Supporting Information:

Effect of conformational disorder on exciton states of an azobenzene aggregate

Evgenii Titov*

Theoretical Chemistry, Institute of Chemistry, University of Potsdam,
Karl-Liebknecht-Straße 24-25, 14476 Potsdam, Germany

E-mail: titov@uni-potsdam.de

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S1
SI1  Details on models, methods, and calculations

The tetramer model was constructed from the B3LYP/def2-TZVP optimized monomer geometry by translating this geometry by 3.5 Å in the direction perpendicular to the molecular plane of trans azobenzene. This tetramer geometry was used as an input geometry for BOMD simulations. The BOMD simulations were performed using B3LYP+D3(BJ) and ωB97X-D functionals with the def2-SV(P) basis set. The velocity Verlet algorithm was used to integrate Newton’s equations of motion. The time step was set to 0.5 fs and the trajectories were run up to 10 ps. The constant-temperature dynamics were simulated using a simple velocity rescaling approach, i.e. multiplying velocities by $\sqrt{\frac{T_{\text{ref}}}{T_{\text{curr}}}}$ at each time step. Here, $T_{\text{ref}}$ is the reference (desired) temperature and $T_{\text{curr}}$ is the current (instantaneous) temperature. The initial velocities were randomly sampled from a uniform distribution and scaled to yield the reference temperature (300 K in our case). The constant-energy trajectories were launched with zero initial velocities. Snapshots selected between 100 fs and 10 ps (with 100 fs period) were subjected to TD-DFT calculations. The excited states were calculated using TD-ωB97X-D and TD-ωB97 with def2-SV(P) basis set. The 20 lowest excited states were requested in the case of the tetramer, and 5 states in the case of the monomer. Transition density matrices were calculated using Multiwfn 3.8.

The calculated stick spectra are broadened with Gaussians as

$$I(E) = \frac{1}{N_s} \sum_{i,\alpha} f_{i,\alpha} \exp \left( -\frac{1}{2\sigma^2} (E - E_{i,\alpha})^2 \right).$$  \hspace{1cm} (S1)

Here, $I$ is intensity, $E$ is excitation energy, $N_s$ is the number of selected snapshots, $E_{i,\alpha}$ and $f_{i,\alpha}$ are the TD-DFT excitation energy and oscillator strength, respectively, for the $S_0 \rightarrow S_i$ transition, for snapshot $\alpha$, and $\sigma$ is a broadening parameter ($\sigma = 0.05$ eV in this work). For each selected snapshot we also calculate the spectra of four monomers forming the aggregate (to calculate the broadened spectrum of the monomer, $4N_s$ instead $N_s$ is used in the denominator of eq. (S1)).
Since the def2-SV(P) basis set was used in dynamics and TD-DFT calculations, we have also considered the tetramer geometry constructed from the B3LYP/def2-SV(P) optimized monomer geometry. The effect of the monomer geometry on the spectra is shown in Fig. S1. While broadened spectra merely show a shift, the stick spectra of the tetramer are quite different. There is a single very intense transition (\(S_0 \rightarrow S_{13}\)) for the def2-SV(P) geometry, and there are two less intense closely lying transitions (\(S_0 \rightarrow S_{12}\) and \(S_0 \rightarrow S_{13}\)) for the def2-TZVP geometry. The FTDM matrices of these transitions are shown in Fig. S2. It is seen that there is a strong admixture of CT excitations for the def2-TZVP geometry.

![Figure S1](image-url)

**Figure S1:** The \(\pi\pi^*\) absorption bands of the monomer (a) and the tetramer (b) calculated with TD-\(\omega\)B97X-D/def2-SV(P) at B3LYP/def2-SV(P) and B3LYP/def2-TZVP geometries.
Figure S2: (a) The $F$ matrix for the $S_0 \rightarrow S_{13}$ transition of the tetramer constructed from the B3LYP/def2-SV(P) optimized monomer geometry. (b,c) The $F$ matrices for the $S_0 \rightarrow S_{12}$ (b) and $S_0 \rightarrow S_{13}$ (c) transitions of the tetramer constructed from the B3LYP/def2-TZVP optimized monomer geometry. Excited states are calculated with TD-$\omega$B97X-D/def2-SV(P).

Figure S3: Instantaneous temperature as a function of time for dynamics starting from the model geometry with zero velocities. The results for B3LYP+D3 and $\omega$B97X-D dynamics are shown.
Figure S4: The $\pi\pi^*$ absorption band of the monomer and the tetramer at $T \approx 300$ K (upper row) and $T < 26$ K (lower row) calculated with TD-$\omega$B97X-D/def2-SV(P) (left column) and TD-$\omega$B97/def2-SV(P) (right column). The monomer-to-tetramer spectral shifts are also shown.
Figure S5: The $n\pi^*$ absorption band of the monomer and the tetramer at $T \approx 300$ K (upper row) and $T < 26$ K (lower row) calculated with TD-$\omega$B97X-D/def2-SV(P) (left column) and TD-$\omega$B97/def2-SV(P) (right column). The monomer-to-tetramer spectral shifts are also shown.
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Figure S7: Distributions of the oscillator strength of the brightest $\pi\pi^*$ transition.
SI2.1 Brightest $\pi\pi^*$ transition

Figure S8: TD-ωB97/def2-SV(P) calculations. Upper row: calculations for single (B3LYP/def2-SV(P)) geometry, for the brightest transition ($S_0 \rightarrow S_{10}$). (a) The $F$ matrix. (b) Diagonal $F_{XX}$ elements as well as the sum of diagonal (LE) and off-diagonal (CT) elements. Lower row: calculations for the ensemble of 100 snapshots (at $T \approx 300$ K), for the brightest transitions. (c) The $F$ matrix averaged over all snapshots. (d) Diagonal $F_{XX}$ elements of the averaged $F$ matrix, as well as the sum of diagonal (LE) and off-diagonal (CT) elements. (e) Averaged highest to lowest diagonal values and their sum (LE). (f) Relative frequency with which monomers have the highest or lowest $F_{XX}$ value.
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Figure S10: TD-ωB97/def2-SV(P) calculations, T < 26 K. (a) The $F$ matrix averaged over all snapshots. (b) Diagonal $F_{XX}$ elements of the averaged $F$ matrix, as well as the sum of diagonal (LE) and off-diagonal (CT) elements. (c) Averaged highest to lowest diagonal values and their sum (LE). (d) Relative frequency with which monomers have the highest or lowest $F_{XX}$ value.
Figure S11: Inverse participation ratios for 100 snapshots.
SI2.2  $S_0 \rightarrow S_5$ transition

![Graphical representation of TD-ωB97X-D/def2-SV(P) calculations for the $S_0 \rightarrow S_5$ transition.](image)

Figure S12: TD-ωB97X-D/def2-SV(P) calculations. Upper row: calculations for single (B3LYP/def2-SV(P)) geometry, for the $S_0 \rightarrow S_5$ transition. (a) The $\mathbf{F}$ matrix. (b) Diagonal $F_{XX}$ elements as well as the sum of diagonal (LE) and off-diagonal (CT) elements. Lower row: calculations for the ensemble of 100 snapshots (at $T \approx 300$ K), for the brightest transitions. (c) The $\mathbf{F}$ matrix averaged over all snapshots. (d) Diagonal $F_{XX}$ elements of the averaged $\mathbf{F}$ matrix, as well as the sum of diagonal (LE) and off-diagonal (CT) elements. (e) Averaged highest to lowest diagonal values and their sum (LE). (f) Relative frequency with which monomers have the highest or lowest $F_{XX}$ value.
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Figure S16: Inverse participation ratios for 100 snapshots.
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Figure S17: TD-ωB97/def2-SV(P) calculations, $T \approx 300$ K. (a) The $F$ matrix averaged over all snapshots. (b) Diagonal $F_{XX}$ elements of the averaged $F$ matrix, as well as the sum of diagonal (LE) and off-diagonal (CT) elements. (c) Averaged highest to lowest diagonal values and their sum (LE). (d) Relative frequency with which monomers have the highest or lowest $F_{XX}$ value.

Figure S18: TD-ωB97X-D/def2-SV(P) calculations, $T < 26$ K. (a) The $F$ matrix averaged over all snapshots. (b) Diagonal $F_{XX}$ elements of the averaged $F$ matrix, as well as the sum of diagonal (LE) and off-diagonal (CT) elements. (c) Averaged highest to lowest diagonal values and their sum (LE). (d) Relative frequency with which monomers have the highest or lowest $F_{XX}$ value.
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Figure S20: Inverse participation ratios for 100 snapshots.
SI3 \( \omega \text{B97X-D dynamics} \)

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Figure S22: The $n\pi^*$ absorption band of the monomer and the tetramer at $T \approx 300$ K (upper row) and $T < 26$ K (lower row) calculated with TD-ωB97X-D/def2-SV(P) (left column) and TD-ωB97/def2-SV(P) (right column). The monomer-to-tetramer spectral shifts are also shown.
Figure S23: Distributions of the brightest $\pi\pi^*$ state label.
Figure S24: Distributions of the oscillator strength of the brightest $\pi\pi^*$ transition.
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Figure S29: Inverse participation ratios for 100 snapshots.
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Figure S34: Inverse participation ratios for 100 snapshots.
### SI3.3 Brightest \(n\pi^*\) transition

Figure S35: TD-\(\omega\)B97X-D/def2-SV(P) calculations, \(T \approx 300\) K. (a) The \(F\) matrix averaged over all snapshots. (b) Diagonal \(F_{XX}\) elements of the averaged \(F\) matrix, as well as the sum of diagonal (LE) and off-diagonal (CT) elements. (c) Averaged highest to lowest diagonal values and their sum (LE). (d) Relative frequency with which monomers have the highest or lowest \(F_{XX}\) value.

Figure S36: TD-\(\omega\)B97/def2-SV(P) calculations, \(T \approx 300\) K. (a) The \(F\) matrix averaged over all snapshots. (b) Diagonal \(F_{XX}\) elements of the averaged \(F\) matrix, as well as the sum of diagonal (LE) and off-diagonal (CT) elements. (c) Averaged highest to lowest diagonal values and their sum (LE). (d) Relative frequency with which monomers have the highest or lowest \(F_{XX}\) value.
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Figure S38: TD-ωB97/def2-SV(P) calculations, T < 26 K. (a) The $F$ matrix averaged over all snapshots. (b) Diagonal $F_{XX}$ elements of the averaged $F$ matrix, as well as the sum of diagonal (LE) and off-diagonal (CT) elements. (c) Averaged highest to lowest diagonal values and their sum (LE). (d) Relative frequency with which monomers have the highest or lowest $F_{XX}$ value.
Figure S39: Inverse participation ratios for 100 snapshots.
Figure S40: Evolution of IPR and $F_H$ in time (for the first 100 fs with time step of 1 fs) along with selected geometrical parameters (bond lengths, intermolecular distances, and dihedral angles defined in the lower right picture). Dotted lines in two upper panels show corresponding values at time 0 fs (for the starting, model (def2-TZVP) geometry). BOMD simulation is done at the B3LYP+D3 level, TD-DFT calculations at the TD-ωB97X-D level. The shown tetramer structure corresponds to 100 fs. Hydrogen atoms in yellow are fixed.