Chemical Science



EDGE ARTICLE

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2017, 8, 6947

Enhancing charge mobilities in organic semiconductors by selective fluorination: a design approach based on a quantum mechanical perspective†

Buddhadev Maiti, (1) ‡a Alexander Schubert, (10) ‡ab Sunandan Sarkar, (10) a Srijana Bhandari, (10) a Kunlun Wang, (10) a Zhe Li, (10) a Eitan Geva, (10) *b Robert J. Twieg (10) *a and Barry D. Dunietz (10) *a

Selective fluorination of organic semiconducting molecules is proposed as a means to achieving enhanced hole mobility. Naphthalene is examined here as a root molecular system with fluorination performed at various sites. Our quantum chemical calculations show that selective fluorination can enhance attractive intermolecular interactions while reducing charge trapping. Those observations suggest a design principle whereby fluorination is utilized for achieving high charge mobilities in the crystalline form. The utility of this design principle is demonstrated through an application to perylene, which is an important building block of organic semiconducting materials. We also show that a quantum mechanical perspective of nuclear degrees of freedom is crucial for a reliable description of charge transport.

Received 3rd June 2017 Accepted 12th August 2017

DOI: 10.1039/c7sc02491f

rsc.li/chemical-science

Organic semiconducting materials have received much attention over the last two decades due to their potential optoelectronic applications, such as organic field-effect transistors (OFETs), ¹⁻³ organic photovoltaics (OPV), ⁴⁻⁷ organic light emitting diodes (OLEDs)⁸⁻¹¹ and sensors. ¹²⁻¹⁴ Charge mobility plays a crucial role in determining device performance. ¹⁵⁻¹⁸ Hence, enhancing charge mobility in organic semiconducting materials is expected to improve device functionality. ^{19,20}

In this paper, we investigate utilizing selective fluorination of conjugated organic molecules as a way of enhancing charge mobilities in organic semiconducting materials.²¹ Fluorinated organic molecules have been previously employed as charge transporting materials.^{22–24} Judicious introduction of polar C–F bonds enhances intermolecular attractive forces, tightening crystal packing, increasing molecular orbital overlap, and thereby affecting the materials optoelectronic properties.^{25–29} At the same time, fluorination is often associated with forming a molecular dipole moment that increases the propensity for charge trapping inhibiting charge mobility.^{30–33} Thus, the

Using dimer models,⁴¹ charge transport (CT) rate constants are calculated within the framework of Fermi's golden rule (FGR) based on inputs obtained from quantum chemical calculations.⁴²⁻⁴⁷ The FGR rate constant is given by:^{42,48,49}

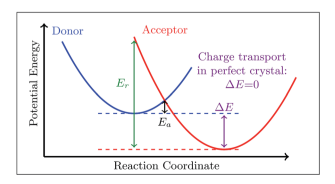


Fig. 1 A schematic view of the charge donor (blue) and acceptor (red) potential energy surfaces. E_r and E_a are the reorganization energy and activation energy, respectively. ΔE is the charge transfer reaction free energy.

challenge is to design structures that maximize charge mobilities while minimizing charge trapping. To this end, we consider a strategy where the local polar C-F bonds are arranged so as to give rise to tightly packed crystals despite having vanishing overall molecular dipoles.³⁴⁻⁴⁰

^aDepartment of Chemistry and Biochemistry, Kent State University, Kent, OH 44242, USA. E-mail: twieg@kent.edu; bdunietz@kent.edu; rtwieg@kent.edu

^bDepartment of Chemistry, University of Michigan, Ann Arbor, MI 48109, USA. E-mail: eitan@umich.edu

[†] Electronic supplementary information (ESI) available: Reorganization energy, orbital-based coupling energies, HRFs, normal mode motions, transfer rates, and atomic coordinates. See DOI: 10.1039/c7sc02491f

[‡] These authors contributed equally to this work.

Chemical Science Edge Article

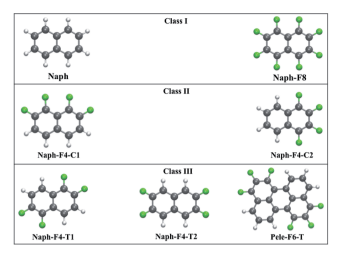


Fig. 2 The three classes of molecules under consideration: Class I consists of nonfluorinated and fully fluorinated naphthalenes; Class II, consists of molecules with asymmetric partial fluorination which gives rise to sizable dipole moments; Class III consists of molecules with symmetrical partial fluorination and vanishing dipole moments.

$$k_{\text{FGR}} = \frac{|\Gamma|^2}{\hbar^2} e^{-\sum_{\alpha} S_{\alpha}(2n_{\alpha}+1)} \times \int_{-\infty}^{\infty} dt \ F_{\text{r}}^{\text{ex}}(t) \exp \left\{ -\frac{i}{\hbar} \Delta E t + \sum_{\alpha} S_{\alpha} \left[(n_{\alpha}+1) e^{-i\omega_{\alpha}t} + n_{\alpha} e^{i\omega_{\alpha}t} \right] \right\}.$$
(1)

Here, Γ is the electronic coupling, ΔE is the energy difference between the two states at their optimized geometries, as illustrated in Fig. 1, $\{\omega_{\alpha}\}$ are the normal mode frequencies, $\{S_{\alpha}\}$ are the Huang–Rhys factors^{49,50} (HRFs), and $n_{\alpha} = \left(\exp\left\{\frac{\hbar\omega_{\alpha}}{k_{\rm B}T}\right\} - 1\right)^{-1}$ are the normal mode's thermal

occupancies. $F_{\rm r}^{\rm ex}(t)=\exp[-k_{\rm B}TE_{\rm r}^{\rm ex}t^2/\hbar^2]$ accounts for outersphere solvation, where $E_{\rm r}^{\rm ex}$ is the corresponding reorganization energy. The inner-sphere reorganization energy is given by $E_{\rm r}^{\rm in}=\sum\hbar\omega_{\alpha}S_{\alpha}$.

 $\frac{1}{\alpha}$

The Marcus rate constant can be obtained from the FGR expression in eqn (1) in the high temperature and short time limits: $^{51-54}$

$$k_{\rm M} = \frac{\left|\Gamma\right|^2}{\hbar} \sqrt{\frac{\pi}{k_{\rm B} T E_{\rm r}}} \exp\left(-\frac{(\Delta E + E_{\rm r})^2}{4k_{\rm B} T E_{\rm r}}\right). \tag{2}$$

here, $E_{\rm r}=E_{\rm r}^{\rm ex}+E_{\rm r}^{\rm in}$ is the overall reorganization energy.

The charge mobility of one-dimensional motion along the molecular stacking axis, η , is evaluated using the Einstein-Smoluchowski equation: ⁵⁵

$$\eta = \frac{ea^2k}{k_{\rm B}T},\tag{3}$$

where, e, a, and k are the electron charge, the distance between the donor and acceptor molecules, and the CT rate constant, respectively (k is set to either $k_{\rm M}$ or $k_{\rm FGR}$).

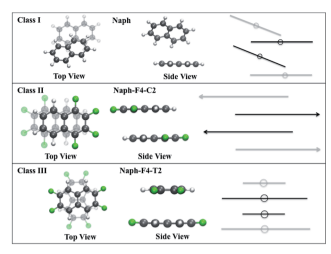


Fig. 3 Class I molecules are arranged in a herringbone-like structure, whereas Class II and III molecules are arranged in stacks due to $\pi-\pi$ interactions. In Class II the monomers are aligned in alternating directions due to their dipole with an in-plane relative shift. In Class III the molecules show a concentric 90° relative in-plane rotation about the stacking axis.

CT rate constants and mobilities were calculated for the following three classes of fluorinated naphthalene molecules (see Fig. 2):

- Class I molecules have vanishing dipole moments^{56–58} that may result in a herringbone-like arrangement in the crystal phase (see Fig. 3).⁵⁹ These are represented by the nonfunctionalized naphthalene molecule (Naph) and the fully functionalized molecule octafluoronaphthalene (Naph-F8), which have both been investigated experimentally.^{60,61} We point out that, while mixtures of Naph/Naph-F8 may form well-packed molecular stacks,⁶¹ in this work we consider only crystals of single compounds.
- Class II molecules include polar C-F bonds that are arranged to produce large molecular dipole moments. The substantial attractive dipole–dipole intermolecular interaction is expected to lead to tightly packed crystals.³⁴⁻³⁷ These are represented by the partially functionalized 1,2,7,8- and 1,2,3,4-tetrafluoronaphthalene molecules (Naph-F4-C1 and Naph-F4-C2, respectively). While Naph-F4-C1 has not been synthesized (to the best of our knowledge), the Naph-F4-C2 crystals were indeed observed to be tightly packed with a partial facial overlap (see Fig. 3).³⁴
- Class III molecules include polar C-F bonds that are arranged to eliminate the molecular dipole moment. Despite of this, substantial attractive intermolecular stacking interactions between those molecules³⁸⁻⁴⁰ give rise to tightly packed crystals (see Fig. 3). These are represented by the partially functionalized 1,2,5,6- and 2,3,6,7-tetrafluoronaphthalene (Naph-F4-T1 and Naph-F4-T2, respectively). Another molecule of this category considered below, is the fluorinated perylene molecule (Pele-F6-T) with a larger conjugated system and a potential to be used in actual optoelectronic applications. To the best of our knowledge, experimental insights into the crystal structure of Class III molecules are not yet available (not reported in the

Edge Article

Cambridge Crystallographic Data Centre, CCDC). The Pele-F6-T molecule is currently targeted by our experimental efforts.

We analyze these molecules using a first-principles approach based on density functional theory (DFT) calculations. The range-separated hybrid (RSH) functional 62 ω B97X-D 63 was employed. This RSH functional addresses well the tendency of local-density-approximation (LDA)-based functionals to underestimate the orbital fundamental gap, $^{64-68}$ and accounts for dispersion interactions. 69 It should be noted that the ω B97X-D functional was shown to yield highly accurate charge reorganization energies with an error of 6.5% in a benchmark study of similar molecules. 70 The polarizable continuum solvation model (PCM) was used to account for interactions with the solid-state host. 68,71,72 Optimal geometries were generated with the 6-31G(d) basis set. To confirm basis set convergence, we compare the 6-31G(d)-based results for Naph-F4-T2 with results obtained with the larger cc-pVTZ basis set (see Table 1).

Electronic coupling coefficients were calculated using a dimer model at the level of configuration interaction with constrained density functional theory (CDFT-CI).⁷³ In the CDFT-CI calculations, each molecule within the ionic dimer is assigned to be either a donor or an acceptor. The two states involved in the CI calculations are then obtained by localizing the charge on one of the molecules M in the ionic dimer calculations. Cation calculations, based on the states M⁺-M and M-M⁺, are used for determining the coupling strength associated with hole mobilities and anion calculations (M⁻-M/M-M⁻) for electron mobilities, respectively.

HRFs $\{S_\alpha\}$ were calculated using the DUSHIN program.^{74,75} Displacement geometries were calculated by comparing the optimized geometries of the neutral (M) and the charged monomer (M⁺). Normal mode frequencies and eigenvectors were obtained using the optimized neutral monomer M. The validity of the harmonic approximation was confirmed by the very good agreement of reorganization energies evaluated using the HRFs with their direct evaluation (see ESI, Table 2†).

Key electronic structure parameters calculated for Class I, Class II, and Class III molecules are shown in Table 1. We confirm that physically relevant Frontier orbital energies, *i.e.* energies of the highest occupied and lowest unoccupied molecular orbitals, HOMO (H) and LUMO (L), are typically within 0.1 eV of the calculated ionization potential (IP) and the electron affinity (EA), respectively.^{68,76} The dipole and quadrupole moments of the different molecules are also provided. As expected, while the molecular dipole moments of Class I and Class III molecules vanish, Class II molecules have substantial dipole moments (3.95 D and 2.90 D). The quadrupole moments of the naphthalene derivatives are observed to increase with the number of fluorine atoms.

The intermolecular separation a for Class I molecules corresponds to the distance between the molecular centers of mass (c.o.m.), and for Class II and III molecules it is set to the distance between the molecular planes. While the intermolecular energies and distances are calculated using the relatively simplified dimer model, we find that the calculated distances are in good agreement with measured crystal structure when available. (In the case of the known compounds Naph, Naph-F8, and Naph-F4-C2, the root mean square deviation (RMSD) of reported crystal structures are 0.134, 0.447, and 0.060 Å, respectively. (34,36,60,61) The intermolecular binding energies $F_{\rm inter}$ of Class I molecules are small with values of up to 0.34 eV, whereas for Class II and III molecules, they are in the range of 0.43–0.59 eV.

In Class II molecules, the intermolecular interactions result in a head-to-tail arrangement that aligns the dipole moments of neighboring molecules in the stack at opposite directions, thereby forming a partial facial overlap, see Fig. 3, middle panel. For Naph-F4-C2, this arrangement is coupled with a lateral displacement of the molecules along the long molecular axis. This shift is in agreement with reported crystal structure and DFT calculations performed on the crystal^{34,36} RMSD between the resolved crystal structure and the calculated dimer structure of Naph-F4-C2 is only 0.06 Å).

Table 1 Physical properties calculated using the ωB97X-D functional. The Frontier orbital energies, $ε_H$ (HOMO) and $ε_L$ (LUMO), are in good agreement with the ionization potential (IP) and the electron affinity (EA), respectively, indicating physically meaningful Kohn-Sham orbitals. The dipole moments (μ) vanish for Class I and Class III molecules and are sizable for Class II molecules. The quadrupole moments (Q) tend to increase with the number of fluorine atoms. The intermolecular separation (q) for Class I molecules was set to the distance between the molecular centers of mass (c.o.m.), and for Class II and III molecules it was set to the distance between the molecular planes. The intermolecular binding energies (F_{inter}), electronic coupling for hole transport (F_{inter}), and hole transport reorganization energy (calculated using dimers, F_{inter}) and monomers, F_{inter} 0 reveal significant differences between the three molecular classes

Cl.	Molecule	$\varepsilon_{\rm H} [{\rm eV}]$	$\varepsilon_{\mathrm{L}}\left[\mathrm{eV}\right]$	IP [eV]	EA [eV]	μ [D]	Q_{xx} [D Å]	Q_{yy} [D Å]	$Q_{zz}\left[\mathrm{D}\;\mathring{\mathrm{A}}\right]$	a [Å]	F _{inter} [eV]	$\Gamma_{\rm h} [{\rm eV}]$	$E_{\mathrm{r}}^{\mathrm{h,d}}\left[\mathrm{eV}\right]$	$E_{\mathrm{r}}^{\mathrm{h,m}}\left[\mathrm{eV}\right]$
6-31G(d)														
I	Naph	-7.73	0.84	7.75	-0.81	0.00	-62.5	-50.2	-49.6	5.00	-0.337	0.001	0.266	0.236
	Naph-F8	-8.38	-0.11	8.32	0.22	0.00	-99.8	-101.1	-89.3	5.99	-0.264	0.005	0.545	0.458
II	Naph-F4-C1	-8.00	0.33	7.98	-0.24	3.95	-75.1	-70.2	-75.9	3.35	-0.562	0.086	0.422	0.362
	Naph-F4-C2	-8.08	0.35	8.05	-0.27	2.90	-75.8	-75.8	-70.1	3.37	-0.588	0.110	0.454	0.380
III	Naph-F4-T1	-8.03	0.27	8.00	-0.20	0.00	-83.5	-67.1	-75.8	3.51	-0.563	0.380	0.424	0.394
	Naph-F4-T2	-8.35	0.27	8.58	-0.16	0.00	-89.7	-66.8	-75.9	3.50	-0.471	0.378	0.335	0.330
	Pele-F6-T	-7.24	-0.71	7.04	0.93	0.03	-151.6	-126.9	-140.7	3.54	-1.183	0.206	0.318	0.299
cc-pVTZ														
	Naph-F4-T2	-8.57	0.03	8.72	-0.15	0.00	-92.0	-68.0	-76.7	3.50	-0.437	0.363	0.331	0.311

Chemical Science Edge Article

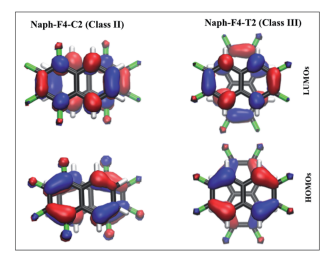


Fig. 4 The dimer orbitals for Naph-F4-C2 (left) and Naph-F4-T2 (right) correspond to a superposition of the monomer Frontier orbitals. Significantly larger overlap between lobes from the HOMOs (lower panels) than for the LUMOs (upper panels) results in an enhanced hole mobility over electron mobilities. The in-plane rotation in Class III molecules (lower right panel) maintains a cofacial arrangement and yields a significantly increased monomer-HOMO overlap as compared to the laterally shifted Class II molecules (lower left panel).

In Class III molecules, a relative in-plane rotation, depending on the overall planar symmetry group, is predicted to enable optimal facial overlap in the stacked structure, see Fig. 3, lower panel. Finally, for the Pele-F6-T molecule, the larger conjugation plane yields an interaction energy of 1.2 eV, which is significantly larger than those found for the naphthalene-based molecules. For this molecule a slight twisting deformation is predicted, induced by H–F interactions.

Electronic couplings for hole transfer, $\Gamma_{\rm h}$, and the corresponding inner sphere reorganization energies, $E_{\rm r}^{\rm h,m}$ and $E_{\rm r}^{\rm h,d}$, calculated using monomer and dimer models,§ respectively, are listed in Table 1. Class III molecules exhibit the largest

electronic coupling for hole transport, \sim 0.2–0.4 eV. Interestingly, the electronic coupling values for electron transport are all small (\lesssim 0.01 eV) except for the case of the larger perylene derivative, where it is 0.11 eV (see ESI, Table 1†). In the following, we will therefore address mobilities for hole transport, unless noted otherwise.

The electronic coupling trends can be understood by considering the relevant orbitals, since coupling values extracted directly from the orbital energies follow the same trend as the couplings obtained *via* CDFT-CI.^{73,77} (Orbitals and orbital energies for all molecules considered in this work can be found in the ESI, Fig. 2–8†). Here, we restrict the discussion to the representative cases of Naph-F4-C2 (Class II) and Naph-F4-T2 (Class III), shown in Fig. 4.

In both cases, the obtained dimer HOMOs, lower panels, (LUMOs, upper panels) resemble a linear combination of two monomer HOMOs (LUMOs). The pair of monomer HOMOs feature substantial overlap within the stacked dimer models, whereas the overlap integral within the pair of LUMOs is either weak (for Class II) or vanishing due to symmetry (for Class III). This can be traced back to the H lobes being oriented along the shifting/rotating direction, whereas L lobes point in the perpendicular direction. Consequently, the dimer state energy splitting between states formed by a superposition of two monomer HOMOs, which serves as a measure for hole coupling, is significantly larger than the dimer state L/L+1 splitting, which serves as a measure for electron coupling, for all naphthalene cases. In the larger perylene-based molecule, on the other hand, mixed orbital lobe orientations occur in the monomer HOMO (see Fig. 5(d)) as well as in the LUMO. Thus, significant splitting related to hole and electron coupling appears, indicating potential ambipolar transport properties.

The larger hole coupling in Class III molecules relative to Class II molecules can be traced back to the displacement of the monomers with respect to each other: expectedly, the lateral displacement in Class II molecules, leads to a loss of the

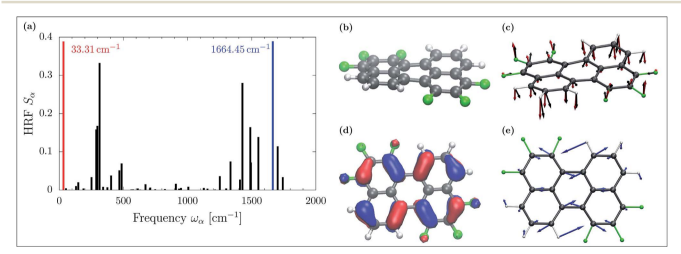


Fig. 5 (a) The Huang–Rhys factors (HRFs) for hole transport in the Pele-F6-T molecule. (b) Fluorination causes an out-of-plane deformation in the Pele-F6-T monomer. (c) Removing an electron from the system relieves that stress, resulting in an relaxation displacement (black), which coincides with the lowest frequency normal mode of frequency 33.31 cm⁻¹ (red). (d) The monomer HOMO is depopulated during the charge transport, causing elongation and contraction of bonds. (e) This movement corresponds to a high frequency normal mode of 1664.45 cm⁻¹.

Edge Article

Table 2 The Marcus rate constants, $k_{\rm M}$, and charge mobilities, $\eta_{\rm M}$, are up to two orders of magnitude lower than $k_{\rm FGR}$ and $\eta_{\rm FGR}$, calculated via Fermi's golden rule. The symmetric selective fluorination results in drastically increased hole mobilities for Class III molecules

Molecule	$k_{\rm M} [{ m s}^{-1}]$	$\eta_{\rm M} \ [{ m cm^2~V^{-1}~s^{-1}}]$	$k_{\rm FGR} \ [{ m s}^{-1}]$	$\eta_{\rm FGR} \ [{ m cm}^2 \ { m V}^{-1} \ { m s}^{-1}]$
6-31G(d)				
Naph	5.52×10^{8}	$0.5 imes 10^{-4}$	2.49×10^{10}	0.0024
Naph-F8	4.75×10^{9}	0.6×10^{-3}	1.48×10^{11}	0.0205
Naph-F4-C1	4.05×10^{12}	0.18	8.46×10^{13}	3.67
Naph-F4-C2	5.21×10^{12}	0.23	1.10×10^{14}	4.83
Naph-F4-T1	5.46×10^{13}	2.60	1.16×10^{15}	55.3
Naph-F4-T2	1.17×10^{14}	5.54	2.14×10^{15}	101
Pele-F6-T	4.81×10^{13}	2.33	5.55×10^{14}	26.9
cc-pVTZ				
	$\textbf{1.37}\times\textbf{10}^{14}$	6.49	2.20×10^{15}	104.23

monomer HOMOs overlap as indicated in Fig. 4, lower left panel. This shift yields a smaller splitting between the H and H–1 dimer orbitals.^{30,78} In contrast, the concentric rotation in Class III molecules achieves an overlap between the two HOMOs, shown in Fig. 4, lower right panel, which is twice as large as in the shifted arrangements. Here the quadrupole-quadrupole attractive interactions appear to be the driving force for maintaining facial overlap. Finally, we stress that in all cases, electronic coupling values are sensitive to the intermolecular arrangement and thus to static and dynamic disorder effects.^{47,79} Further analysis of the electronic coupling trends and their disorder sensitivity is provided in the ESI.†

The reorganization energies $E_{\rm r}$ as listed in Table 1 are within the range of 0.2–0.6 eV. Reorganization energies calculated using ionic dimer CDFT models are reproduced rather well by the simpler ionic monomer models, within 0.1 eV in all cases. We therefore proceed to calculate displacement geometries and HRFs using monomer models.

CT is dominated by modes with large HRFs. The HRFs for Pele-F6-T are presented in panel (a) of Fig. 5 (the HRFs for naphthalene-based molecules and illustration of the key modes are provided in the ESI Fig. 9 and 10,† respectively). The dominant low frequency modes originate from the fluorination-induced bay region distortion, resulting in a slight twist of about 10.5° in the otherwise flat perylene molecule as shown in panel (b) of Fig. 5. Removing an electron from the monomer during hole transport, partially relieves that stress, initiating a twisting motion shown in panel (c), black arrows, which essentially coincides with the low frequency normal mode of 31.31 cm⁻¹ (red). Such dominant low frequency modes do not appear in the smaller and more rigid naphthalene-based molecules without bay regions.

The high frequency normal mode of 1664.45 cm^{-1} provides the dominant contribution to E_r^{in} . Similar to the cases of the naphthalene molecules, such modes are caused by a slightly deformed C=C stretch along the longitudinal molecular axis. This in-plane mode is illustrated with blue arrows in panel (e) of Fig. 5. Its dominant contribution to hole transport can be understood in terms of molecular orbitals: depopulating the

HOMO, depicted in Fig. 5 panel (d), yields weakened π bonds, whereas bonds of low HOMO density contract.

Hole transport FGR rate constants and hole mobilities, as well as their values at the semi-classical Marcus limit obtained at $T=300\,$ K (ref. 80 and 81) are presented in Table 2. Comparison of the FGR rate constants to their Marcus limit reveals that while the Marcus rate constants follow a similar trend, they are one to two orders of magnitude smaller than the FGR rate constants. This emphasizes the deficiency of Marcus theory^{43,79,82} in describing charge transfer (and transport) processes. An important finding of our work is that even in the case of charge transport, where the vanishing reaction free energy guarantees a setting in the Marcus-normal region, nuclear quantum effects should not be neglected.

For estimating the transport rates we have to consider also the external reorganization. However, the outer sphere reorganization energy is unknown for most of the molecules considered here. External reorganization energies are expected to be about a tenth of the internal reorganization energies due to the predominant contribution of low frequency modes. Below, we adopted the naphthalene crystal outer-sphere reorganization energy, $E_{\rm r}^{\rm ex}=35$ meV, from ref. 84, for all compounds. An analysis of the rate constant sensitivity with respect to $E_{\rm r}^{\rm ex}$ can be found in the ESI, Tables 3 and 4,† revealing only marginal influence. We point out that similar values for the $E_{\rm r}^{\rm ex}$ have been reported for other oligoacenes. Larger values of up to 300 meV have been found in few cases and only in the context of photoinduced backbone distortions, e.g. intermolecular torsional or shifting motions, 87,88 which are not expected to occur in the case of charge transport.

Class I molecules show the lowest charge mobilities, which can be traced back to the loose packing and weak coupling. Class II molecules exhibit higher mobilities, which can be traced back to the tighter stacking and stronger coupling. However, the shift between adjacent units reduces the monomer-orbital overlap. Class III molecules exhibit the highest mobilities. This can be traced back to the tight stacking and lack of charge traps due to the vanishing dipole moments. In these molecules, a concentric 90° torsion in the cofacial arrangement maintains large orbital overlap between the monomers.

Overall, fluorination may enhance the mobility by more than three to four orders of magnitude in the case of the naphthalene-based molecules. However for Class III molecules, the stronger electronic coupling values implies that the perturbative FGR approach may not be suitable. Strong electronic coupling may lead to delocalization, and thereby to band-like transport that could further enhance mobilities. So, We also point out that the diffusive Einstein–Smoluchowski equation, eqn (3), assumes hopping-like transport (although tunneling between adjacent molecules is accounted for. In addition, entering the femtosecond regime, transient nuclear non-equilibrium effects might arise, which are not accounted for by the FGR approach used in this work. However, in a recent analysis we find that such transient effects are only of minor influence on the overall transition timescales.

The molecule Pele-F6-T is predicted to exhibit a hole mobility of about $30 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. For reference, the calculated mobility in nonfluorinated perylene in herringbone configuration (as in Class I) is $0.07 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. Thus, fluorination, in the case of the

Chemical Science

perylene derivative, is seen to enhance the mobility by more than two orders of magnitude. It should be noted that the measured mobilities in perylene are 0.017 cm2 V-1 s-1 (electrons) and 0.02 cm² V⁻¹ s⁻¹ (holes), ⁹² while even higher electron mobilities (2.0 cm² V⁻¹ s⁻¹) have been reported under ultrapure conditions.⁹³

In conclusion, the results reported in this paper shed light on the circumstances under which selective fluorination of organic semiconducting molecules can enhance charge mobility. High mobility is found when the functionalization is done so that the overall molecular dipole moment vanishes. The quadrupole, which is the leading electrostatic pole in such cases, can still lead to tight packing, without risking the emergence of charge traps. We also show that while Marcus rates can capture basic trends, they differ quantitatively from the corresponding FGR rates.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

B. D. D. is grateful to support by a NSF-CHEM grant CHE-1362504. E. G. is grateful to support by a NSF-CHEM grant CHE-1464477. B. D. D. and E. G. also acknowledge support from DOE award No. DE-SC0016501. A. S. was awarded an Institute of Complex Adaptive Matter (ICAM) fellowship by the Kent State and University of Michigan ICAM branches. We are also grateful to generous resource allocations on the Ohio Supercomputer Center94 and the Kent State University, College of Arts and Sciences Computing Cluster.

Notes and references

§ In the monomer model, DFT energy calculations yield $E_r^{h,m} = |E_M^M - E_{M^+}^M| + |E_{M^+}^{M^+}|$ $E_{\rm M}^{\rm M^+}|$. Here, the superscript denotes the state, and the subscript identifies the geometry. The dimer based $E_{\rm r}^{\rm h,d}=E_{\rm M^+M}^{\rm M^+M}-E_{\rm M^+M}^{\rm MM^+}$ are obtained via CDFT for a cationic dimer state.

- 1 G. Horowitz, Adv. Mater., 1998, 10, 365-377.
- 2 M. Malachowski and J. Zmija, Opto-Electron. Rev., 2010, 18, 121-136.
- 3 Z. Bao, A. J. Lovinger and A. Dodabalapur, Appl. Phys. Lett., 1996, 69, 3066-3068.
- 4 K. M. Pelzer and S. B. Darling, Mol. Syst. Des. Eng., 2016, 1, 10 - 24.
- 5 K. Gao, J. Miao, L. Xiao, W. Deng, Y. Kan, T. Liang, C. Wang, F. Huang, J. Peng, Y. Cao, F. Liu, T. P. Russell, H. Wu and X. Peng, Adv. Mater., 2016, 28, 4727-4733.
- 6 D. M. DeLongchamp, Semiconductor Materials for Solar Photovoltaic Cells, Springer, 2016, pp. 169-196.
- 7 K. A. Mazzio and C. K. Luscombe, Chem. Soc. Rev., 2015, 44, 78-90.
- 8 A. Dodabalapur, Solid State Commun., 1997, 102, 259-267.

- 9 S. Reineke, F. Lindner, G. Schwartz, N. Seidler, K. Walzer, B. Lüssem and K. Leo, *Nature*, 2009, **459**, 234–238.
- 10 S. Lamansky, P. Djurovich, D. Murphy, F. Abdel-Razzaq, H.-E. Lee, C. Adachi, P. E. Burrows, S. R. Forrest and M. E. Thompson, J. Am. Chem. Soc., 2001, 123, 4304-4312.
- 11 C.-Y. Kuei, W.-L. Tsai, B. Tong, M. Jiao, W.-K. Lee, Y. Chi, C.-C. Wu, S.-H. Liu, G.-H. Lee and P.-T. Chou, Adv. Mater., 2016, 28, 2795-2800.
- 12 L. E. Kreno, K. Leong, O. K. Farha, M. Allendorf, R. P. Van Duyne and J. T. Hupp, Chem. Rev., 2011, 112, 1105-1125.
- 13 J. Kong, N. R. Franklin, C. Zhou, M. G. Chapline, S. Peng, K. Cho and H. Dai, Science, 2000, 287, 622-625.
- 14 B. V. Harbuzaru, A. Corma, F. Rey, P. Atienzar, J. L. Jordá, H. García, D. Ananias, L. D. Carlos and J. Rocha, Angew. Chem., Int. Ed., 2008, 47, 1080-1083.
- 15 A. Gheno, S. Vedraine, B. Ratier and J. Bouclé, Metals, 2016, 6,
- 16 M. Saliba, S. Orlandi, T. Matsui, S. Aghazada, M. Cavazzini, J.-P. Correa-Baena, P. Gao, R. Scopelliti, E. Mosconi, K.-H. Dahmen, F. De Angelis, A. Abate, A. Hagfeldt, G. Pozzi, M. Graetzel and M. K. Nazeeruddin, Nat. Energy, 2016, 1, 1-7.
- 17 Z. Yu and L. Sun, Adv. Energy Mater., 2015, 5, 1500213.
- 18 H. Tsuji, K. Sato, Y. Sato and E. Nakamura, J. Mater. Chem., 2009, 19, 3364-3366.
- 19 J. Halls, C. Walsh, N. Greenham, E. Marseglia, R. Friend, S. Moratti and A. Holmes, Nature, 2000, 376, 498-500.
- 20 C. J. Brabec, C. Winder, M. C. Scharber, N. S. Sariciftci, J. C. Hummelen, M. Svensson and M. R. Andersson, J. Chem. Phys., 2001, 115, 7235-7244.
- 21 J. T. Welch, Selective fluorination in organic and bioorganic chemistry, American Chemical Society, 1991.
- 22 B. Winkler, F. Meghdadi, S. Tasch, R. Müllner, R. Resel, R. Saf, G. Leising and F. Stelzer, Opt. Mater., 1998, 9, 159.
- 23 Q. Lia, Y. Duana, H.-Z. Gaob, Z.-M. Sua and Y. Genga, J. Mol. Graphics Modell., 2015, 59, 50.
- 24 J. Calvo-Castro, G. Morris, A. R. Kennedy and C. J. McHugh, Cryst. Growth Des., 2016, 16, 5385.
- 25 M. Broadhurst, G. Davis, A. DeReggi, S. Roth and R. Collins, Polymer, 1982, 23, 22-28.
- 26 A. Iyer, J. Bjorgaard, T. Anderson and M. E. Köse, Macromolecules, 2012, 45, 6380-6389.
- 27 N. Koch, A. Vollmer, S. Duhm, Y. Sakamoto and T. Suzuki, Adv. Mater., 2007, 19, 112-116.
- 28 E. Di Donato, R. P. Fornari, S. Di Motta, Y. Li, Z. Wang and F. Negri, J. Phys. Chem. B, 2010, 114, 5327-5334.
- 29 K. Fuchibe, T. Morikawa, K. Shigeno, T. Fujita and J. Ichikawa, Org. Lett., 2015, 17, 1126-1129.
- 30 V. Coropceanu, J. Cornil, D. A. da Silva Filho, Y. Olivier, R. Silbey and J.-L. Brédas, Chem. Rev., 2007, 107, 926-952.
- 31 J. H. Yun, S. Park, J. H. Heo, H.-S. Lee, S. Yoon, J. Kang, S. H. Im, H. Kim, W. Lee, B. Kim, J. M. Ko, D. S. Chung and H. J. Son, Chem. Sci., 2016, 6649-6661.
- 32 O. L. Griffith, X. Liu, J. A. Amonoo, P. I. Djurovich, M. E. Thompson, P. F. Green and S. R. Forrest, Phys. Rev. B: Condens. Matter Mater. Phys., 2015, 92, 085404-085409.
- 33 W.-T. Park, G. Kim, C. Yang, C. Liu and Y.-Y. Noh, Adv. Funct. Mater., 2016, 26, 4695-4703.

Edge Article

- 34 I. Y. Bagryanskaya, Y. V. Gatilov, A. M. Maksimov, V. E. Platonov and A. V. Zibarev, *J. Fluorine Chem.*, 2005, 126, 1281–1287.
- 35 B. W. Gung, X. Xue and H. J. Reich, *J. Org. Chem.*, 2005, **70**, 3641–3644.
- 36 F. Cozzi, S. Bacchi, G. Filippini, T. Pilati and A. Gavezzotti, *Chem.-Eur. J.*, 2007, **13**, 7177–7184.
- 37 S. A. Arnstein and C. D. Sherrill, *Phys. Chem. Chem. Phys.*, 2008, 10, 2646–2655.
- 38 X.-H. Ju and X. Liao, Int. J. Mater. Res., 2013, 104, 109-113.
- 39 T. Kato and T. Yamabe, J. Chem. Phys., 2004, 121, 2356-2366.
- 40 D. Kovaček, Z. B. Maksić and I. Novak, *J. Phys. Chem. A*, 1997, **101**, 1147–1154.
- 41 B. Engels and V. Engel, *Phys. Chem. Chem. Phys.*, 2017, **19**, 12604–12619.
- 42 M. H. Lee, B. D. Dunietz and E. Geva, *J. Phys. Chem. C*, 2013, 117, 23391–23401.
- 43 M. H. Lee, E. Geva and B. D. Dunietz, *J. Phys. Chem. C*, 2014, **118**, 9780–9789.
- 44 M. H. Lee, B. D. Dunietz and E. Geva, *J. Phys. Chem. Lett.*, 2014, 5, 3810–3816.
- 45 A. K. Manna and B. D. Dunietz, *J. Chem. Phys.*, 2014, **141**, 121102.
- 46 D. E. Wilcox, M. H. Lee, M. E. Sykes, A. Niedringhaus, E. Geva, B. D. Dunietz, M. Shtein and J. P. Ogilvie, *J. Phys. Chem. Lett.*, 2015, 6, 569–575.
- 47 M. H. Lee, E. Geva and B. D. Dunietz, *J. Phys. Chem. A*, 2016, **120**, 2970–2975.
- 48 J. Jortner, J. Chem. Phys., 1976, 64, 4860-4867.
- 49 A. Nitzan, Chemical Dynamics in Condensed Phases: Relaxation, Transfer and Reactions in Condensed Molecular Systems, Oxford University Press, 2006.
- 50 K. Huang and A. Rhys, *Proc. R. Soc. London, Ser. A*, 1950, **204**, 406–423.
- 51 R. A. Marcus, J. Chem. Phys., 1956, 24, 966-978.
- 52 R. A. Marcus, J. Chem. Phys., 1956, 24, 979-989.
- 53 R. A. Marcus and N. Sutin, *Biochim. Biophys. Acta*, 1985, **811**, 265–322.
- 54 R. A. Marcus, Rev. Mod. Phys., 1993, 65, 599-610.
- 55 G. L. Hornyak, J. Dutta, H. F. Tibbals and A. Rao, *Introduction to nanoscience*, CRC Press, 2008.
- 56 S. Abrahams, J. M. Robertson and J. White, *Acta Crystallogr.*, 1949, 2, 233–238.
- 57 S. Abrahams, J. M. Robertson and J. White, *Acta Crystallogr.*, 1949, 2, 238–244.
- 58 D. Cruickshank, Acta Crystallogr., 1957, 10, 504-508.
- 59 J. Oddershede and S. Larsen, *J. Phys. Chem. A*, 2004, **108**, 1057–1063.
- 60 J. Potenza and D. Mastropaolo, Acta Crystallogr., Sect. B: Struct. Crystallogr. Cryst. Chem., 1975, 31, 2527.
- 61 J. C. Collings, K. P. Roscoe, R. L. Thomas, A. S. Batsanov, L. M. Stimson, J. A. K. Howard and T. B. Marder, *New J. Chem.*, 2001, 25, 1410.
- 62 R. Baer and D. Neuhauser, *Phys. Rev. Lett.*, 2005, **94**, 043002–
- 63 J.-D. Chai and M. Head-Gordon, *Phys. Chem. Chem. Phys.*, 2008, **10**, 6615–6620.

- 64 E. Livshits and R. Baer, *Phys. Chem. Chem. Phys.*, 2007, 9, 2932–2941.
- 65 T. Stein, L. Kronik and R. Baer, J. Am. Chem. Soc., 2009, 131, 2818–2820.
- 66 L. Kronik, T. Stein, S. Refaely-Abramson and R. Baer, *J. Chem. Theory Comput.*, 2012, **8**, 1515–1531.
- 67 H. Phillips, E. Geva and B. D. Dunietz, *J. Chem. Theory Comput.*, 2012, **8**, 2661–2668.
- 68 H. Phillips, Z. Zheng, E. Geva and B. D. Dunietz, *Org. Electron.*, 2014, **15**, 1509–1520.
- 69 R. Sure and S. Grimme, *J. Chem. Theory Comput.*, 2015, **11**, 3785–3801.
- 70 C. Bruckner and B. Engels, *J. Comput. Chem.*, 2016, 37, 1335–1344.
- 71 J. Tomasi, B. Mennucci and R. Cammi, *Chem. Rev.*, 2005, 105, 2999–3094.
- 72 J.-M. Mewes, Z.-Q. You, M. Wormit, T. Kriesche, J. M. Herbert and A. Dreuw, *J. Phys. Chem. A*, 2015, **119**, 5446–5464.
- 73 Q. Wu, B. Kaduk and T. Van Voorhis, J. Chem. Phys., 2009, 130, 034109.
- 74 J. R. Reimers, J. Chem. Phys., 2001, 115, 9103-9109.
- 75 J. Tang, M. T. Lee and S. H. Lin, *J. Chem. Phys.*, 2003, **119**, 7188–7196.
- 76 E. J. Baerends, O. V. Gritsenko and R. van Meer, *Phys. Chem. Chem. Phys.*, 2013, 15, 16408–16425.
- 77 Q. Wu and T. V. Voorhis, J. Chem. Phys., 2006, 125, 164105.
- 78 J.-L. Brédas, J. P. Calbert, D. A. da Silva Filho and J. Cornil, *Proc. Natl. Acad. Sci. U. S. A.*, 2002, **99**, 5804–5809.
- 79 S. Jang and A. Montoya-Castillo, *J. Phys. Chem. B*, 2015, **119**, 7659–7665.
- 80 A. Troisi, Adv. Mater., 2007, 19, 2000-2004.
- 81 C. R. Kagan and C. B. Murray, Nature, 2015, 10, 1013-1026.
- 82 S. Jang and M. D. Newton, *J. Phys. Chem. B*, 2006, **110**, 18996–19003.
- 83 D. Beljonne, J. Cornil, L. Muccioli, C. Zannoni, J.-L. Brédas and F. Castet, *Chem. Mater.*, 2011, 23, 591–609.
- 84 V. Stehr, R. F. Fink, M. Tafipolski, C. Deibel and B. Engels, WIREs Computational Molecular Science, 2016, 6, 694–720.
- 85 J. E. Norton and J.-L. Brédas, *J. Am. Chem. Soc.*, 2008, **130**, 12377–12384.
- 86 D. P. McMahon and A. Troisi, *J. Phys. Chem. Lett.*, 2010, 1, 941–946.
- 87 V. Lemaur, M. Steel, D. Beljonne, J.-L. Brédas and J. Cornil, *J. Am. Chem. Soc.*, 2005, **127**, 6077–6086.
- 88 R. F. Fink, J. Seibt, V. Engel, M. Renz, M. Kaupp, S. Lochbrunner, H.-M. Zhao, J. Pfister, F. Würthner and B. Engels, *J. Am. Chem. Soc.*, 2008, **130**, 12858–12859.
- 89 F. Ortmann, F. Bechstedt and K. Hannewald, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2009, **79**, 235206.
- 90 S. Jang, J. Chem. Phys., 2011, 135, 034105.
- 91 X. Sun and E. Geva, *J. Chem. Theory Comput.*, 2016, **12**, 2926–2941.
- 92 Y. Maruyama, T. Kobayashi, H. Inokuchi and S. Iwashima, *Mol. Cryst. Liq. Cryst.*, 1973, **20**, 373–380.
- 93 W. Warta, R. Stehle and N. Karl, Appl. Phys. A, 1985, 36, 163-170.
- 94 Ohio Supercomputer Center, http://osc.edu/ark:/19495/f5s1ph73, 1987.