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Cavity Manipulation of Attosecond Charge Migration in Conjugated Dendrimers

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migration modes and the character of photoexcited wave function can be significantly influenced by the strong light-matter interaction in the cavity. This presents a new avenue for modulating initial ultrafast charge dynamics and subsequently controlling coherent energy transfer in dendritic nanostructures.

INTRODUCTION

Dendrimers are branched tree-like polymers,¹ which have drawn significant interest as photosensitizers² and photocatalysts,³ due to their distinct structure and versatile functionality.¹ The crux of a dendrimer's utility lies in its unique branched tree-like architecture, which shows a high degree of surface functionality coupled with internal pockets and enables them to engage with a diverse array of molecular entities effectively. In photodynamic therapy, for instance, dendrimers serve as vehicles to transport photosensitizers to specific body regions, offering a targeted approach to cancer treatment.⁴ Similarly, in photovoltaics, conjugated denstimers potentially enhance solar cell efficiency by facilitating improved light absorption and directed carrier transport utilizing antenna-like cascading principles.^{5–8}

diffraction. Our simulations demonstrate that the dendrimer charge

Extensive efforts have been dedicated to improving optoelectronic and light harvesting functions of dendrimers through the optimization of their topologies, structural rigidity, and electronic-vibrational (vibronic) couplings.^{9–11} For instance, initial investigations revealed the pivotal role of building block connectivity and topology in dendritic structures.¹² Here connectivity through a meta-position of a central benzene ring often constrains electronic communications leading to appearance of Frenkel-like excitonic states¹³ followed by a picosecond Förster resonance energy transfer (FRET). In contrast, para- and ortho- type linkages facilitate the formation of more delocalized electronic states approaching the Wannier exciton limit,¹³ potentially resulting in more

efficient energy transfer with coherent characteristics.^{14,15} It is established that photoexcited dynamics in dendrimers can be well controlled using pulse-shaping techniques.¹⁰ It is intriguing to explore the early stages of electronic dynamics that shape the initial photoexcited wavepacket in dendrimers.

Modern attosecond science makes it possible to probe charge migration, which involves ultrafast charge oscillations caused by the electronic coherence when a molecule is photoexcited or photoionized into a superposition of electronic states. These dynamics occur on time scales of a few femtoseconds when nuclei remain essentially frozen.^{16–20} Charge migration is the early stage of many charge and energy transfer processes and charge-induced reactions, playing an important role in the efficiency of these photophysical and photochemical processes.^{21–23} Investigating charge migration in dendrimers is of particular interest, given the complex interplay between their branched structure and electronic states.^{10–12}

Optical cavities provide potent means to manipulate charge migration in dendrimers. Molecules coupled to an optical cavity can interact with the vacuum field of the cavity photon

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mode even without external driving fields, forming hybrid light-matter states known as polaritons.^{24–31} Optical cavities can be characterized by the effective radiation matter coupling strength $\kappa = \mu \sqrt{\hbar \omega_c / 2 \varepsilon_0 V_c}$, where μ is the transition dipole moment, ε_0 is the vacuum electric permittivity, V_c is the cavity mode volume and ω_c is the cavity frequency. The cavity mode can be characterized by the ratio of the cavity frequency to the resonance width, known as the quality factor (*Q*). A larger *Q* indicates higher quality and a longer cavity photon lifetime. Theoretical and experimental studies have shown that photophysical and photochemical processes such as chemical reactivity, reaction rates and long-range energy transfer can be altered by creating polaritons.^{32–43}

Here, we investigate the effects of optical cavities on charge migration in conjugated dendrimer molecules that are photoexcited into a superposition of electronic states. Our target systems include meta-connectivity between dendritic branches that weakens the π electronic conjugation. In this case electronic states are spatially localized,¹³ which brings barriers to electron and hole migration between branches of the bare dendrimer. We find that interbranch exciton passage becomes possible by placing the dendrimer in an optical cavity that facilitates mixing of the electronic states and enables transitions between them. We further simulate the heterodynedetected time-resolved X-ray diffraction (TR-XRD) signal for monitoring this process.^{44,45} Polariton effects are clearly observed. These findings suggest a novel approach to control the initial charge dynamics in conjugated dendrimers, offering potential applications across photochemistry and photophysics. This could potentially enhance the light harvesting efficacy, elevate the performance of optoelectronic devices, and bolster sensor capabilities.

PROBING ELECTRONIC DYNAMICS BY TIME-RESOLVED X-RAY DIFFRACTION

Our molecules are described by the Hamiltonian $H_{\text{Mol}} = \sum_i \hbar \omega_i |i\rangle \langle i|$, where index *i* labels ground (i = 0) and relevant excited (i > 0) states. The molecule is then photoexcited into a superposition of electronic states. The excited state dynamics are computed through numerical solutions of the time dependent Schrödinger equation. The time-dependent electronic charge density is given by

$$\sigma_{\text{total}}^{E}(\boldsymbol{r}, t) = \sum_{i} \rho_{ii} \sigma_{ii}^{E}(\boldsymbol{r}) + \sum_{i \neq j} \rho_{ij}(t) \sigma_{ij}^{E}(\boldsymbol{r})$$
(1)

where \mathbf{r} is the spatial coordinate and $\sigma_{ii}^{E}(\mathbf{r})$ and $\sigma_{ij}^{E}(\mathbf{r})$ are the state and the transition charge density between electronic states *i* and *j*, respectively. In this study, the nuclei are assumed frozen over the attosecond charge migration time scale. The electronic populations ρ_{ii} become time-independent after the pump excitation, while the electronic coherences $\rho_{ij}(t)$ oscillate at frequency corresponding to the energy difference of states *i* and *j*.

The total Hamiltonian of the molecule coupled to a single cavity mode is given by

$$H = H_{\rm Mol} + H_{\rm Cav} + H_{\rm MolCav} \tag{2}$$

where the cavity Hamiltonian is $H_{\text{Cav}} = \hbar \omega_c \left(a^{\dagger} a + \frac{1}{2} \right)$ and *a* and a^{\dagger} are the cavity mode annihilation and creation operators, respectively. The interaction between the molecule and the

cavity is characterized within the electronic dipole approximation $H_{MolCav} = -\mu \cdot E_c(r)$, where

$$\mathbf{E}_{\mathbf{c}}(\mathbf{r}) = i\sqrt{\hbar\omega_{\mathbf{c}}/2\varepsilon_{0}V_{c}}ae^{i\mathbf{k}_{c}\cdot\mathbf{r}}\mathbf{e}_{\mathbf{c}} + H. \ c. \tag{3}$$

where k_c is the cavity-mode wave vector, e_c is the field polarization, and *H.c.* stands for the Hermitian conjugate. The cavity coupling strength is $g = \sqrt{\hbar \omega_c / 2 \varepsilon_0 V_c}$. The polariton states $\{|P_k\rangle\}$ are calculated by diagonalizing the polaritonic Hamiltonian using the direct product basis set $|i,n\rangle = |i\rangle \otimes |n\rangle_o$ where $\{|n\rangle_c\}$ is the Fock states of the cavities photons.

The attosecond charge migration is monitored by a TR-XRD movie. The heterodyne-detected attosecond X-ray diffraction signal is given by

$$S_{\rm het}(\boldsymbol{q}) \propto 2\Im \int \mathrm{d}t \boldsymbol{A}_{\rm het}(t) \boldsymbol{A}_{X}(t) \sigma(\boldsymbol{q}, t) \tag{4}$$

where $A_{het}(t)$ and $A_X(t)$ are envelopes of two coincident X-ray pulses with a controlled relative phase. The diffraction signal time resolution is therefore determined by the duration of the attosecond X-ray probe pulses. $\sigma(q,t)$ is the spatial Fourier transform of the charge density:

$$\sigma(\boldsymbol{q}, t) = \int \mathrm{d}\boldsymbol{r} e^{-i\boldsymbol{q}\cdot\boldsymbol{r}} \sigma(\boldsymbol{r}, t)$$
(5)

where q is the scattering vector. The charge migration process can be visualized as the time-dependent difference charge density, which is the difference between the time-dependent charge density of the excited molecule and the ground state $\sigma_{\text{diff}}^{E}(\mathbf{r}, t) = \sigma_{\text{total}}(\mathbf{r}, t) - \sigma_{\text{total}}^{g}(\mathbf{r})$. Thanks to the linear dependence of the heterodyne XRD signal on the charge densities, the corresponding difference signal can be obtained by $S_{\text{het}}(\sigma_{\text{diff}}^{E}) = S_{\text{het}}(\sigma_{\text{total}}) - S_{\text{het}}(\sigma_{\text{total}}^{g})$.

RESULTS AND DISCUSSION

We first consider the conjugated dendrimer $C_{30}H_{18}$ shown in Figure 1a. We adapt the electronic structure description provided by the NEXMD package as detailed in Method section. Charge migration is commonly triggered by ionizing the systems into cations. In this study, instead of photoionization, we photoexcite the molecule into a superposition of electronic states and investigate the following electron and hole dynamics. A linearly polarized Gaussian pulse $E(t) = A \overrightarrow{e_0} e^{-i\omega_0 t} e^{-(t^2/2\sigma^2)}$ with a peak amplitude $A = 3.60 \times$ 10^7 V/cm, $\sigma = 1.0$ fs duration, central frequency $\omega_0 = 3.30$ eV is used for initialization. These parameters were selected to ensure effective population of the relevant excited states (with largest absorption cross sections) while minimizing the ionization and other nonlinear effects that could complicate the desired charge migration dynamics. Specifically, the central frequency ω_0 was tuned to the average energy of the first and second excited states of the system ($e_1 = 3.23$ eV, $e_2 = 3.56$ eV), ensuring efficient coupling to these states. The pulse duration and amplitude were optimized to provide a balance between efficient excitation and minimal perturbation of the system while demonstrating maximal contrast between there and polaritonic states (see Figure S2). We include the nine relevant excited states and numerically solve the time dependent Schrödinger equation using the fourth-order Runge-Kutta method with 0.03 fs time step. The simulated 6 fs electronic population dynamics triggered by the 1.0 fs pump excitation is shown in Figure 1b. Owing their strong



Figure 1. (a) Structure of the dendrimer $C_{30}H_{18}$. (b) Population dynamics of $C_{30}H_{18}$ excited by pump pulse of energy $\omega_0 = 3.30$ eV and duration $\sigma = 1.0$ fs. (c) Transition density from ground state $|0\rangle$ to excited state $|1\rangle$. (d) Transition density from ground state $|0\rangle$ to excited state $|2\rangle$.

transition dipole moments, two excited states are significantly populated. $|1\rangle$ becomes dominant with a population of 0.8, whereas state $|2\rangle$ population reaches 0.1. To display the charge differences of these two states with respect to the ground state, the transition charge densities $\sigma_{ij}^{E}(\mathbf{r})$ between the ground state | 0 \rangle and $|1\rangle$ and $|2\rangle$ are calculated through $\langle 0|\hat{\sigma}|i\rangle$, where $\hat{\sigma}$ is the density operator. As shown in Figure 1c, $\sigma_{01}^{E}(\mathbf{r})$ spans the long dendrimer branch. In contrast, $\sigma_{02}^{E}(\mathbf{r})$ is localized at the short branch as illustrated in Figure 1d. Different charge migration modes involving states $|1\rangle$ and $|2\rangle$ lead to distinct charge dynamics along different branches.

We have first simulated the charge dynamics of the bare dendrimer molecule and visualized it by the time-dependent difference charge density. Figure 2a-f shows the difference charge densities at selected times t. The charge only migrates along the long branch and there is no significant migration to the short branch. This is consistent with previous studies, that showed that meta-substitution can block interbranch electronic communications.¹³ Figure 2m-r shows the difference charge densities of the molecule in an optical cavity with coupling strength g = 0.103 V/Å and $\omega_c = 3.559 \text{ eV}$ at selected times t. In contrast to the bare molecule, the charge dynamics of the short branch is now significant. Charge migration enabled by polaritonic effects exhibits long-range character spanning the entire molecule. We systematically varied the coupling strength to explore its modification of the charge migration pathways, which change gradually with the coupling strength. As shown in Figure 2g-l, with g = 0.0514 V/Å, the charge migration

process does not significantly vary by the cavity (Figure 2a-f). As the coupling strength is increased, remarkable changes in the charge migration pathway emerge with the short branch of the dendrimer activated. (Figure 2m-x). Note that the coupling strengths employed here are about 1%-3% of the cavity frequency, which are experimentally feasible. Recent publications have reported the realization of single-molecule strong coupling using plasmonic nanocavities with coupling strengths that are commensurate with those we used in our simulations.⁴⁶ Plasmonic cavities often exhibit high losses and broad line widths. Their Q factors typically range between 5 and 20.47 To further explore the loss rate influence of optical cavities on the attosecond charge migration, charge dynamics of cavity modified dendrimers with different Q factors were simulated and presented in Figure S9. We emphasize that the charge migration process is not disturbed even with a Q factor lower to 10 due to the ultrafast time scale of pure electronic dynamics. Another factor that may influence the charge migration is the coupling of high photon cavity modes. Simulations of the charge dynamics of photoexcited dendrimer with the two-photon mode included in our Hamiltonian are presented in Figure S10. The charge dynamics is identical to those of the single-photon mode calculations. We demonstrate that these effects can be neglected in our simulations because of the off resonance of the high photon modes with the dendrimer electronic states.

To further interpret these findings, we have calculated the population dynamics of the electronic states for different coupling strengths. As shown in Figure 3a, upon pump excitation, the populations of all the states remain constant in the bare molecule. Since the nuclei are assumed frozen, charge dynamics shown in Figure 2a-f only arises from the electronic coherence. In an optical cavity, all polariton states mix. The original electronic coherence turns into a polaritonic coherence. Subsequently, the electronic populations become time-dependent and new electronic states may become active. As the coupling strength increases, the populations of states | 0, $|1\rangle$ and $|2\rangle$ start to significantly oscillate in time (Figure 3bd). Some inactive states such as $|7\rangle$ and $|8\rangle$ also show up in the charge dynamics bringing significant modifications to the corresponding charge migration processes. Separate contributions of each state are provided in Figure S3. The contribution of population dynamics to charge migration is a new feature induced by the optical cavity and does not exist in the bare molecule. The polaritonic charge dynamics arises from both the cavity modifications to electronic coherence and population dynamics.

Figure 4 depicts the TR-XRD signals of the bare and the cavity-embedded dendrimer with a g = 0.103 V/Å coupling strength. The signals are similar at short times (Figure 4a,b and g,h). Notable differences between the bare and cavity-embedded molecules emerge after 5.2 fs. This is consistent with the time-resolved difference charge densities shown in Figure 2. New charge migration features result in new strong signals (Figure 4j,k) that are absent in the bare molecule (Figure 4d,e). Cavity modification of the charge dynamics can thus be clearly imaged by TR-XRD signals.

To explore the generality of cavity manipulation of charge migration, we have applied the same approach to a larger dendrimer $C_{54}H_{30}$ with three branches (see Figure S4). A 3 to 8 fs time scale is selected for both the $C_{30}H_{18}$ (two-branched) and $C_{54}H_{30}$ (three-branched) dendrimers according to the ultrafast nature of the attosecond charge migration. The pure

-_y(Å)

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Figure 2. (a–f) Time-dependent charge density difference of excited bare dendrimer $C_{30}H_{18}$. Polaritonic time-dependent charge density differences of this dendrimer coupled with cavity with coupling strengths g = 0.0514 V/Å (g–l), 0.103 V/Å (m–r), and 0.154 V/Å (s–x). The cavity frequency $\omega_c = 3.559 \ eV$. All density differences are integrated in the *z*-axis direction.

electronic charge dynamics should disappear at longer time scales due to the participation of nuclear dynamics. Figure 5a-d shows the difference charge densities of the bare dendrimer at selected times *t*. Charge migration mainly occurs between two branches and there is no significant charge migration into the third one. Figure 5e-h shows the difference charge densities of $C_{54}H_{30}$ coupled to an optical cavity with coupling strength $g = 0.0514 \ V/\text{\AA}$ and $\omega_c = 3.559 \ \text{eV}$ at selected times *t*. The polaritonic charge dynamics is significantly modulated

compared to Figure 5a-d. Global charge migration mode involving all branches is now activated. Figure 6 compares the TR-XRD signals of the bare and the cavity-embedded $C_{54}H_{30}$ dendrimer. The differences between the charge dynamics can be clearly tracked in the momentum space.

Our results demonstrate that polaritons in optical cavities can activate the excited-state charge dynamics in dendritic structures across meta-substitutions. Many theoretical and experimental studies have shown that photoinduced dynamics



Figure 3. Population dynamics with various coupling strengths. The cavity frequency of (a-d) is 3.559 *eV*, and the coupling strengths are 0, 0.0514, 0.103, 0.154 *V*/Å, respectively.

of conjugated dendrimers are localized within segments connected by benzene rings substituted at the metaposition.^{11,13,48} Ultrafast absorption also shows that metasubstitutions of disubstituted benzene exhibit the most hindered excited-state charge dynamics, compared to orthoand para-substitutions.⁴⁹ As shown in Figure 2a–f and 5a–d, our dynamical simulations of charge evolution in two bare interstitial dendrimer molecules agree with these findings. However, in an optical cavity, the suppressed long-range electronic delocalization and charge migration can be activated. Recent study implies that ultrafast charge dynamics does not show effects extended over many molecules.⁴³ Thus, cavity modification of molecular charge migration cannot be enhanced by collective strong coupling. In this study, we show that the charge migration processes in conjugated dendrimers can be significantly modified with a relatively weak cavity coupling strength (2% of the cavity frequency), which is achievable by a single molecule strong coupling with no cooperativity. Polaritons in optical cavities thus have the potential to alter the charge dynamics in a broad range of conjugated species.

CONCLUSIONS

Our computational study explores the initial stages of coherent electronic dynamics in conjugated dendrimers subjected to polaritonic effects. We have shown that attosecond charge migration in molecules can be significantly manipulated by coupling to optical cavities. While meta-linkages suppress spatial electronic delocalization and charge migration mode in a bare molecule, our polaritonic simulations of charge dynamics in dendrimers demonstrate that this limitation is alleviated and charge migration can be significantly amplified. This important observation is substantiated by population analysis of the electronic states to explicitly show how the charge migration process is affected by light-matter coupling. Representative applications to two dendrimer molecules with two and three branch geometries emphasize generality of the phenomenon. Simulating time-resolved X-ray diffraction signals to monitor the modulated ultrafast charge dynamics in momentum space provide direct link to potential attosecond experiments. Our modeling suggests to potential novel use of optical cavities to control molecular photochemistry and photophysics by manipulating the initial coherent charge





Figure 4. (a–f) Time-reolved XRD of bare excited dendrimer $C_{30}H_{18}$. (g–l) Time-resolved XRD of this dendrimer embedded in a cavity, with frequency $\omega_c = 3.559 \text{ eV}$, the coupling strength g = 0.103 V/Å. The plane of molecule $C_{30}H_{18}$ is vertical to the *z*-axis.

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Figure 5. (a–d) Time-dependent charge density difference of excited dendrimer integrated in the *z*-axis direction $C_{54}H_{30}$. (e–h) Polaritoncalculated time-dependent charge density difference of dendrimer ($C_{54}H_{30}$) coupled with cavity integrated in the *z*-axis direction, and the cavity frequency $\omega_c = 3.559 eV$, the coupling strength g = 0.0514 V/Å.





Figure 6. (a–d) Time-resolved XRD of bare excited dendrimer $C_{54}H_{30}$. (e–h) Time-resolved XRD of this dendrimer embedded in a cavity, with frequency $\omega_c = 3.559 eV$, the coupling strength g = 0.0514 V/Å. The plane of molecule $C_{54}H_{30}$ is vertical to the *z*-axis.

dynamics, which can be probed by contemporary attosecond spectroscopy techniques.

METHODS

Semiempirical Approach and Dendrimer Optimization. The electronic structure calculations in this work are based in the Collective Oscillator Approach⁵⁰ interfaced with the Austin Model 1 semiempirical approach⁵¹ as implemented in the NEXMD code.^{52,53} The analytical gradients⁵⁴ implemented in NEXMD were used to optimize the dendrimer structures in the ground state without the optical cavity interaction. The electronic structure level of theory implemented in NEXMD has shown to describe efficiently and accurately a wide range of conjugated molecules^{55–57} and in particular dendrimers and dendrimer building blocks.^{58–64}

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.4c06727.

Transition densities and population dynamics of $C_{30}H_{18}$ and $C_{54}H_{30}$, charge dynamics with different dissipation of $C_{30}H_{18}$, and charge dynamics of $C_{30}H_{18}$ embedded with single/two-mode cavities (PDF)

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Notes

The authors declare no competing financial interest.

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Supporting Information of Cavity Manipulation of Attosecond Charge Migration in Conjugated Dendrimers

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Figure S1 provide the transition densities $\langle 0 | \hat{\sigma} | i \rangle$ between the ground state $|0\rangle$ and various excited states $|i\rangle$ of the bare molecule, states reflecting the density differences of those states from ground state. The energies of the excited states are $e_1 = 3.23$ eV, $e_2 = 3.56$ eV, $e_3 = 3.67$ eV, $e_4 = 3.72$ eV, $e_5 = 3.84$ eV, $e_6 = 3.84$ eV, $e_7 = 3085$ eV, $e_8 = 4.08$ eV, and $e_9 = 4.14$ eV, respectively.



Figure S1: Transition densities between the ground state $|0\rangle$ and various excited states of $C_{30}H_{18}$. States $|1\rangle$ through $|9\rangle$ correspond to the lowest nine excited states, respectively. $\hat{\sigma}$ is the density operator.



Figure S2: (a-c) Population dynamics excited by pump pulse of energy $\omega_0 = 3.30$ eV, duration $\sigma = 0.5$ fs, 1.0 fs, 1.5 fs, and amplitude $A = 2.60 \times 10^7$ V/cm. Same for the amplitude 3.60×10^7 V/cm. (g-i) Same for the amplitude 4.60×10^7 V/cm.



Figure S3: Separate contributions of population dynamics of all relevant excited states to the charge dynamics in Figure 3(m-r).



Figure S4: Geometric structure of $C_{54}H_{30}$

Figure S5 provides the transition densities $\langle 0 | \hat{\sigma} | i \rangle$ between the ground state $| 0 \rangle$ and various excited states $| i \rangle$ of the bare molecule.



Figure S5: Transition densities between the ground state and various excited states of $C_{54}H_{30}$. $|0\rangle$ represents the ground state. States $|1\rangle$ through $|9\rangle$ correspond to lowest nine excited states, respectively. $\hat{\sigma}$ is the density operator.



Figure S6: Population dynamics of $C_{54}H_{30}$ excited by a pump pulse with frequency $\omega_0 = 3.56 \ eV$ and duration $\sigma = 1.5 \ fs$.



Figure S7: Population dynamics with coupling strength of $C_{54}H_{30}$. The cavity frequency of (a-d) is 3.559 eV, and the coupling strengths are 0, 0.0514, 0.103, 0.154 V/Å, respectively.



Figure S8: Separate contributions of population dynamics of all relevant excited states to the charge dynamics in Figure 5(e-h).



Figure S9: (a-f) Polaritonic time-dependent charge density difference of $C_{30}H_{18}$ coupled with cavity integrated in the z-axis direction, and the cavity frequency $\omega_c = 3.559 \ eV$, the coupling strength $g = 0.103 \ V/Å$, the quality factor Q = 1000. (g-l) Same for the quality factor Q = 100. (m-r) Same for the quality factor Q = 10.



Figure S10: (a-f) Polaritonic time-dependent charge density difference of this dendrimer coupled with single-photon mode of cavity integrated in the z-axis direction, and the cavity frequency $\omega_c = 3.559 \ eV$, the coupling strength $g = 0.103 \ V/\text{\AA}$. (g-l) Same for the two-photon mode of cavity included.