

Ultrafast Vibronic Dynamics of Functional Organic Polymer Materials: Coherence, Confinement, and Disorder

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Acknowledgments & Collaborations

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Topics

- 1 Photoinduced Energy & Charge Transfer in Functional Organic Materials
 - Goal: First-Principles Approach to Organic Photovoltaics
 - Electron-Hole Lattice Models & Vibronic Coupling
 - Quantum Dynamics in Many Dimensions

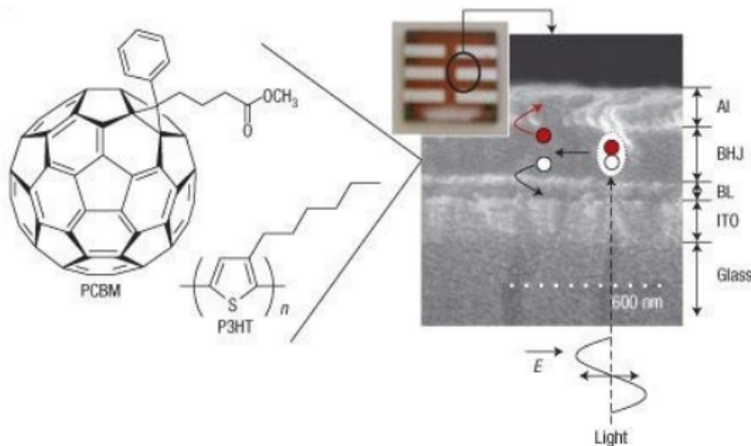
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- 2 Case Studies I: Exciton Migration Guided by Conformational Dynamics
Torsion-Induced Intra-Chain Exciton Migration
Ultrafast, Coherent Exciton-Polaron Dynamics
Do Semiclassical and Ehrenfest-Langevin Dynamics Work?

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Charge Transfer Excitons in Neat Polythiophene
Collective Vibronic Effects
Charge Transfer Excitons and Interfacial Charge Separation

Role of Electron-Phonon Coupling in Organic Photovoltaics



Kim et al., Nature Materials, 5, 197 (2006)

elementary steps:

- creation of electron-hole pairs (excitons)
- **exciton migration** to donor-acceptor interface area
- **exciton dissociation** at donor-acceptor junctions (here, PCBM-P3HT)¹
- capture of charge carriers at electrodes
- potentially competing process: electron-hole recombination

¹PCBM = phenyl-C₆₁-butyric acid methyl ester, P3HT = poly(3-hexylthiophene)

What is the Best Nano-Morphology?

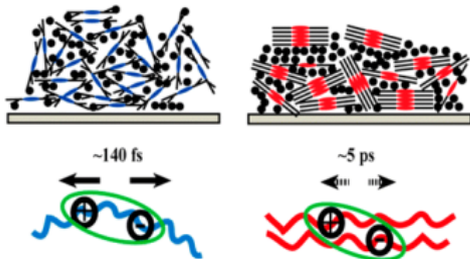
Highly ordered nanostructured domains (typically sub-10 nm) are thought to

- facilitate exciton diffusion
- favor exciton dissociation
- facilitate free carrier transport

Nanostructured domains can be achieved by

- self-assembly properties of D/A oligomers
- thin film processing methods (e.g., nanoimprint lithography)

However, the role of nanoscale ordering is controversial:

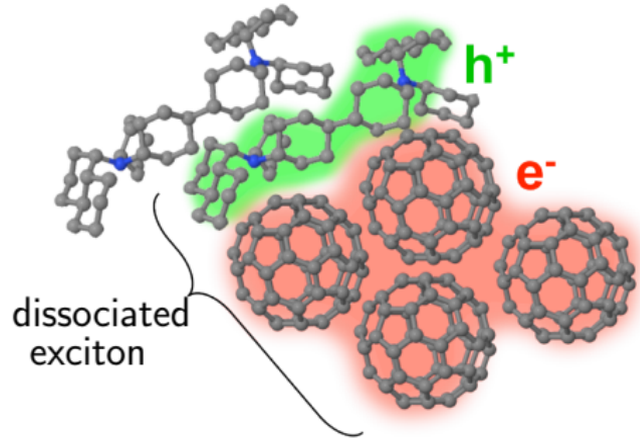


Guo et al., JACS 136, 10024 (2014)

- e.g., in a recent study of DA copolymer:fullerene systems, it is shown that the charge separation energetics changes **unfavorably** upon formation of crystalline domains

Which Methods, Even for a Minimal Model?

- tens to hundreds of electronic states
- aggregate-type systems
- charge transfer and excitonic couplings^(*) required
- delocalized excitations
- strong electron-phonon coupling
- non-Markovian dynamics
- non-exponential transfer
- coherent wavepacket dynamics
- standard rate theories (Förster / Marcus) not necessarily valid



<http://phys.org/news/2014-02-result-cheaper-efficient-solar-cells.html>

(*)excitonic coupling = transition density interaction:

$$V_{DA} = \frac{1}{4\pi\epsilon_0} \int d\mathbf{r}_D d\mathbf{r}_A \frac{\rho_D^{(eg)}(\mathbf{r}_D) \rho_A^{(ge)}(\mathbf{r}_A)}{|\mathbf{r}_D - \mathbf{r}_A|} \longrightarrow \text{limiting case: transition dipole interaction}$$

Two Types of Approaches

approximate electron-nuclear dynamics:
e.g., time-dep. Kohn-Sham equation

$$i \frac{\partial}{\partial t} \phi_i(r, t) = \left(-\frac{\nabla^2}{2} + v_{\text{KS}}(r, t) \right) \phi_i(r, t)$$

expand in adiabatic KS basis,

$\phi_i(r, t) = \sum_k c_{ik}(t) \tilde{\phi}_k(r; R)$ such that

$$i \frac{dc_{ik}}{dt} = \sum_l c_{il}(t) (\epsilon_l \delta_{kl} + d_{kl} \cdot \dot{R})$$

Ehrenfest or Surface Hopping dynamics

e.g., Craig, Duncan, Prezhd, PRL 95, 163001 (2005)

pro's: no pre-computed potentials

con's: possibly poor description of excited states and nuclear dynamics

parametrized model Hamiltonian
+ multi-state quantum nuclear dynamics

$$i \frac{\partial}{\partial t} \psi(R, t) = \hat{H} \psi(R, t)$$

with a multi-state/site Hamiltonian

$$\hat{H} = \sum_{mn} (\hat{h}_{mn}^e + \hat{h}_{mn}^{e-ph}(R)) |m\rangle \langle n| + \hat{H}_0^{ph}(R)$$

and $|\psi(R, t)\rangle = \sum_n c_n(t) \Phi_n(R, t) |n\rangle$

use (approximate) quantum dynamics

e.g., Kondov et al., JPCC 111 (2007), Tamura et al., JACS 135 (2013)

pro's: immediate physical interpretation

con's: restricted number of coordinates, electronic couplings *via* diabatisation

Road Map: Model Hamiltonians & Quantum Dynamics

e-h lattice models + non-perturbative *e-ph* interaction + quantum dynamics

- electron-hole (*e-h*) lattice models including vibronic interactions
- *ab initio* (typically CC2, ADC(2)) and/or TD-DFT parametrization
- **diabatization** procedures to generate electronic couplings
- compute **spectral densities** and effective-mode decomposition
- efficient high-dimensional **nonadiabatic quantum dynamics** using multi-configurational methods (MCTDH) or reduced dynamics (HEOM) + semiclassical/quantum-classical approaches (SQC/MM, Ehrenfest)
- “molecular aggregate” perspective: parametrization for small fragments & dynamics for larger systems Polkehn, Eisenbrandt, Tamura, Burghardt, Int. J. Quant. Chem. 118:e25502 (2018)

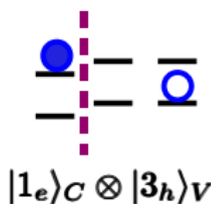
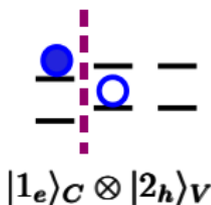
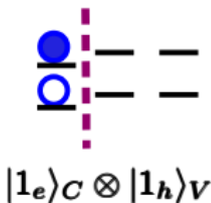
CC2 = Second-Order Approximate Coupled-Cluster

ADC(2) = Second-Order Algebraic-Diagrammatic Construction (ADC(2)) scheme

MCTDH = Multi-Configuration Time-Dependent Hartree Beck et al., Phys. Rep. 324, 1 (2000)

HEOM = Hierarchy of Equations of Motion Tanimura, J. Phys. Soc. Jpn. 75, 082001 (2006)

Electron-Hole Lattice Model



- electron-hole ($e-h$) configurations:

$$|\mathbf{n}\rangle = |n_e n'_h\rangle = |n_e\rangle_C \otimes |n'_h\rangle_V$$

- Hamiltonian in this basis:

$$\hat{H} = \sum_{\mathbf{m}\mathbf{n}} (\hat{h}_{\mathbf{m}\mathbf{n}}^{eh} + \hat{h}_{\mathbf{m}\mathbf{n}}^{eh-ph}(\mathbf{x})) |\mathbf{m}\rangle \langle \mathbf{n}| + \hat{H}_0^{ph}(\mathbf{x})$$

Merrifield, J. Chem. Phys. 34, 1835 (1961)

Wang and Mukamel, Chem. Phys. Lett. 192, 417 (1992)

Karabunarliev and Bittner, J. Chem. Phys. 118, 4291 (2003)

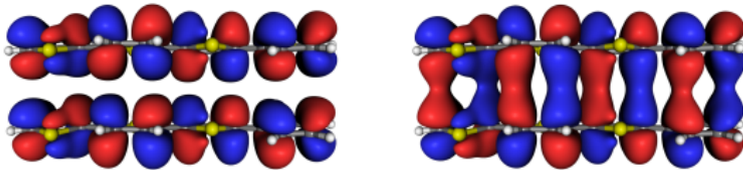
Binder, Wahl, Römer, Burghardt, Faraday Discuss, 163, 205 (2013)

- includes Frenkel-type exciton (XT) states and charge transfer (CT) states
- oligomer (fragment) *ab initio* or TDDFT calc's: on-site energies, diabatic couplings
- vibronic couplings from Franck-Condon gradients, geometry optimization, PES cuts

Special Case: Frenkel Exciton Model

- Frenkel model ($n_e = n'_h = n$) often a good approximation to describe exciton
- exact analytic mapping of oligomer PES's to Frenkel model

Binder, Römer, Wahl, Burghardt, J. Chem. Phys. 141, 014101 (2014)



stacked oligothiophene (OT4)₂: "HJ aggregate"

- **J-aggregate**: end-to-end alignment of monomer units; lowest state of the exciton manifold is the bright state
- **H-aggregate**: plane-to-plane stacked geometry; highest state of the exciton manifold is the bright state
- **HJ-aggregate**: combination of both, as in stacked oligomers

Yamagata, Spano, JCP **136**, 184901 (2012)

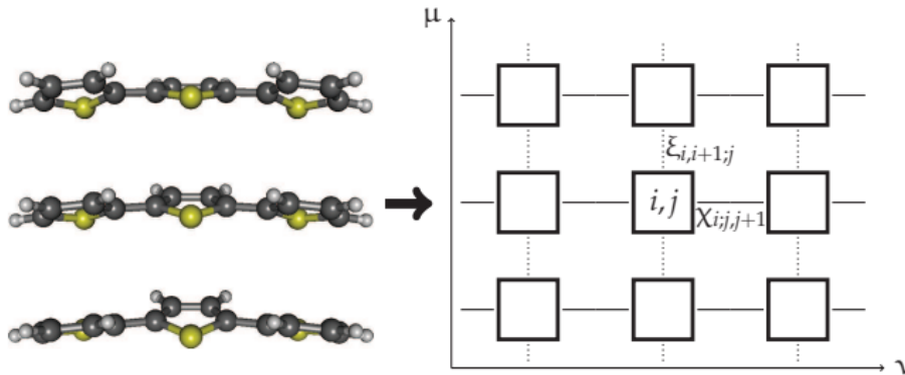
- delocalized states

$$|\Psi_{\text{exciton}}\rangle = \sum_n^{N_{\text{exc}}} c_n |n\rangle$$

where $N_{\text{exc}} \sim 5-10$; $|n\rangle =$ configuration with single excitation on n th monomer

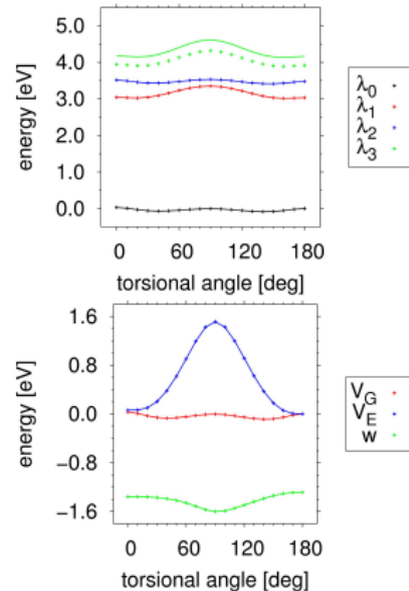
- trapping due to exciton-phonon interactions

HJ-Aggregate: Vibronic Lattice Model



Binder, Römer, Wahl, Burghardt, J. Chem. Phys. 141, 014101 (2014)

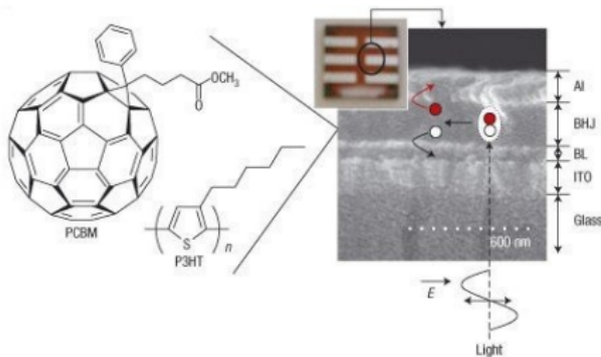
Binder, Polkehn, Ma, Burghardt, Chem. Phys. 482, 16 (2017)



- here: analytic mapping of oligomer PES onto Hückel type model in 1D or 2D: solution to an inverse eigenvalue problem

(NB.: V_G/V_E : monomer potentials, w : site-to-site coupling)

System-Bath Models



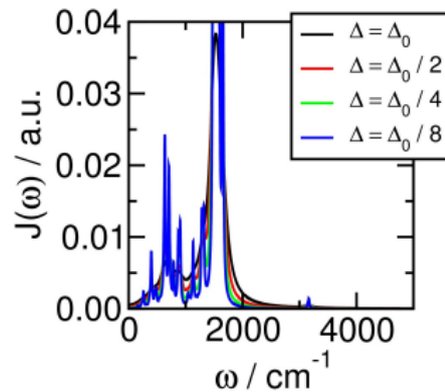
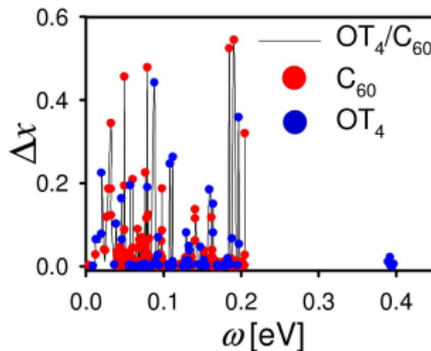
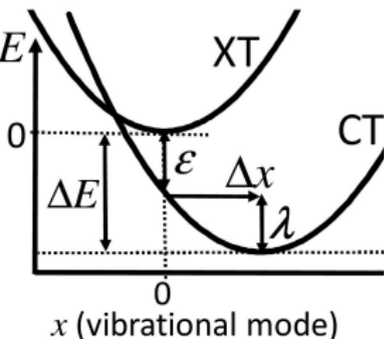
S region: e.g., electronic degrees of freedom (electron-hole states)

B region: all vibrations (phonons) mapped to harmonic oscillator model

$$\hat{H}_B + \hat{H}_{SB} = \sum_n \frac{1}{2} (\hat{p}_n^2 + \frac{1}{2} \omega_n^2 \hat{x}_n^2) + \hat{s} \sum_n c_n \hat{x}_n$$

$$J(\omega) = \pi/2 \sum_n c_n^2 / \omega_n \delta(\omega - \omega_n)$$

spectral density

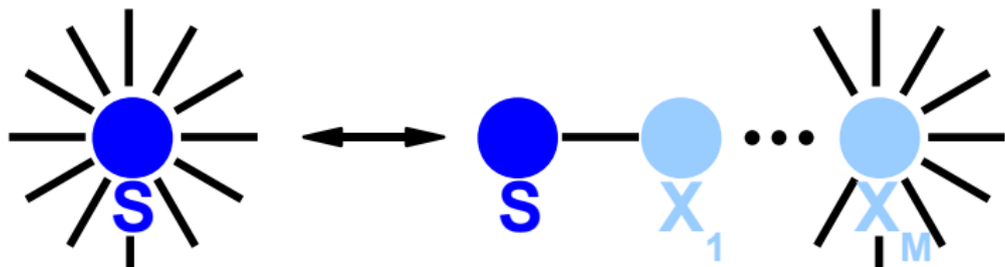
Spectral Densities from Electronic Structure Calculations^(*)

$$J(\omega) = \frac{\pi}{2} \sum_n^N \frac{c_n^2}{\omega_n} \delta(\omega - \omega_n) \simeq \frac{\pi}{2} \sum_n^N \frac{c_n^2}{\pi} \frac{\Delta}{(\omega - \omega_n)^2 + \Delta^2}$$

Tamura, Martinazzo, Ruckebauer, Burghardt, J. Chem. Phys., 137, 22A540 (2012)

^(*)NB. Alternatively: obtain SD's from correlation functions (MD, CPMD, ...)

Bath Dynamics in Less Dimensions: Effective-Mode Chains



Martinazzo, Vacchini, Hughes, Burghardt, J. Chem. Phys. 134, 011101 (2011), Hughes, Christ, Burghardt, J. Chem. Phys. 131, 024109 (2009)
 Tamura, Bittner, Burghardt, J. Chem. Phys. 126, 021103 (2007), Gindensperger, Köppel, Cederbaum, J. Chem. Phys. 126, 034106 (2007)
 Cederbaum, Gindensperger, Burghardt, Phys. Rev. Lett., 94, 113003 (2005), Garg, Onucic, Ambegaokar, J. Chem. Phys. 83, 4491 (1985)

$$\hat{H}_{SB} + \hat{H}_B = \hat{s} \sum_i c_n \hat{x}_n + \hat{H}_B \longrightarrow D \hat{s} \hat{X}_1 + d_{12} \hat{X}_1 \hat{X}_2 + \dots + \hat{X}_M \text{-residual bath}$$

- orthogonal coordinate transformation $\hat{X} = \mathbf{T}\hat{x}$
- short-time dynamics captured by first few effective modes
- truncate hierarchical chain to define approximate, reduced-dimensional model
- (quasi-)Markovian closure
- connection to TD-DMRG!

Unitary Propagation vs. Master Equations

- ① explicit, multidimensional dynamics for the full system + bath space:
 wavefunction $\psi_{SB}(t)$ or density operator $\hat{\rho}_{SB}(t) = \sum_n p_n |\psi_{n,SB}(t)\rangle \langle \psi_{n,SB}(t)|$
 → typically (ML-)MCTDH

Meyer, Manthe, Cederbaum, Chem. Phys. Lett. **165**, 73 (1990), Beck et al., Phys. Rep. **324**, 1 (2000)

- ② reduced dynamics (master equation) methods: $\hat{\rho}_S(t) = \text{Tr}_B \hat{\rho}_{SB}(t)$
 → typically Hierarchy of Equations of Motion (HEOM)

Tanimura, J. Phys. Soc. Jpn. **75**, 082001 (2006)

- ③ intermediate methods: explicit treatment of subsystem + effective-mode (E) part of the bath + master equation for residual (B') bath:²

$$\frac{\partial \hat{\rho}_{SE}}{\partial t} = -\frac{i}{\hbar} [\hat{H}_{SE}, \hat{\rho}_{SE}(t)] + \hat{L}_{\text{diss}}^{(B')} \hat{\rho}_{SE}(t) \quad ; \quad \hat{\rho}_{SE}(t) = \text{Tr}_{B'} \hat{\rho}_{SEB'}(t)$$

²e.g., Caldeira-Leggett: $\hat{L}_{\text{diss}}^{(B')} \hat{\rho}_{SE} = -i\frac{\gamma}{\hbar} [\hat{X}_E, [\hat{P}_E, \hat{\rho}_{SE}]_+] - \frac{2\gamma M kT}{\hbar^2} [\hat{X}_E, [\hat{X}_E, \hat{\rho}_{SE}]]$

Unitary Dynamics: MCTDH

$$\Psi(r, t) = \sum_J A_J(t) \Phi_J(r, t) \equiv \sum_{j_1=1}^{n_1} \dots \sum_{j_N=1}^{n_N} A_{j_1 \dots j_N}(t) \prod_{\kappa=1}^N \phi_{j_\kappa}^{(\kappa)}(r_\kappa, t)$$

- **Multi-Configuration Time-Dependent Hartree**: tensor approximation scheme
Meyer, Manthe, Cederbaum, Chem. Phys. Lett. **165**, 73 (1990), Beck et al., Phys. Rep. **324**, 1 (2000)
- EoM's from the Dirac-Frenkel variational principle: $\langle \delta\Psi | \hat{H} - i\frac{\partial}{\partial t} | \Psi \rangle = 0$
- MCTDH takes one to **50-100 modes**; exponential scaling alleviated
- restriction on the form of the potential: sums over products
- related multi-layer variant (**ML-MCTDH**) goes up to **1000 modes**
Wang, Thoss, J. Chem. Phys. **119**, 1289 (2003), Manthe, J. Chem. Phys. **128**, 164116 (2008), Vendrell, Meyer, *ibid* **134**, 044135 (2011)
- related **MCTDH-F** (fermion) and **MCTDH-B** (boson) methods
Kato, Kono, Chem. Phys. Lett. **392**, 533 (2004), Nest, Klamroth, Saalfrank, J. Chem. Phys. **122**, 124102 (2005)
Alon, Streltsov, Cederbaum, Phys. Lett. A **362**, 453 (2007)
- **density matrix** variant
Raab, Burghardt, Meyer, J. Chem. Phys. **111**, 8759 (1999)
- **hybrid** approaches: e.g., Gaussian-based variant (**G-MCTDH**, **vMCG**)
Burghardt, Meyer, Cederbaum, J. Chem. Phys. **111**, 2927 (1999), Worth, Burghardt, Chem. Phys. Lett. **368**, 502 (2003)

Multi-Layer(ML)-MCTDH: Hierarchical Tensor Form

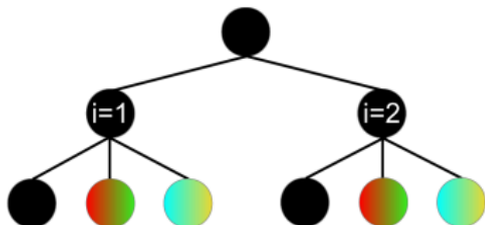
$$\Psi(r,t) = \sum_J A_J(t) \Phi_J(r,t) = \sum_J A_J(t) \prod_{\kappa=1}^M \varphi_{j_\kappa}^{(\kappa)}(r_\kappa, t)$$

where the 1st-layer SPFs $\varphi_{j_\kappa}^{(\kappa)}$ are now built as superpositions of 2nd-layer SPFs,

$$\varphi_{j_\kappa}^{(\kappa)}(r_\kappa, t) = \sum_L B_{j,L}^{(\kappa)}(t) \Phi_L^{(\kappa)}(r_\kappa, t) = \sum_L B_{j,L}^{(\kappa)}(t) \prod_{\mu} \varphi_{l_\mu}^{(\kappa,\mu)}(r_{\kappa\mu}, t)$$

... and so on ...

- intra-SPF correlations *via* MCTDH form
- continue to higher orders: ML-MCTDH
- “hierarchical Tucker format”
- recent Gaussian-based variant Römer, Ruckebauer, Burghardt, J. Chem. Phys. 138, 064106 (2013)

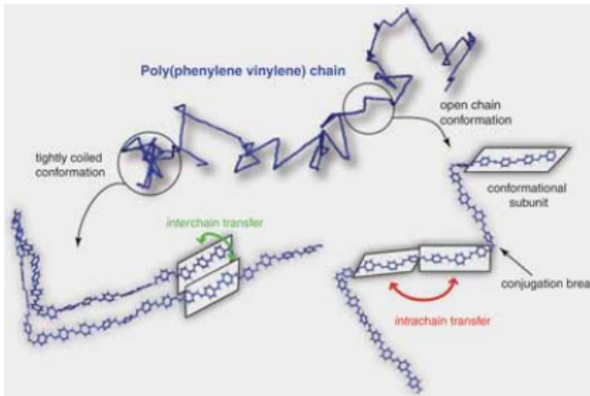


Wang, Thoss, J. Chem. Phys. 119, 1289 (2003), Manthe, J. Chem. Phys. 128, 164116 (2008), Vendrell, Meyer, J. Chem. Phys. 134, 044135 (2011)

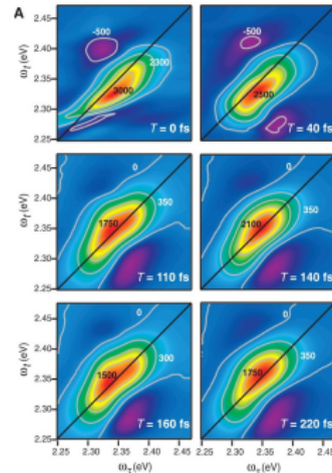
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Charge Transfer Excitons and Interfacial Charge Separation

Exciton Dynamics in Organic Semiconducting Polymers



Collini, Scholes, Science 323, 369 (2009)



~ 0.1 -1 ps: coherent intra-chain excitation energy transfer (EET) dynamics

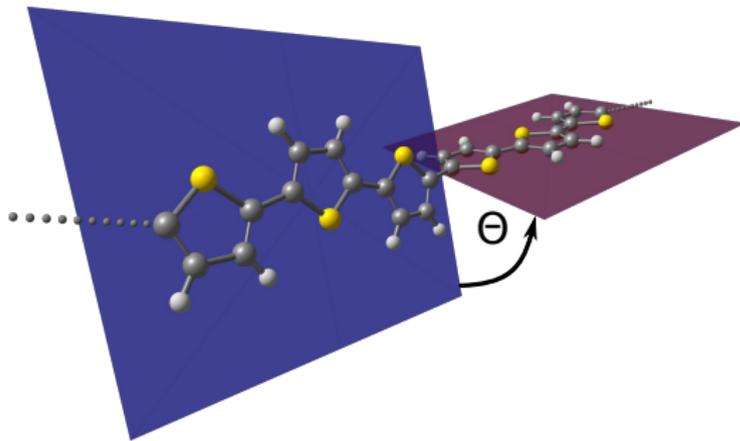
~ 0.1 -1 ps: self-trapped exciton-polaron states

~ 0.1 -few ps: torsional geometry relaxation interfering with EET

~ 1 -10 ps: inter-chain EET

\sim ps-ns: thermally assisted hopping

Exciton Migration Guided by Conformational Dynamics



- full quantum dynamical study for small oligomers (5-20 units)
- Oligothiophene (OT) and Oligo-Phenylenevinylene (OPV)
- ML-MCTDH (up to 50 states, 100 vibrational modes)
- monomer-based, *ab initio* parametrized Hamiltonian

Binder, Wahl, Römer, Burghardt,

Faraday Discuss 163, 205 (2013)

Panda, Plasser, Aquino, Burghardt, Lischka

J. Phys. Chem. A, 117, 2181 (2013)

Wahl, Binder, Burghardt

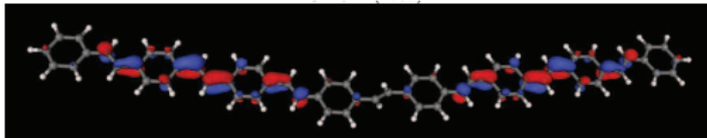
Comp. Theor. Chem. 1040, 167 (2014)

- Is the transfer dynamics on ultrafast time scales **coherent** or of hopping type?
- Is a trapped **exciton-polaron** generated and if so, on which time scale?
- Is the **spectroscopic unit** concept valid?

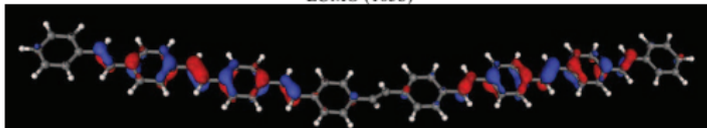
see also: Tretiak, Saxena, Martin, Bishop, Conformational Dynamics of Photoexcited Conjugated Molecules, Phys. Rev. Lett. 89, 097402 (2002)

Electronic Structure: Trapping in OPV Oligomers

Collaboration with H. Lischka, F. Plasser (Vienna/Texas Tech/Tianjin University)

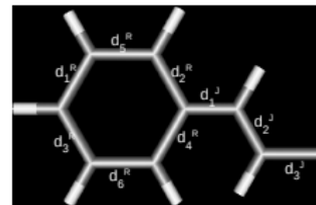
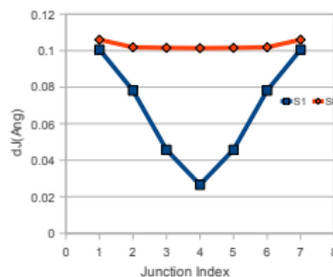
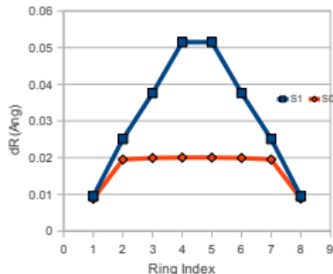


LUMO (105b)



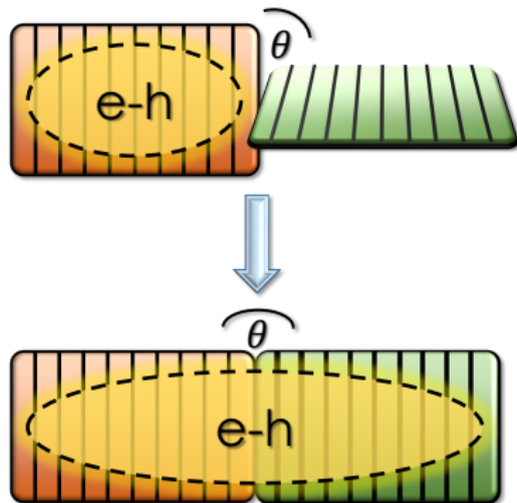
HOMO (106a)

- high-level electronic structure methods (ADC(2), CC2, MRCI)
- exciton trapping, due to BLA modes, described correctly



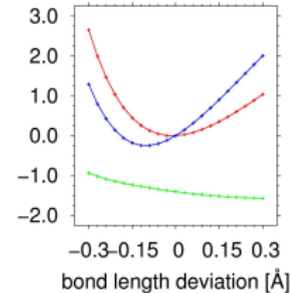
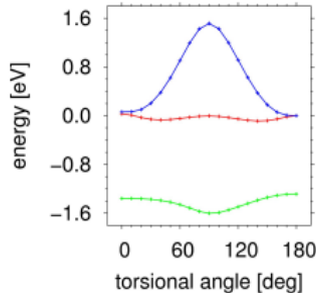
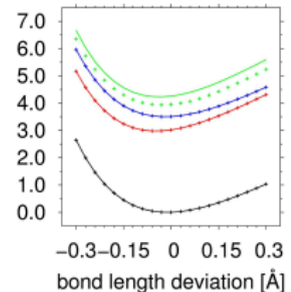
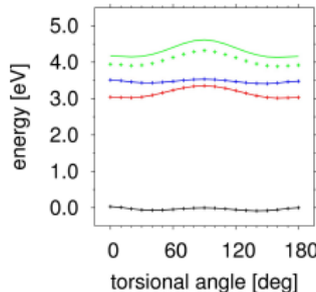
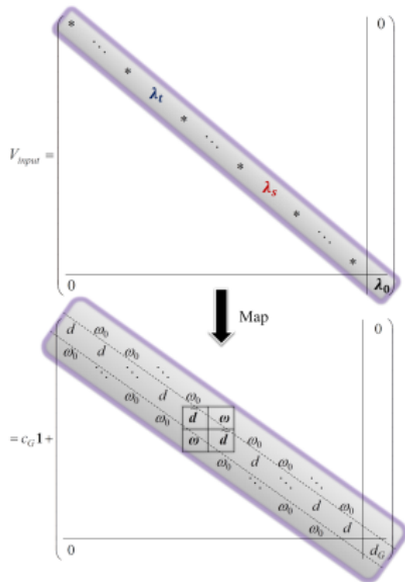
Dynamics: Test Case OT-20

- Do we see trapped exciton-polarons in the dynamics?
- How exactly does the exciton migrate as the conjugation break “heals”?
- How does the spatial extension of the exciton change as a function of conformational (torsional) fluctuations?



Monomer representation:
most unbiased picture to answer
these questions!

Relevant Coordinates: Torsions, CC Stretch, Ring Modes



- analytical, pointwise mapping of oligomer PES's onto a Frenkel model
- diabaticization in terms of solution to an inverse eigenvalue problem
- applicable to “extended Hückel systems” of J / H / HJ-aggregate type

Ab initio Frenkel-Holstein Hamiltonian

$$\hat{H} = \sum_{n,n'=1}^N (\delta_{n,n'} \hat{T} + \delta_{n,n'} \hat{V}_n^{\text{site}} + \hat{V}_{n,n'}^{\text{exc}}) |n\rangle \langle n'| + \hat{H}_{\text{bath}} \hat{1}$$

- kinetic energy in curvilinear coordinates (using TNUM code):

$$\hat{T} = \frac{1}{2} \left(\sum_{n=1}^N G_{xx} \hat{p}_{x_n}^2 + \sum_{n=1}^{N-1} (G_{yy} \hat{p}_{y_{n,n+1}}^2 + G_{\theta\theta} \hat{p}_{\theta_{n,n+1}}^2) + 2 \sum_{n=1}^N G_{xy} \hat{p}_{x_n} (\hat{p}_{y_{n,n+1}} + \hat{p}_{y_{n,n-1}}) \right)$$

- site energies:

$$\hat{V}_n^{\text{site}}(\{\hat{x}, \hat{y}, \hat{\theta}\}) = \hat{V}_0(\{\hat{x}, \hat{y}, \hat{\theta}\}) + \hat{\Delta}_n(\hat{x}_n, \hat{y}_{n,n\pm 1}, \hat{\theta}_{n,n\pm 1})$$

with the difference potential

$$\hat{\Delta}_n(\hat{x}_n, \hat{y}_{n,n\pm 1}, \hat{\theta}_{n,n\pm 1}) = c_E + \hat{v}_E(\hat{x}_n, \hat{y}_{n,n\pm 1}, \hat{\theta}_{n,n\pm 1}) - \hat{v}_G(\hat{x}_n, \hat{y}_{n,n\pm 1}, \hat{\theta}_{n,n\pm 1})$$

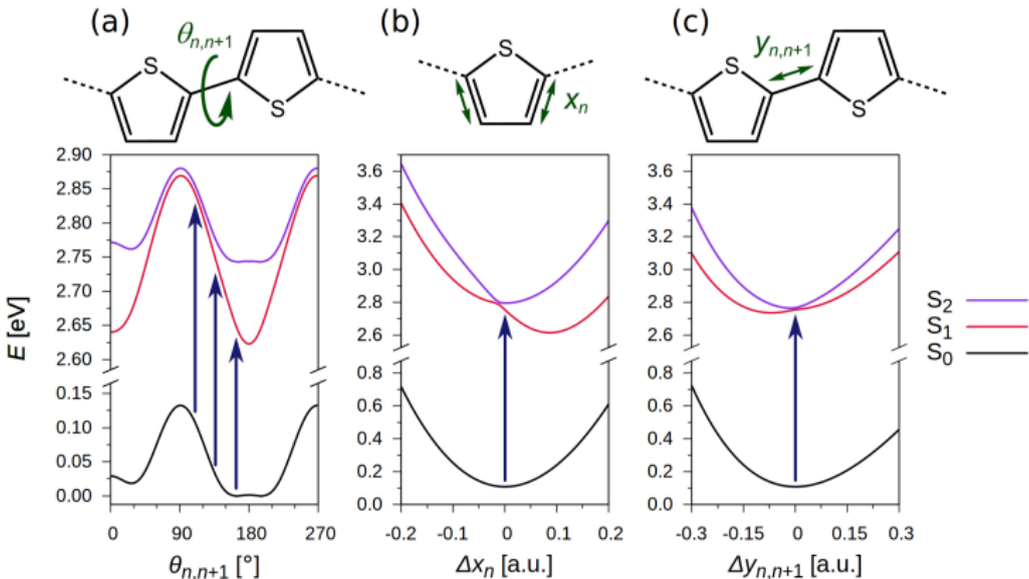
- excitonic coupling:

$$\hat{V}_{n,n\pm 1}^{\text{exc}}(\hat{\theta}_{n,n\pm 1}) = \hat{w}(\hat{\theta}_{n,n\pm 1}) \quad \leftarrow \text{large (of the order of 1 eV)}$$

reduces to standard Frenkel-Holstein model if

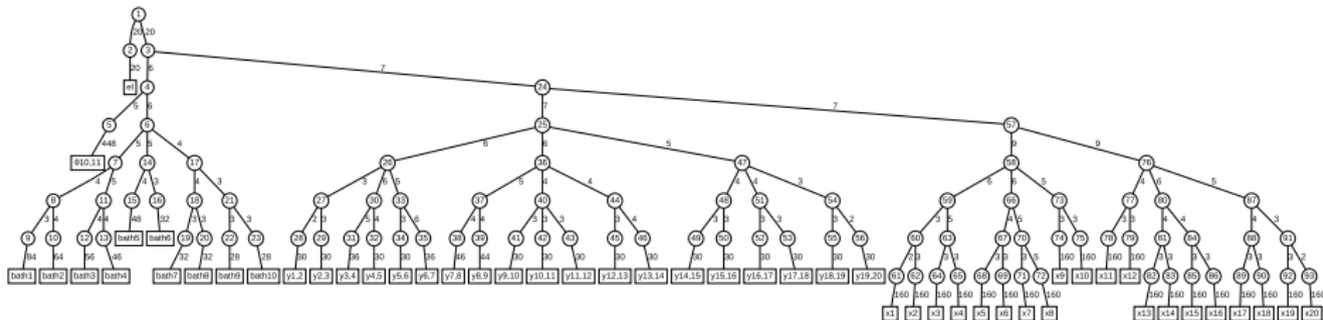
- the excitonic coupling is constant
- a single *site-local* mode per site is included
- the monomer potentials conform to shifted harmonic-oscillator potentials

Relevant Coordinates: Torsions, CC Stretch, Ring Modes



- high-dimensional PES as a function of site-local and site-correlated modes
- 20 monomer sites, 50 phonon modes

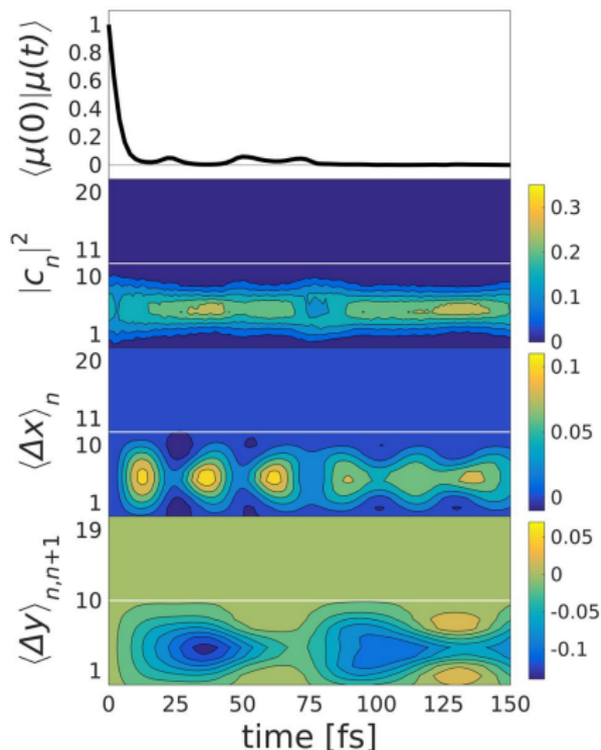
Quantum Dynamics: Hierarchical Multi-Layer MCTDH Tree



- **ML-MCTDH** (20 states, 50 modes)
- **7 layers**, 2-10 single-particle functions (SPFs) – highly correlated system!
- combined electronic particle (here, 20 states)
- active torsional mode as isolated particle (long DVR grid)
- delocalized initial condition prepared by imaginary time propagation

Binder, Lauvergnat, Burghardt, Phys. Rev. Lett. 120, 227401 (2018), Supp. Mater.

Quantum Dynamics: 20-Site J-Aggregate with Central Torsion



Binder, Lauvergnat, Burghardt, Phys. Rev. Lett. 120, 227401 (2018)

- ML-MCTDH calculations: 20 states, 50 modes (6-layer set-up)
- **earliest time scale**: exciton trapping (contraction by ~ 3 sites)
- high-frequency modes adapt to exciton: **quasi-stationary trapping**
- LEGS = local exciton ground state: nodeless left-localized exciton

Tozer, Barford, JPCA 116:10310 (2012)

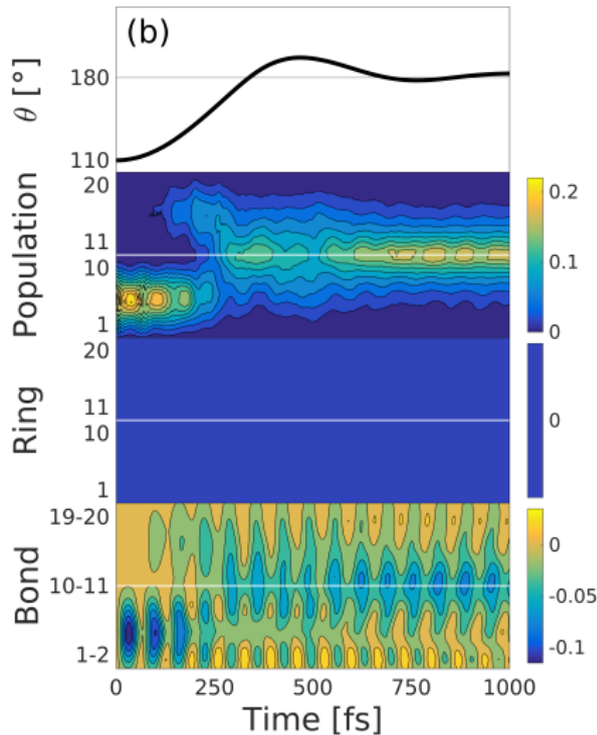
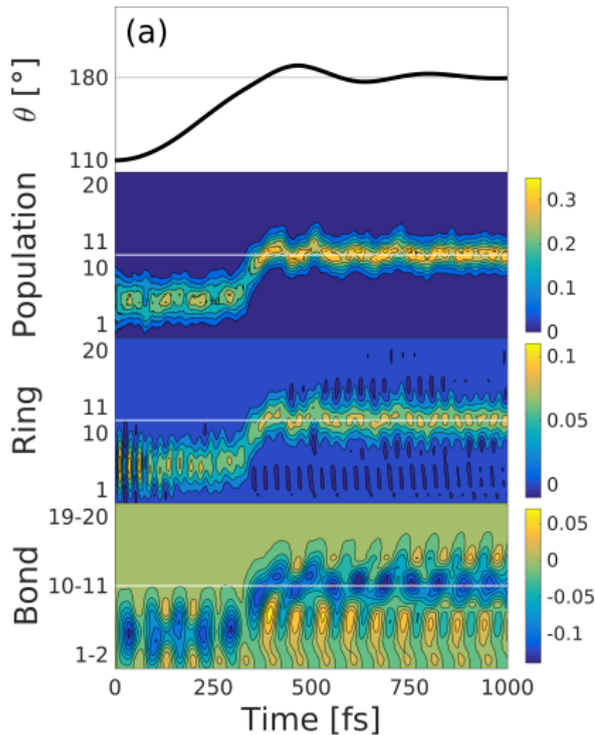
- ultrafast decay of transition dipole autocorrelation function (~ 10 fs): $\langle \mu(0)\mu(t) \rangle = |\mu|^2 \langle \psi_{\text{exc}}(0) | \psi_{\text{exc}}(t) \rangle$
- relates to anisotropy decay: ~ 40 fs

Grage et al., Phys. Rev. B, 67, 205207 (2003)

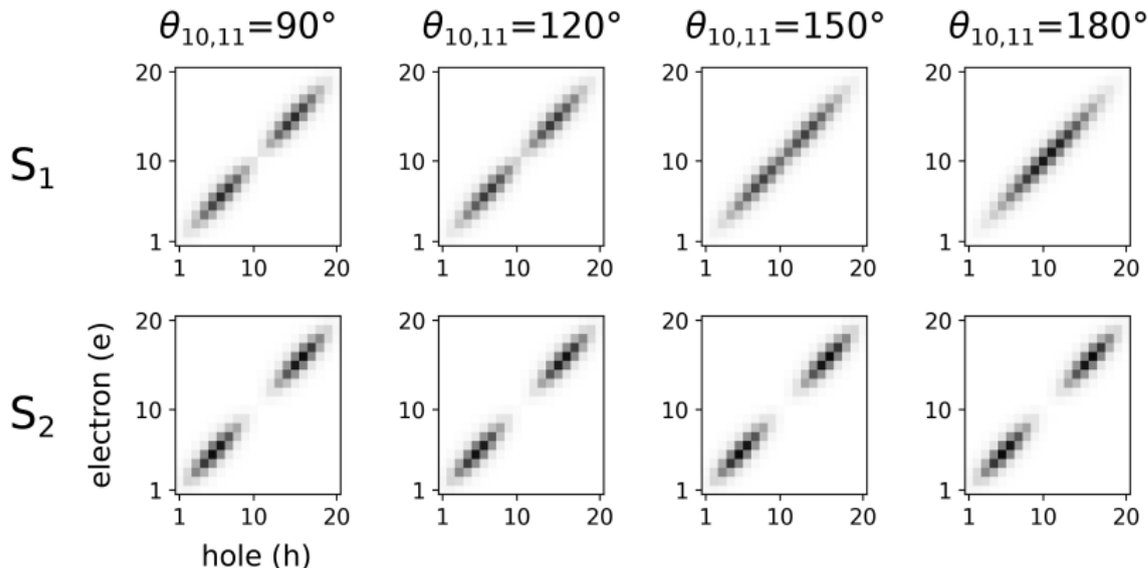
Quantum Dynamics: 20-Site J-Aggregate with Central Torsion

C-C inter-monomer mode + local C=C + torsion + bath

C-C inter-monomer mode + torsion + bath



Electronic Structure – Torsion Dependence



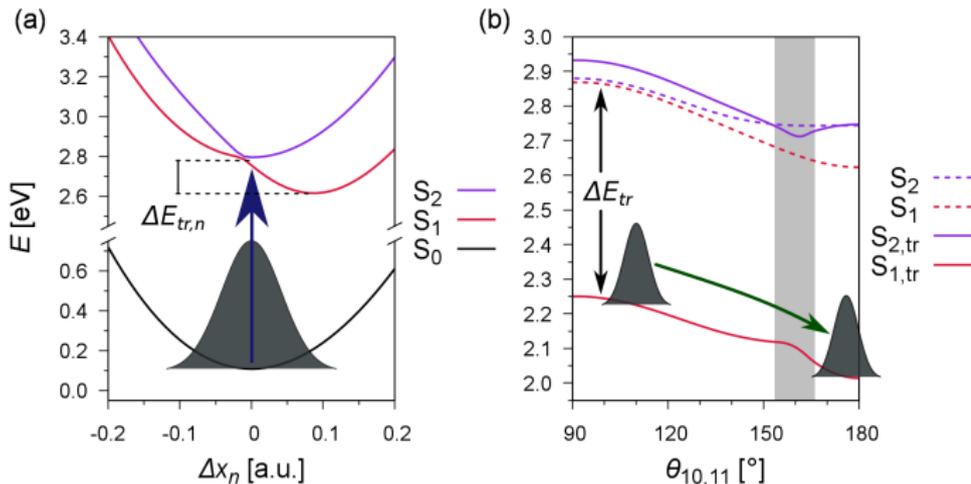
- transition density analysis

Panda, Plasser, Aquino, Burghardt, Lischka, JPCA (2013), Binder, Lauvergnat, Burghardt, Phys. Rev. Lett. 120, 227401 (2018)

- particle-in-the-box type e - h states
- marked dependence on torsion
- LEGS = local exciton ground state: nodeless S_1 exciton @ 180°

Tozer, Barford, JPCA 116:10310 (2012)

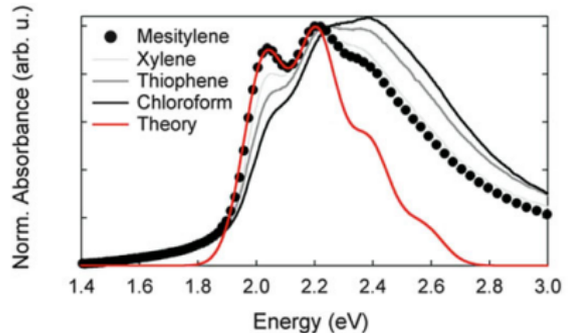
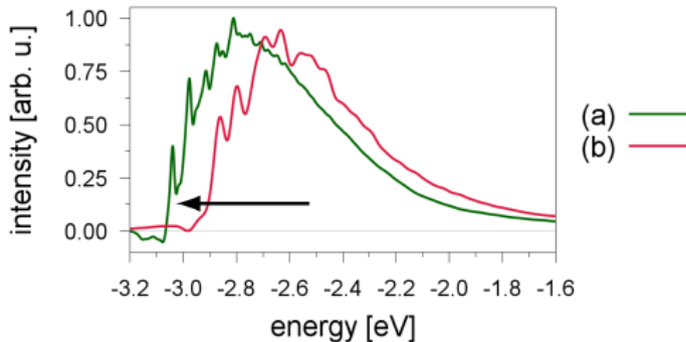
Two Time-Scale Process



- dynamics is essentially happening on coupled S₁/S₂ surfaces
- effective torsion potentials including stabilization due to trapping (S_{1,tr}/S_{2,tr})¹
- initial left/right localized state = superposition of S₁/S₂
- energy loss due to external bath acting on torsional mode
- **exciton-polaron**: quasi-stationary trapping at all stages of the dynamics

¹i.e., "polaron transformed" potentials

Initial vs. Final States: Absorption Spectra

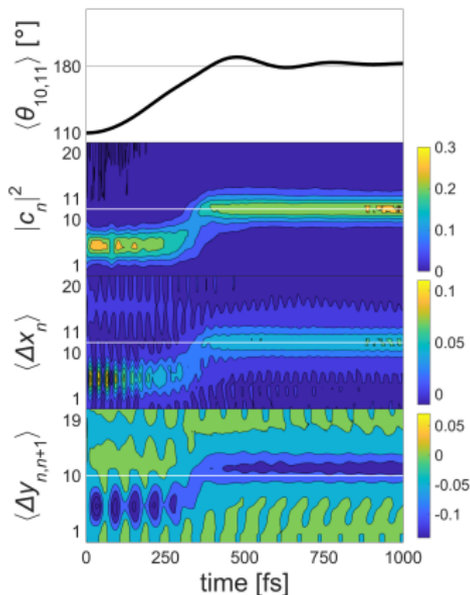


Tremel, Ludwigs, Adv. Polym. Sci. 265, 39 (2014)

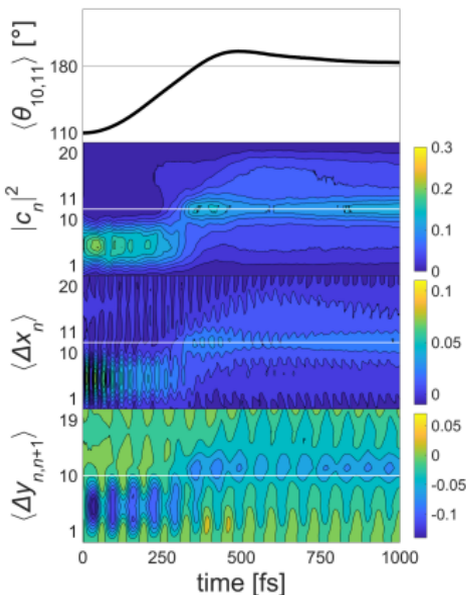
- spectra calculated from Fourier transform of $C(t) = \langle \psi(0) | \psi(t) \rangle$:
 - (a) full 20-mer system: torsionally relaxed, stationary state
 - (b) left-localized 10-mer fragment for initial defect structure
 (a) ← (b): downhill energy transfer
- dominant vibronic signature of ring-breathing mode

Semiclassical SQC/MM dynamics (T=0K)

SQC/MM – single trajectory



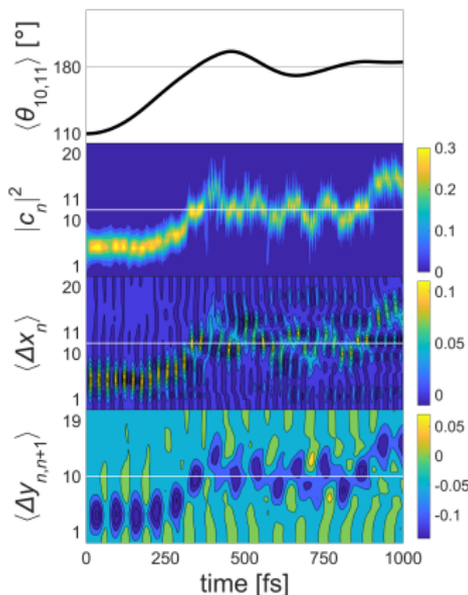
SQC/MM – Wigner average



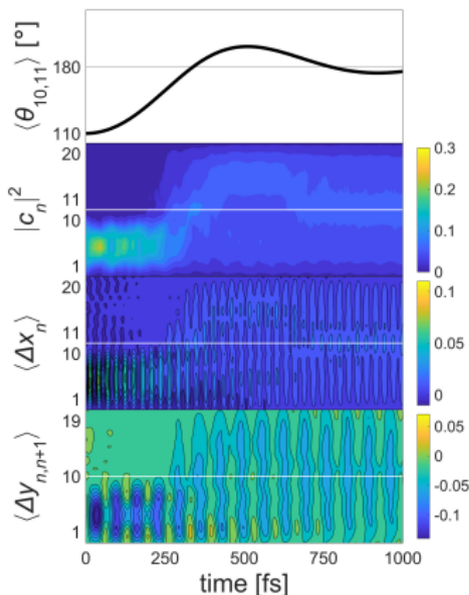
- SQC/MM = Symmetrical Quasi-Classical / Meyer-Miller model
- single-trajectory result close to MCTDH, but Wigner average “fuzzy”

Ehrenfest/Langevin dynamics (T=0K)

Ehrenfest – single trajectory



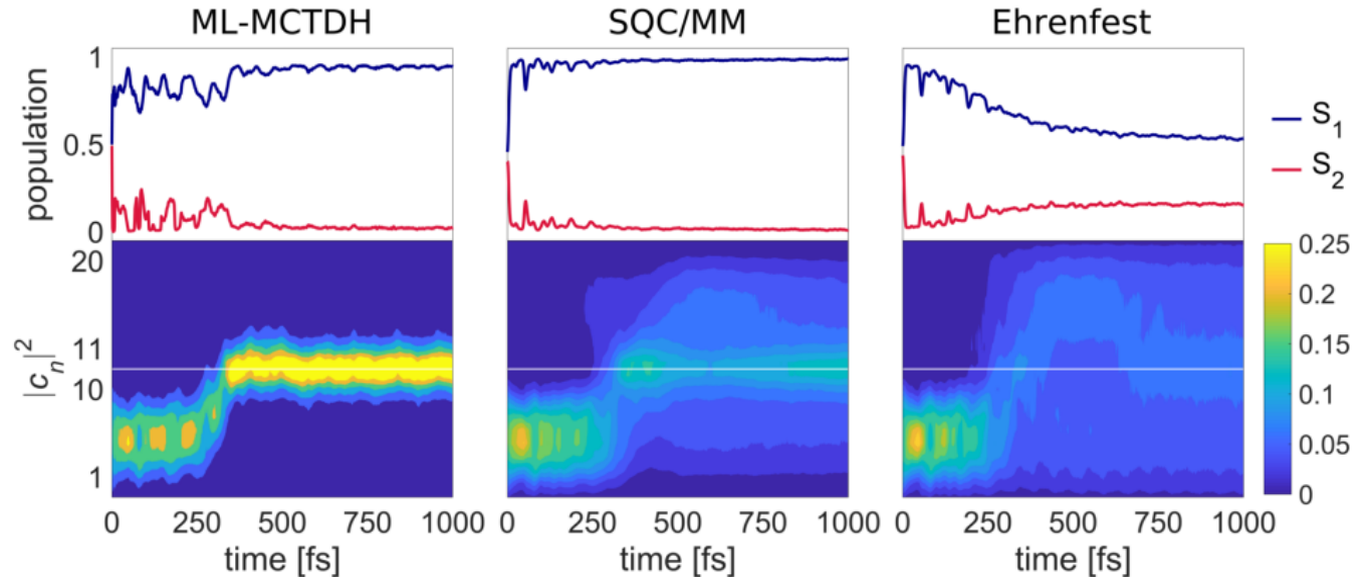
Ehrenfest – Wigner average



- single-trajectory dynamics exhibits fluctuations
- Wigner average “fuzzy” – due to Wigner sampling of high-frequency modes

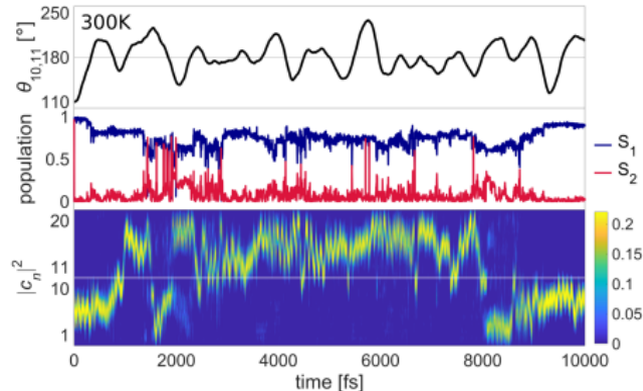
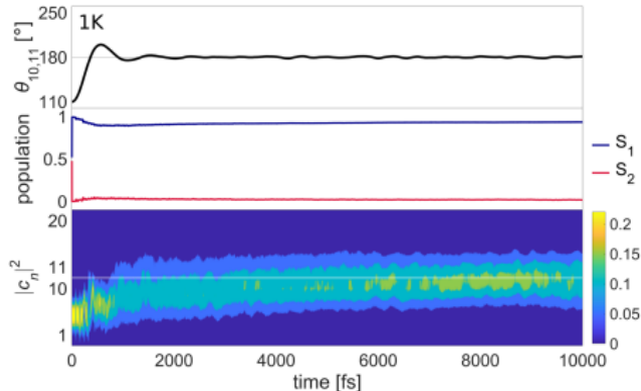
Liang, Cotton, Binder, Hegger, Burghardt, Miller, J. Chem. Phys., in press (2018)

Adiabatic Populations (T=0K)



- SQC/MM reproduces the adiabatic populations quite accurately
- Ehrenfest shows severe shortcomings (related to detailed-balance problem)
- both methods incur problems due to ZPE of high-frequency modes

Temperature Effects: Ehrenfest/Langevin dynamics



- single-trajectory simulation, with ZPE of high-frequency modes removed
- exciton migration at higher T is related to repeated non-adiabatic events
- interplay of torsional fluctuations and trapping explains observations
- quantum benchmark simulations needed (*via* random-phase wavefunctions, or thermofield method, combined with MCTDH)

Wahl, Hegger, Binder, Burghardt, in preparation

Polarons in π -Conjugated Polymers: Anderson or Landau?

William Barford,^{*,1} Max Marcus,^{1,2} and Oliver Robert Tozer^{1,3}

"We show that the high-frequency C-C bond oscillation only causes Landau polarons for a very narrow parameter regime; generally we expect disorder to dominate and Anderson polarons to be a more applicable description."

J. Phys. Chem. A 120, 615 (2016)

Excitons in conjugated polymers: Do we need a paradigm change?

Wichard J. D. Beenken

"The fact that we could not find partition of excitons by structural defects – except of rare gauche defects and accidental chemical defects – leads us to the conclusion that we have to search for new mechanisms."

Phys. Status Solidi A 206, 2750 (2009)

Our interpretation: Exciton-polarons driven by defects and fluctuations

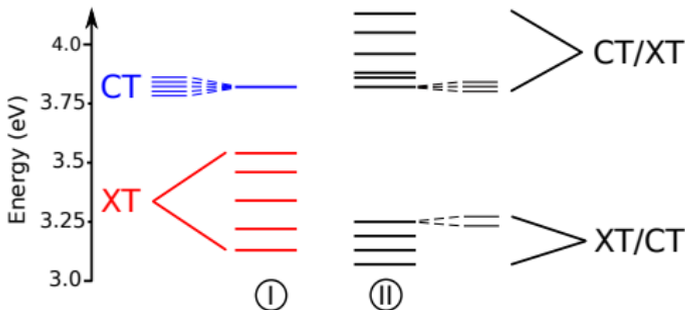
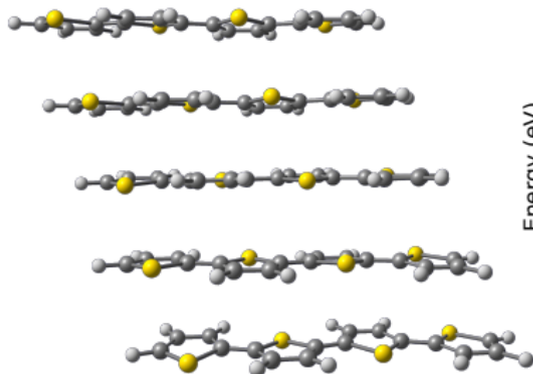
- exciton-polaron species: "exciton dressed by a cloud of local oscillators"
- typical delocalization length: 5-10 units (in line with experiment)
- Landau polaron (or Holstein "large" polaron) subject to disorder
- torsional defects confine excitons to sublattices ("spectroscopic units")
- hopping-type transition between exciton-polaron states induced by torsion
- elementary step is of "coherent hopping" type: highly correlated dynamics!

Topics

- 1 Photoinduced Energy & Charge Transfer in Functional Organic Materials
Goal: First-Principles Approach to Organic Photovoltaics
Electron-Hole Lattice Models & Vibronic Coupling
Quantum Dynamics in Many Dimensions
- 2 Case Studies I: Exciton Migration Guided by Conformational Dynamics
Torsion-Induced Intra-Chain Exciton Migration
Ultrafast, Coherent Exciton-Polaron Dynamics
Do Semiclassical and Ehrenfest-Langevin Dynamics Work?
- 3 Case Studies II: Charge Transfer Excitons & Charge Separation
Charge Transfer Excitons in Neat Polythiophene
Collective Vibronic Effects
Charge Transfer Excitons and Interfacial Charge Separation

Charge Transfer Excitons in Neat Regioregular Polythiophene

experiment: Reid et al., Chem. Mater. 26, 561 (2014), De Sio et al., Nature Comm. 7, 13742 (2016)

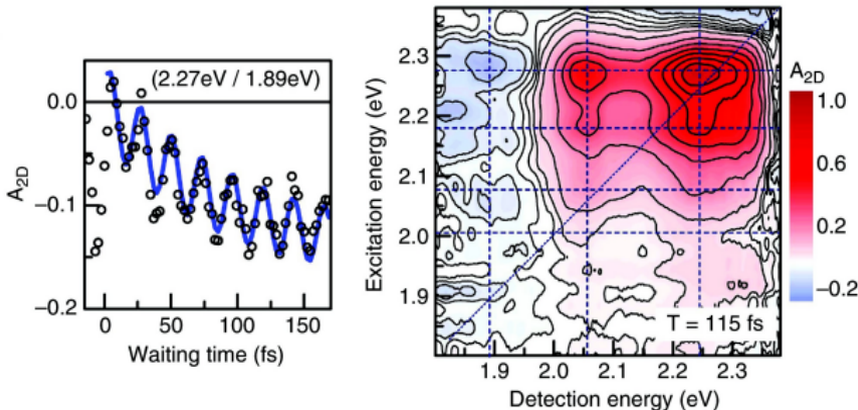


I: local adiabatic, II: full adiabatic

Popp, Polkehn, Tamura, Burghardt, to be submitted

- inter-chain CTX states favored in PT (as compared with, e.g., PPV)
- electronic structure (ADC(2), TDDFT): low-energy inter-chain CT states
- representative quantum dynamics calculations for $(OT)_n$, $n = 3, 5$
- diabaticization + Linear Vibronic Coupling (LVC) model
- ML-MCTDH for up to 13 electronic states, 196 modes

Charge Transfer Excitons in Neat Regioregular Polythiophene



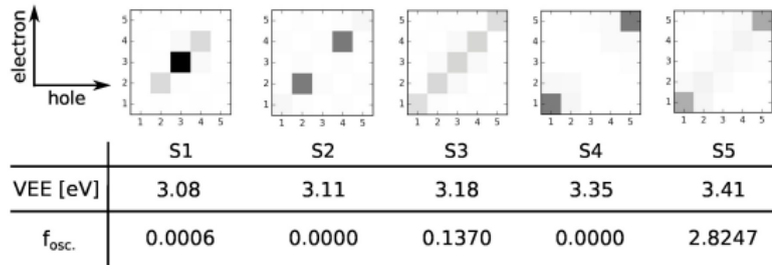
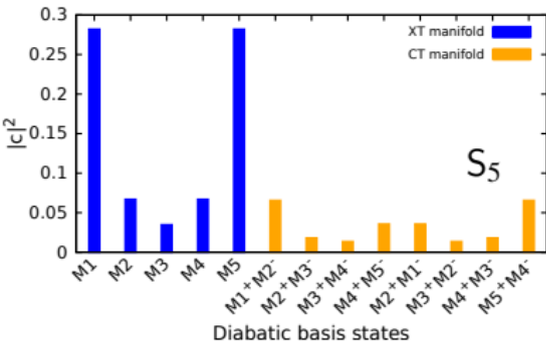
2D electronic spectroscopy
P3HT (thin film)

De Sio et al., Nature Comm. 7, 13742 (2016)

- polaron cross-peak signal assigned to inter-chain CTX states
- sustained high-frequency oscillations (23 fs periodicity)
- simulated using 2-state/1-mode model
- similar observations by Scholes & collaborators
- **What is the origin of the persistent oscillations?**

Song et al., JCP 142, 212410 (2015)

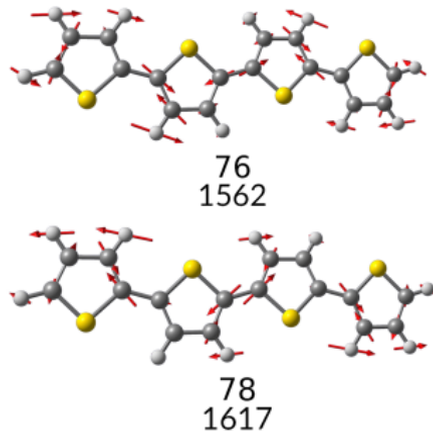
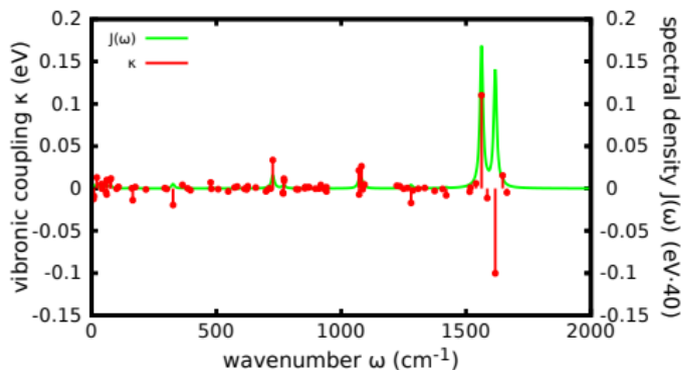
Modified H-Aggregate: What's the Bright State?



Popp, Polkehn, Tamura, Burghardt, to be submitted

- (OT)₅ stack: significant mixing of XT and CT states
- bright state (S_5) looks distinctly different from typical H-aggregate (inverted curvature of nodeless exciton wavefunction) Hestand and Spano, J. Chem. Phys. 143, 244707 (2015)
- transition densities for H-type dimer *via* TheoDORÉ program (F. Plasser)
- very good agreement between ADC(2) and TDDFT/ ω B97XD results
- in line with benchmark study by Lischka & collaborators JCTC 10, 3280 (2014)

Spectral Densities (SD's)

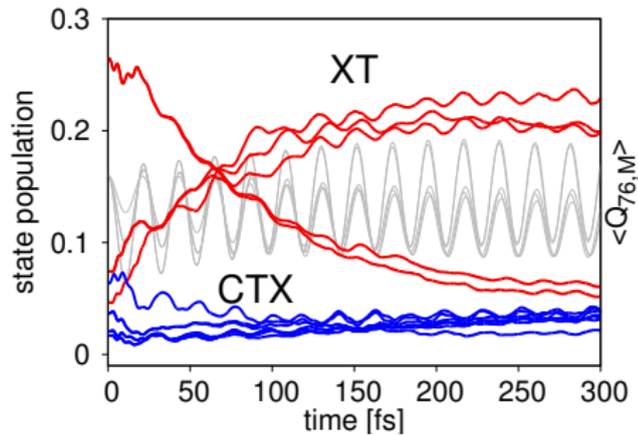
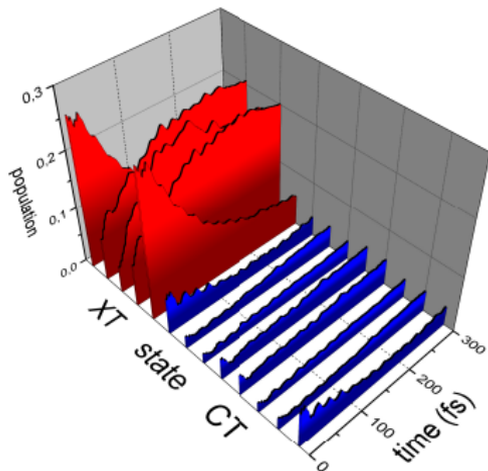


- SD's calculated from state-specific Frank-Condon gradients
- SD's show large amplitude (\sim Huang-Rhys factor) for CC stretch modes
- SD's similar for various electronic states
- use SD's to parametrize Linear Vibronic Coupling (LVC) Hamiltonian
- ML-MCTDH: 13 states/196 modes (or 78 effective modes)

LVC Model – Full Quantum Dynamics (13 States, 196 modes)

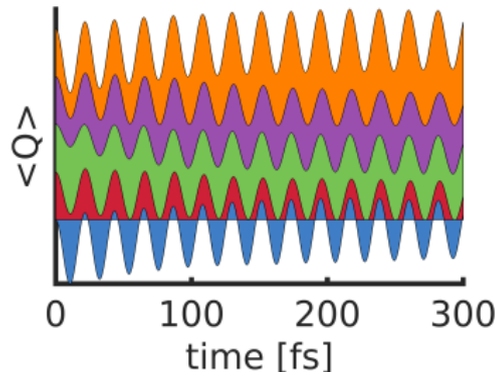
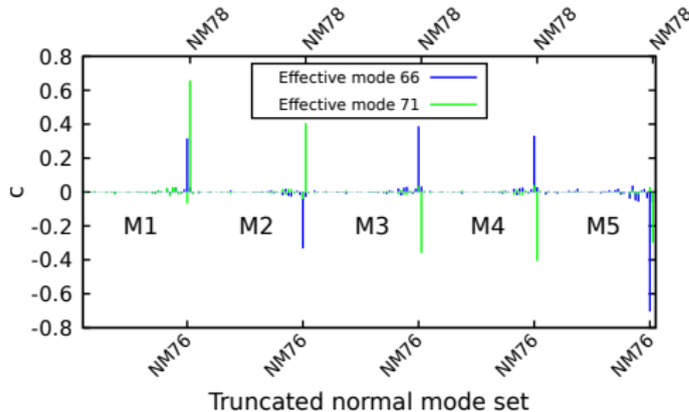
$$\hat{H} = \hat{H}^{\text{on-site}} + \hat{H}^{\text{coup}} + \hat{H}^{\text{e-ph}}$$

$$\hat{H}^{\text{coupl}} = j \sum_{n=1}^N |\text{XT}_n\rangle \langle \text{XT}_{n+1}| + \kappa_1 \sum_{n=1}^{N-1} |\text{XT}_n\rangle \langle \text{CTX}_{n+1,n}| + \kappa_2 \sum_{n=1}^{N-1} |\text{XT}_n\rangle \langle \text{CTX}_{n,n+1}| + h.c.$$



- two diabatisation schemes Tamura, JPCA, 120, 9341 (2016), Blancafort, Voityuk, JCP 140, 095102 (2014)
- pronounced oscillatory signature in all state populations

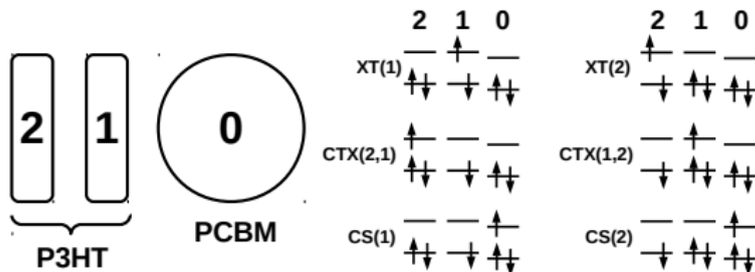
Collective Vibronic Response



- high-frequency normal modes of OT fragments are evolving in phase
- amplification of vibronic effects due to exciton delocalization
- effective modes: linear combinations of dominant normal modes
- generalized effective-mode transformation (correlated XT/CTX modes)

Popp, Polkehn, Tamura, Burghardt, to be submitted

Do CTX States Affect Interfacial Charge Generation?

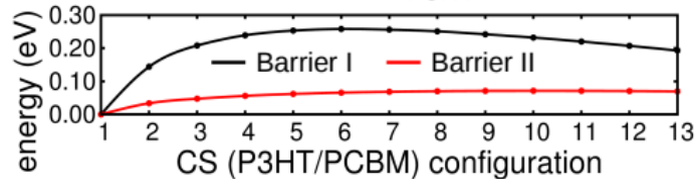
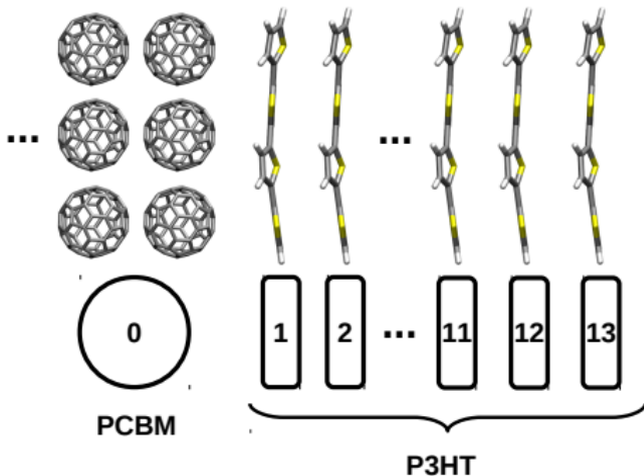


	XT(1)	XT(2)	CTX(2,1)	CTX(1,2)	CS(1)	CS(2)
XT(1)	0.100	0.100	0.357	0.139	0.200	0.007
XT(2)		0.100	0.139	0.357	0.014	0.013
CTX(2,1)			0.280	0.001	0.005	0.002
CTX(1,2)				0.230	0.019	0.165
CS(1)					0.000	0.102
CS(2)						0.140

Polkehn, Tamura, Burghardt, J. Phys. B, 51, 014003 (2018)

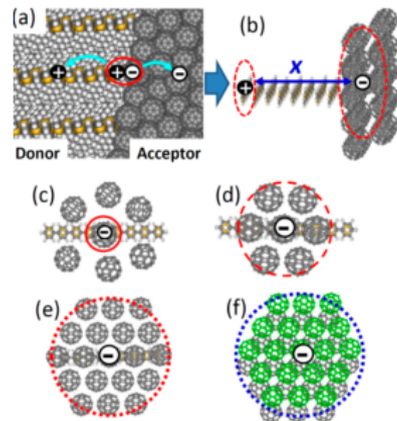
- CTX-to-CS transfer can circumvent interfacial XT-to-CS transfer step
- here: parameter determination via diabaticization by projection onto reference wavefunctions Tamura, JPCA 120, 9341 2016, Polkehn, Tamura, Burghardt, J. Phys. B, 51, 014003 (2018)

Interfacial Charge Generation in P3HT:PCBM Type Systems



Tamura, Burghardt, JACS (Communication) 135, 16364 (2013)

Huix-Rotllant, Tamura, Burghardt, J. Phys. Chem. Lett., 6, 1702 (2015)

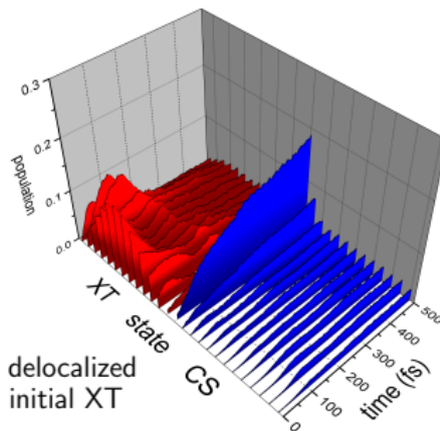
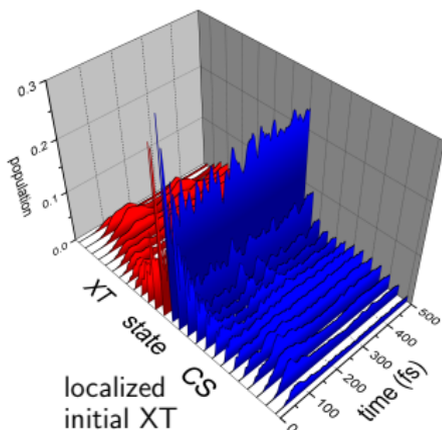


factors favoring ultrafast *e-h* separation:

- electron delocalization over fullerene aggregates: **strong decrease of barrier**
- hole delocalization on oligothiophenes
- XT delocalization in H-aggregate donor
- exciton (XT) excess energy: **“Hot CT”** mechanism

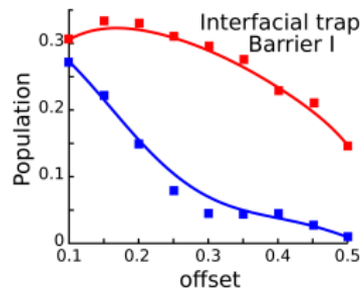
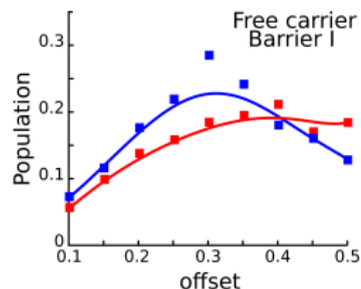
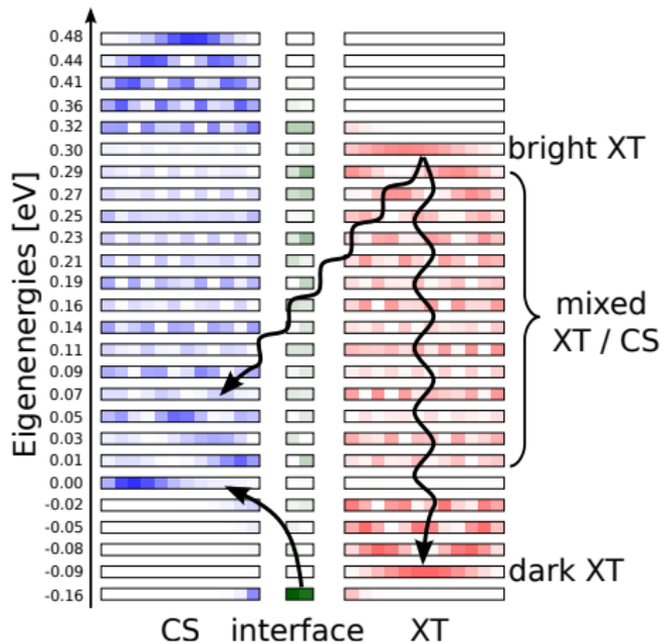
Interfacial Charge Separation in the Absence of CTX States

$$\hat{H} = \hat{H}_{\text{XT}_1-\text{CT}}(\mathbf{x}) + \sum_n \hat{H}_{\text{CS}}^{(n)}(\mathbf{x}) |\text{CS}_n\rangle \langle \text{CS}_n| + t(\mathbf{x}) (|\text{CS}_1\rangle \langle \text{CT}| + \sum_{nm'} |\text{CS}_n\rangle \langle \text{CS}_{n'}| + h.c.) \\ + \sum_n \hat{H}_{\text{XT}}^{(n)}(\mathbf{x}) |\text{XT}_n\rangle \langle \text{XT}_n| + j(\mathbf{x}) \sum_{n'} (|\text{XT}_n\rangle \langle \text{XT}_{n'}| + h.c.)$$



- CT/CS generation depends on exciton (de)localization
- ML-MCTDH calculations: 26 states/120 modes (barrier II)

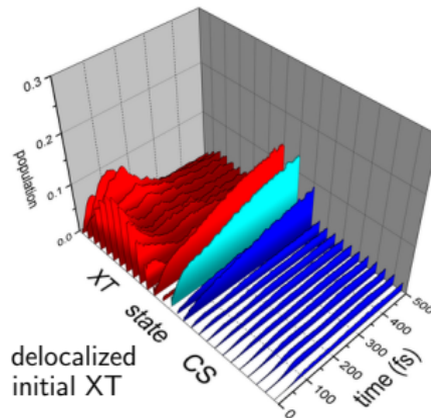
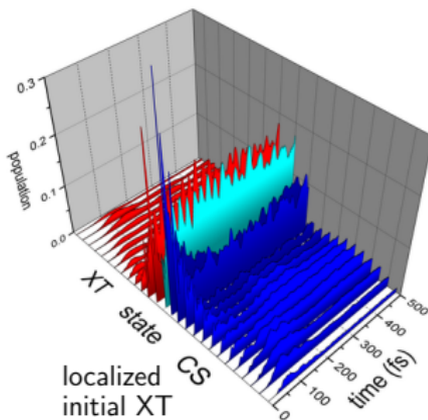
Electronic Eigenstate Picture



- interplay of delocalization, internal conversion, and charge transfer
- de/localized initial condition (blue/red) reduces/enhances interfacial trapping

Huix-Rotllant, Tamura, Burghardt, J. Phys. Chem. Lett., 6, 1702 (2015)

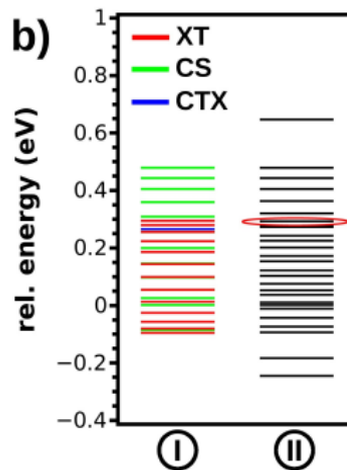
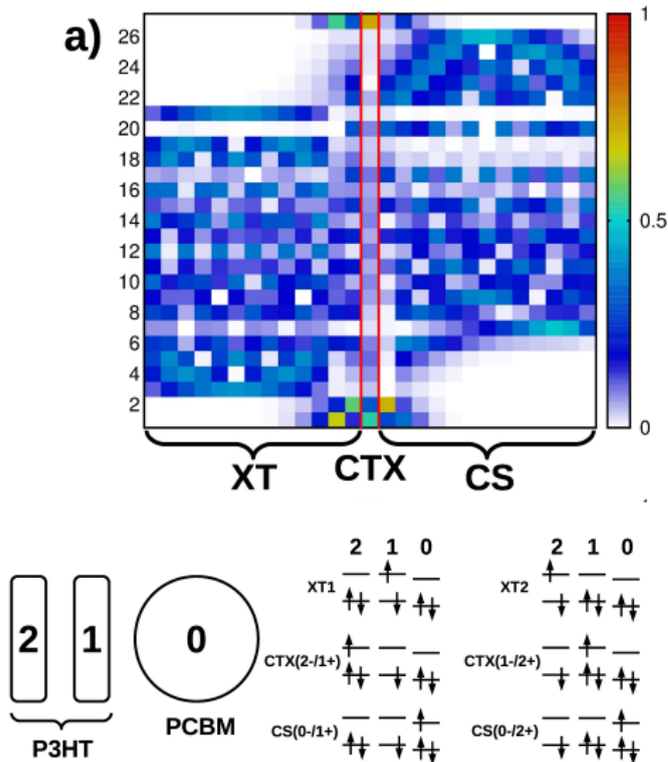
Interfacial Charge Separation in the Presence of CTX States



- CTX (turquoise) states emerge prominently as additional energetic traps
- ML-MCTDH calculations up to 182 states and 112 modes
- for a single CTX state: CS formation slightly reduced
- for larger models (50/182 states): reduction of CS yield is significant
- results depend in a sensitive fashion upon energetics and electronic couplings

Polkehn, Tamura, Burghardt, J. Phys. B, Special Issue "Light Energy Conversion, Light Harvesting", 51, 014003 (2018).

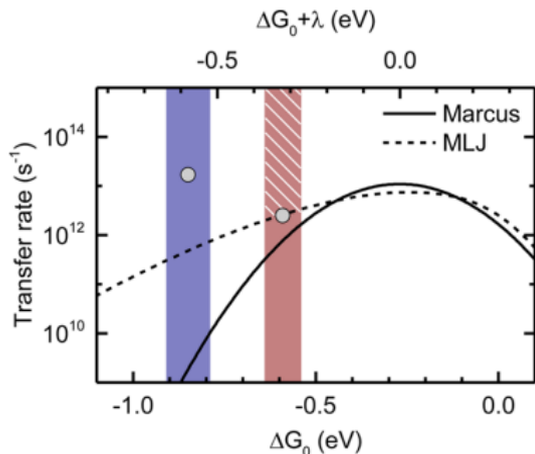
P3HT/PCBM Model Including CTX States



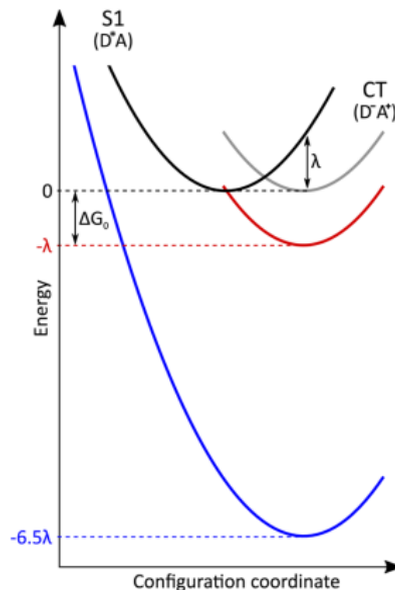
- 182 states/112 modes
- OT charge transfer excitons open new pathways for carrier formation
- but also act as energetic traps

Polkehn, Tamura, Burghardt, J. Phys. B, 51, 014003 (2018)

Marcus Theory Doesn't Work for Ultrafast Exciton Break-Up



Unger et al., J. Phys. Chem. C 121, 22739 (2017)



- Marcus (M) & Marcus-Levich-Jortner (MLJ) rates:

$$k_M = \frac{|V|^2}{\hbar} \sqrt{\frac{\pi}{\lambda k_B T}} \exp\left[-\frac{(\lambda + \Delta G_0)^2}{4\lambda k_B T}\right] ; \quad k_{MLJ} = \frac{|V|^2}{\hbar} \sqrt{\frac{\pi}{\lambda_0 k_B T}} \sum_{v=0}^{\infty} \frac{e^{-S} S^v}{v!} \exp\left[-\frac{(\lambda + \Delta G_0)^2}{4\lambda k_B T}\right]$$

Summary

① Molecular-Level Approach to Organic Photovoltaics

- $e-h$ lattice model: highlights fragment properties + molecular packing
- extensions to conical intersections, multi-exciton states¹ etc.
- accurate on-the-fly dynamics remains highly challenging

¹See our work on singlet fission: Tamura, Huix-Rotllant, Burghardt, Olivier, Beljonne, Phys. Rev. Lett. 115, 107401 (2015)

Summary

① Molecular-Level Approach to Organic Photovoltaics

- $e-h$ lattice model: highlights fragment properties + molecular packing
- extensions to conical intersections, multi-exciton states¹ etc.
- accurate on-the-fly dynamics remains highly challenging

② Coherent Exciton Migration

- ultrafast formation of quasi-stationary exciton-polaron states
- elementary exciton-polaron migration step is coherent
- strongly correlated exciton-phonon states

¹See our work on singlet fission: Tamura, Huix-Rotllant, Burghardt, Olivier, Beljonne, Phys. Rev. Lett. 115, 107401 (2015)

Summary

① Molecular-Level Approach to Organic Photovoltaics

- e - h lattice model: highlights fragment properties + molecular packing
- extensions to conical intersections, multi-exciton states¹ etc.
- accurate on-the-fly dynamics remains highly challenging

② Coherent Exciton Migration

- ultrafast formation of quasi-stationary exciton-polaron states
- elementary exciton-polaron migration step is coherent
- strongly correlated exciton-phonon states

③ Role of Charge Transfer Excitons in Regioregular Domains

- charge-transfer excitons in regioregular oligothiophene phases
- coherent formation of charge transfer excitons
- impact on charge separation at donor-acceptor interface

¹See our work on singlet fission: Tamura, Huix-Rotllant, Burghardt, Olivier, Beljonne, Phys. Rev. Lett. 115, 107401 (2015)

Summary

① Molecular-Level Approach to Organic Photovoltaics

- e - h lattice model: highlights fragment properties + molecular packing
- extensions to conical intersections, multi-exciton states¹ etc.
- accurate on-the-fly dynamics remains highly challenging

② Coherent Exciton Migration

- ultrafast formation of quasi-stationary exciton-polaron states
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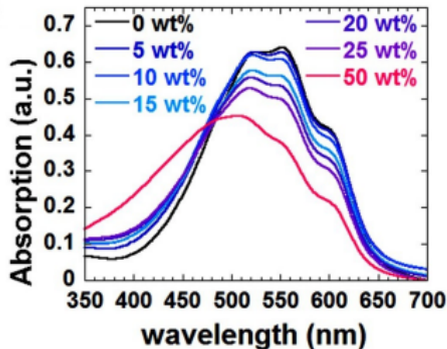
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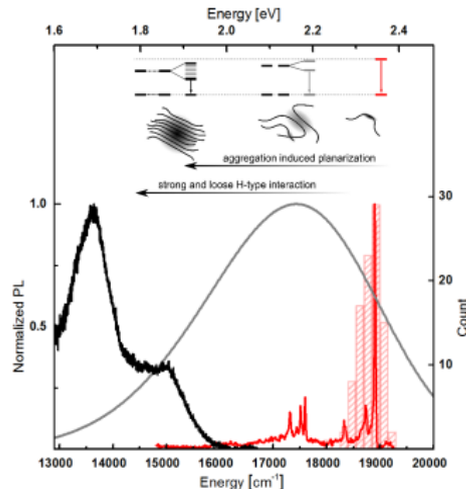
¹See our work on singlet fission: Tamura, Huix-Rotllant, Burghardt, Olivier, Beljonne, Phys. Rev. Lett. 115, 107401 (2015)

Insights from Spectroscopy

absorption spectrum: regioregular P3HT
with admixtures from regiorandom P3HT



Vohra et al., Appl. Phys. Lett. (2012)

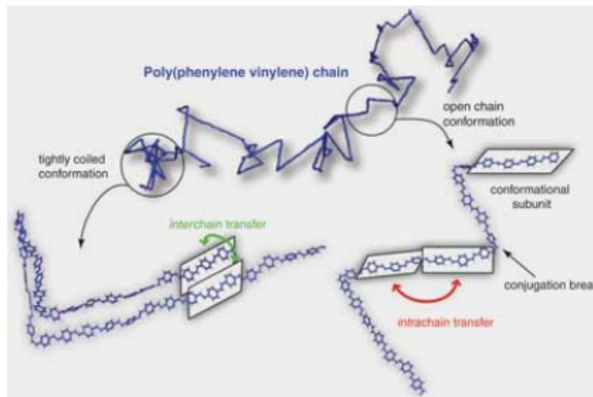


Raithel et al., Macromolecules 49, 9553 (2016)

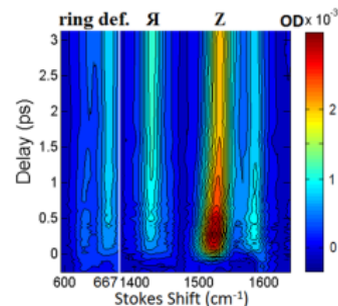
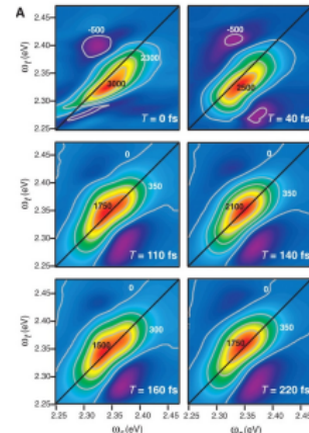
- P3HT: inhomogeneous broadening across 200 nm
- huge red shift between solution and bulk: ordered low-energy conformations
- signatures of vibronic structure due to high-frequency modes ($\sim 1500 \text{ cm}^{-1}$)
- single-molecule spectroscopy unravels signals from single chromophores

Thiessen, Vogelsang, Adachi, Steiner, Vanden Bout, Lupton, PNAS E3550 (2013)

Time-Resolved Spectroscopies



Collini, Scholes, Science 323, 369 (2009)



- 2D electronic spectroscopy monitors ultrafast energy and charge transfer

Anna, Song, Dinshaw, Scholes, Pure. Appl. Chem. 85, 1307 (2013)

De Sio, Lienau, PCCP 19, 18813 (2017)

- time-resolved Raman spectroscopy provides signatures of ultrafast structural dynamics

Zhou, Yu, Bragg, J. Phys. Chem. Lett. 6, 3496 (2015)

Acknowledgments & Collaborations

Group Frankfurt:

- M. Polkehr
- P. Eisenbrandt
- R. Binder
- J. von Cosel
- K. Falahati
- D. Picconi
- W. Popp
- M. Bonfanti
- K. Schwinn

Former members:

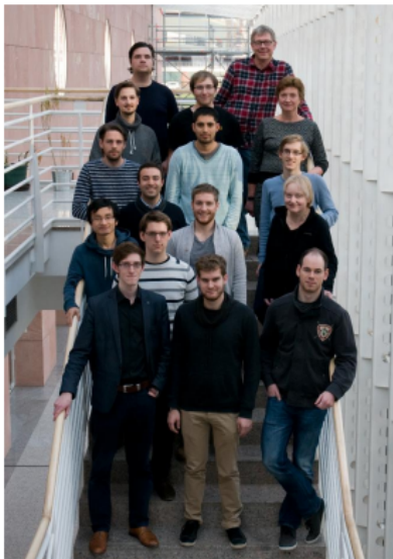
- J. Wahl
- M. Huix-Rotllant

Collaborations:

- H. Tamura (Tokyo, Japan)
- W. H. Miller, S. Cotton, R. Liang (Berkeley)
- K. H. Hughes (Bangor, UK)
- R. Martinazzo (Milano, Italy)
- H. Lischka, A. Aquino (TTU, USA)
- F. Plasser (Vienna, Austria)
- D. Lauvergnat (Orsay, France)
- S. Haacke, S. Méry (Strasbourg, France)
- G. d'Avino (Institut Néel, Grenoble)
- G. A. Worth (London, UK)
- A. Panda (IIT Guwahati, India)
- D. Beljonne, Y. Olivier (Mons, Belgium)

Thanks to: **DFG / ANR (France)** for financial support



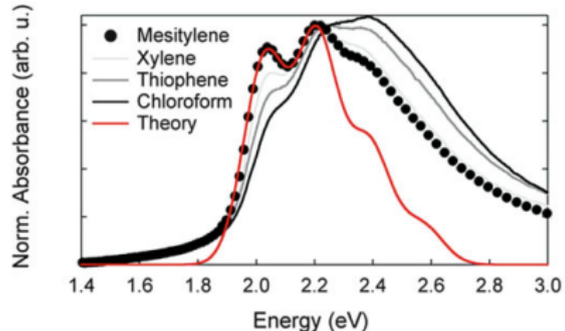
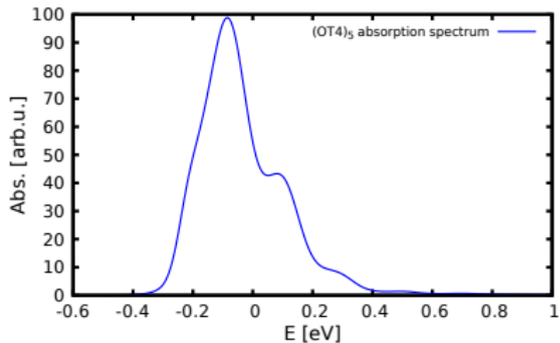


Theoretical Chemistry
of Complex Systems

AK Burghardt



Absorption Spectra

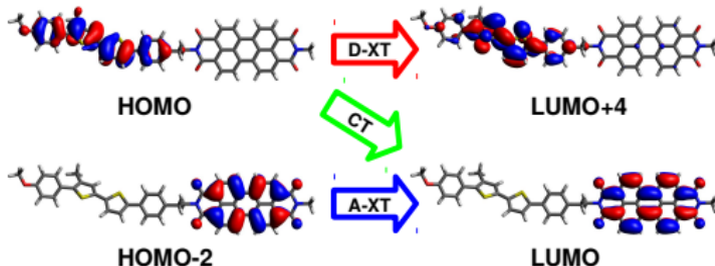
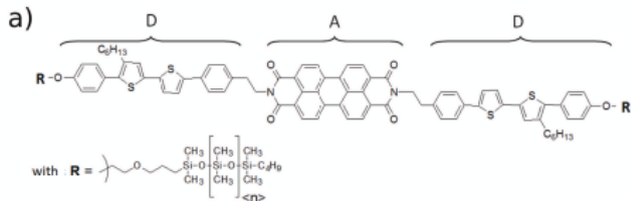


Tremel, Ludwigs, Adv. Polym. Sci. 265, 39 (2014)

- spectra calculated from Fourier transform of $C(t) = \langle \psi(0) | \psi(t) \rangle$ for OT H-aggregate (6 layers)

Highly Ordered DA Assemblies: Liquid Crystalline Material

collaboration with S. Haacke, S. Méry (Strasbourg)

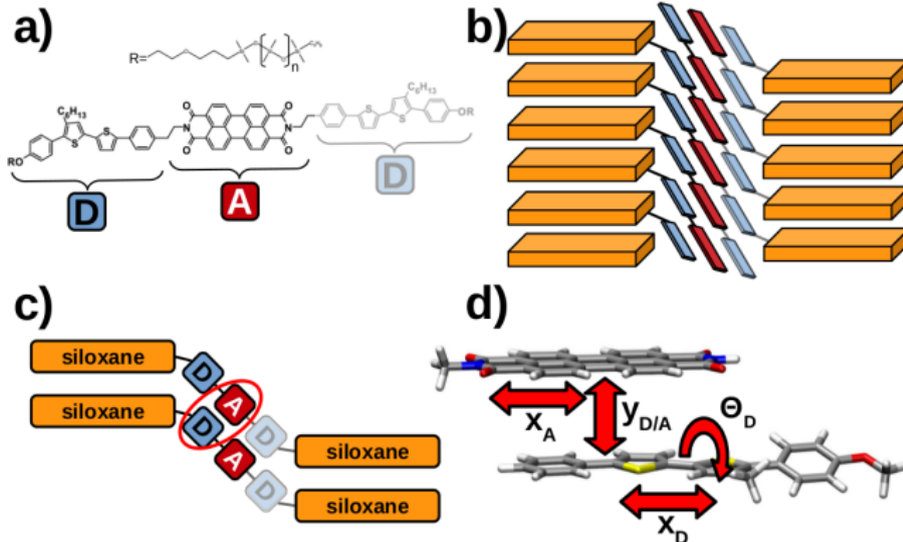


Roland, Ramirez, Léonard et al., PCCP, 14, 273 (2012)

Wenzel, Dreuw, Burghardt, PCCP, 15, 11704 (2013)

- competing ultrafast energy transfer (EET) and charge transfer (CT) processes
- in chloroform: EET in 130 fs, followed by CT in 2.7 ps
- in liquid crystalline phase: CT in 60 fs!
- relatively fast recombination (50 ps) – material doesn't really work well ...

What is Happening in the First-Generation Material?



Polkehn et al., J. Phys. Chem. Lett., 7, 1327 (2016)

- first-generation material: liquid crystalline smectic mesophase
- idea: D/A stacks serve as “quantum wells” for carrier transport
- much faster charge transfer in film than solution (~ 50 fs vs. ~ 3 ps)
- calculations suggest unexpected inter-chain D-A interactions

$$\hat{H} = \hat{H}_{\text{on-site}} + \hat{H}_{\text{coupl}} + \hat{H}_{\text{e-ph}}$$

$$\hat{H}_{\text{on-site}} = \epsilon_D \sum_{i=1}^{N_D} |D_i^{XT}\rangle \langle D_i^{XT}| + \epsilon_A \sum_{i=1}^{N_A} |A_i^{XT}\rangle \langle A_i^{XT}| + \sum_{i=1}^{N_D} \sum_{j=1}^{N_A} \epsilon_{D_i^+ A_j^-} |D_i^+ A_j^-\rangle \langle D_i^+ A_j^-|$$

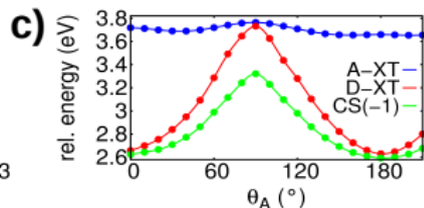
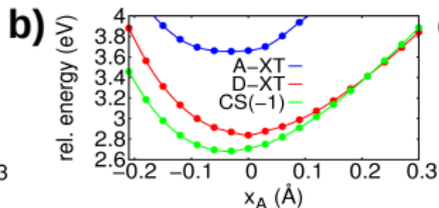
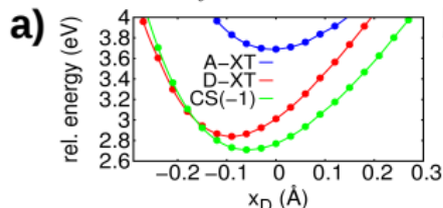
$$\hat{H}_{\text{coupl}} = J_D \sum_{i=1}^{N_D} \sum_{j=1}^{N_D} (|D_i^{XT}\rangle \langle D_j^{XT}| + \text{h.c.}) + J_A \sum_{i=1}^{N_A} \sum_{j=1}^{N_A} (|A_i^{XT}\rangle \langle A_j^{XT}| + \text{h.c.}) + J_{DA} \sum_{i=1}^{N_D} \sum_{j=1}^{N_A} (|D_i^{XT}\rangle \langle A_j^{XT}| + \text{h.c.})$$

$$+ \kappa_D \sum_{i=1}^{N_D} \sum_{j=1}^{N_A} (|D_i^{XT}\rangle \langle D_i^+ A_j^-| + \text{h.c.}) + \kappa_A \sum_{i=1}^{N_A} \sum_{j=1}^{N_D} (|A_i^{XT}\rangle \langle D_j^+ A_i^-| + \text{h.c.})$$

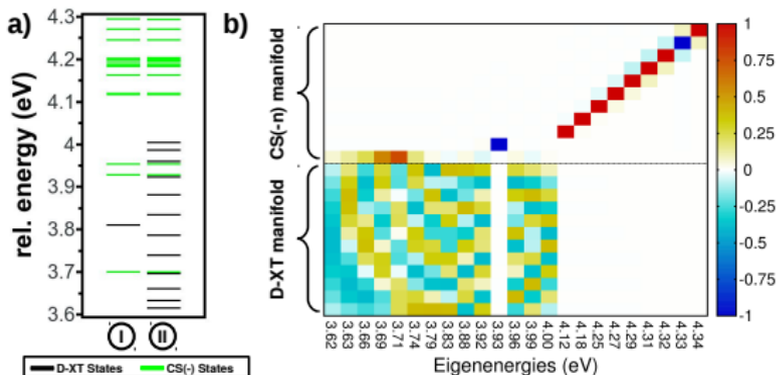
$$+ t_e \sum_{i=1}^{N_D} \sum_{j=2}^{N_A-1} (|D_i^+ A_j^- \rangle \langle D_i^+ A_{j\pm 1}^-| + \text{h.c.}) + t_h \sum_{i=2}^{N_D-1} \sum_{j=1}^{N_A} (|D_i^+ A_j^- \rangle \langle D_{i\pm 1}^+ A_j^-| + \text{h.c.})$$

$$\hat{H}_{\text{e-ph}} = \hat{T}_{\text{ph}}(\{\zeta^D\}, \{\zeta^A\}, \{\zeta^{DA}\}) + \sum_i \hat{V}_D^{XT}(\{\zeta_i^D\}, \{\zeta_i^{DA}\}) |D_i^{XT}\rangle \langle D_i^{XT}| + \sum_i \hat{V}_A^{XT}(\{\zeta_i^A\}, \{\zeta_i^{DA}\}) |A_i^{XT}\rangle \langle A_i^{XT}|$$

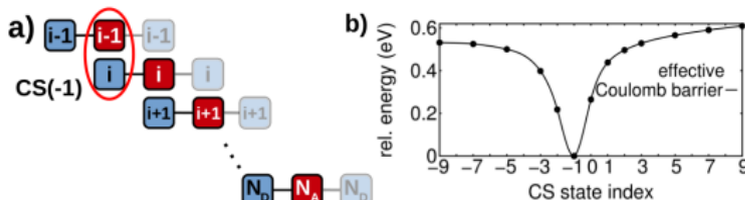
$$+ \sum_i \sum_j \hat{V}_{DA}(\{\zeta_i^D\}, \{\zeta_j^A\}, \{\zeta_{ij}^{DA}\}) |D_i^+ A_j^- \rangle \langle D_i^+ A_j^-|$$



Liquid Crystalline Phase – Energetics



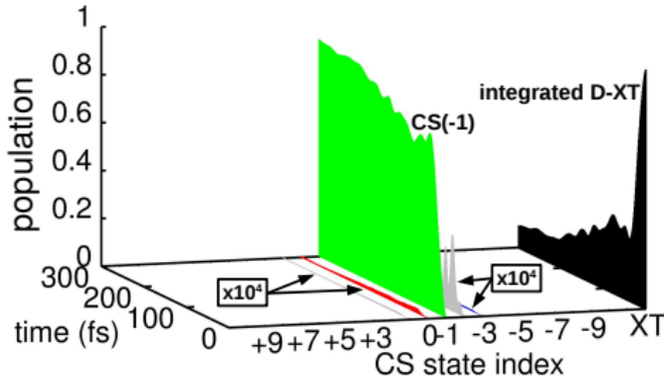
- energetics at Franck-Condon geometry
- state mixing: excitonic manifold and CS(-1) state
- but higher charge separated states barely accessible



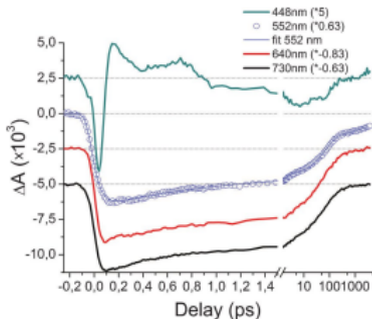
- on-site energies computed from ADC(2) and TDDFT
- internal field: $20 \text{ V}/\mu\text{m}$
- CS(-1) state strongly stabilized

Polkehn, Tamura, Eisenbrandt, Haacke, Méry, Burghardt, J. Phys. Chem. Lett., 7, 1327 (2016)

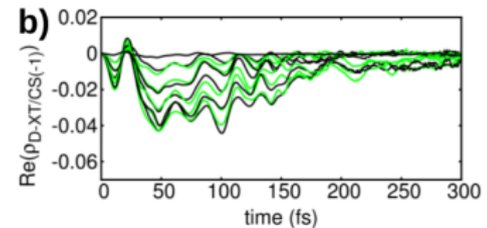
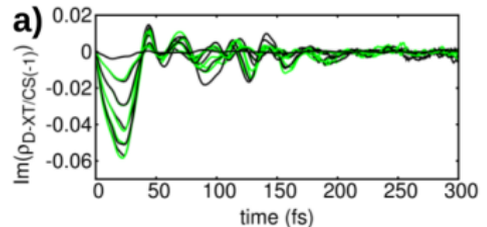
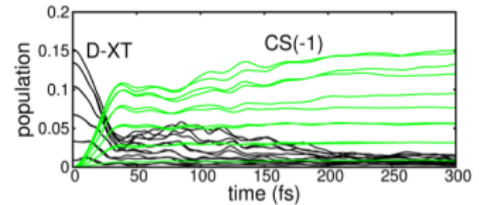
Liquid Crystalline Phase – Dynamics



Polkehn et al., JPCL (2016)



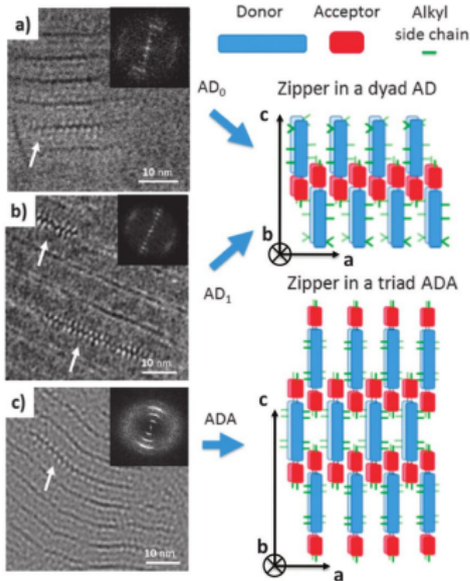
Roland et al., PCCP (2012)



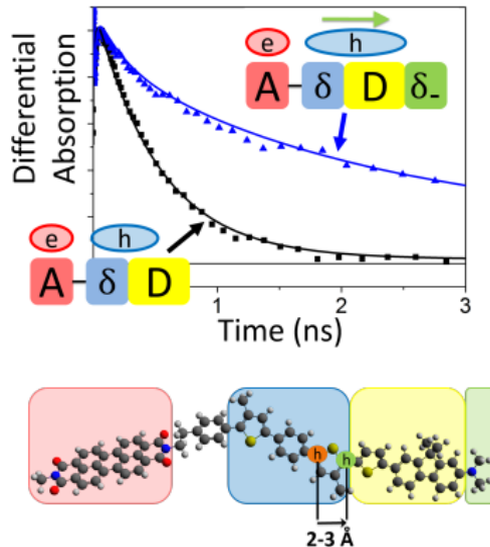
- transition to CS(-1) states (~ 50 fs) explains transient absorption experiments (Haacke)

- ML-MCTDH simulations for 156 states/48 modes

Second Generation Material: Zipper-like Molecular Packing



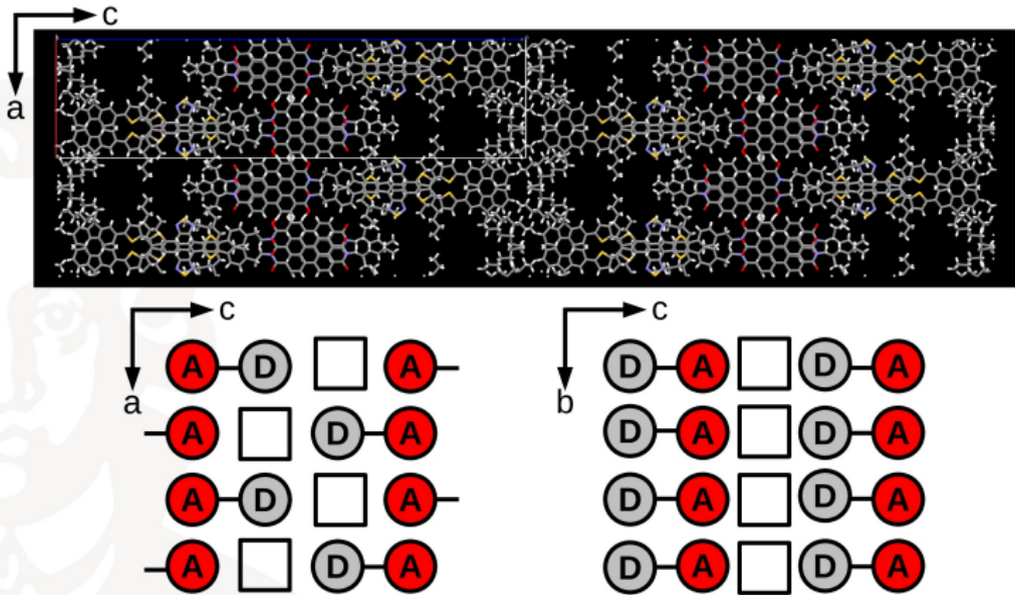
Biniak et al., J. Mater. Chem. C 3, 3342 (2015)



Liu, Eisenbrandt et al., PCCP 18, 18536 (2016)

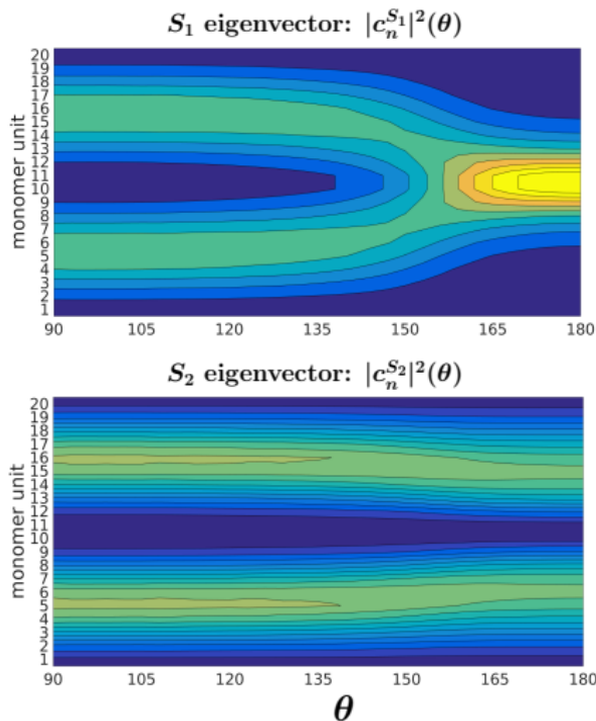
- tunable donor species: alternating thiophene/fluorene/benzothiadiazole units; electrodeficient bridge to the perylene acceptor – **chemical design!**
- organization in lamellae (both DA and ADA – but not DAD)
- comparatively slow CT formation (tens of ps) - **and less recombination**

Second Generation Material: In Progress ...

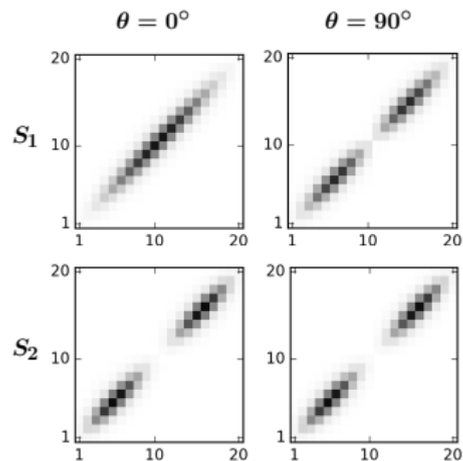


- coarse-grained model for Kinetic Monte Carlo (KMC) simulations
- microelectrostatics calculations (collaboration with G. d'Avino, Grenoble)
- multi-scale modeling needed!

Electronic Structure – Torsion Dependence



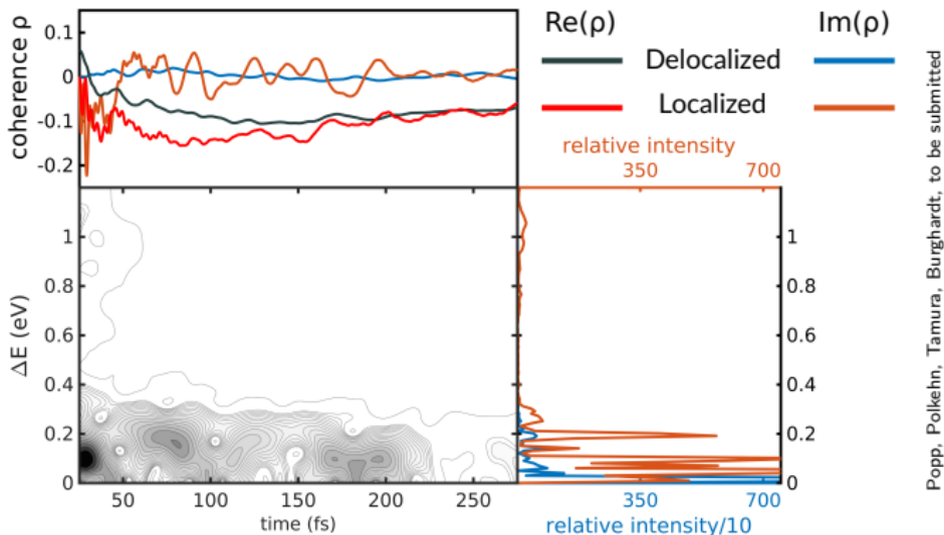
NB. polaronic (trapped) S_1/S_2 states



- transition density analysis
Panda, Plasser, Aquino, Burghardt, Lischka, JPCA (2013)
- particle-in-the-box type $e-h$ states
- marked dependence on torsion
- LEGS = local exciton ground state:
nodeless S_1 exciton @180°

Tozer, Barford, JPCA 116:10310 (2012)

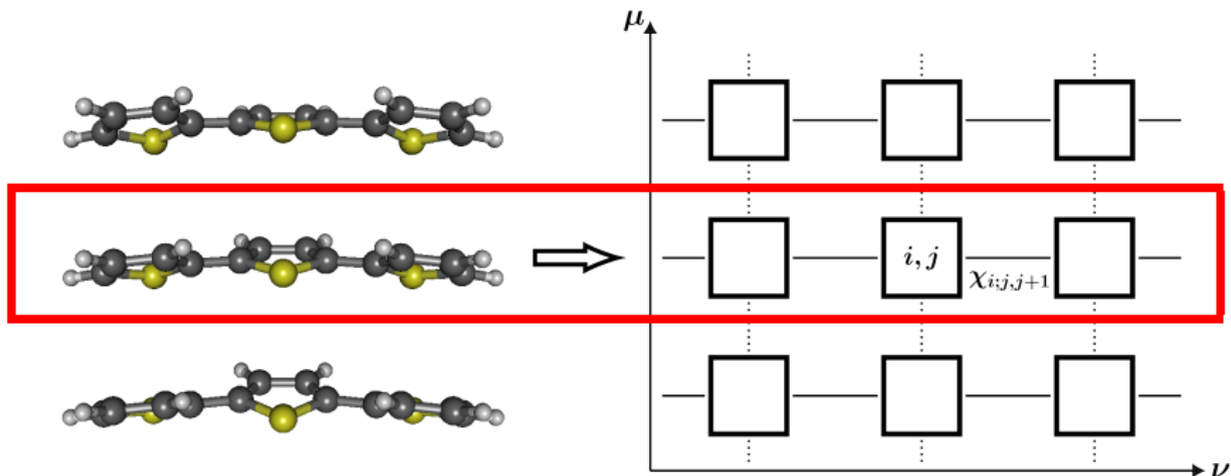
Signatures of Excitonic Coherence



Popp, Polkehn, Tamura, Burghardt, to be submitted

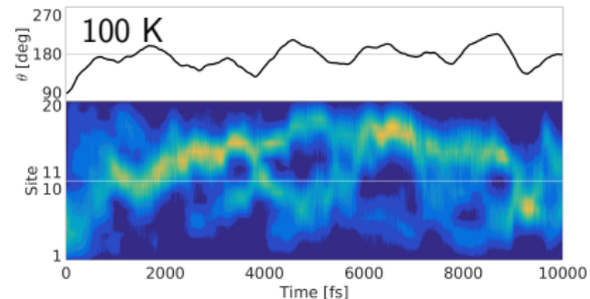
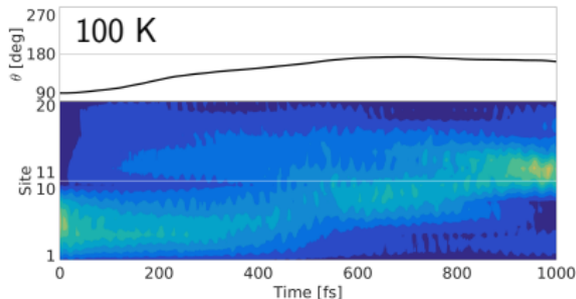
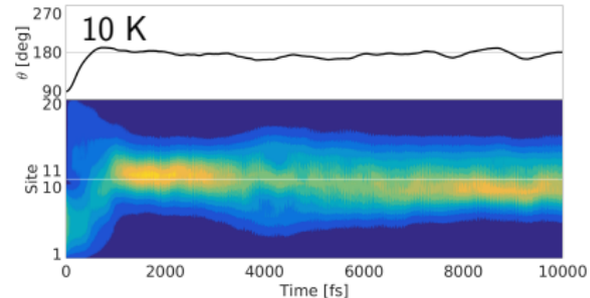
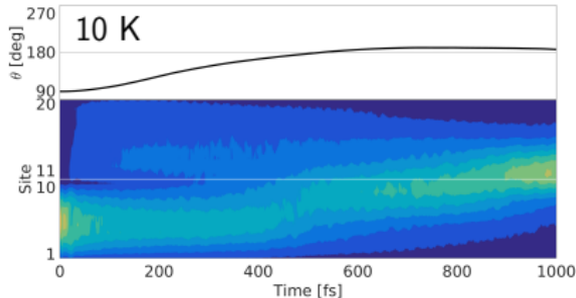
- windowed Fourier transform of time-dependent XT-XT electronic coherence
- localized initial condition produces beatings within excitonic manifold
- very similar frequency as dominant vibrations ($\sim 0.1-0.2$ eV)
- both vibrational and excitonic coherence could contribute

HJ Aggregates: Intra- & Inter-Chain Interactions



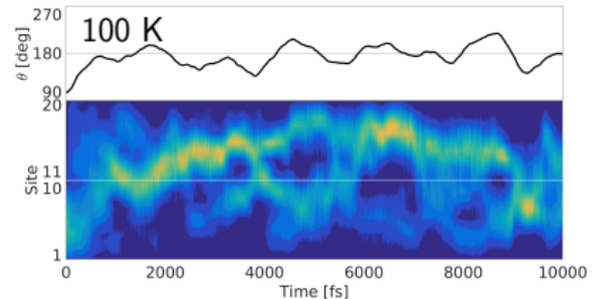
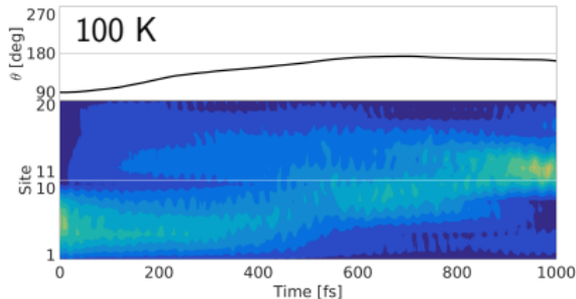
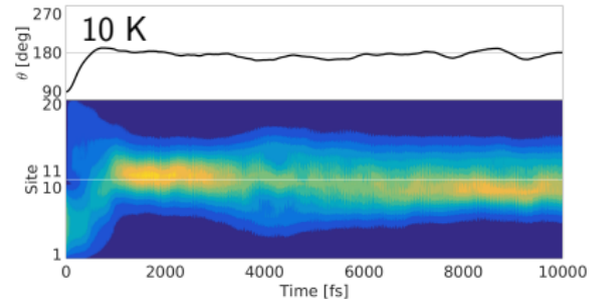
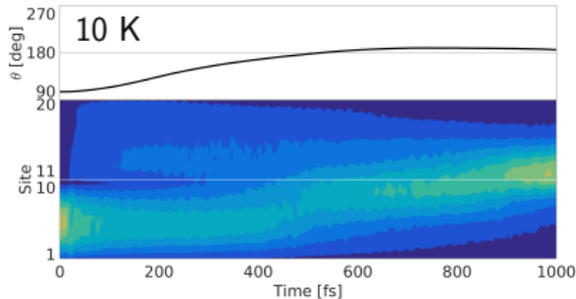
- partitioning either monomer-based or oligomer-based
- **monomer-based partitioning:** on-chain exciton dynamics
- **oligomer-based partitioning:** interchain exciton and charge transfer dynamics

Temperature Effects: Ehrenfest/Langevin dynamics



- ultrafast transients not correctly reproduced by Ehrenfest dynamics
- at increasing temperatures, fluctuations start driving exciton migration

Temperature Effects: Ehrenfest/Langevin dynamics



- ultrafast transients not correctly reproduced by Ehrenfest dynamics
- at increasing temperatures, fluctuations start driving exciton migration

Quantum Coherence Plays a Non-Negligible Role!

$$|\psi(t)\rangle = c_0(t)|0\rangle|\phi_0(t)\rangle + c_1(t)|1\rangle|\phi_1(t)\rangle$$

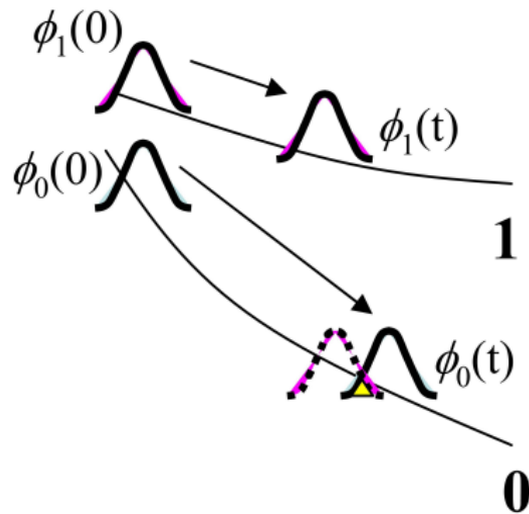
electronic coherence:

$$\begin{aligned}\rho_{01}(t) &= \text{Tr}[|0\rangle\langle 1|\hat{\rho}(t)] \\ &= \langle 1|\hat{\rho}(t)|0\rangle = c_1^*(t)c_0(t)\langle\phi_1(t)|\phi_0(t)\rangle\end{aligned}$$

- coherence \propto **overlap of nuclear wavefunctions**
- typical decoherence times: tens to hundreds of fs or more (estimate from $\tau_{\text{dec}} \sim \tau_g(6k_B T/\lambda)^{1/2}$ or $\tau_{\text{dec}} \sim \gamma^{-1}(\lambda_T/\Delta x)^2$)

Prezdho, Rosicky, PRL 81, 5294 (1998)

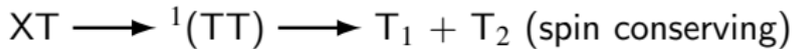
- loss of coherence not captured by classical trajectory picture



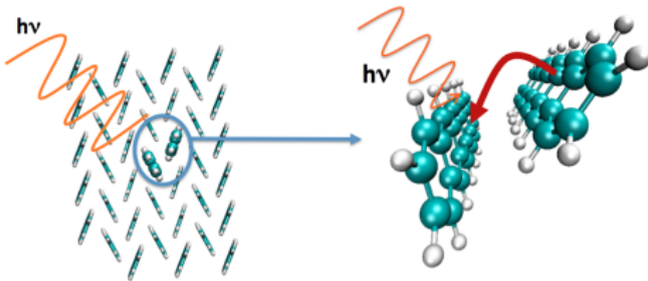
picture: P. Rosicky et al.

Singlet Fission: Route To Carrier Multiplication

(collaboration with H. Tamura (Sendai/Tokyo), D. Beljonne (Mons))



possibly overcome Shockley-Queisser limit

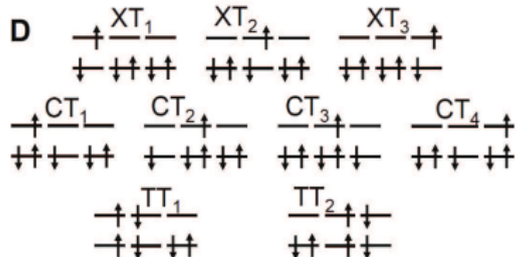
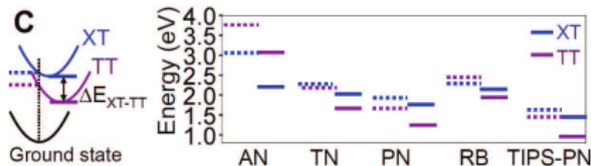


<http://sites.lsa.umich.edu/zimmerman-lab/wp-content/uploads/sites/52/2014/03>

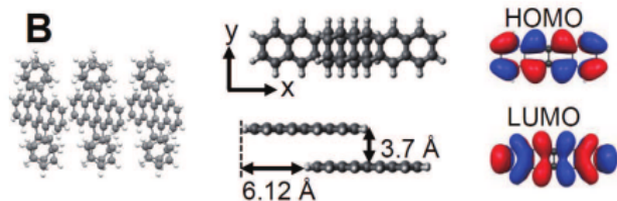
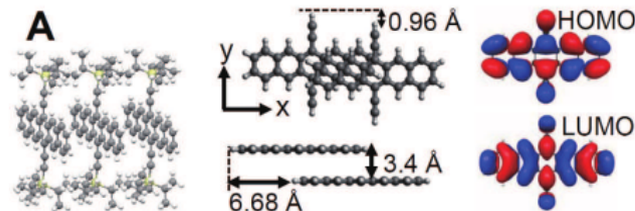
- driving force $\Delta E_{XT-TT} < 0$ doesn't explain the whole picture
- possible involvement of intermediate CT states (superexchange)
- vastly different time scales for different materials

- discovered in 1965: anthracene
- reviews by Smith & Michl, *Annu. Rev. Phys. Chem.* 2013, 64, 361, *Chem. Rev.* 110, 6891 (2010)

Molecular Packing: Energetics & Electronic Couplings



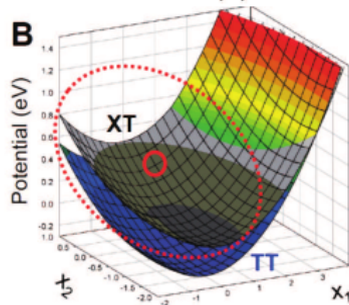
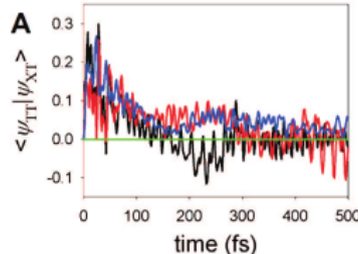
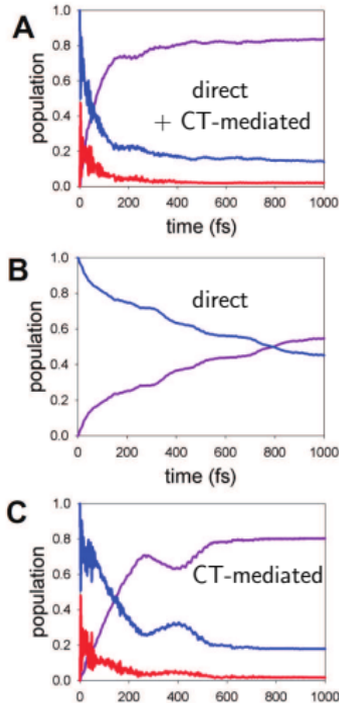
- energetics of acene series (C)
- trimer model: 9 states (D)



- TIPS-pentacene (A): slip-stacked
- rubrene (B): C_{2h} symmetric

Electronic coupling @ equil. non-zero for TIPS-pentacene but vanishing for rubrene!
 (MRMP2 calculations for dimers, el. couplings via diabatisation protocol)

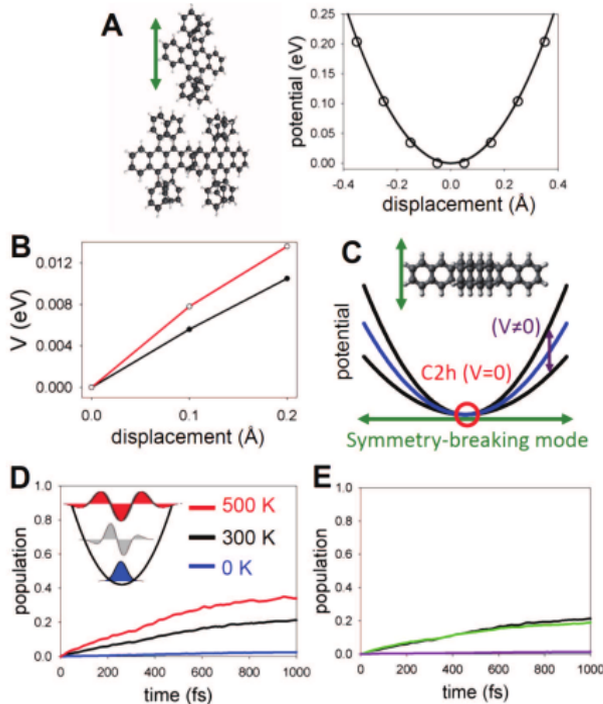
TIPS-Pentacene: Ultrafast Singlet Fission



- ultrafast, coherent SF
 - slip-stacked geometry: avoided crossing
 - interfering direct and CT-mediated pathways (electronic coupling via CT's dominates)
 - vibrational coherence effectively transferred between XT and TT states (see $\langle \psi_{TT} | \psi_{XT} \rangle$)
- Musser et al., Nature Phys. 11, 352 (2015)
- MRMP2 calculations + diabaticization

Tamura, Huix-Rotllant, Burghardt, Olivier, Beljonne, Phys. Rev. Lett., 115, 107401 (2015)

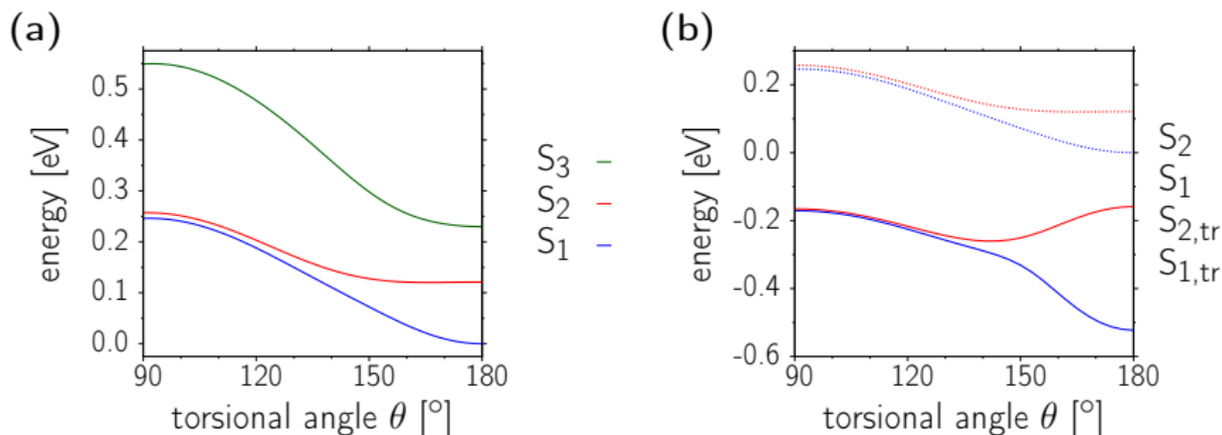
Rubrene: Thermally Activated Singlet Fission



- driving force slightly exergonic (like tetracene), but thermally activated SF (picosecond scale)
- C_{2h} crystal geometry: electronic coupling vanishes – conical intersection
- electronic couplings depend on symmetry-breaking coordinate: $V(X) = \lambda X$, $\lambda \sim 10^{-3}$ eV
- slow, incoherent dynamics
- key influence of molecular packing

Tamura, Huix-Rotllant, Burghardt, Olivier, Beljonne, Phys. Rev. Lett., 115, 107401 (2015)

Exciton-Polaron Dynamics: Adiabatic Picture



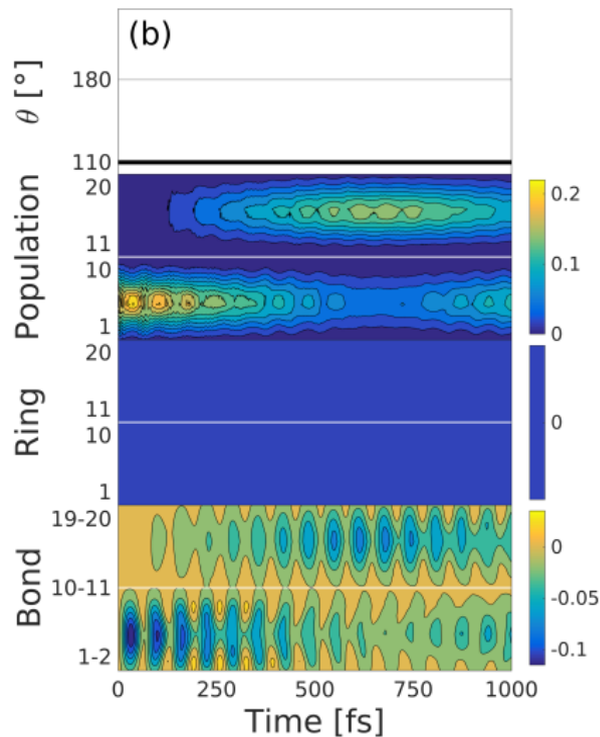
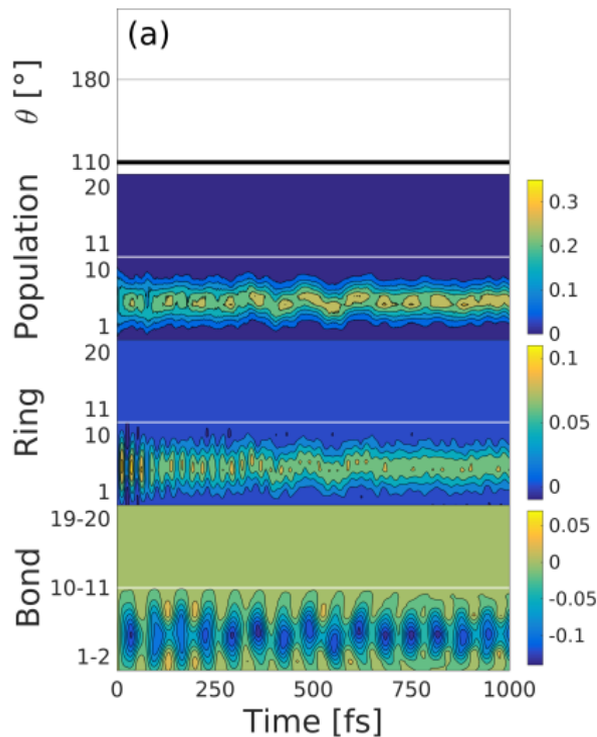
- dynamics is essentially happening on coupled S_1/S_2 surfaces
- effective torsion potentials including stabilization due to trapping ($S_{1,tr}/S_{2,tr}$)²
- initial left/right localized state = superposition of S_1/S_2
- energy loss due to external bath acting on torsional mode
- exciton-polaron trapping persists at all stages of the dynamics

²i.e., “polaron transformed” potentials

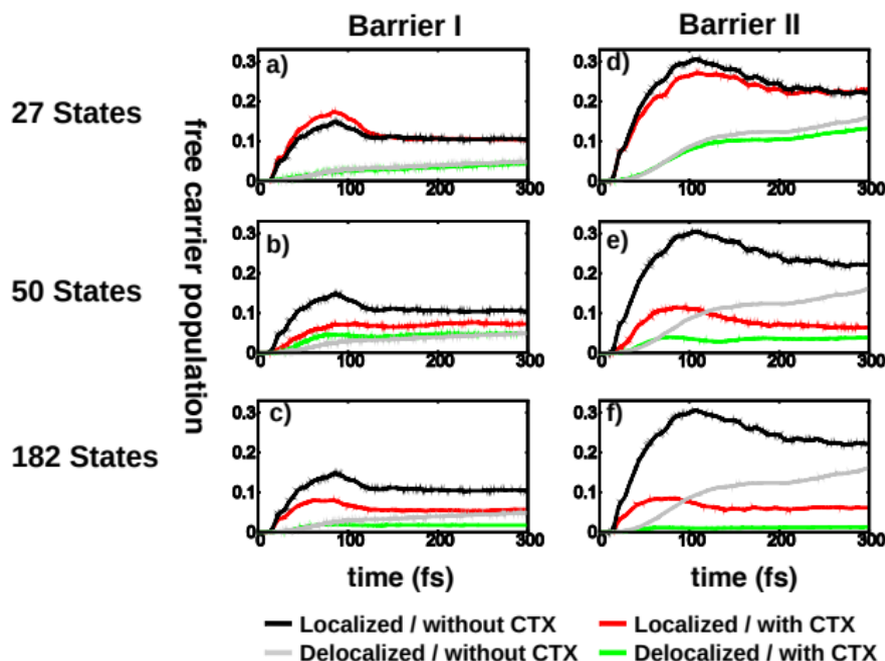
Quantum Dynamics: 20-Site J-Aggregate with Central Torsion

C-C inter-monomer mode + local C=C + torsion + bath

C-C inter-monomer mode + torsion + bath



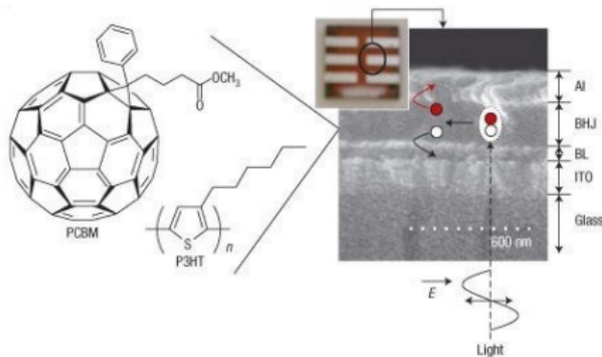
Interfacial Charge Separation in the Presence of CTX States



- time-dependent free carrier populations

Polkehn, Tamura, Burghardt, J. Phys. B, Special Issue "Light Energy Conversion, Light Harvesting", submitted (2017)

System-Bath Models



S region: e.g., electronic degrees of freedom (electron-hole states)

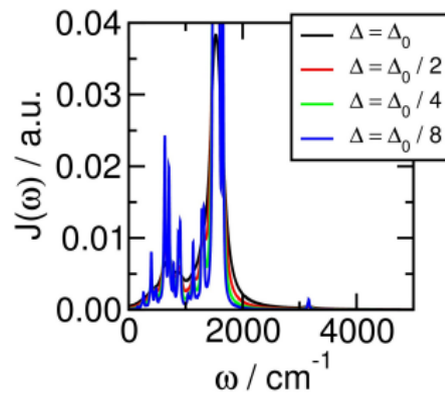
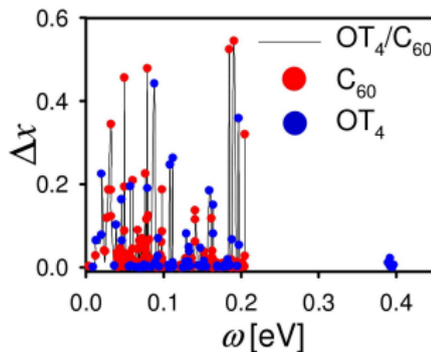
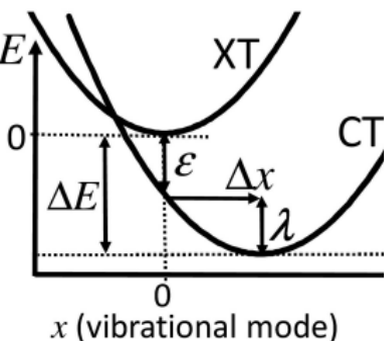
B region: all vibrations (phonons) mapped to harmonic oscillator model

$$\hat{H}_B + \hat{H}_{SB} = \sum_n \frac{1}{2} (\hat{p}_n^2 + \frac{1}{2} \omega_n^2 \hat{x}_n^2) + \hat{s} \sum_n c_n \hat{x}_n$$

$$J(\omega) = \pi/2 \sum_n c_n^2 / \omega_n \delta(\omega - \omega_n)$$

spectral density

Spectral Densities from Electronic Structure Calculations^(*)



$$J(\omega) = \frac{\pi}{2} \sum_n^N \frac{c_n^2}{\omega_n} \delta(\omega - \omega_n) \simeq \frac{\pi}{2} \sum_n^N \frac{c_n^2}{\pi} \frac{\Delta}{(\omega - \omega_n)^2 + \Delta^2}$$

Tamura, Martinazzo, Ruckebauer, Burghardt, J. Chem. Phys., 137, 22A540 (2012)

^(*)NB. Alternatively: obtain SD's from correlation functions (MD, CPMD, ...)