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# *Ab initio* investigation of hot electron transfer in CO<sub>2</sub> plasmonic photocatalysis in the presence of hydroxyl adsorbate†

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Photoreduction of carbon dioxide (CO<sub>2</sub>) on plasmonic structures is of great interest in photocatalysis to aid selectivity. While species commonly found in reaction environments and associated intermediates can steer the reaction down different pathways by altering the potential energy landscape of the system, they are often not addressed when designing efficient plasmonic catalysts. Here, we perform an atomistic study of the effect of the hydroxyl group (OH) on CO<sub>2</sub> activation and hot electron generation and transfer using first-principles calculations. We show that the presence of OH is essential in breaking the linear symmetry of CO<sub>2</sub>, which leads to a charge redistribution and a decrease in the O=C=O angle to 134°, thereby activating CO<sub>2</sub>. Analysis of the partial density of states (pDOS) demonstrates that the OH group mediates the orbital hybridization between Au and CO<sub>2</sub> resulting in more accessible states, thus facilitating charge transfer. By employing time-dependent density functional theory (TDDFT), we quantify the fraction of hot electrons directly generated into hybridized molecular states at resonance, demonstrating a broader energy distribution and an 11% increase in charge-transfer in the presence of OH groups. We further show that the spectral overlap between excitation energy and plasmon resonance plays a critical role in efficiently modulating electron transfer processes. These findings contribute to the mechanistic understanding of plasmon-mediated reactions and demonstrate the importance of co-adsorbed species in tailoring the electron transfer processes, opening new avenues for enhancing selectivity.

## Introduction

Subwavelength metallic nanoparticles that support plasmonic resonances represent a versatile platform for nanoscale light

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## New concepts

The development of plasmonic photocatalysts for CO<sub>2</sub> reduction is slowed down by a lack of atomistic understanding of the CO<sub>2</sub> behaviour on metallic structures under optical bias. Illuminated metallic nanoparticles generate hot electrons which can be transferred to adsorbate molecular species, thereby initiating a chemical reaction. While it is accepted that co-adsorbed species and/or reaction intermediates can influence the reaction pathways, this aspect is often overlooked. We report a first-principles investigation of plasmon-driven CO<sub>2</sub> reduction in the presence of a co-adsorbed hydroxyl group (OH) on a gold cluster. By using time-dependent density functional theory, we investigate the dynamics of plasmon excitation and decay and we analyse the energy distribution of hot electrons and quantify their transfer to the molecular adsorbates as a function of the distance, excitation frequency and an increasingly non-monochromatic frequency distribution. We demonstrate the importance of co-adsorbed species in changing the energetics of the system, showing that the OH group mediates the orbital hybridization between gold and CO<sub>2</sub> resulting in more accessible states, thus facilitating charge transfer. These findings contribute to the mechanistic understanding of plasmon-mediated reactions and demonstrate the importance of co-adsorbed species in tailoring the electron transfer processes, opening new avenues for enhancing selectivity.

manipulation. These structures hold great potential in nanophotonic applications that benefit from enhanced near-field light-matter interactions, such as plasmon-enhanced biosensing,<sup>1–3</sup> surface-enhanced Raman scattering<sup>4,5</sup> and photocatalysis.<sup>6–8</sup> Harnessing the hot electrons (HE)<sup>9</sup> generated by the non-radiative decay of illuminated metal nanoparticles before they thermalise is an attractive way to initiate and selectively drive chemical reactions,<sup>10–13</sup> and has recently attracted significant interest in the heterogeneous catalysis community.<sup>14–18</sup> Hot electrons can be rapidly transferred to the orbitals of nearby molecules, thereby driving catalytic chemical reactions. The energy and momentum distribution of the hot carriers depend on multiple factors, including material, particle morphology, crystal orientation, defects and environs,<sup>19,20</sup> influencing many aspects of heterogeneous catalysis, including the reaction rate, selectivity, turnover frequency, yield, and reaction order.<sup>21</sup> To date, the significant role

of HEs provided by photoexcited plasmons has been attributed to many reactions, including water splitting,<sup>22</sup> bond cleavage<sup>23</sup> or formation,<sup>24</sup> and carbon dioxide reduction.<sup>25</sup>

Once generated, HEs can be transferred to nearby molecules directly or indirectly.<sup>26,27</sup> In the former process, HEs are directly excited from the metal into the unoccupied molecular orbitals of interacting acceptor adsorbates. The latter process comprises HEs generated within metal nanoparticles, subsequently being scattered to empty states of surface-adsorbed molecules with suitable overlaps in energy and momentum distribution. The direct injection of HEs is a faster and more efficient process compared to the indirect transfer,<sup>28</sup> and offers great potential for enabling high selectivities to adsorbed chemical species and their reaction paths.<sup>28–30</sup> High selectivities can be achieved by precisely tuning the plasmon resonance and the energy distribution of HEs to align with targeted unoccupied adsorbate states, thereby selectively promoting a specific reaction pathway.<sup>6,17,30</sup> The design and engineering of the optoelectronic properties of plasmonic architectures are thus important in particular to achieve selectivity in multiproduct reactions such as CO<sub>2</sub> reduction.<sup>31</sup>

In the context of plasmon-enabled chemical transformations, reaction pathways and activation energies are often being investigated using ground-state density functional theory (DFT) methods.<sup>32–35</sup> Although these analyses provide important information on the systems and contribute to the understanding of the action mechanisms, they cannot accurately capture the physics of excited-states and the dynamics of the photogenerated hot carriers. Instead, time-dependent DFT (TDDFT) approaches have been increasingly employed to shed light on plasmon dynamics and hot carrier generation<sup>36–38</sup> by providing atomic-scale insights into excitation and transfer processes of electrons. In particular, Rossi *et al.* have developed a real-time TDDFT methodology to study the dynamics of plasmon-molecule systems,<sup>39–43</sup> demonstrating that the generated HEs have an energetic and spatial distribution that depends on the atomic structure, with lower-coordinated atoms exhibiting a

higher proportion of HEs.<sup>41</sup> These can be directly transferred to adjacent semiconductors<sup>40</sup> or molecules,<sup>29,42</sup> even when the molecule is not chemisorbed to the plasmonic nanoparticle:<sup>42</sup> using Ag–CO as a model system, Fojt *et al.* demonstrated that HEs can be directly generated on the molecule at distances up to 5 Å.

We extend the approach of Rossi *et al.* to investigate the energy required to activate carbon dioxide (CO<sub>2</sub>) on a small Au cluster in the presence of co-adsorbed environmental species and probe the carrier dynamics. It is widely accepted that the reaction environment and intermediate species can steer the reaction pathway<sup>44–46</sup> by altering the potential energy landscape of the chemical transformation. However, a mechanistic study of the effect of intermediate adsorbates on HE generation and transfer in CO<sub>2</sub> reduction is critical to enhance our understanding of the fundamental processes involved. Here, we focus specifically on the role of adsorbed hydroxyl groups, as they take part in the CO<sub>2</sub>RR and are crucial for enhanced stability, activity and selectivity.<sup>47–49</sup> By using TDDFT calculations, we study the HE transfer process as a function of distance, excitation frequency and an increasingly non-monochromatic frequency distribution, extending the fundamental understanding of HE transfer across nanoparticle–molecule interfaces and their impact on the overall CO<sub>2</sub> reduction reaction.

## Results and discussion

### System description

We consider CO<sub>2</sub> as a reactant species because its reduction is of significant industrial and societal importance: artificial synthesis of hydrocarbons and alcohols from CO<sub>2</sub>, using water to provide hydrogen and sunlight to drive the reaction, has the potential to provide renewable fuels and chemicals at scale.<sup>17,50</sup> A considerable challenge in converting CO<sub>2</sub> into organic products lies in its activation, due to the notably high reaction barrier linked to the initial electron transfer (−1.9 V *vs.*

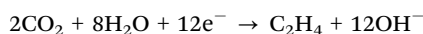
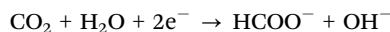


**Fig. 1** (a) Activation of CO<sub>2</sub> on a catalytic surface. The free molecule has a linear configuration with an O=C=O bond angle of 180°; upon interaction with the surface atoms, the bond angle reduces, thus activating CO<sub>2</sub>. (b) Spatial scheme of the valence orbitals of a gold atom (bright grey sphere) on the catalytic surface: colour-coding represents the spherical 6s (grey), d<sub>xy</sub> (yellow), d<sub>x<sup>2</sup>-y<sup>2</sup></sub> (red and green), d<sub>xz</sub> (purple), d<sub>yz</sub> (cyan) and d<sub>z<sup>2</sup></sub> (blue). The colours refer to AO lobes aligned to the x-/y-/z-axis (red/green/blue) with superimposed colours indicating bi-axial orientation; xy shown by yellow, xz shown by magenta, and yz shown by cyan. (c) Schematic representation of the system analysed in this work, comprising of a Au<sub>87</sub> cluster with an OH group and CO<sub>2</sub>.

standard hydrogen electrode). Adsorption of CO<sub>2</sub> on catalytic surfaces forms partially charged CO<sub>2</sub><sup>δ-</sup> species which – due to charge redistribution – results in a change in the O=C=O bond angle and leads to a bent configuration,<sup>51,52</sup> as shown in Fig. 1(a). As a consequence, the lowest unoccupied molecular orbital (LUMO) shifts to lower energies, reducing the energy barrier required for electron transfer and promoting a CO<sub>2</sub> reduction reaction.<sup>53</sup>

We use gold as a standard plasmonic material because of its unique electronic configuration which allows it to efficiently mediate catalytic reactions.<sup>31,54</sup> The electronic configuration of Au is shaped considerably by its atomic orbital arrangement. Gold has a 5d<sup>10</sup>6s<sup>1</sup> valence orbital configuration which consists of five filled 5d atomic orbitals that enable hybridization with various reactant molecules, and a partially filled 6s orbital. The five 5d orbitals are strongly lobed and are overlaid by the diffused spherical 6s orbital which does not hybridize with the 5d orbitals. The large nuclear charge of gold leads to a more pronounced nuclear-electron attraction of the valence electron shell: due to relativistic effects, the s-orbitals are found to contract in response to the large nuclear charge, while the d-orbitals are in fact expanded, increasing their involvement in the chemical and physical properties of gold.<sup>55,56</sup> These specific atomic orbital lobes allow for various combinations of hybridisation and formation of complexes with molecular species and ligands, making gold a promising material for catalytic applications and HE transfer. It is worth noting that this versatility may pose challenges to spatial selectivity, as gold can couple with numerous molecular species. However, the resulting higher spatial probability of electron transfer due to an increased overlap integral of the atomic and/or molecular orbitals involved promotes catalytic reactions. Selectivity for a specific reaction can be achieved by carefully tuning and aligning the energy distribution of HEs with the energy of the lowest unoccupied molecular orbitals (LUMOs) of the reactant species,<sup>28</sup> as discussed at the end of this section.

This is particularly relevant for multibranching chemical transformation like CO<sub>2</sub> reduction. Most of the high performing experimental literature on CO<sub>2</sub> reduction uses aqueous media that are either near-neutral or slightly alkaline.<sup>57–59</sup> While there are recent reports on CO<sub>2</sub> reduction under acidic conditions,<sup>60,61</sup> neutral and alkaline electrolytes are preferred because they provide a better CO<sub>2</sub> activation capacity, improve the suppression of the competing hydrogen evolution reaction and help in further promoting the C–C coupling efficiency.<sup>62,63</sup> An example of CO<sub>2</sub> reduction to typical products like formate and ethylene in alkaline media can be represented by the following equations:<sup>64,65</sup>



Under these conditions, the competing hydrogen evolution reaction can be drastically suppressed and the generated OH<sup>-</sup> intermediate can be an active participant in the reaction,

speeding up the activation of CO<sub>2</sub> and providing a means for further electron transfer.

### Electronic ground state properties

We start by investigating the adsorption and activation of a CO<sub>2</sub> molecule on a Au<sub>87</sub> cluster in presence and absence of an adsorbed hydroxyl group through structural relaxation. The two model systems are Au<sub>87</sub> + CO<sub>2</sub> and Au<sub>87</sub> + OH + CO<sub>2</sub>, with the CO<sub>2</sub> initially placed at a varying distance of 2 Å < *d* < 5 Å on a bridge-site, as this is the most catalytically active site for CO<sub>2</sub> reduction on gold surfaces,<sup>66</sup> while the OH group is adsorbed on an on-top site.

Fig. 2(a) shows the O=C=O bond angle when the systems reached equilibrium (forces < 0.05 eV per atom). In the absence of the OH group, CO<sub>2</sub> maintains a linear symmetry with a final equilibrium O=C=O angle which fluctuates around 178°, independent of the distance from the Au cluster (red curve). Contrarily, the presence of an OH group perturbs the charge symmetry of the molecule, leading to structural transformation for *d* < 3 Å. This results in a bent configuration with a final equilibrium O=C=O angle of 133°, in line with the previous theoretical reports on activated CO<sub>2</sub> on catalysts' surfaces.<sup>51,52</sup> Similar results are also obtained on periodic gold slabs (see ESI,† S1). Simultaneously, as CO<sub>2</sub> approaches the Au cluster (*d* < 3 Å), a redistribution of the charge density occurs, as can be seen by the different LUMO shapes of the system (Fig. 2(b)). A Bader charge analysis<sup>67,68</sup> was performed to investigate the charge redistribution which induces a charge dipole on the CO<sub>2</sub> molecule, with negative charges on the oxygen atoms (Fig. S2, ESI†). This in turn breaks its linear symmetry and results in an increased likelihood of adsorption on the Au cluster. These results suggest the importance of investigating the effects of co-adsorbed hydroxyl groups – and possibly other environmental species – in the overall activation of CO<sub>2</sub>, as well as the role of such species in HE transfer for efficiently driving catalytic processes.

When CO<sub>2</sub> approaches the Au and Au + OH clusters, the strength of orbital hybridization increases.<sup>42</sup> This can be demonstrated by analysing the projected density of states (pDOS) of the adsorbed molecules as a function of their distance to the Au surface for the different systems (Fig. 2(c) and (d)). At distances *d* ≥ 4 Å for both Au + CO<sub>2</sub> and Au + OH + CO<sub>2</sub> systems, the pDOS converges to the molecular density of states (DOS) of the isolated molecules. As *d* decreases, the pDOS eventually splits into several branches due to hybridisation with Au orbitals.<sup>42</sup> The addition of the OH group qualitatively shows a similar trend with a branched pDOS (Fig. 2(b)), but with an increased number of electronic states. In other words, the OH group mediates the orbital hybridization between Au and CO<sub>2</sub>, allowing for more accessible states.

We anticipate that these states are of significant importance for the further HE transfer process;<sup>31</sup> in fact, a HE transfer to the adsorbed molecule requires the presence of acceptor states at suitable energies, *i.e.* where the pDOS is present,<sup>42</sup> thus having more unoccupied levels which can lead to an increase of catalytic efficiency.



**Fig. 2** (a) CO<sub>2</sub> bond angle as a function of the distance during the adsorption process on Au and Au + OH clusters, represented by red and black curves, respectively. The LUMO wave functions of the systems are also shown at the same isosurface value of 0.05 a.u. (b) Magnification of the top-view of the LUMO at the same distances of (a). At  $d = 2.5$  Å, when transitioning from weak to strong hybridization (from 3 Å to 2.5 Å), we observe a phase flip in the molecular orbital (MO) lobes. This is due to the spin eigenfunction of the MO changing state, *i.e.* from spin-up to spin-down. When a phase flip happens, it means that the energy difference between both spin states of the MO is very small ( $\leq k_B T$ ) so that both electrons can swap their spin eigenfunction (spinor) within the MO. The spin-flip occurrence when going from 3 Å to 2.5 Å and then back to the original spin configuration shows that one spin configuration of the LUMO has a slightly higher  $E_{\text{LUMO}}$  compared to the  $E_{\text{LUMO}}$  at  $d = 2$  Å. (c) and (d) Map of the projected density of states as a function of the distance and energy for Au + CO<sub>2</sub> and Au + CO<sub>2</sub> + OH.

Additionally, this adsorption process is also accompanied by a shift towards a lower binding energy of the LUMO, decreasing from  $\sim 4.9$  eV ( $d \gtrsim 4$  Å) to  $\sim 4.6$  eV ( $d \sim 3.5$  Å) in line with the previous literature,<sup>14</sup> eventually splitting up into 4.7 eV (LUMO+1) and 4.5 eV (LUMO) for  $d < 3$  Å due to exchange interactions rising with inverse inter-atomic distance (with the overlap integral of associated orbitals forming a bond). This is another indication that the OH group acts as a “bridge”

between CO<sub>2</sub> and Au in terms of electronic coupling, which can then enable electron transfer from a considerably larger distance.

The total DOS of the different systems is shown in Fig. S3 (ESI<sup>†</sup>). The introduction of CO<sub>2</sub> and OH induces only minor alterations in the total DOS due to the dominant contribution of states originating from the Au cluster. Notably, in the presence of the molecules, the Au cluster exhibits a slightly

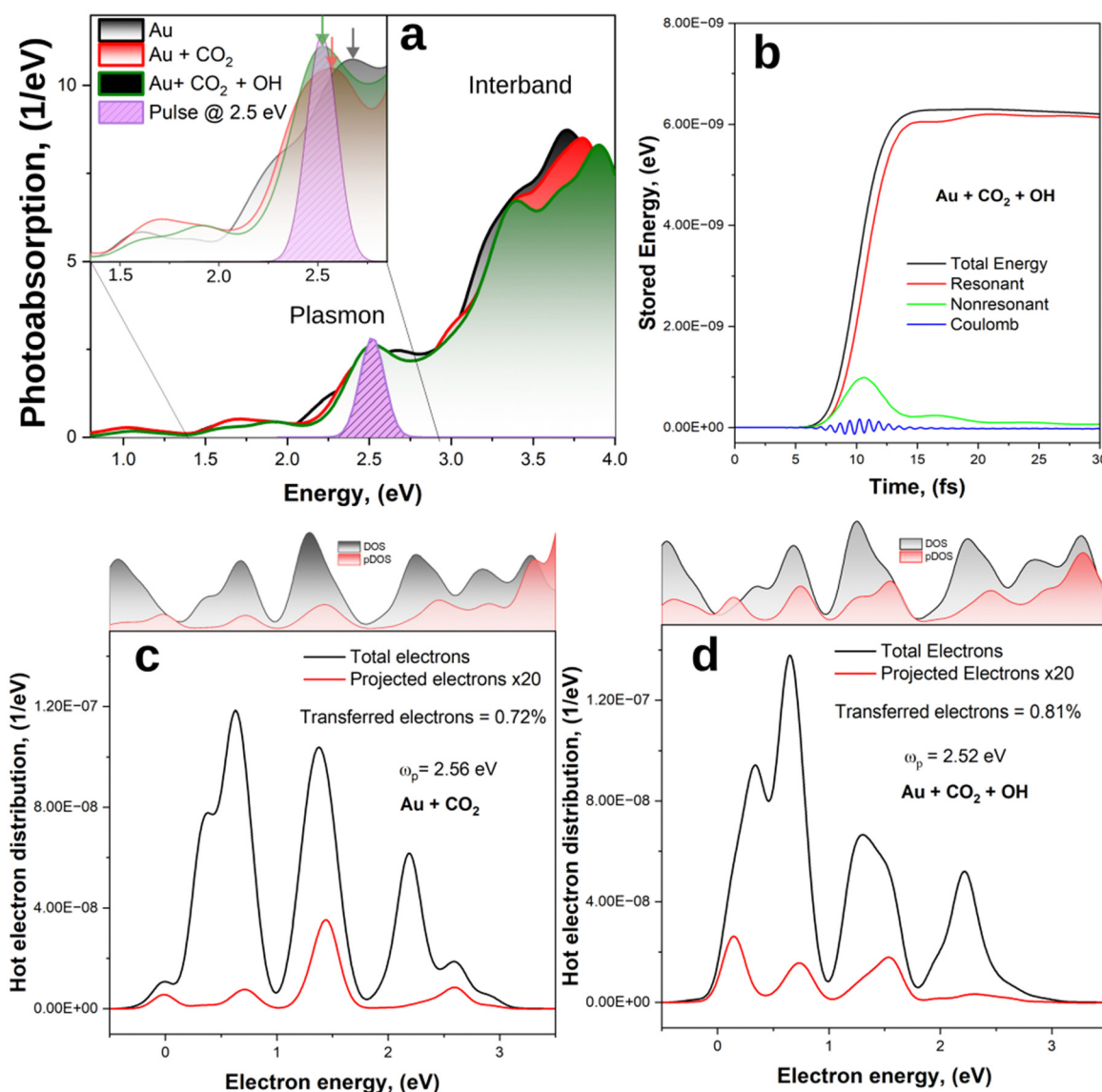


positive charge, which becomes more pronounced when the OH group is involved. The shift of the HOMO to a higher energy is  $\sim 0.1$  eV for Au + CO<sub>2</sub> (red curve) and  $\sim 0.16$  eV for Au + CO<sub>2</sub> + OH (green curve). This indicates that both the adsorbed OH and the CO<sub>2</sub> attract negative charges from the Au nanoparticle, which becomes apparent when comparing electronegativities, ionization energies, and electron affinities of C and O vs. Au.<sup>69</sup> Consequently, an electron transfer should occur from the Au NP to the molecular species. The CO<sub>2</sub> molecule induces a significantly increased shift of the HOMO to higher binding energy as compared to the shift when the OH group is included in the system. This dominant energy shift due to the CO<sub>2</sub> molecule suggests that an electron transfer from the Au cluster will eventually reside with the CO<sub>2</sub> molecule rather than with

the OH group, the latter presenting merely a transient charge path for transferring an electron to the CO<sub>2</sub> molecule.

### Plasmon dynamics and hot electron (HE) generation

We now study the time evolution of the Au, Au + CO<sub>2</sub> and Au + CO<sub>2</sub> systems under light illumination; when present, CO<sub>2</sub> is placed at a distance of 2 Å from the Au surface. First, the photoabsorption spectra of the system are calculated by applying an initial Dirac pulse ( $\delta$ -kick) in the linear-response regime along the  $z$ -direction (and perpendicular to the CO<sub>2</sub> molecule, when present). As the Fourier transform of the  $\delta$ -kick contains all frequencies with equal proportion, this kick equally excites all the optically allowed transitions and allows us to extract the photoabsorption spectrum. Next, we analyse the energy



**Fig. 3** (a) Photoabsorption spectra of Au, Au + CO<sub>2</sub> and Au + CO<sub>2</sub> + OH. The purple curve represents a typical driving laser pulse in the frequency domain, centered at the LSPR of the Au + CO<sub>2</sub> + OH. The colour coded arrows represent the LSPR of the different systems. (b) Time evolution of the stored energy for Au + CO<sub>2</sub> + OH, highlighting the different energy contributions. (c) and (d) Generated and transferred HEs energy distribution for the Au + CO<sub>2</sub> and Au + CO<sub>2</sub> + OH systems, respectively, averaged between 15 and 30 fs.

evolution of the systems after excitation with ultra-short optical pulses at the plasmonic resonance energy, which allows us to calculate the energy distribution of hot carriers generated by the decay of the plasmon. This analysis allows us to estimate the plasmon dephasing time after which most of the energy has been transferred to hot carriers.

The photoabsorption spectra (Fig. 3(a)) are characterized by a smoothly growing intensity starting at around 1 eV, with a plasmon peak at  $\sim 2.5$  eV, followed by a wide modulation up to 4 eV. The latter is in line with previous works on the optical properties of small gold clusters<sup>70,71</sup> and can be ascribed to interband transitions, which have a larger probability and a stronger contribution to the photoabsorption spectra than the plasmon excitation.<sup>72</sup> As expected, the plasmon peak is relatively weak, in line with the previous theoretical works on small gold clusters<sup>73–75</sup> which showed that a clear plasmon emerges only when the cluster size exceeds 2 nm.<sup>76</sup> For the combined systems, we observe a slight red-shift (green and red curves in Fig. 3(a)) of the plasmon peak from 2.66 eV for the single Au cluster, to 2.56 eV and 2.52 eV for Au + CO<sub>2</sub> Au + CO<sub>2</sub> + OH, respectively. This indicates the presence of additional decay channels for the plasmon due to formation of hybridized cluster-molecule states.

Next, we excite the systems with a Gaussian pulse  $-E(t) = E_0 \cos(\omega_p(t - t_0))\exp(-\sigma^2(t - t_0)^2/2)$  – with  $\omega_p$  set to be in resonance with the systems ( $\omega_p = 2.66$  eV for Au, 2.56 eV for Au + CO<sub>2</sub> and 2.52 eV for Au + CO<sub>2</sub> + OH), centred at 10 fs and with a temporal FWHM of 5 fs (Fig. 3(a), purple peak). In accord with the Heisenberg's uncertainty principle  $-\Delta t \times \Delta E \geq \hbar/2$  – a FWHM of  $\Delta t = 5$  fs corresponds to an uncertainty in the photon energy range of  $\Delta E = 0.41$  eV, presenting a broadband excitation as intended for plasmonic excitation with solar irradiation.

The energy of the light pulse is absorbed by the electrons in the system, promoting them into excited states. The electronic energy is not equally distributed, and can be divided into Coulomb energy, and electron-hole transition energy contributions<sup>41,77</sup> as shown for the Au + OH + CO<sub>2</sub> system in Fig. 3(b). The latter can be further separated into the energy of transitions resonant with the excitation pulse (*i.e.* with frequency  $\omega = \omega_p \pm 2\sigma$ ), constituting hot carriers, and non-resonant transitions, attributed to transitions from d-states in the metal nanoparticle which screen the plasmon. The evolution of the different energy contributions is similar for all the investigated systems (Fig. S4, ESI<sup>†</sup>), as the plasmon dephasing process is not affected on a significant scale by a single molecular species. Initially, the plasmon is excited *via* Coulomb interactions<sup>78</sup> and non-resonant contributions carry most of the energy during the plasmon excitation. After  $\sim 15$  fs, as the plasmon decays, the total energy is redistributed into electron-hole excitations that are resonant with the pulse.

Having established that after  $\sim 15$  fs the energy of the plasmon is mostly stored in hot carriers, we proceed to analyse the energy distribution of the photogenerated HEs, as well as the fraction of such HEs directly excited to states localised on the CO<sub>2</sub> molecule for the Au + CO<sub>2</sub> and Au + CO<sub>2</sub> + OH

systems. The HE energy distributions are shown in Fig. 3(c) and (d), where we have averaged the energies of the HEs between 15 fs and 30 fs as the resonant transitions have already reached their steady state. Therefore, HEs possess one temperature in accord with Fermi–Dirac statistics relative to the energy of their steady state as given by the convolution of the initial HE population with the total DOS of the system in the relevant energy range. For both the analysed systems, the generated HEs (black lines) can have energies up to the laser pulse frequency ( $E_F + \hbar\omega_p$ ) due to the conservation of energy. In addition, their energy distributions are not uniform within this range but show occupation probabilities that match the total DOS of the systems; electrons can only be excited to existing unoccupied states. The total DOS for Au + CO<sub>2</sub> and Au + CO<sub>2</sub> + OH are similar as they are dominated by the Au contributions. It follows that all systems have similar total HE energy distributions, presenting major peaks at  $\sim 0.7$  eV, 1.3 eV and 2.2 eV.

Conversely, the transferred electron energy distributions (red lines), which represent the fraction of HEs directly excited to states localised on the CO<sub>2</sub>, mirror the pDOS and exhibit differences between the systems. In particular, in the absence of the OH group, the fraction of HEs generated on the CO<sub>2</sub> is 0.72%, with most of them having an energy of 1.4 eV. The addition of the OH group increases the density of electronic states (see also Fig. 2(d)), thus opening more channels for electron transfer and effectively improving the charge transfer to 0.81%, which corresponds to an 11% increase with respect to the case without OH.

In the next section, we investigate the effect of the excitation parameters – pulse frequency ( $\omega_p$ ) and width (FWHM)– on the direct excitation of HEs to CO<sub>2</sub> orbitals. The pulses are visualised in the time domain as shown in Fig. S5 (ESI<sup>†</sup>) and have varying  $\omega_p$  from 1.6 eV to 2.8 eV, and FWHM from 4 fs ( $\Delta E = 0.52$  eV) to 15 fs ( $\Delta E = 0.14$  eV). The energy range was chosen to investigate the HE generated from the plasmon excitation as well as from interband excitation, noting that the threshold energy for interband transition (5d to 6sp) for gold is about 2.3 eV.<sup>79,80</sup> We have also investigated a pulse at 3.75 eV with a FWHM of 5 fs to match the maximum of the interband transitions within the analysed region (see Fig. 3(a)). The energy distributions of the HEs excited to states localised on the CO<sub>2</sub> molecule are reported in Fig. 4(a)–(d) as a function of pulse energy; the distributions for the total amount of HEs in both systems are given in Fig. S6 (ESI<sup>†</sup>). The amount of generated HEs increases with the increase of pulse width. For FWHM = 4 fs, we obtain a maximum HE density when  $\omega_p \sim 2.8$  eV, while for FWHM = 15 fs, the maximum occurs when  $2.3 < \omega_p < 2.7$  eV for all systems. In this energy range, the pulse excitation approaches the plasmon resonance of the systems (Fig. 3(a)), thus emphasizing the critical role of spectral overlap in efficiently modulating electron transfer processes.<sup>81,82</sup> Furthermore, below  $\sim 2$  eV, photoabsorption decreases, thus resulting in a negligible generation of HEs within the molecule.

At energies above the plasmon resonance, hot electrons are being generated *via* interband transitions.<sup>72</sup> While the total amount of HEs the systems is higher at the interband

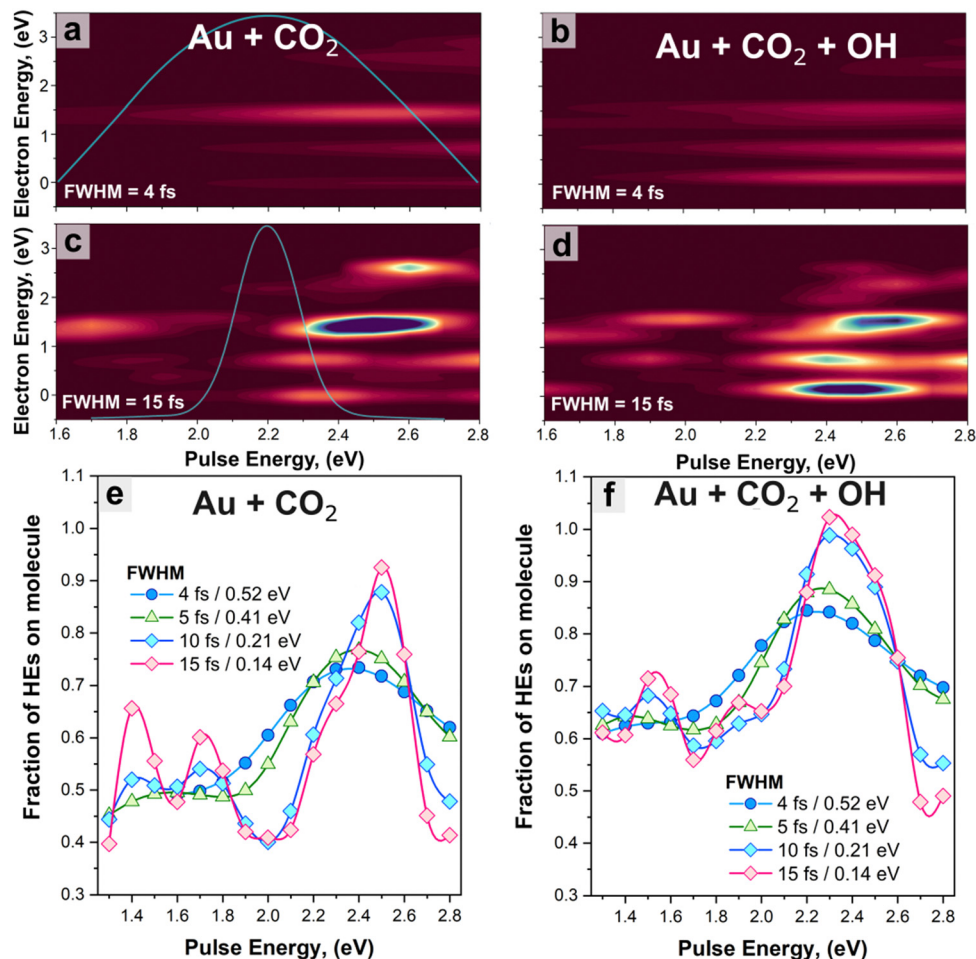


Fig. 4 (a) and (b) Amount of HEs generated on CO<sub>2</sub> and their energy distribution as a function of the pulse frequency (with a temporal FWHM of 4 fs (0.52 eV)) for Au + CO<sub>2</sub> and Au + CO<sub>2</sub> + OH, respectively. (c) and (d) Same as a and b, but the excitation has a FWHM of 15 fs (0.14 eV). The light blue lines in (a) and (c) represent a typical laser pulse in the frequency domain. (e) and (f) Fraction of carriers transferred to the molecule as a function of the excitation frequency for multiple FWHM values.

transitions (because of the larger photoabsorption<sup>72,80</sup>), the fraction of those HEs transferred to the CO<sub>2</sub> molecule is slightly smaller than at the plasmon resonance (Fig. S7, ESI†). This suggests that using the interband transition to activate and subsequently reduce CO<sub>2</sub> has a little advantage over the plasmon absorption in this specific system. In addition, exciting such transitions would come at the cost of a narrow excitation range and in-phase oscillation, preventing practical applications: using a laser to excite such interband transitions would preclude any use of sustainable energy such as solar irradiation for CO<sub>2</sub> reduction. This finding suggests that engineering the photoabsorption, specifically the plasmon resonance energy, should be considered to efficiently leverage photogenerated HEs for tailored reactions.

In the absence of the OH group (Fig. 4(a) and (c)), the energy distribution of HEs exhibits a prominent single peak centred around ~1.4 eV for pulse energies of 2.4–2.6 eV. Notably, approximately 55% (75%) of the HEs assume this specific energy level when subjected to pulse durations with the FWHM of 4 fs (15 fs) and under resonance conditions. Conversely, the

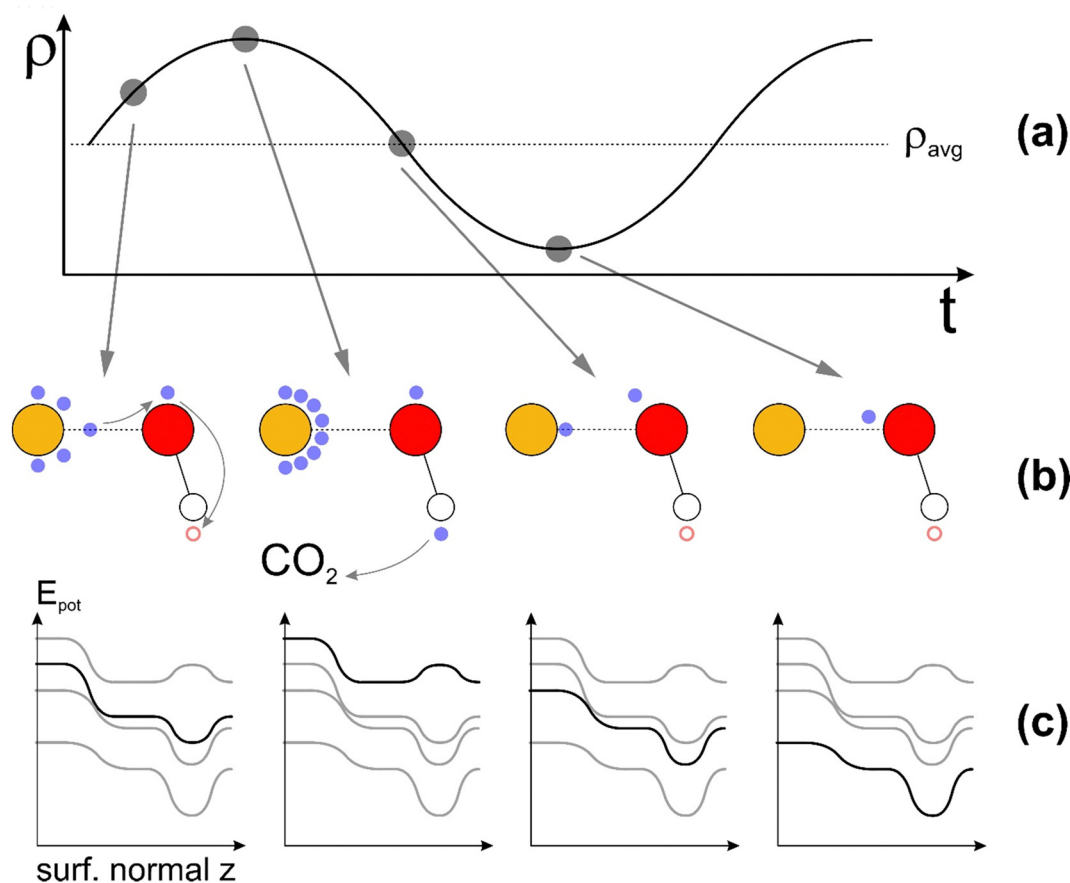
presence of the OH group results in a broader electron energy distribution. This is attributed to the presence of more accessible hybridized states within the energy range of 0–2 eV, which can be populated by HEs. The broadened energy distribution as facilitated by the OH group provides more channels for HEs transfer and promotes the system to a different excited level, potentially leading to a different product distribution.<sup>6</sup> This demonstrates that the presence of the OH group in the CO<sub>2</sub> photoreduction process alters the energy landscape and transfer behaviour of HEs, thereby possibly affecting the reaction pathway, and highlight the importance of taking into account the influence of environmental and intermediate species for selective HE catalysis.

Furthermore, the excitation width also affects the HE distributions. Narrow band pulses (FWHM = 15 fs;  $\Delta E = 0.14$  eV) lead to an overall increase in the generation of HEs on the molecule; conversely, with 'broadband' excitation (FWHM = 4 fs;  $\Delta E = 0.52$  eV), the pulse energy is redistributed to a wider range of possible transitions in the photoabsorption spectrum, leading to a broader but less intense (less resonant) electron

transfer. The transfer probability is given in Fig. 3(f) and 4, calculated as the ratio of the number of HEs generated on the molecule to the total HEs in the systems, and exhibits consistently higher values when the OH group is present, indicating a more efficient transfer. In particular, the addition of the OH group leads to an increase of  $\sim 10$  to 15% in the transfer probability, depending on pulse frequencies and FWHMs. Interestingly, when transitioning from the narrowband to broadband excitation (FWHM = 0.14 eV to 0.52 eV), despite a reduced transfer probability, the electron transfer to acceptor hybridized states remains high, with a maximum probability of *ca.* 0.84% for Au + CO<sub>2</sub> + OH at  $\omega_p = 2.3$  eV. This result suggests that HE transfer can still occur under broadband solar illumination with minor reductions of the yield, which is critical for the practical operation of photocatalysts.

While it is beyond the scope of this work to delve into the quantitative details of Landau damping, it is important to acknowledge the potential contribution of Landau damping from plasmons generated in gold (Au). Plasmons are collective longitudinal electron density waves which align electrons at their wave front to the resonant energy and  $k$ -vector of the

plasmon wave, thereby providing energy and momentum to electrons below the resonant energy or depleting energy and momentum from electrons with energies above the resonant energy and  $k$ -vector.<sup>83</sup> Such a resonant HE alignment, in the compound with the adsorbed OH group enabling the HE transfer to CO<sub>2</sub> species, may present a resonant charge pump system as briefly outlined below and illustrated schematically in Fig. 5. We note that Au is excellent for plasmon propagation along its cluster surface in an arbitrary direction due to its wide-angle coverage of surface states originating from its hybrid-AO configuration (Fig. 1(b)). With the plasmon wavefront increasing the HE density at the Au atom with an adsorbed OH group, HEs get pushed into OH groups where they get weakly localized, mainly at the outermost tip (towards positively charged H atom). Further advancement of the plasmon wavefront will result in the crest of the HE density wave, providing a dynamic Coulomb force to the HE localized at the end of the OH group, likely facilitating the HE transfer from the OH group to the CO<sub>2</sub> molecule, and providing the dynamic energy to overcome the weak localization at the OH group. With the passing of the HE density wave, its back end will



**Fig. 5** (a) Schematic of the evolution of electron density  $\rho$  over time  $t$ , featuring a surface plasmon. The grey points refer to specific situations in the charge transfer dynamics brought about by the plasmon. (b) Snapshots of charge distributions at key points of plasmonic oscillation. The gold-yellow sphere represents the Au surface atom, and the red and white spheres represent the O and H atom, respectively, of the adsorbed OH group. The small blue particles represent electrons, and the red empty circle represents the unoccupied valence state of the H atom. (c) Schematic evolution of the potential energy landscape over the Au surface atom with an adsorbed OH group, whereby the black line is the energy landscape referring to the molecule charge snapshot above.



dynamically deepen the electron localization potential of the OH group, reaching maximum localization ability at the density minimum of the plasmon. This ability sets the best possible condition to localize another HE in the next oscillation cycle, thereby repeating the HE transfer process.

In addition to the resonant HE transfer in the energy spectrum discussed earlier, the charge pump process described above may provide an additional resonance on the time spectrum. While it is beyond the scope of our work to delve into the quantitative details of such a Landau damping-mediated HE pump in the time regime, we point out that the dynamic electric dipole moment of the OH group as the adsorbed species may play a prominent role in further enhancing the HE charge transfer to the CO<sub>2</sub> molecule. Furthermore, the Landau damping-mediated resonance will likely have an impact on the selectivity of chemical species to receive the HE, thereby being reduced to a specific hydrocarbon or hydroxycarbon compound. With the CO<sub>2</sub> molecule and H<sub>2</sub> or H<sub>2</sub>O present as reactants, the testing of analogous adsorbed species to OH groups such as –O–Ag or –O–Li as stable adsorbents with suitable ionization energy and electron affinity of the cation (Ag, Li) provides a clear direction for future research into the ongoing optimization of adsorbent- and plasmon-mediated catalytic reduction of CO<sub>2</sub>.

This work contributes to the development of a mechanistic and kinetic picture of illuminated nanoparticle systems, which has led to active debate on the role of hot carriers in observed reaction rates.<sup>84–87</sup> We note that the quantitative results are specific for the systems considered. However, the fundamental physics and general conclusions elucidated here remain translatable to other systems and co-adsorbed species, as recently shown for electron transfer from Ag and Cu clusters to CO molecules.<sup>42</sup> We have studied both CO and CO<sub>2</sub> as co-adsorbed species (Fig. S8, ESI†) and performed ground state calculations to investigate the hybridization between the Au cluster and the molecules. However, these species do not appear to hybridize – and thus promoting resonant charge transfer to CO<sub>2</sub> molecules – to the same extent as adsorbed OH groups. Importantly, our findings highlight the critical role of OH adsorbates in determining (i) the adsorption and activation of CO<sub>2</sub>, (ii) metal-molecule hybridization, (iii) importance of matching energy levels to efficiently transfer hot electrons to hybridized states. Consequently, the presence of OH species can promote the CO<sub>2</sub> reduction reaction to a different potential energy surface, possibly leading to different product distributions and reaction speeds. It is, therefore, very important to consider adsorbed species during the design of plasmonic catalysts to achieve high efficiency and selectivity for specific reactions, particularly when dealing with plasmonic structures functionalized with ligands.

## Conclusions

We investigated the mechanisms of plasmon-driven CO<sub>2</sub> reduction in the presence of a co-adsorbed hydroxyl group *via*

real-time time-dependent density functional theory. Our findings show that the OH group induces a redistribution of charges in CO<sub>2</sub>, leading to favourable absorption and activation on small Au clusters. Critically, adding an OH group significantly increased hybridization between molecular states of CO<sub>2</sub> and electronic states of Au as mediated by the OH group, enabling and improving charge transfer. This coupling is distance-dependent, reaching its maximum when CO<sub>2</sub> is close to the Au cluster, and this behavior can be predicted by analysing the branching of the partial density of states of the molecules.

By applying TDDFT, we are able to study the carrier dynamics of these systems at an atomic level, whereby the quantification of the hot electron (HE) energy distribution is critical. While the total HE distribution is largely unaffected by the presence of the OH group, the additional hybridized states facilitate the transfer of HEs to the CO<sub>2</sub> molecule, thereby increasing the resonant direct HE transfer by 11%. Furthermore, our findings confirm the importance of spectral overlap between the excitation energy and the plasmon resonance, and show that direct HE transfer is only slightly decreased (~1.05% to 0.84%) when the excitation is changed from a narrowband to a broadband. This finding shows that a significant chemical reaction rate still occurs under broadband excitation such as solar light. In addition, Landau dampening of the plasmon wave in the compound with the OH group may present another resonance mechanism on the time scale. This mechanism may further enhance HE transfer to CO<sub>2</sub> for specific reductive reactions, whereby the dynamic dipole moment of the OH group coupling the plasmon to a CO<sub>2</sub> molecule *via* HE transfer is a key property.

In summary, this investigation elucidates the role of adsorbed hydroxyl groups in the adsorption and activation of CO<sub>2</sub> on plasmonic surfaces. It highlights the importance of considering adsorbed molecules to enhance direct HE transfer, ultimately enabling the design of highly efficient and selective plasmonic photocatalysts with polychromatic excitation.

## Conflicts of interest

There are no conflicts to declare.

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