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Automatic discovery of photoisomerization mechanisms with nanosecond machine learning photodynamics simulations†

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Photochemical reactions are widely used by academic and industrial researchers to construct complex molecular architectures *via* mechanisms that often require harsh reaction conditions. Photodynamics simulations provide time-resolved snapshots of molecular excited-state structures required to understand and predict reactivities and chemoselectivities. Molecular excited-states are often nearly degenerate and require computationally intensive multiconfigurational quantum mechanical methods, especially at conical intersections. Non-adiabatic molecular dynamics require thousands of these computations per trajectory, which limits simulations to \sim 1 picosecond for most organic photochemical reactions. Westermayr *et al.* recently introduced a neural-network-based method to accelerate the predictions of electronic properties and pushed the simulation limit to 1 ns for the model system, methylenimmonium cation (CH_2NH_2^+). We have adapted this methodology to develop the Python-based, Python Rapid Artificial Intelligence *Ab Initio* Molecular Dynamics (PyRAI²MD) software for the *cis-trans* isomerization of *trans*-hexafluoro-2-butene and the 4π -electrocyclic ring-closing of a norbornyl hexacyclodiene. We performed a 10 ns simulation for *trans*-hexafluoro-2-butene in just 2 days. The same simulation would take approximately 58 years with traditional multiconfigurational photodynamics simulations. We generated training data by combining Wigner sampling, geometrical interpolations, and short-time quantum chemical trajectories to adaptively sample sparse data regions along reaction coordinates. The final data set of the *cis-trans* isomerization and the 4π -electrocyclic ring-closing model has 6207 and 6267 data points, respectively. The training errors in energy using feedforward neural networks achieved chemical accuracy (0.023–0.032 eV). The neural network photodynamics simulations of *trans*-hexafluoro-2-butene agree with the quantum chemical calculations showing the formation of the *cis*-product and reactive carbene intermediate. The neural network trajectories of the norbornyl cyclohexadiene corroborate the low-yielding *syn*-product, which was absent in the quantum chemical trajectories, and revealed subsequent thermal reactions in 1 ns.

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1. Introduction

Photochemistry provides access to highly strained molecular architectures,¹ photoswitches,² organic photovoltaics,³ and solar fuel materials,^{4,5} which are characterized by mild conditions and high atom economy. Photochemical reactions typically consist of a series of molecular transformations that occur in excited molecules after light absorption. Unlike ground state processes, where spectroscopy and crystallography reveal structural information, light-promoted excited-state reactions involve short-lived femto-to picosecond (10^{-15} to 10^{-12} s) molecular excited states and reactive intermediates. This ultrafast processes involve relaxation to the excited-state minima for fluorescence or non-radiative transition to the ground-state through a state crossing point or seam, which plays essential roles in the chemoselectivity of a photochemical reaction.^{6,7}



Unravelling the origin of chemoselectivity and stereoselectivity in organic photochemical reactions is challenging because of the short-lived molecular excited states. Quantum chemical calculations offer insight into the bonding changes that occur along a reaction coordinate and non-adiabatic molecular dynamics (NAMD) simulations to gain mechanistic insights and develop structure–reactivity relationships in complex photochemical reactions. The nuclei-electron coupling and time-dependent terms increase the complexity of Hamiltonian in NAMD and, thus, computation time. Multiple methods developed in the last two decades simplified the time-dependent molecular wave functions, *e.g.* *ab initio* multiple spawning (AIMS),^{8,9} and fewest switches surface hopping (FSSH).^{10–12} However, computing high dimensional PESs with the requisite multiconfigurational methods is extremely resource-intensive. For example, most quantum chemical software packages encode an upper limit to active space of 16 electrons and 16 orbitals for complete active space self-consistent field (CASSCF) calculations. A typical NAMD experiment requires thousands of such calculations, resulting in a maximum simulation time on the order of 1 picosecond, at a computational cost of approximately 10^1 – 10^4 wall-clock hours. Unfortunately, many direct excitation photochemical reactions have low quantum yields and long excited-state lifetimes. This results in prohibitively expensive NAMD simulations, thus limiting broad understanding of excited-state structure–reactivity relationships in photochemistry.

An increasing number of studies reported that fitting machine learning (ML) potentials could substantially accelerate the NAMD simulation.^{13–15} Aleotti *et al.* have parameterized *ad hoc* force fields for a 10 ps dynamic simulation of azobenzene.¹⁶ Westermeyr *et al.* have trained multilayer feedforward neural networks (NNs) to enable 1 ns simulation of methyl-enimmonium cation (CH_2NH_2^+) in 59 days.¹⁷ Later, they have extended the application of the NN model in predicting excited-states electronic properties for other small molecules, such as SO_2 and CSH_2 employing a deep continuous-filter convolutional-layer neural network, SchNet, combined with SHARC.¹⁸ Recently, Ha *et al.* have trained the SchNet model to study the excited-state dynamics of penta-2,4-dieniminium cation.¹⁹ Applying ML-based NAMD on complex molecules continues to challenge theorists because of additional conformational flexibility and increase in available degrees of freedom. Training ML models becomes increasingly difficult with the rapidly growing required size of training datasets, especially when based on atom-wise molecular representations.

Here, we address the need for our combined ML and NAMD approach (ML-NAMD) that expands the scope of organic photochemical reactions. We demonstrate the high accuracy in the trained NN and low computational cost in the ML-NAMD simulations. Our trajectory statistics support that the ML-NAMD predicted photochemical product distribution is in good agreement with quantum chemical results. Our initial ML-NAMD simulations focused on the first such photodynamics of hexafluoro-2-butene (**1**), which is mechanistically well understood due to its characteristic vertical $\pi\pi^*$ excitation from HOMO to LUMO (the π and π^* -orbital). Hexafluoro-2-butene is

a nontoxic and not flammable industrial working fluid used as a refrigerant and a foam-blown agent.²⁰ We also applied the ML-NAMD simulations to the 4π -electrocyclic ring-closing of norbornyl cyclohexadiene (**3**), a previously unknown and complex photochemical system.

2. Computational methods

2.1 Quantum chemical calculations

We performed quantum chemical calculations with OpenMolcas 19.11²¹ to generate reference and training data. We chose the less resource-intensive CASSCF method because it produced consistent potential energy surfaces with CASPT2 calculations, tested by the interpolated reaction coordinate diagrams of the model reactions (Fig. S11 and S17†). The converged orbitals and optimized geometries are available in the ESI.†

In the *cis*–*trans* isomerization of **trans-1**, we selected an active space of 2 π -electrons and 2 π -type orbitals (*i.e.* (2,2)) for the C=C bond. The geometries of **trans-1** and **cis-1** were optimized with the cc-pVDZ basis set.²² A vibrational analysis confirmed only positive frequencies. We located two minimum energy crossing points (MECPs) corresponding to the *trans* \rightarrow *cis* (**MECP-trans-1**) and the reverse reaction (**MECP-cis-1**). The NAMD simulations with CASSCF(2,2)/cc-pVDZ used the fewest switches surface hopping algorithm (FSSH)^{10–12} in Hammes-Schiffer/Tully (HST) scheme¹⁰ with decoherence correction of 0.1 Hartree²³ implemented in OpenMolcas 19.11.²¹ We generated 1500 initial geometries and associated velocities near the equilibrium geometry of **trans-1** with Wigner sampling at 300 K. The NAMD trajectories were propagated at 300 K (Nosé–Hoover thermostat²⁴) from the S_1 Franck–Condon region for \sim 500 fs with 0.5 fs timesteps. It is important to note that applying a thermostat can bias the excited-state dynamics because the excited-state lifetime of a few hundred femtoseconds often does not suffice for thermalization. We used a thermostat to reduce the kinetic energy gained from surface hopping, which compensates for the overestimated excitation energy of CASSCF calculation. We collected 1371 of the 1500 trajectories that reached the ground-state within 500 fs, whereas the others remained in the excited-state or showed molecular structures with unexpected broken bonds resulting from incorrectly converged CASSCF calculations.

The 4π -electrocyclic ring-closing of **3** used 4 π -electrons and 3 π -type orbitals active space (*i.e.*, (4,3)) for the conjugated bonds. As Martinez and co-workers suggested in a previous study on cyclohexadiene,²⁵ we removed the π^* -orbital in A_2 symmetry to ensure consistent CASSCF state ordering with CASPT2 calculations. The geometries of **3** and possible ladderene products were optimized with the ANO-S-VDZP basis set.^{26–29} Frequency calculations showed all positive values. We set up the same NAMD simulations for **3** with CASSCF(4,3)/ANO-S-VDZP while the simulations are done in 1000 fs. We only propagated 250 trajectories at 300 K because the NAMD simulation time increased significantly to 17 days. 240 of the 250 trajectories were completed at the ground-state and were used for analysis.



2.2 Machine learning model

We have implemented multilayer feedforward NN as the primary ML model in our ML-NAMD approach. Our NN was inspired by those used in SchNarc;¹⁸ energies and forces are trained together with a combined loss function. Non-adiabatic couplings (NACs) are computed by the difference between NN predicted state energies and interstate couplings, which has shown to be more suitable for NN training.^{30,31} The forces and interstate couplings are trained as the derivatives of the energies and a virtual potential, respectively.¹⁸ We used an atom-specific virtual potential for more accurate NACs predictions. The phase of NACs is internally corrected with a phase-less loss function.¹⁸ We include a detailed discussion on force and NAC prediction in ESI.† The NN receives an inverse distance-based^{17,18} feature representation. **1** has 12 atoms ($N = 12$), which leads to 66 unique entries in the inverse distance matrix. The NN predicts two energy values (ground- and excited-state; $k = 2$) and use their derivatives with respect to the input coordinates to compute the $k \times 3N = 72$ force components. $3N = 36$ NAC components are simultaneously predicted. For **3** ($N = 25$), the input feature size is 300, with 2 energies, 150 force, and 75 NAC components. We chose a leaky soft plus activation function for hidden layers. We have optimized the hyperparameters to predict energies, forces, and NACs with a grid-search over 864 NNs. The training was done using the Adam optimizer³² with a stepwise decrease of the initial learning rate from 10^{-3} to 10^{-5} on validation error plateaus (energies and forces: 2700 epochs; NACs: 1600 epochs). To evaluate the prediction uncertainty in ML-NAMD simulation, we used the standard deviation of an ensemble of NNs. We picked two sets of the most accurate and efficient NNs (each set has one NN for energies and forces together, and another one for NACs) with distinct architectures. We implemented all NNs using the TensorFlow/Keras (v2.3) packages³³ for Python. Information about grid search and NN training are available in ESI.†

2.3 Initial training set generation

Accurate NN predictions require robust training data; thus, the training data must include a broad sampling of essential and relevant conformations of the target molecule. Westermayr *et al.* recently reported a comprehensive review including the initial training set generation techniques that sample the conformations of small molecules from cost-efficient ground- and excited-state molecular dynamics, Wigner sampling, normal mode scan, and optimized critical geometries.^{34,35} None of these individual techniques can adequately sample the relevant potential energy surfaces of a photochemical reaction as the molecular structure become more complex and the degrees of freedom increase. For example, Wigner sampling is limited to the configurations immediately related to equilibrium geometries. Conversely, a series of multiconfigurational NAMD is prohibitively expensive for training. In this work, we invoke a composite scheme that generates compact yet relevant training data for ML-accelerated excited-state dynamics (Fig. 1).

Our initial training set generation scheme combines *Wigner sampling*, *geometrical interpolation*, and *trajectories*. The *Wigner*

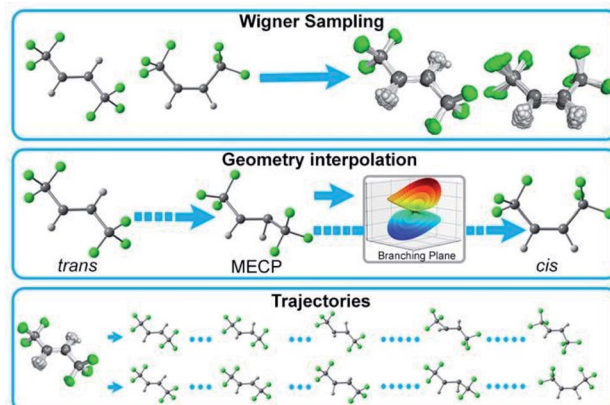


Fig. 1 Three initial training set generation approaches in PyRAI²MD. *Wigner sampling* explores the conformational space near the equilibrium geometries. *Geometry interpolation* collects the data along with reaction coordinates from reactant to product. *Trajectories* samples the data from quantum chemical trajectories. The 2D PES data can also supplement the data sampling. Detailed information about the initial training set generation is available in the ESI.†

sampling approach generates training data by sampling reactant and product structures. The *geometry interpolation* approach accesses the reaction coordinate diagram with the optimized reactant, product, and minimum energy crossing point (MECP) geometries. It systematically varies the reactant geometry to that of the minimum energy crossing point and then to that of the product in equal increments of Z-matrix coordinate parameters. We expanded the range of sampled structures in the training data by applying the Wigner sampled geometrical perturbations to the interpolated reaction coordinate diagram. The *trajectories* approach samples conformations from short-time quantum chemical NAMD trajectories (50–100 fs). For instance, we included every 10th snapshot of the first 50 fs of 132 CASSCF NAMD trajectories of **1** corresponding to the *trans* → *cis* isomerization.

2.4 Adaptive sampling

The initial training set represents a chemically intuitive conformational space to fit the NN potential for the investigated photochemical reactions. The NN potential becomes less reliable when trajectories frequently encounter the structures outside the data set. We efficiently expanded the initial training set to cover important structures on the PESs using an active learning approach, adaptive sampling. It was first introduced by Behler *et al.*³⁶ then adapted to ground- and excited-state dynamics by Marquetand and co-workers.^{17,37} The adaptive sampling propagates ML-NAMD trajectories to explore the conformational space and sample the uncertain geometries based on the standard deviation between two sets of NNs. The collected geometries are recomputed with quantum chemical calculations and then join the initial training set to improve the NN potentials iteratively. In the *cis*–*trans* isomerization of **trans-1**, the adaptive sampling included 250 ML-NAMD trajectories in 500 fs with a 0.5 fs timestep at 300 K. The adaptive sampling



stopped at 28 iterations with 6207 data points (98% of the trajectories completed without finding new geometries). In the 4π -electrocyclic ring-closing of **3**, the adaptive sampling ran 250 ML-NAMD trajectories in 1000 fs with a 0.5 fs timestep. We increased the temperature to 1200 K to access high energy non-equilibrium geometries along the reaction coordinates. The temperature was reset to 300 K in the production ML-NAMD simulations. We obtained a final set of 6272 data points after 100 iterations (96% of the trajectories completed normally). Detailed information about the adaptive sampling is available in the ESI.[†]

3. PyRAI²MD

We developed the Python Rapid Artificial Intelligence *Ab Initio* Molecular Dynamics program (PyRAI²MD) to integrate ML models and the NAMD algorithms. It is an open-source code to enable ML-NAMD simulations for unprecedented molecular complexity and timescales. Fig. 2 illustrates the computational architecture of PyRAI²MD, which requires ML and NAMD kernels.

The workflow in the NAMD kernel starts with the Wigner sampled initial conditions. The following procedures, red arrows in Fig. 2, iteratively compute the energies, forces, NACs, and surface hopping probability, propagating the NAMD trajectory. We generalized the input and output format of computing energies, forces, and NACs in the NAMD kernel. This feature enables efficient communication between the ML kernel and our chosen external quantum chemical program, OpenMolcas 19.11.²¹ The workflow in ML kernel reads a training set or an existing model. It provides a convenient interface to train models and make predictions marked by blue arrows in Fig. 2.

The green arrows in Fig. 2 illustrate the adaptive sampling workflow incorporating the NAMD and ML kernels. It first loads two trained ML models to run multiple trajectories in the NAMD kernel. The trajectories explore the unsampled conformational space while the two ML models evaluate the prediction uncertainty on-the-fly, as described in Section 2.4 and ESI.[†] The sampled data were added into the initial set to train a new model.

4. Results and discussion

4.1 *Cis-trans* isomerization model for *trans*-hexafluoro-2-butene

As the first demonstration of our ML-NAMD approach, we chose an emblematic photochemical reaction, an isomerization of *trans*-hexafluoro-2-butene, **trans-1**. We have characterized the photodynamics of **trans-1** with 1371 CASSCF(2,2)/cc-pVQZ trajectories. 946 trajectories correspond to the *cis-trans* isomerization, whereas 425 trajectories undergo an intramolecular hydrogen abstraction to form hexafluoro-2-butene carbene, **2** shown in Scheme 1.

This pathway is consistent with prior theoretical studies on ethylene intramolecular hydrogen migration reactions.³⁸⁻⁴⁰ To test whether the NN-driven adaptive sampling method can discover the missing data for the alternative carbene pathway without prior knowledge or human bias, the initial training set only included the information about the *trans* \rightarrow *cis* isomerization of **trans-1**.

4.2 ML potential for the *cis-trans* isomerization of *trans*-hexafluoro-2-butene

We trained four NNs for the *cis-trans* isomerization of **trans-1** with PyRAI²MD. On a single CPU, the predictions of the energies, forces, and NACs take 0.00991 seconds, whereas the equivalent computations at CASSCF(2,2)/cc-pVQZ require 336 seconds. These results represent an NN acceleration of 3.4×10^4 -fold relative to ground-truth CASSCF calculations. We summarized the NN architecture, mean absolute error (MAE), and coefficient of determination (R^2) of the predicted energies, forces, and NACs in Table 1.

The MAE of predicted energies is 0.023–0.025 that meets the ‘chemical accuracy’ threshold (1 kcal mol⁻¹ = 0.043 eV). The R^2 values are 0.9984–0.9989 in NN1 and NN2, suggesting an almost linear correlation between the NN prediction and QC reference. The MAE and R^2 in force calculation are 0.14–0.15 eV Å⁻¹ and 0.9357–0.9536. These values are consistent with previous reports on SO₂ and CH₂NH₂⁺ photochemistry.¹⁸ The MAE and R^2 in the interstate coupling term of the NACs are 0.170–0.182 eV

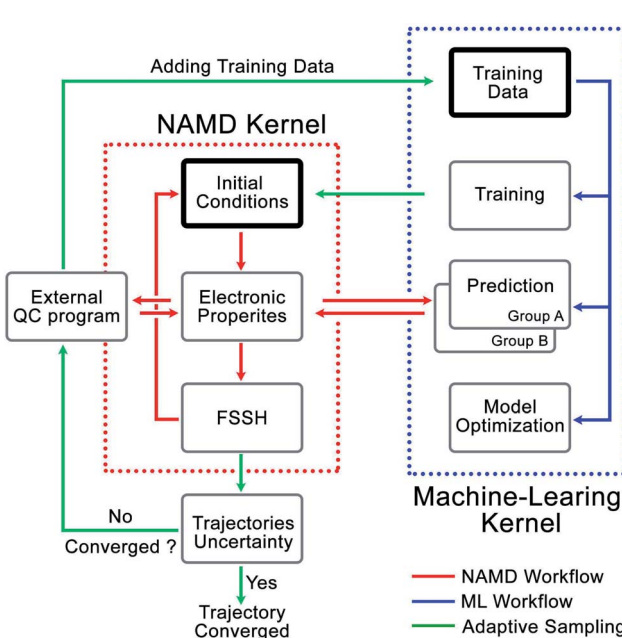
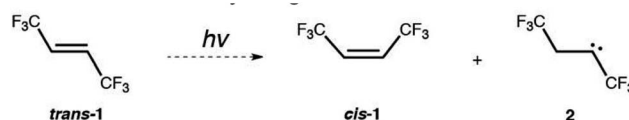


Fig. 2 The computational architecture of PyRAI²MD. The bold boxes are the initial stage in NAMD and ML kernel.



Scheme 1 *Cis-trans* isomerization and hydrogen abstraction of hexafluoro-2-butene.



Table 1 Selected NN hyperparameters and the mean absolute errors with R^2 of predicted energies (eV), forces (eV Å $^{-1}$), and interstate coupling term of NACs (eV Å $^{-1}$) trained on 6207 data points of 1. Additional details on hyperparameters and training statistics are available in ESI

Hyperparameters	Energies, forces		NACs ^a	
	NN1	NN2	NN1	NN2
Hidden layers	3	5	5	3
Neurons/layer	400	300	300	600
Batch size	64	128	128	128
MAE	0.023(0.14) ^b	0.025(0.15) ^b	0.182	0.170
R^2	0.9989(0.9357) ^b	0.9984(0.9536) ^b	0.7137	0.7423

^a Here the NAC only represents the interstate coupling term. ^b The MAE and R^2 of forces are shown in the parenthesis.

Å $^{-1}$ and 0.7137–0.7423, respectively. The low R^2 relative to energies and forces suggests possible overfitting of NACs (the training MAE are 0.014–0.015 eV Å $^{-1}$ and the training R^2 are 0.9969–0.9970) due to the indefinite term of the antiderivative of NACs and unavailable training data for it.

4.3 ML-NAMD simulations for the *cis*–*trans* isomerization of *trans*-hexafluoro-2-butene

Because the predicted interstate terms of NACs have relatively larger errors than energies and forces, the behavior of NN surface hopping events can be very different from them in QC trajectories. We compared the FSSH and Zhu–Nakamura theory of surface hopping (ZNSH)^{41–45} to determine the NACs influence on the NN trajectories. The ZNSH method is independent on NACs that has been successfully applied to excited-state dynamics of other systems^{46,47} and used to train machine learning trajectories.^{13,14} Propagating a 500 fs NN FSSH trajectory only needs 23 seconds in contrast to 33 hours of running an equivalent trajectory with CASSCF(2,2)/cc-pVDZ. The ZNSH method can further reduce the cost of NN trajectory to 13 seconds because it only computes surface hopping probability near crossing regions (S₁/S₀ gap < 0.5 eV). We have collected 5573 and 5820 NN trajectories with FSSH and ZNSH methods,

respectively. Fig. 3a shows the state population dynamics of *trans*-1 in 500 fs.

The state populations of QC trajectories (Fig. 3a) suggest the S₁ half-life is 33.5 fs. The NN FSSH trajectories depending on predicted NACs show rather slower decay of the S₁ state. The S₁ half-life is 71.0 fs and the MAE in population between the NN FSSH and QC trajectories is 0.105. The NN ZNSH trajectories using NN energies and forces predict that the S₁ half-life is 29.0 fs with the MAE in population of 2.5%. The good agreement between the NN ZNSH and QC trajectories indicates the NN energies and forces are sufficiently accurate to detect surface hopping events. The overestimated S₁ lifetime in NN FSSH trajectories is attributed to the errors in the NAC model. It seems to underestimate the surface hopping probability, which keeps trajectories at S₁ in a longer time until it finds a smaller energy gap for a hop to S₀. The S₁/S₀ gap is similar for QC FSSH and the NNs ZNSH (0.51 eV and 0.46 eV, respectively) while the NN FSSH is 0.12 eV. Due to the closer match, we focused on the ZNSH trajectories in the subsequent discussions.

We tested the limits of our newly established ML-NAMD model by running unprecedented 10 ns simulations with a 0.5 fs time step. We obtained 89 NN ZNSH trajectories and the average state populations are shown in Fig. 3b. The state populations in the first 500 fs also include the 5820 trajectories mentioned above. The non-radiative S₁/S₀ relaxation completes within 1 ps. The flat population curve after 10³ fs indicates all trajectories stayed in S₀ up to 10 ns, without surface hopping up to the S₁ state. The total 2 × 10⁷ iterations were accomplished in an average of 50 hours on a single CPU. The 10 ns simulation with CASSCF(2,2)/cc-pVDZ would otherwise require approximately 58 years of wall time. We have included the ML-NAMD trajectory video in ESI.†

To quantify the difference between the photochemical reaction pathways enumerated by the QC and NN trajectories, we plot the trajectories corresponding to the *trans* → *cis* isomerization of *trans*-1 and their energies with respect to the \angle H-C-C-H and \angle C-C-C-C dihedral angles in Fig. 4.

Fig. 4a illustrates the trace of the QC trajectories that bifurcates toward *cis*-1 and *trans*-1. All trajectories started with a narrow spreading in the range of \angle C-C-C-C = 150–180° and \angle H-C-C-H = 0–60° (upper right corner). 248 surface hopping

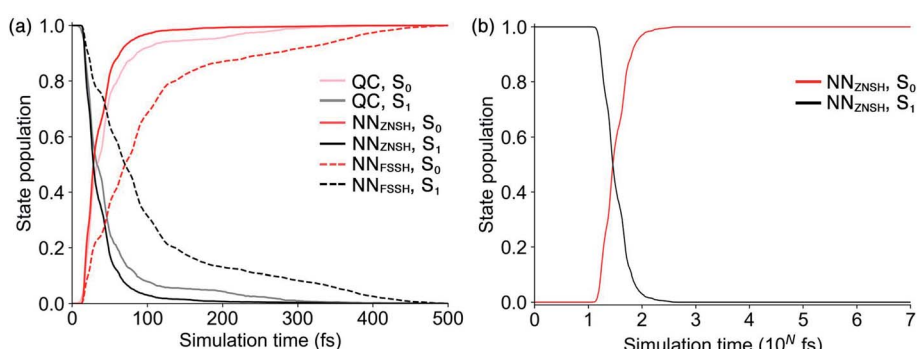


Fig. 3 (a) The QC vs. NN state populations for *trans*-1 at 300 K. The QC state populations average 1371 trajectories. The NN state populations average 5573 FSSH trajectories and 5820 ZNSH trajectories. (b) The NN state population of 89 ZNSH trajectories in 10 ns.



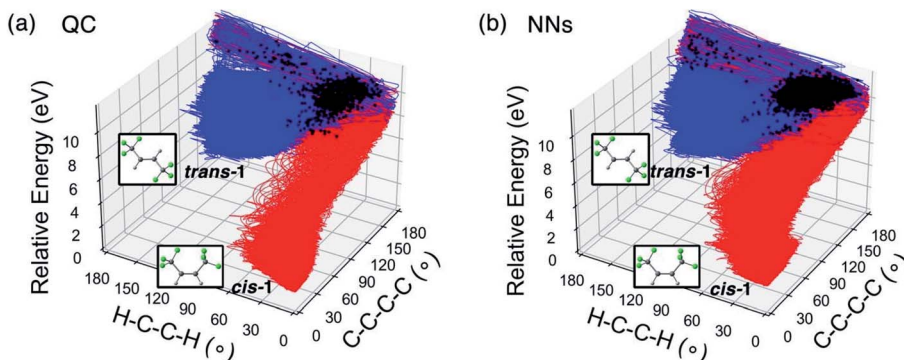


Fig. 4 The NAMD trajectories of *trans* → *cis* isomerization of **1**, computed with (a) CASSCF(2,2)/cc-pVDZ and (b) NN in 500 fs simulations at 300 K. The black dots represent the last surface hopping point in each trajectory.

events turned the trajectories along the direction of the \angle H-C-C-H axis toward *cis*-**1**. 698 trajectories took the same path returning to *trans*-**1**. The *trans* : *cis* ratio is 2.8 : 1 in the QC trajectories. The NN predicted trajectories in Fig. 3b recreated the topology of the reference. The NN predicted 772 trajectories transformed to *cis*-**1** and 2680 trajectories back to *trans*-**1**, resulting in a corresponding ratio of 3.5 : 1. The NN and QC trajectories show virtually identical evolution of the average \angle C-C-C-C and \angle H-C-C-H angles during the 500 fs simulation (Fig. S14†).

The QC and NN trajectories undergoing the *trans* → carbene pathway are shown in Fig. 5a and b with 5 snapshots from a QC trajectory in Fig. 5c, including the surface hopping point.

425 QC trajectories formed **2** in a *trans* : carbene ratio of 1.6 : 1. The trajectories in Fig. 5a display that the transformation shared the path with *trans* → *cis* isomerization from the Franck-Condon region to the S₁/S₀ crossing region. The S₁/S₀ surface hopping happened in the area of \angle C-C-C-C = 150–180° and \angle H-C-C-H = 0–60°. In Fig. 5c, the hydrogen migrated to the other carbon atom from 58 to 70 fs. The C=C π-bond broke resulting in a nonplanar geometry (\angle C-C-C-C = 164° and \angle H-C-C-H = 113° at 75 fs). We intentionally omitted the *trans* → carbene pathway data in the initial training set, yet the adaptive sampling located 347 structures with significantly broken C-H bonds (>1.40 Å) resembling **2** (5.6% in the final

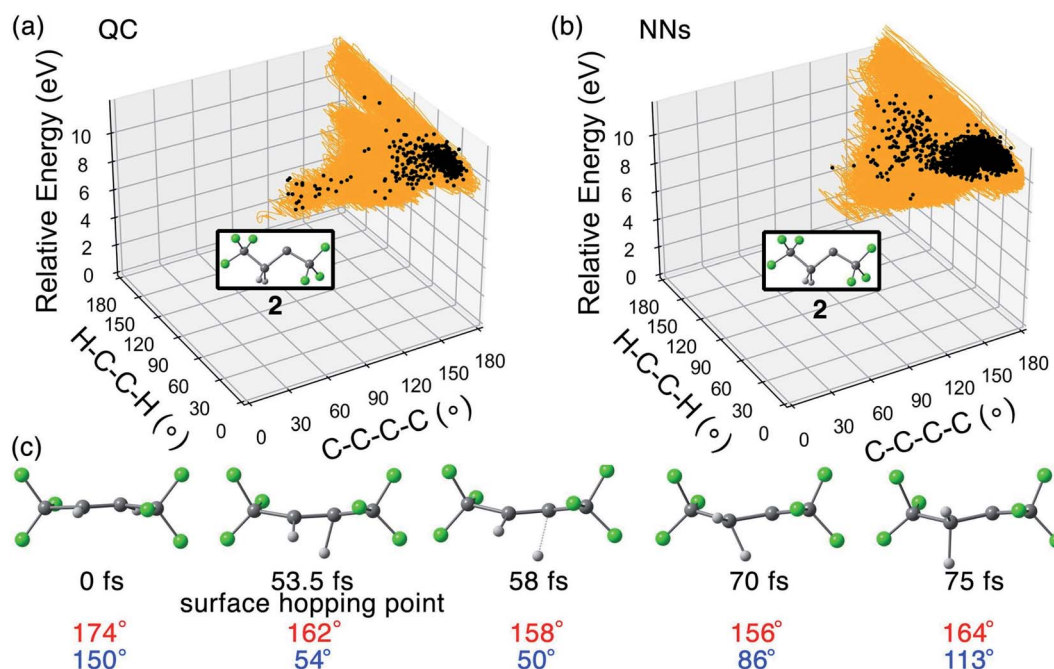


Fig. 5 The NAMD trajectories of *trans* → carbene isomerization of **1** computed with (a) CASSCF(2,2)/cc-pVDZ and (b) NN in 500 fs simulations at 300 K. The black dots represent the last surface hopping point in each trajectory. (c) Snapshots of the formation of **2** in CASSCF(2,2)/cc-pVDZ trajectories. The \angle H-C-C-H and \angle C-C-C-C angles are in blue and red, respectively.



training set). Fig. 5b shows the 2368 NN trajectories toward 2. The predicted *trans* : carbene ratio is 1.1 : 1.

We analyzed the crossing region and surface hopping points for the QC and NN trajectories. Fig. 6 shows the spatial distribution of the latest S_1/S_0 surface hopping point in each trajectory and examples of surface hopping geometries along the S_1/S_0 crossing seam.

Fig. 6a projects the latest S_1/S_0 surface hopping points from the CASSCF(2,2)/cc-pVDZ trajectories into the 2D conformational space of **1**. The dense areas represent the S_1/S_0 crossing region as defined by the two reaction coordinates. In the density map (Fig. 6a, right panel), the surface hopping points accumulated at two regions, $\angle C-C-C-C = 170^\circ$; $\angle H-C-C-H = 60^\circ$ and $\angle C-C-C-C = 180^\circ$; $\angle H-C-C-H = 40^\circ$, which have larger $\angle C-C-C-C$ angles than **MECP-trans-1**. Beside the hopping points near **MECP-trans-1**, Fig. 6c shows two pyramidalized surface hopping geometries at **A** ($\angle C-C-C-C = 180^\circ$ and $\angle H-C-C-H = 80^\circ$) and **B** ($\angle C-C-C-C = 180^\circ$ and $\angle H-C-C-H = 20^\circ$). We noted C-H stretching surface hopping points at **D** ($\angle C-C-C-C = 180^\circ$ and $\angle H-C-C-H = 180^\circ$); the C-H distance is 2.24 Å. The surface hopping geometries at **E** ($\angle C-C-C-C = 110^\circ$ and $\angle H-C-C-H = 140^\circ$) have shown the formation of **2**, which suggests that excited-state hydrogen abstraction is theoretically possible. Fig. 6b depicts the distribution and density of NN predicted S_1/S_0 surface hopping points. These surface hopping events occurred closely at $\angle C-C-C-C = 180^\circ$; $\angle H-C-C-H = 40^\circ$ agreeing with those observed in QC trajectories. The significant increase in the number of NN trajectories provided an increased statistical significance of our crossing region analysis. The NNs located a twisting surface hopping geometry

at **C** ($\angle C-C-C-C = 120^\circ$ and $\angle H-C-C-H = 90^\circ$). The density map (Fig. 6b, right panel) suggests it is a relatively rare surface hopping event as few points were located at **C**. The NN could detect the rare event because of the substantially increased number of trajectories (5820) with a nearly negligible increase in computational cost.

4.4 4π electrocyclic ring-closing model of norbornyl cyclohexadiene

To demonstrate our ML-NAMD method on a more complex reaction, we studied the photochemical behavior of norbornyl cyclohexadiene **3** (Scheme 2). Burns and Lopez hypothesized that irradiation of **3** would initiate 4π -electrocyclic ring-closing to produce diastereoisomeric ladderene products **anti-4** and **syn-4**, in line with known cyclohexadiene photochemistry.⁴⁸ Irradiation of **3** with 254 nm light was performed for 0.5, 1, 2, or 4 hours and provided several hydrocarbon products. The anticipated major reaction pathway produced ladderene diastereomers **anti-4** (28%) and **syn-4** (4%). Under these conditions we also observed an alternative reaction pathway to **anti-4'** (8%) via intermediate diene **3'** (see ESI† for more details). We then decided to study the major reaction pathway of **3** to form products **anti-4** and **syn-4** using our ML-NAMD method.

4.5 ML potential for the 4π -electrocyclic ring-closing of norbornyl cyclohexadiene

We trained NNs for **3** as described in Sections 2.3 and 2.4. The NN predict energies, forces, and NACs of **3** with comparable efficiency (0.0118 seconds) to **1** (0.00991 seconds). However, the marked increase in electrons and active space size of **3** results in

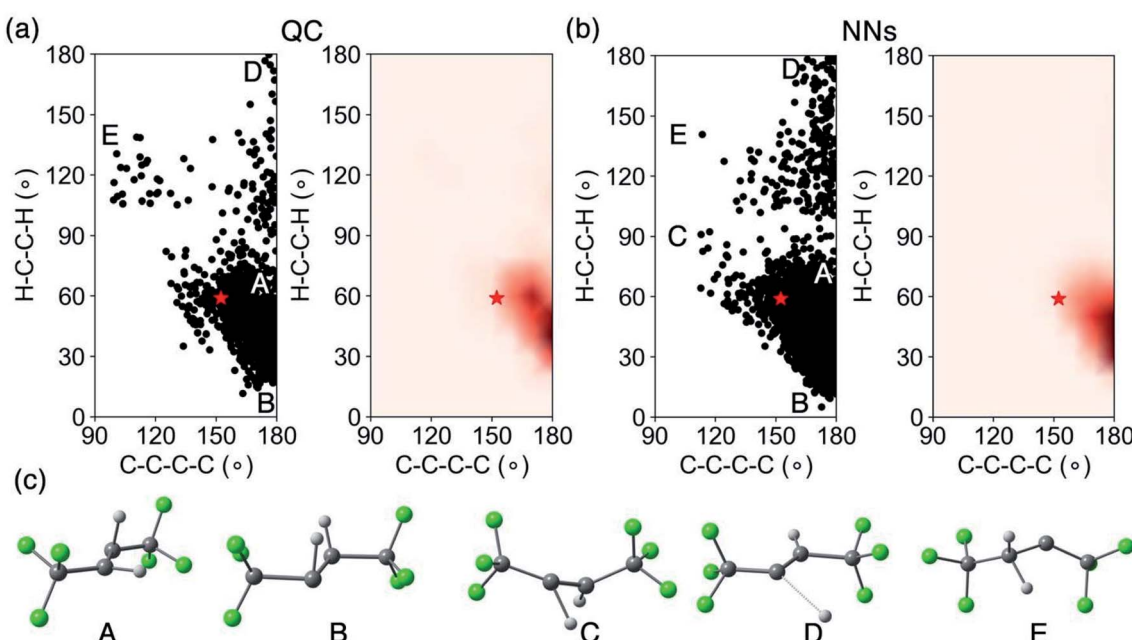
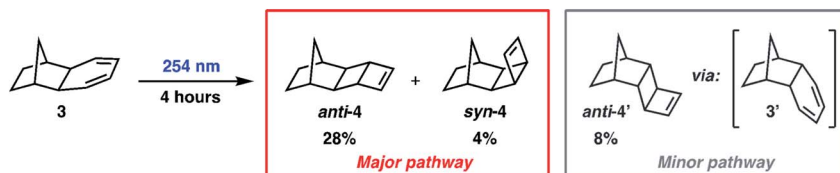


Fig. 6 Scatter plots of the latest S_1/S_0 surface hopping point geometries in (a) CASSCF(2,2)/cc-pVDZ and (b) NN predicted trajectories in 500 fs simulation at 300 K. The left and right panel illustrate the position and the density of the geometries. The density is defined as the fraction of geometries in an interval of 10° . The magnitude ranges from 0.00 to 0.12 and is colored from light pink to dark red. The red star marks **MECP-trans-1**. (c) Examples of five surface hopping point structures defining the S_1/S_0 crossing seam.





Scheme 2 Photochemical reaction of norbornyl cyclohexadiene.

considerably longer wall time of a single point CASSCF(4,3)/ANO-S-VDZP calculation (2480 seconds) than **1** (336 seconds). Note the ANO-S-VDZP and cc-pVDZ basis sets are the same size. The NN benefit from a 2.1×10^5 -fold acceleration compared to the CASSCF calculations. We summarized the NN architecture, MAE, and R^2 of predicted energies, forces, and NACs in Table 2.

The MAE of the energy predictions for compound **3** are 0.027–0.031 eV, which approaches chemical accuracy (1 kcal mol⁻¹ or 0.043 eV). The R^2 values are 0.9991–0.9996 and demonstrate a good correlation between the predicted and the target energies. The MAE of the forces ranges from 0.12–0.13 eV Å⁻¹, with R^2 values ranging from 0.9991–0.9996. They are 0.02 eV Å⁻¹ smaller than in case of compound **1**, suggesting improved fitting of forces. The MAE of the S₁/S₀-coupling are 0.078–0.081 eV Å⁻¹, which are smaller than 50% of that of **1**. Nevertheless, possible overfitting of the S₁/S₀-coupling still occurs as the R^2 values (0.5723–0.7956) are relatively low in the validation set (the MAE and R^2 are 0.036–0.040 eV and 0.9217–0.9452 in the training set).

4.6 ML-NAMD simulations for the 4π-electrocyclic ring-closing of norbornyl cyclohexadiene

The photodynamics study on **3** requires computationally intensive NAMD simulations. We have obtained 240 CASSCF(4,3)/ANO-S-VDZP trajectories of the 4π-electrocyclic ring-closing of **3**. The single trajectory of 1000 fs simulations required an average of 17 days of computation time. With our ML-NAMD method, we propagated 3910 NN FSSH trajectories and 3954 NN ZNSH trajectories, requiring just 56 and 38 seconds, respectively.

Table 2 Selected NN hyperparameters and the mean absolute errors and R^2 of predicted energies (eV), forces (eV Å⁻¹), and interstate coupling term of NACs (eV Å⁻¹) trained on 6267 data points of **3**. Additional details on hyperparameters and training statistics are available in ESI

Hyperparameters	Energies, forces		NACs ^a	
	NN1	NN2	NN1	NN2
Hidden layers	3	4	3	4
Neurons/layer	400	300	500	300
Batch size	128	128	128	128
MAE	0.027(0.12) ^b	0.031(0.13) ^b	0.078	0.081
R^2	0.9996(0.9934) ^b	0.9991(0.9858) ^b	0.7956	0.5732

^a Here the NAC only represents the interstate coupling term. ^b The MAE and R^2 of forces are shown in the parenthesis.

We compared the QC and NN state populations to assess the errors of our trained NN in predicting excited-state dynamics of **3** (Fig. 7).

Fig. 7 shows that the S₁ half-life of the QC trajectories is 108 fs. The NN FSSH trajectories overestimated it to be 175 fs. The resulting MAE in population between NN FSSH and QC trajectories is 14.8% in the first 500 fs. In contrast, the NN ZNSH trajectories agree better with the QC trajectories. The predicted S₁ half-life is 105 fs and the MAE in population is 4.5%. The delayed surface hopping events in the NN FSSH trajectories resulted from the underestimation of NACs as discussed in Section 4.3. This causes the NN FSSH trajectories to spend longer times on approaching regions with smaller energy gaps. The average surface hopping S₁/S₀ energy gap is 0.22, which is notably smaller than 0.45 eV in the QC trajectories (0.26 eV in the NN ZNSH trajectories). Thus, the following discussion of the ML-NAMD simulations of **3** will focus on the NN ZNSH trajectories.

The QC and NN trajectories are characterized with the distance R between the two carbon atoms closing the cyclohexadiene ring and the bending angle θ of the cyclohexadiene plane, shown in Fig. 8a and b. The optimized geometries of observed products and intermediates are shown in Fig. 8c.

The QC and NN trajectories show two types of pathways. We distinguished the *syn*- and *anti*-pathways with the bending angle of the cyclohexadiene plane, $\theta = 120\text{--}150^\circ$ and $\theta = 210\text{--}240^\circ$, respectively. 2 of 240 QC trajectories formed *anti*-**4** in 1 ps corresponding to a low yield of 0.8%, while a pathway to *syn*-**4** is

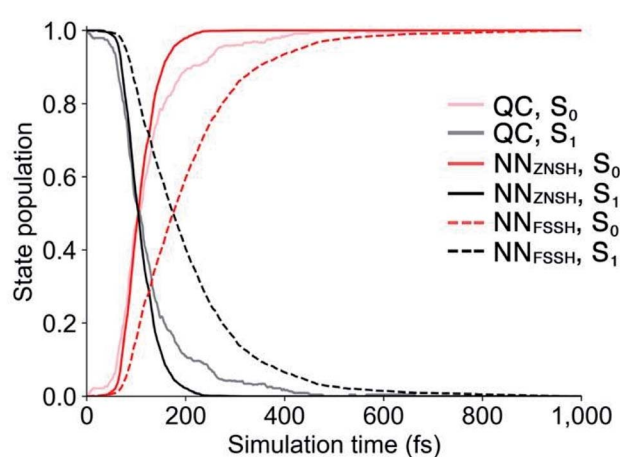


Fig. 7 The QC vs. NN state populations of **3** at 300 K. The QC state populations average is based on 240 trajectories. The NN state population is computed from 3910 FSSH trajectories and 3954 ZNSH trajectories, respectively.



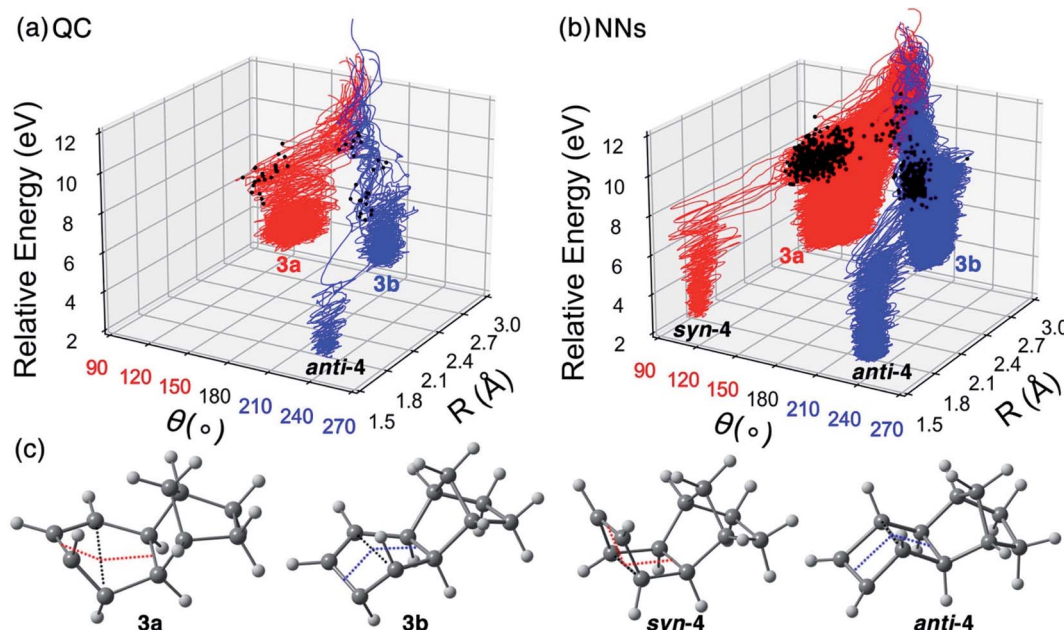


Fig. 8 The NAMD trajectories of 4π -electrocyclic ring-closing of **3** computed with (a) CASSCF(4,3)/ANO-S-VDZP and (b) NNs in 1000 fs simulations at 300 K. The black dots represent the last surface hopping point in each trajectory. R is defined as the distance between the two carbon atoms closing the cyclohexadiene ring. θ is defined as the bending angle of the cyclohexadiene plane. $\theta = 0\text{--}180^\circ$ represents a *syn*-configuration and $\theta = 180\text{--}360^\circ$ represents an *anti*-configuration. (c) The CASSCF(4,3)/ANO-S-VDZP optimized geometries of **3a**, **3b**, *syn-4* and *syn-4*. The red dotted lines highlight the angle θ in **3a** and *syn-4*; the blue dotted lines highlight the angle θ in **3b** and *anti-4*. The black dotted lines show the distance R in all compounds.

absent due to the insufficient number of trajectories. Our calculations suggest that the major outcome of the photoreaction of **3** return to the reactant (Fig. S19†). The higher experimental yield of *anti-4* (28%) and *syn-4* (4%) is because the reactant can be re-excited to have another chance of ring-closing during the 4 hours of irradiation. Our QC-NAMD simulations also suggest a less direct pathway involving reactive intermediates, **3a** and **3b** (Fig. 8a and c). The QC trajectories suggest a yield of **3a** and **3b** are 13% and 7% (Fig. 8a). Among those non-productive trajectories (Fig. S19†), 44% of the trajectories involved *syn*-configurations, and 35% formed *anti*-configuration before they reverted to **3**. Collectively, from the S_1 -FC regions, 57% of the trajectories followed the *syn*-pathway, and 43% of the trajectories pursued the *anti*-pathway. The preferential *syn*-pathway in the QC-NAMD simulations agree with the minimum energy path (MEP) calculations of **3** (Fig. S17†) where the *syn*-pathway is steeper descent than the *anti*-pathway.

The increase in NN trajectories in Fig. 8b identified a pathway to *syn-4*, with a 0.2% yield. The yield of *anti-4*, **3a**, and **3b** are 0.7%, 11%, and 5%, respectively, resulting in 84% non-productive trajectories reverting to **3**. The formations of *syn-4* and *anti-4* start near 100 fs and complete near 200 fs (Fig. S20†). We compare the yield of *syn-4* and *anti-4* in the NN trajectories and the experiment by zooming in the 100–200 fs region and normalizing the yield and time scale. The predicted yield curves show excellent agreement with the experiment data (Fig. S21†). Among the non-productive trajectories, 53% preferred the *syn*-pathway, and 31% went through the *anti*-pathway (Fig. S19†). Overall, the NN trajectories are consistent with QC trajectories

and MEP calculations that the *syn*-pathway (64%) is preferred over the *anti*-pathway (36%). It is also worth mentioning that we had not provided any information about the intermediates **3a** and **3b** in the initial training set since we did not recognize them until running QC-NAMD. The NN effectively collected the necessary training data through adaptive sampling.

We then analyzed the QC and NN surface hopping geometries to understand the behavior of NN trajectories in the crossing regions. Fig. 9a and b plot the distributions and density of the last S_1/S_0 surface hopping geometries in the QC and NN trajectories. Fig. 9c illustrates examples of three surface hopping geometries.

The 2D projection of surface hopping geometries from the QC trajectories shows three distinct regions (Fig. 9a). The crossing region around F and H resemble the MECP geometries along the *syn*- and *anti*-pathway, **MECP-syn-3** and **MECP-anti-3** (Fig. S16†). These geometries show shortened C–C distances ($R = 2.2\text{--}2.6 \text{ \AA}$) and bent cyclohexadiene plane ($\theta = 140\text{--}160^\circ$ near F and $\theta = 210\text{--}230^\circ$ near H) that are distorted toward the 4π -electrocyclic ring-closing. The region F is denser than H indicating the *syn*-pathway is preferred. Another crossing region G corresponds to cyclohexadiene-like surface hopping geometries ($R = 2.8\text{--}3.2 \text{ \AA}$, $\theta = 150\text{--}210^\circ$). These surface hopping events occur *via* the stretching of the cyclohexadiene plane. Fig. 9b plots surface hopping geometries in the NN trajectories. The distributions are similar in the three crossing regions. According to the density map, the NN tend to predict more surface hopping at the ring-closing regions (F and H) than the stretching region (G).

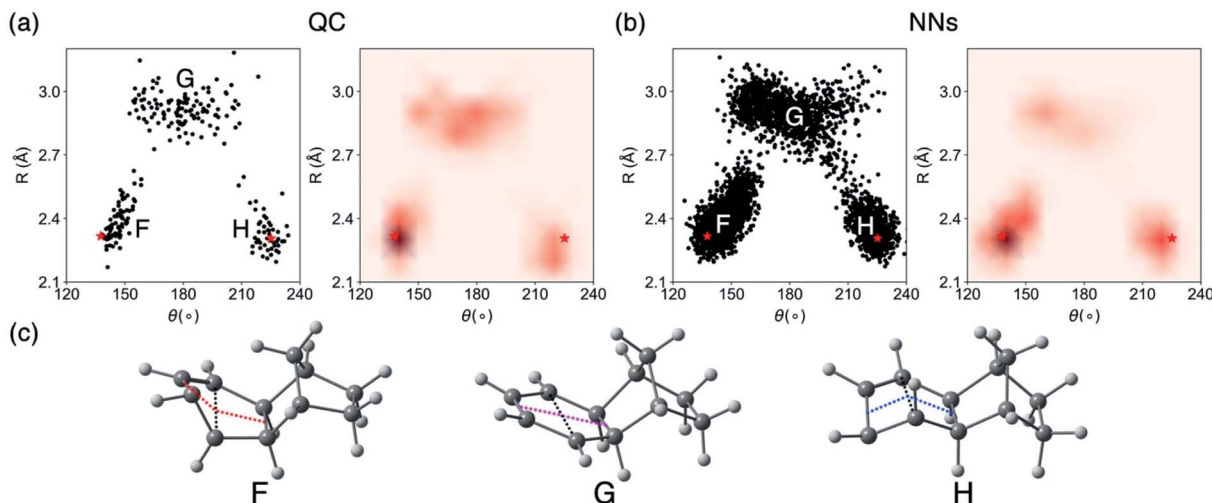


Fig. 9 Scatter plots of the surface hopping geometries in (a) CASSCF(4,3)/ANO-S-VDZP and (b) NN trajectories. The left and right panel show the distribution and the density of the geometries. The density is defined as the fraction of geometries in intervals of 10° . The magnitude ranges from 0.00 to 0.12 and is colored from light pink to dark red. The red stars mark the CASSCF(4,3)/ANO-S-VDZP optimized minimum energy crossing points geometries. (c) Examples of three surface hopping geometries. The dotted lines highlight the distance R (black) between the two carbon atoms closing the cyclohexadiene ring and the bending angle θ of the cyclohexadiene plane: $\theta = 0-180^\circ$ (red) represents a *syn*-configuration and $\theta = 180-360^\circ$ (blue) represents an *anti*-configuration (pink for $\theta = 180^\circ$).

In the non-productive trajectories, we observed thermal conversions from **3a** and **3b** to **3**. Fig. 10 depicts the QC trajectories snapshots showing the conversion from **3a** to **3** after it hops to the ground-state.

The snapshot in Fig. 10 shows a first appearance of **3a** at 122.5 fs. It completely transformed to **3** at 331.5 fs suggesting a lifetime of 209 fs. The trajectory statistics concludes that 0.8% of the QC trajectories undergo **3a** \rightarrow **3** conversion and 0.8% go for **3b** \rightarrow **3**. The NN trajectories confirm the **3a** \rightarrow **3** and **3b** \rightarrow **3** thermal conversions in 5% and 3% of the trajectories, respectively. The remaining **3a** and **3b** at the end of the NAMD simulations suggest the lifetime of **3a** and **3b** can be longer than 1 ps. That encourages us to explore subsequent thermal reactions of the photodynamics of **3** in nanosecond scale.

We continued to propagate 984 NN trajectories from 1 ps to 1 ns, which completes in just 4.8 hours per trajectory. We monitored the interconversion from **3a** and **3b** to **3** with the $\angle H-C-H$ dihedral angles α , as shown in Fig. 11.

The top panel in Fig. 11 illustrates trajectories that formed the aforementioned short-lived **3a** and **3b**, which convert to **3** within 1 ps. The dihedral angle α is below 60° in the initial 100

fs. The formation of **3a** and **3b** rotate the $C=C$ bond leading to $\alpha = 180^\circ$ and the continued conversion to **3** recovery the planar cyclohexadiene with $\alpha < 30^\circ$. The bottom panel in Fig. 11 shows the trajectories involved with a long-lived **3a** and **3b**. The trajectories retain the dihedral angle $\alpha = 180^\circ$ at 1 ps. By tracking reaction coordinate α , we note that the earliest conversion occurred right after 1 ps and the last one appeared at 0.9 ns. In the end of 1 ns simulation, the yield of **3a** reduced to 1% and the **3b** completely converted to **3**. Measuring the ~ 0.9 ns lifetime of these reactive intermediates would not be possible without ML-NAMD simulations; this dynamic effect offers

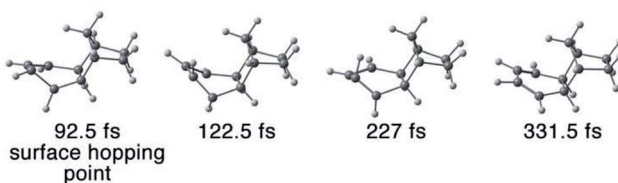


Fig. 10 Snapshots of the formation of transient intermediate **3a** in a CASSCF(4,3)/ANO-S-VDZP trajectory. The trajectory first formed **3a** at 122.5 fs and then converted to **3** at 331.5 fs.

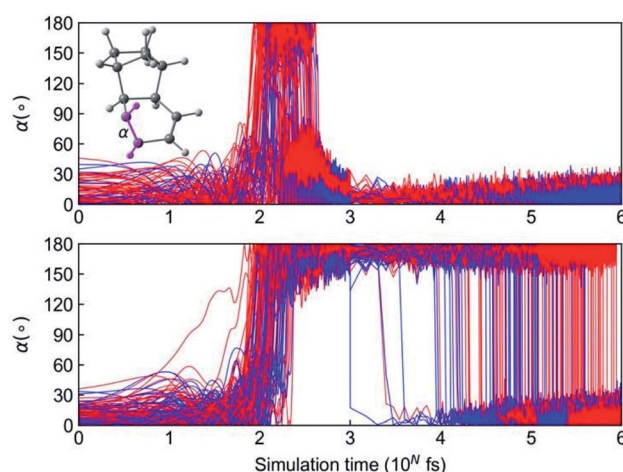


Fig. 11 The NN trajectories showing the thermal conversion from **3a** (red) and **3b** (blue) to **3** in 1 ns. The top panel shows the conversions occurred within 1 ps; the bottom panel shows the conversions after 1 ps. α is defined as the $\angle H-C-C-H$ dihedral angles ($0-180^\circ$) marked in pink. $\alpha = 0^\circ$ in **3**; $\alpha = 180^\circ$ in **3a** and **3b**.



important mechanistic insight into photochemical ladderene formation reactions.

5. Concluding remarks

We have applied and expanded a ML-NAMD approach using neural networks that overcome current limitations in photodynamics simulations, including simulation time and molecular complexity. We used a composite generation scheme to effectively sample the initial training set, which combines the Wigner sampling, geometrical interpolations, and short-time quantum chemical trajectories. We implemented an adaptive sampling workflow to automatically expand the initial training set with the important data in the unexplored conformational space using a query of committee model. The final data set of the *cis-trans* isomerization and 4π -electrocyclic ring-closing models has 6207 and 6267 data points, respectively. The training error in energy achieved chemical accuracy with a mean absolute error of 0.023–0.032 eV. The forces and NAC interstate couplings errors were 0.12–0.15 eV Å⁻¹ and 0.078–0.182 eV Å⁻¹ respectively.

For the *cis-trans* isomerization of **trans-1**, the NNs effectively accelerated the computations of energies, forces, and NACs for **trans-1** in 3.4×10^4 -fold compared to the CASSCF calculations. The 10 ns ML-NAMD simulations from the S₁ Franck–Condon regions of the Wigner sampled initial conditions at 300 K on a single CPU completed in 2 days 2 hours on average of 89 trajectories, which would have approximately required 58 years for running the CASSCF(2,2)/cc-pVDZ calculations. The analysis of 5820 NN trajectories in 500 fs shows consistent results with the QC trajectories, predicting the formation of **cis-1** (*cis* : *cis* = 3.5 : 1) and 2 (*trans* : carbene = 1.1 : 1). The NN acceleration becomes more significant (2.1×10^5 -fold) for complex organic molecules such as **3** because of the considerably increasing cost of CASSCF calculations with more electrons and larger active space. Our results of 3954 NN trajectories in 1 ps confirmed the low-yield of **syn-4** (0.2%) and **anti-4** (0.7%) that support our experimental results. The continued 1 ns NN photodynamics simulations revealed almost complete thermal conversion from **3a** and **3b** to **3**, which explain their absence in experiment.

We have demonstrated that the PyRAI²MD ML-NAMD approach is able to not only reproduce experimental results, but can be used to identify new light-induced mechanistic pathways. The identification of reactive intermediates such as the carbene **2** and isomers **3a** and **3b** help to explain experimental results. Our study provides a path forward to quantify the lifetimes of reactive long-lived excited-states and reactive intermediates currently inaccessible with quantum chemical methods alone. We anticipate that these PyRAI²MD-enabled simulations will enable other researchers to understand photochemical reactivities and selectivities with thousands of nanosecond trajectories, facilitating connections between excited- and ground-state mechanistic pathways.

Author contributions

Jingbai Li developed Python Rapid Artificial Intelligence *Ab Initio* Molecular Dynamics (PyRAI²MD) codes, performed

machine learning and quantum chemical calculations, and prepared the initial draft of the manuscript and ESI. Patrick Reiser implemented the TensorFlow/Keras neural networks interface for PyRAI²MD, prepared the machine learning section in the ESI, and helped review and edit the draft. Benjamin R. Boswell performed the photochemistry experiment of norbornyl cyclohexadiene, and helped prepare the experimental section in the draft and ESI. André Eberhard tested Gaussian Process regression model. Noah Z. Burns supervised the photochemistry experiments, and helped review and edit the draft. Pascal Friederich supervised the implementation of TensorFlow/Keras neural networks interface, advised the discussions in the machine learning data, and helped review and edit the draft. Steven A. Lopez supervised the development of PyRAI²MD, managed the project administration, reviewed and edited the manuscript revisions, and responded to the submission process.

Conflicts of interest

There are no conflicts of interest to declare.

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