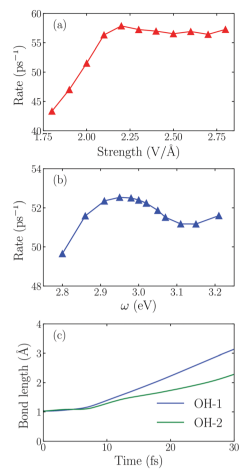


Fig. 3 (a) The reaction rate varies with the intensity of the incident laser with ω_{0} = 3.07 eV. (b) The reaction rate varies with the frequency of the incident laser with $E_{\text{max}}^0 = 2.00 \text{ V Å}^{-1}$. (c) The time evolution of two OH bonds under the laser field with $E_{\rm max}^0$ = 2.80 V Å⁻¹ and ω_0 = 3.07 eV.

bands confirms that multiphoton absorption occurs. This nonlinear coupling between the system and laser field definitely affects the dynamics of water splitting and may lead to the water fragmentation.

Fig. 3b shows that the splitting rate varies with the laser frequency. The maximum rate appears at $\omega_0 = 2.95$ eV, not at 3.20 eV, indicating that more hot electrons are transferred when the frequency of the incident field matches the energy gap between the Fermi-level and AB orbital, namely, the hot electron injection mechanism in the water splitting reaction follows the direct one.11,25

3.3 Electronic/nuclear dynamics of nanosized tetrahedron@H2O

To simulate the plasmon-induced water splitting reaction with an Au particle at the nanometer scale with a mixed TDDFT/ FDTD method, we firstly calculate the time-dependent spatial inhomogeneous scattering field generated by the truncated Au tetrahedron via the FDTD++ package, then describe the time evolution trajectory dynamics of the Au₂₀@H₂O system under the intensive near field by the TDDFT/Ehrenfest scheme. With this treatment, the hot electron injection is assumed to always appear around the interface between the metal cluster and the adsorbate, and the different sized NPs provide the near field with different intensity to which Au₂₀@H₂O is subjected, as Fig. 4 shows.

Fig. 5 shows the time evolution of the OH bond length in Au₂₀@H₂O under the intensive near field which is generated by the truncated Au tetrahedrons with different sizes. Here the laser field with $E_{\rm max}^0$ = 0.50 V Å⁻¹ and ω_0 = 3.07 eV is set. We observe that one of the OH bonds breaks at 25.12 fs under the field enhancement by the truncated Au tetrahedron with L =2.88 nm. In this case, the disassociation reaction takes place at

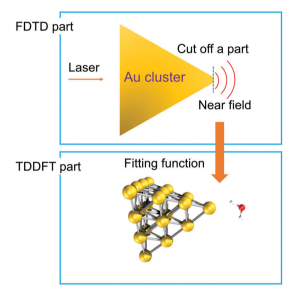


Fig. 4 A schematic diagram showing the mixed FDTD/TDDFT scheme for the PDWS reaction with large-sized NPs.

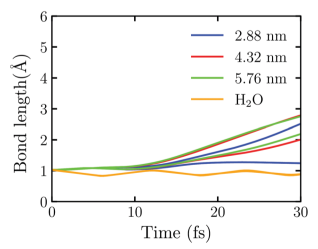


Fig. 5 Time evolution of OH bonds in Au₂₀@H₂O interact with the intensive near field generated by the truncated Au tetrahedron with different side lengths of 2.88 nm, 4.32 nm, and 5.76 nm, respectively. The orange line represents the time evolution of OH bonds in the isolated H₂O interacting with the near field generated by the whole Au tetrahedron of L = 2.88 nm (without the truncation of Au_{20}).

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 $E_{\rm max}^0 < 1.80 \text{ V Å}^{-1}$. Furthermore, we observe that the larger the NP's size, the faster the disassociation reaction. For example, at L = 4.32 nm, the two OH bond dissociation times are 21.47 fs and 27.80 fs while at L = 5.76 nm, they are 21.03 fs and 25.88 fs, respectively. For comparison, we also calculate the splitting dynamics of an isolated H₂O molecule which interacts with the near field generated by the whole Au tetrahedron of L = 2.88 nm obtained by FDTD. In this case, no electron is injected to H₂O, and only the field enhancement effect of Au NPs is involved. As Fig. 5 shows, the OH bond oscillates at a certain frequency, and there is no sign of splitting. These numerical simulations suggest that the near field can enhance the reaction rate of water splitting and one can adjust the size of a metal NP to control the plasmon-assisted photochemistry via the field enhancement effect. However, without the injection of hot electrons, the water splitting reaction will not take place, which highlights the decisive factor to drive the water splitting.

To show the field enhancement effect, we calculate the absorption spectra and near fields of the truncated Au tetrahedrons with different side lengths of 2.88 nm, 4.32 nm, and 5.76 nm in Fig. S5 and S6 (ESI†). The major low-energy absorption bands locate at 1.92-2.16 eV, deviating largely from the major low-energy absorption peak of Au₂₀, manifesting the quantum size effect of the optical responses of NPs with a few nanometers in size. The larger the NP's size, the stronger the absorption. And then, we calculate the strong scattered field generated by the truncated Au tetrahedrons. The incident laser is set to match the one used in TDDFT calculations. Fig. S6 (ESI†) clearly shows the strong field enhancement effect near the 'tip' of the tetrahedron, and the 'hot spots' are formed close to the Au particles. In addition, it is clear that the field enhancement ratio increases with the size of NPs.

The near field decays rapidly with the surface separation as Fig. S6 (ESI†) shows. To have a clear picture on the effect of the near field's inhomogeneity on the water splitting, we perform the TDDFT/Ehrenfest dynamics of Au₂₀@H₂O interacting with the real near field generated by the truncated Au tetrahedron with L = 2.88 nm and the uniform electric field taken from the center of mass of Au₂₀@H₂O generated by the same NP, respectively. In this case, a very weak laser field with ω_0 = 3.07 eV and $E_{\text{max}}^0 = 0.65 \text{ V Å}^{-1}$ is applied. As Fig. S7 (ESI†) shows, the duration of the scattered field at the center of mass of Au₂₀@H₂O is much longer than the incident field, and the scattered field's amplitude is magnified about three-fold so that the OH bond disassociates even with a weak laser field. Fig. 6 displays the evolution of OH bonds in Au₂₀@H₂O which is interacting with the two kinds of fields, respectively. In the laser activity window, in the case with a uniform field, the change of OH bond length shows an "upward arc", while in the real near field case, it shows as a "downward arc". The near field generated from LSPR is a very short-range electromagnetic field with a strong intensity gradient which may generate a gradient force to affect the molecule. 62,63

To explain why the field's spatial distribution can affect the evolution of OH bonds with time, we compare the forces acting on the H₂O molecule in Fig. S8 and S9 (ESI†). During the laser

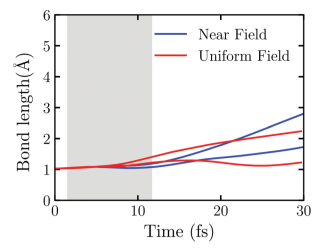


Fig. 6 Time evolution of the OH bonds of Au₂₀@H₂O interacting with the real near field and the uniform electric field, respectively. Here the laser field with $E_{\rm max}^0 = 0.65 \text{ V Å}^{-1}$ and $\omega_0 = 3.07 \text{ eV}$ is applied.

activity window (from 6.60 fs to 6.84 fs), the results with the uniform field and the near field are quite dissimilar. In the uniform field, the force vectors in the XZ plane acting on the H2O molecule are affected by the field and electron injection, and its direction and magnitude change rapidly. While in the near field, the force vectors in the XZ plane always point to the 'tips' of Au₂₀, especially the force acting on the O atom. It is known that this direction is also the negative gradient direction of the field. We thus suggest that the inhomogeneous near field affects the water splitting reaction through the gradient force and causes a different reaction during the field active window.

4 Concluding remarks

We have presented a theoretical study on the PDWS via TDDFT/ Ehrenfest nonadiabatic dynamics. By comparing the calculated results of four model systems: H2O and Au20@H2O separately interacting with the laser field and the near fields, we clearly reveal the microscopic mechanism of PDWS and the interface electron transfer, and the correlation of the reaction rate with the laser field frequency and intensity, the NP sizes, and the field's spatial distribution. A multiscale scheme has been applied to describe PDWS dynamics. For H₂O and Au₂₀@H₂O systems, its electronic degrees of freedom were treated quantum mechanically within the real-space real-time implementation of TDDFT, while their ions were handled classically. For the system with a larger sized NP, we divided this NP into Au₂₀ + residue, where Au₂₀ was described by TDDFT and the truncated NP was described by FDTD.

The main conclusions are summarized as follows:

- (1) The electrons populated on the AB orbitals of H₂O are mandatory to drive the OH bond breaking. The strong orbital hybridization between Au₂₀ and H₂O creates the condition for photo-induced direct electron injection.
- (2) The dynamic results under different laser intensities unveil that the linear dependence of the reaction rate of PDWS

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and the incident field amplitude holds only at a relatively weak field, however the linear correlation breaks down by the participation of other reactions such as water fragmenting in the intensive field regime. The splitting rate varies with the laser frequency, and the maximum rate appears when the laser frequency matches the energy gap between the metal Fermilevel and AB orbital.

- (3) To describe the effect of field enhancement induced by large-sized Au NPs, the mixed FDTD/TDDFT method is used. By dividing the large sized NPs into the Au₂₀ + residue, the requirement of near-field enhancement and hot-electron injection in the water splitting reaction can be simultaneously satisfied. Setting the water near the apex of the large-sized tetrahedron, OH bonds can be disassociated by a very weak laser field.
- (4) The intensive electric field can make two OH bonds in the Au₂₀@H₂O system break successively, attributed to the nonlinear coupling between the system and the applied field. When the intensity of the incident field reaches a certain degree, the field couples with the system nonlinearly, opening the possibility of multiphoton absorption. These hot electrons excited by multiphoton absorption can give important contributions in H₂O splitting reactions.
- (5) The influence of the field's spatial distribution on water splitting is significant. The gradient force caused by the strong intensity gradient of the near field makes the distinction of force vectors in the XZ plane compared to those in the uniform field case, leading to the different OH bond evolution dynamics.

This work is useful for understanding hot electron induced reactions at ambient conditions by plasmonic excitations and can provide a reference for the development of a related mixed quantum-classical method in describing large-sized plasmonic systems.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The financial support from the National Natural Science Foundation of China with Grant No. 21833006 and 22173074 is appreciated.

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