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Charge-Transfer Probes for Molecular Recognition via Steric Hindrance in Donor-Acceptor Pairs

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Abstract

Molecular association of various aromatic hydrocarbons (D, including sterically hindered donors) with a representative group of diverse acceptors $(A = \text{quinone}, \text{trinitrobenzene}, \text{densar}$ tetracyanoethylene, tropylium, tetranitromethane, and nitrosonium) is visually apparent in solution by the spontaneous appearance of distinctive colors. Spectral (UV−vis) analyses of the colored solutions reveal their charge-transfer origin (λ_{CT}) , and they provide quantitative

information of the intermolecular association in the form of the K_{DA} and ε_{CT} values for the formation and visualization, respectively, of different [D,A] complexes. Importantly, such measurements establish charge-transfer absorption to be a sensitive analytical tool for evaluating the steric inhibition of donor−acceptor association. For example, the steric differences among various hindered aromatic donors in their association with quinone are readily dramatized in their distinctive charge-transfer (color) absorptions and verified by X-ray crystallography of the charge-transfer crystals and/or QUANTA molecular modeling calculations of optimum intermolecular separations allowed by van der Waals contacts.

Introduction

Molecular recognition and preassociation are conceptually vital to catalytic stereospecificity and other contemporaneous topics in organic chemistry like self-assembly and organization, supramolecular (host−guest) chemistry, etc.¹⁻³Of the various measures available for the quantitative evaluation of intermolecular interactions in solution $-$ especially weak nonbonded ones $-$ the appearance of new spectral bands arising from the charge-transfer (CT) transitions between electron donors and electron acceptors is especially useful and easy to apply.⁴⁻⁶ Indeed, the ubiquitous CT absorptions are diagnostic of a very wide spectrum of intermolecular electron donor−acceptor (DA) interactions arising in extremely stable, isolable 1:1 complexes on one hand,^{7} to highly transient complexes (with collisional lifetimes) at the other extremum.⁸ From a structural point of view, however, it is not at all clear *a priori* what the critical donor−acceptor encounter (distance) must be for the relevant charge-transfer absorptions to be in evidence $\frac{9}{2}$

In order to establish the limits to which charge-transfer is applicable as an analytical probe for intermolecular interactions, we employ in this study four classes of aromatic donors (Chart 1), in which the essential benzenoid (donor) π -chromophore is sterically encumbered to various degrees by increasing alkyl bulk. Thus, the first (simply methylated) member in each class represents the sterically most accessible donor, and all members in each class are of comparable donor strengths.¹⁰

Chart 1

In a complementary way, we identify two classes of sterically graded electron acceptors in Chart 2. As such, the planar π-acceptors are presented in the order of their increasing size from tetracyanoethylene to 1,3,5-trinitrobenzene. Among the σ-acceptors, the diatomic nitrosonium cation and dibromine are the smallest and least subject to steric hindrance, especially in comparison with the larger, tetrahedral acceptors tetranitromethane and carbon tetrabromide. The relative acceptor strengths in Chart 2 are indicated by the trend in the reduction potentials (E° _{red} V vs SCE).^{11,12}

Donor-acceptor pairs in solution are quantitatively monitored in this study by UV−vis spectral changes, and the relevant charge-transfer interactions identified in X-ray crystal structures and compared with molecular modeling calculations. Steric effects are not only exploited in the structural requirements for the intermolecular formation of 1:1 donor−acceptor complexes but also to achieve intramolecular selectivity in biaryls and in a tethered donor containing more than one aromatic center.

Chart 2

Results and Discussion

I. Visual Detection of Intermolecular Donor−Acceptor Associations. Exposure of the various methylbenzenes in Chart 1 to the prototypical π-acceptor chloranil (**CA**) spontaneously resulted in brightly colored dichloromethane solutions. The colorations progressively evolved from yellow (mesitylene) to orange (durene) to purple (hexamethylbenzene) in line with the decreasing ionization potentials of the aromatic donors.¹⁰ Similar vivid colorations were also observed when the same methylbenzenes were mixed with other electron acceptors such as 1,3,5-trinitrobenzene (**TNB**), and further red-shifts in colors occurred with the stronger πacceptor tetracyanoethylene (**TCNE**) as well as the tropylium cation (**TR**+) in Chart 2.

II. Quantification of Donor−Acceptor (π−π) Associations. The UV−vis spectral changes in Figure 1 typically show the monotonic growth of the diagnostic charge-transfer absorbance with its visible maximum at λ _{CT} = 520 nm upon the incremental addition of hexamethylbenzene (**HMB** in Chart 1) to a solution of chloranil (**CA**) in dichloromethane at 24 °C. This well-resolved (featureless) absorption band was ascribed to the intermolecular donor−acceptor association, *i.e*.

 $HMB + CA \stackrel{K_{DA}}{\Longleftrightarrow} [HMB, CA]$ (1)

in which the characteristic color derives from the charge-transfer transition, as originally formulated by Mulliken.⁴ Such a structural assignment was readily verified by the isolation of dark purple crystals of the 1:1 complex in >95% yield from an equimolar solution of hexamethylbenzene and chloranil, simply by the very slow removal of dichloromethane *in vacuo*. X-ray crystallography of the charge-transfer crystal revealed the hexamethylbenzene to be directly juxtaposed on the chloranil and separated by an interplanar distance of $d = 3.51$ Å, as illustrated in the top and side perspectives A_T and A_S (all hydrogens omitted for clarity).^{13,14} Indeed, the observed donor−acceptor separation in the purple crystal is remarkably close to the calculated distance of $d = 3.57$ Å by energy minimization of the intermolecular van der Waals contacts between hexamethylbenzene and chloranil. The predicted structure based on the QUANTA molecular modeling analysis¹⁵ is shown in the space-filling representation **B** below.

For the quantitative analysis of the donor−acceptor association in solution, the spectrophotometric absorbance changes in Figure 1 were treated by the Benesi-Hildebrand procedure[,16](javascript:void(0);) *i.e*.

$$
\frac{[CA]}{A_{\text{CT}}} = \frac{1}{\epsilon_{\text{CT}}} + \frac{1}{K_{\text{DA}}\epsilon_{\text{CT}}} \frac{1}{[D]}
$$
(2)

where A_{CT} is the molar absorbance and ϵ_{CT} is the extinction coefficient of the charge-transfer band at the monitoring wavelength (generally close to λmax). For hexamethylbenzene (D) concentrations much greater than that of chloranil, a plot of [**CA**]/ACT *vs* the reciprocal donor concentration was linear, and the least-squares fit produced a correlation coefficient of greater than 0.999 in the inset of Figure 1. From the slope [*K*_{DA}ε_{CT}] 1and the intercept [ε_{CT}] 1, the values of the association constant and the extinction coefficient were readily extracted as K_{DA} $= 2.8$ M⁻¹ and $\epsilon_{520} = 2800$ M⁻¹ cm⁻¹, respectively. Such a limited magnitude of *K*_{DA} for hexamethylbenzene and chloranil indicated that the donor−acceptor interaction is described as weak at best (Δ*G*°_{DA} = −0.6 kcal mol⁻¹), as typical for the spontaneous formation of electron donor−acceptor complexes of quinones (and other π-electron acceptors) with various types of other electron donors (such as alkenes, enol ethers, sulfides, etc.). $\frac{17}{12}$

Figure 1 Spectral (UV−vis) changes attendant upon the incremental addition of hexamethylbenzene to chloranil in dichloromethane. Inset:  Benesi−Hildebrand plot.

The donor−acceptor interactions of the homologous methylbenzenes in Chart 1 with chloranil showed a progressive red-shift of the charge-transfer band λ _{CT} for **HMB** > **DUR** > **MES**, as listed in Table 1 (column 4). The same trend was observed in the association constants K_{DA}, but the magnitude of the change was somewhat limited (see column 5). Weak but distinctive donor−acceptor interactions of hexamethylbenzene were also indicated by the comparison of the charge-transfer absorptions (λ_{CT}) and the magnitudes of the formation constants (K_{DA}) with the other π-acceptors including trinitrobenzene, tropylium, and tetracyanoethylene, as listed in Tables 1 and 4. For comparative purposes, the intensities of the CT (color) absorptions of the various donor−acceptor complexes in Table 1 are given by the values of K_{DA}ε_{CT} in column 7.¹⁸

Table 1. Donor−Acceptor Association of Various Methylbenzenes with Different π-Acceptors*^a*

^a In dichloromethane containing 4−6 mM acceptor and 50−500 mM aromatic donors at 24 °C.

III. Complete Steric Inhibition of Donor−Acceptor (π−π) Association. Analogous exposure of hexaethylbenzene (**HEB**) to chloranil led to no coloration, and no new CT absorption band was observed in the colorless dichloromethane solution even in the presence of a large excess of **HEB**. Furthermore, many attempts to isolate charge-transfer crystals of the **HEB** complex with chloranil in various molar ratios from dichloromethane, chloroform, acetone, ethyl acetate, etc. were all unsuccessful, and low-temperature crystallization merely led to phase separation of the individual (pure) components. Such a striking difference between hexaethylbenzene and hexamethylbenzene (*vide supra*) was not restricted to chloranil. Thus, all other π-acceptors including trinitrobenzene, tropylium, and tetracyanoethylene showed distinctive CT colorations ranging from yellow to orange to green when mixed with hexamethylbenzene, whereas no (or very faint) colorations were detected with hexaethylbenzene at even higher concentrations. The UV−vis spectral changes in Figure 2 confirmed that neither chloranil nor trinitrobenzene participated in charge-transfer association with hexaethylbenzene, and the smallest acceptors tetracyanoethylene and tropylium showed (at best) very weak CT interactions with **HEB** (Table $2)$. $19,20$

The comparative charge-transfer behaviors notwithstanding, the *intrinsic* electron-donor properties of **HEB** are even somewhat better than those of **HMB** insofar as their relative oxidation potentials of E_{ox}° = 1.59 and 1.62 V *vs* SCE, respectively.¹⁰ In order to clarify this anomalous variation (of λCT *vs E*ox°) between **HMB** and **HEB**, we synthesized the novel hybrid triethylmesitylene (**TEM**) with an intermediate oxidation potential of *E*ox° = 1.61 V *vs* SCE. In

fact, the exposure of **TEM** to chloranil immediately led to the characteristic purple coloration of the charge-transfer association (λ _{CT} = 516 nm), and dark purple crystals of the EDA complex were readily isolated in high yields from an equimolar mixture of triethylmesitylene and chloranil. Indeed, the charge-transfer parameters of the purple [**TEM**, **CA**] were essentially identical to those of the **HMB** analogue. Further comparisons of the **TEM** and **HMB** complexes with trinitrobenzene, tropylium, and tetracyanoethylene in Tables 1 and 2 established the same similarities. However, the association constants of the **TEM** complexes (Table 2, column 5) were always roughly half the value of K_{DA} for the corresponding **HMB** complex (Table 1) in dichloromethane. The latter was consistent with a statistical factor of 0.5 for only half the faces available for association, as established by the unique conformation of **TEM** in the chloranil complex shown below in structure **C** by X-ray crystallography. Such a tripodal arrangement of all three ethyl groups on the opposite face of **TEM** is directly related to the conformation of the ethyl groups on *both* faces of **HEB** with quasi *D*³*^d* symmetry in structure **D** which was previously established by Mislow and co-workers,21 *i.e*.

^a See Table 1.*^b* Not determined.*^c* No new absorption band.*^d* Estimated values, see Experimental Section.

From such a comparative behavior of **HMB**, **TEM**, and **HEB** with chloranil as well as the other π-acceptors **TNB**, **TR**+, and **TCNE** it is easy to conclude that a group of three (1,3,5) ethyl substituents is sufficient for the complete steric inhibition of the face of a benzenoid (donor) chromophore for intermolecular association by a π-acceptor. Moreover, the comparable steric inhibition is achieved by tris-annulations at the α-carbons, as presented in the bicyclic structure **TET** (see Table 2, last 3 entries) and the homologous **TMT**. Energy minimization between chloranil and **HEB** was predicted with the aid of QUANTA molecular modeling calculations to occur at an interplanar distance of $d = 4.50 \text{ Å}$ in structure **E**, and the predicted separation of chloranil and **TET** was $d = 4.51$ Å in structure **F**. As such, the difference of 0.9 Å between the observed separation of 3.6 Å in [**HMB**, **CA**] and the predicted separation of 4.5 Å in [**HEB**, **CA**] could represent a "gray" area in which very weak, but visually (color) and spectrally (CT) observable association may be apparent. In order to pursue this possibility, we synthesized a series of unsymmetrical aromatic donors designed to cover the benzenoid face only partially.

IV. Steric Modulation of Donor−Acceptor (π−π) Associations. Mesitylene (**MES**) yielded a bright yellow solution when exposed to chloranil, but 1,3,5-tri-*tert*-butylbenzene (**TTB**) under the same conditions, as expected, led to no coloration. However, intermediate behavior was shown by the homologous *tert*-butylxylene (**TXY**) which afforded a very pale yellow solution with chloranil, and the coloration with di-*tert*-butyltoluene (**DTT**) was barely discernible. The quantitative effects of these color (intensity) changes are given by the values of *K*DAεCT in Table 3 (column 6),²⁰ which were obtained from the spectrophotometric analysis of the chargetransfer absorptions (Figure 3A) attendant upon the incremental additions of these aromatic donors to chloranil in dichloromethane. The qualitative trend of the color intensity followed the monotonic decrease in the association constant K_{DA} with increasing number of *tert*-butyl groups.²² However, the latter had no significant effect on the intrinsic donor strength, since the values of the oxidation potentials *E*ox° tabulated in column 2 were uniformly invariant. This observation, taken together with the constancy of the charge-transfer transition (λ _{CT} in column 3), indicated that the interplanar separations (*d*) between the chloranil and **MES**, **TXY**, and **DTT** in the donor−acceptor associations were all comparable,²³ despite the decreasing strength of the interaction (ΔG_{DA}). In order to identify the origin of the difference, we carried out the QUANTA molecular modeling analysis of the nonbonding interactions in all four donor−acceptor associations. It is particularly noteworthy that energy minimization was predicted to occur at essentially the same interplanar separation of $d = 3.4$ Å in the chloranil complex with **MES**, **TBX**, and **DTT** in structures **G**, **H**, and **J**, respectively, but at a significantly larger separation of *d* = 4.5 Å for tri-*tert*-butylbenzene in structure **K**. Such a donor−acceptor association in the mono- and di-*tert*-butyl-substituted donors was achieved by a small parallel shift of chloranil away from the *tert*-butyl group(s).²⁴

Figure 2 Steric hindrance in the CT absorption bands of **HMB** (thick line), **TEM** (dashed line), and **HEB** (thin line) complexed with (A) chloranil, (B) trinitrobenzene, and (C) tropylium obtained by spectral subtraction of the uncomplexed acceptor from the experimental UV−vis spectra for (A) 6 mM **CA** with 50 mM **HMB**, 50 mM **TEM**, and 500 mM **HEB** in CH2Cl2, (B) 5 mM **TNB** with 100 mM **HMB**, 100 mM **TEM**, and 500 mM **HEB** in CH2Cl2, and (C) 9 or (4.5) mM **TR**⁺ BF4- with 22 mM **HMB**, 22 mM **TEM**, and (22 mM) **HEB** in CH3CN.

Figure 3 Partial steric hindrance in donor−acceptor association of (A) 4 mM **CA** with 250 mM **MES**, **TXY**, and **DTT**, (B) 4 mM **CA** with 240 mM **ME**, **MEA**, and **MEA2**, and (C) 1.7 mM **TCNE** with 102 mM **PET** and **HEB** in dichloromethane, obtained as difference spectra analogous to those in Figure 2.

Indeed, such limited lateral displacements along the aromatic planes are not expected to be important factors in the charge-transfer transitions to significantly alter the values of λ $\rm{c\tau.}^{25}$

Table 3. Donor−Acceptor Association of the Class II, III, and IV Aromatic Donors with Chloranil*^a*

^a From 4 mM **CA** and 0.01−1 M donor in dichloromethane at 24 °C.*^b* -*^d*See Table 2.

Among the class II donors (based on durene in Chart 1), the tetramethyl derivative **TMA** showed the same "partial" steric behavior relative to the completely hindered octamethyl analogue **OMA** (and the bicyclic version **DMA**) at the other extremum. For example, the results in Table 3 (entry 6) point to the strongly diminished donor−acceptor interaction of **TMA** relative to **DUR** in its association with chloranil,26 in a manner similar to the differentiation of **TXY** and **MES** in the *tert*-butyl series (class III). Similarly, the bis-annulated donors **OMA** and **DMA** were subject to complete steric inhibition, much like tri-*tert*-butylbenzene in class III and **TET** and **TMT** in class I.

The replacement of a pair of methyl groups in **HMB** with methoxy groups render the class IV aromatic ethers to be the best π-donors by virtue of the low E_0x° values in Table 3.²⁷ As such, these methyl ethers were more tolerant to steric encumbrance $-$ the bis-annulated **MEA**₂ being

much less subject to partial steric hindrance than its counterpart **DMA** in class II, as shown by a direct comparison of *K*DA and *K*DAεCT in entry 11 with those in entry 8 in Table 3. Repeated attempts to grow single crystals of the weak yellow complex [**MEA**2,**CA**] for X-ray crystallography were unsuccessful. However, from the significantly blue-shifted value of λ _{CT} = 452 nm in column 4, we tentatively conclude that donor−acceptor association of the strong donor **MEA**² and chloranil probably occurs at an intermediate interplanar separation of *d* greater than 3.5 Å (but less than 4.5 Å).²⁸

A close inspection of the charge-transfer absorption of the small and rather powerful πacceptor tetracyanoethylene (**TCNE**) with high concentrations of the highly hindered hexaethylbenzene revealed a weak but distinctive absorbance at $\lambda_{\text{CT}} = 540$ nm (Table 4).²⁰ In order to determine how hexaethylbenzene could be subject to partial steric hindrance, we synthesized the hybrid pentaethyltoluene (**PET**) to establish evidence for donor−acceptor association with **TCNE**. UV−vis spectral analysis (Figure 3c) indeed revealed a blue complex to be formed in CH₂Cl₂ with values of K_{DA} and ε_{CT} which were substantially less than those evaluated for either **HMB** or **TEM**. In particular, the value of $\epsilon_{CT} = 300$ M⁻¹ cm⁻¹ for the [PET, **TCNE**] was sharply diminished from $\epsilon_{CT} = 5200 \text{ M}^{-1} \text{ cm}^{-1}$ for [HMB, TCNE], and such a significant difference could result from the reduced π-orbital overlap of **PET** and **TCNE**. Although repeated attempts to grow single crystals of [**PET**, **TCNE**] suitable for X-ray crystallography failed, QUANTA molecular modeling studies showed that donor−acceptor association was possible at an interplanar separation of $d = 3.5 \text{ Å}$ by a parallel shift along the aromatic plane, much in the manner previously described in structures **H** and **J**. ²⁴ It is noteworthy that with the exception of the lateral slippage of ∼2 Å, structure **L** for [**PET**, **TCNE**] is akin to the centrosymmetric structure of [HMB, $TCNE$] with $d = 3.35 \text{ Å}$ (established by X-ray crystallography[\)29](javascript:void(0);) as illustrated in the top perspective **M**. If so, the donor−acceptor association observed between the highly hindered **HEB** and tetracyanoethylene, albeit weak (Table 4, entry 4),²⁰ may be achieved by a conformational change of a single ethyl group in hexaethγlbenzene by a 180° rotation about the aromatic−Cα bond,^{21b} so as to effect the partial steric hindrance analogous to the [**PET**, **TCNE**] structure in **L**. ²⁴ Indeed the complete absence of any donor−acceptor association of the conformationally rigid tris-annulated donor **TET** with tetracyanoethylene (Table 4, entry 5) lends a certain credence to this possibility. Be that as it may, the enhanced donor−acceptor association of the tetracyanoethylene with **HEB** (compared to that with chloranil) was in line with its smaller size and increased acceptor strength. In order to consider these factors further, we next asked how molecular shape could influence the selectivity in donor−acceptor associations by utilizing (a) the powerful diatomic cation NO⁺ with E_{red}° = 1.28 V *vs* SCE as well as the uncharged tetranitromethane (TNM) as rather small σ -acceptors¹² and (b) the substituted biaryls and a tethered aromatic system to serve as bichromophoric donors, as follows.

Table 4. Donor−Acceptor Association of Hexaalkylbenzenes with Tetracyanoethylene*a*

^a From 2 mM **TCNE** and 10−100 mM donor in dichloromethane at 24 °C.*^c* ,*^d*See Table 2.

V. Shape Selectivity of Aromatic Donors. A. With σ-Acceptors. Exposure of nitrosonium (NO+) tetrafluoroborate to hexamethylbenzene in acetonitrile immediately resulted in an intense red coloration, and the UV−vis spectral analysis of the bright red solution revealed a well-resolved CT absorption band (λ _{CT} = 337 nm) with a characteristic low-energy band extending beyond 600 nm, as shown in Figure 4A. Although the extinction coefficient of ϵ _{CT} = 3100 M-1 cm-1 in Table 5 was in line with those evaluated for the **HMB** complexes with the πacceptors in Table 1, the association constant of $K_{DA} = 31000 \text{ M}^{-1}$ was more than four orders of magnitude larger – indicative of an exceptionally strong donor-acceptor association of NO⁺ with **HMB**. [30](javascript:void(0);)

Figure 4 Steric effects in charge-transfer absorptions from σ-acceptors with aromatic donors:  (A) 5 mM NO⁺BF₄ with 1 mM **HMB**, HEB, and TET (as indicated); NO⁺BF₄ alone (\cdots) in acetonitrile and (B) 2 M **TNM** with 5 mM **HMB**, **HEB** and **TET** as indicated; **TNM** alone (···) in dichloromethane.

Table 5. Donor−Acceptor Association of Hexaalkylbenzenes with Nitrosonium Tetrafluoroborate*a*

^a In acetonitrile containing 1 mM NOBF⁴ and 5−10 mM donor at 24 °C.

Surprisingly, the same red color (with comparable intensity) was observed when hexaethylbenzene was treated with NO⁺BF₄⁻ under identical (concentration) conditions, and the UV−vis spectrum in Figure 4A confirmed the mostly unaltered charge-transfer absorption. More surprising were the results in Table 5 which showed that the highly hindered trisannulated donors **TET** and **TMT** enjoyed undiminished donor−acceptor association with NO+, the measured association constants in all cases being uniformly large, with $K_{DA} > 3 \times 10^4$ M⁻¹. In order to account for such an unexpected stability, we grew red crystals of the **HEB** and **TET** complexes with NO⁺SbCl₆ for X-ray crystallographic analysis (see Experimental Section). Indeed, the excellent quality of both single crystals allowed an unambiguous assignment of the nitrogen terminus of the slightly tilted NO⁺ to be centrally situated over the benzenoid (donor) chromophore at a nonbonded distance of 2.08 Å in structures **N** and **P**. The donor−acceptor association, characterized as the very close encounter of NO+ to the benzenoid centers of both hexaethylbenzene and **TET** (inside van der Waals distance), is achieved by significant incursion within the "picket fence" formed by three (alternating) ethyl groups in structure **N** and three ethano bridges in structure **P**. The tight fit of NO+ within the van der Waals cavity in the **HEB** complex (shown in **N**) is sufficient to severely dampen its librational (crystallographic) disorder.31 It is particularly noteworthy that such a donor−acceptor interaction derives from the intrinsic donor properties of **TET** and **HEB** that are akin to that in the electron-rich **HMB**, [33](javascript:void(0);) as shown by the constant values of E_{ox} ° in Table 4. As such, the nitrosonium association in the **TET** and **HEB** complexes (which occurs in the teeth of the potentially repulsive interactions with the ethyl and ethano substituents) is allowed by the dimensions of the (van der Waals) cavity sufficient to accommodate the diatomic acceptor. 34 The three-dimensional requirements for the nestling of NO+ is graphically illustrated by the space-filling representations **R** and **S**. For comparison, the molecular structure of the unhindered hexamethylbenzene complex^{32b} is presented in **Q**.

As a further elaboration of the shape-selectivity of aromatic donors, we examined the donor−acceptor interaction with the tetrahedral σ-acceptor tetranitromethane (**TNM**). Thus, the exposure of a colorless solution of **TNM** to hexamethylbenzene in dichloromethane immediately resulted in a dark red solution, the UV−vis spectrum of which showed an intense (unresolved) CT absorption that extended to well beyond 600 nm (Figure 4B). 35 The quantitative treatment of the absorbance data according to the Benesi−Hildebrand procedure yielded the values of K_{DA} and ε_{CT} for the donor−acceptor association in Table 6 which were substantially less than those for the NO⁺ complex. It is particularly noteworthy that the association of **TNM** with the hindered **HEB** resulted in a blue-shift of the nondescript CT tail absorption in Figure 4B, but the quantitative (UV−vis) spectral analysis indicated that the association constant of $K_{DA} = 1.5 M^{-1}$ was only slightly less than that with **HMB** (by about a factor of 3). Furthermore, the highly hindered tris-annulated donors **TET** and **TMT** were not significantly different from **HEB** insofar as their association with **TNM**. We interpret the rather invariant values of the extinction coefficients in Table 6 (column 5) to be consistent with optimal CT coupling of the small-sized **TNM** irrespective of steric hindrance from the aromatic donor. Although attempts to grow single crystals of the **TNM** complexes were unsuccessful, the QUANTA molecular modeling calculations revealed rather large donor−acceptor separations of a single NO2 group of **TNM** to the **HMB** centroid in structure **T** and to the trisannulated **TET** in structure **U**. [36](javascript:void(0);)

^a In dichloromethane containing 0.2 M **TNM** and 5−50 mM donor at 24 °C.

The mixture of hexamethylbenzene with carbon tetrabromide (colorless) resulted in a pale yellow solution which showed a weak tailing UV−vis absorbance without a discernible absorption maximum arising from the blue-shifted charge-transfer band. Such an overlap of the CT absorption with the local absorption of CBr4 precluded a quantitative evaluation of the donor−acceptor association.

B. In Bichromophoric Systems. Exposure of *p*,*p*'-dimethylbiphenyl (**T-T**) to chloranil in dichloromethane was accompanied by an immediate color change to a bright purple solution. UV-vis spectral analysis of the well-resolved CT absorption band (λ _{CT} = 502 nm in Table 7) indicated a very weak association of **T-T** to chloranil with $K_{DA} = 0.09$ M⁻¹ and a value of $\epsilon_{CT} =$ 6000 $M⁻¹$ cm⁻¹ in line with the other chloranil complexes with methylbenzenes in Table 1. By analogy with the structure **V** (top perspective) of the biphenyl/tetracyanobenzene complex previously established by X-ray crystallography, 37 a similar centrosymmetric structure for the intermolecular association of the bitolyl **T-T** with chloranil was indicated. In marked contrast, the unsymmetrically permethylated homologue pentamethylphenyl-*p*-toluene (**PM-T**) afforded a dark purple solution. The UV-vis spectral analysis of the charge-transfer absorption (λcτ = 510 nm) indicated a substantially larger association with $K_{DA} = 0.99$ M⁻¹ but significantly diminished value of ϵ_{CT} = 700 M⁻¹ cm⁻¹, which we interpreted as a shape-selective association

of chloranil with the pentamethylphenyl moiety. Indeed, X-ray crystallography of the purple [**PM-T**,**CA**] confirmed this spectral assignment and revealed the interplanar separation of *d* = 3.41 Å with significant slippage along the pentamethylphenyl plane to avoid the orthogonal tolyl moiety as shown in the side and top perspectives W_S and W_T.

By comparison, the unsymmetrically hindered pentaethylphenyl-*p*-toluene (**PE-T**) with chloranil afforded a yellow solution. The charge-transfer absorption λ _{CT} = 437 nm and ϵ _{CT} = 70 M⁻¹ cm⁻¹ in Table 7 were indicative of an analogous slipped structure for [**PE-T,CA**], but with the chloranil complexed to the less hindered tolyl moiety.³⁸

Table 7. Donor−Acceptor Association of Various Biphenyls and the Tethered Aromatic Donor with Chloranil*^a*

DONOR	E_{ox}^o	λ_{CT}		K_{DA} ε_{CT}	$K_{\text{DA}}\varepsilon_{\text{CT}}$
	$(V)^b$	(nm)		(M^{-1}) $(M^{-1}$ cm ⁻¹)	
	$T-T = 1.64$	502	0.09	6,000	500
PM-T 1.62 510			0.99	700	700
PE-T	1.67	437	1.2	70	90
OMe $HMB-MEA_2$ 1.62, 514 1.0	1.11			2,600	2,600

^a See Table 1.

A qualitative view of steric hindrance was also useful in predicting the site selectivity of donor−acceptor association in bifunctional aromatic donors that were not directly connected, as in the bichromophoric **HMB**-**MEA**² in which hexamethylbenzene (**HMB**) was tethered to the bis-annulated ether **MEA**2. Thus the treatment of the tethered donor **HMB**-**MEA**² with chloranil afforded a dark purple solution in which UV−vis spectral analysis yielded the charge-transfer parameters in Table 7 (entry 4) that were essentially identical with those obtained for the mononuclear **HMB** in Table 1 (entry 1). Indeed, X-ray crystallographic analysis of the [**HMB**-**MEA**2, **CA**] complex provided unambiguous support for the predicted structure **Y**.

Summary and Conclusions

Visualization (color) and attendant charge-transfer absorption are reliable and sensitive analytical probes for monitoring the donor−acceptor (DA) association of the various aromatic hydrocarbons in Chart 1 with the different types of π- and σ-acceptors in Chart 2. Thus, the facile molecular association of the polymethylbenzenes, including hexamethylbenzene **HMB**, with all the π-acceptors is immediately apparent by the vividly colored solutions, the UV-vis spectral analysis of which reveals new charge-transfer absorption bands (λ_{CT}) for the quantitative evaluation of the association constant (K_{DA}) as well as the CT color (ϵ _{CT}). In marked contrast, the homologous congener hexaethylbenzene **HEB** shows neither color change nor charge-transfer absorption when exposed to the same π-acceptors under identical conditions. Such unfavored molecular associations of **HEB** are readily ascribed to its unique conformational structure **D**, in which the alternant ethyl groups sterically protect both benzenoid faces from the close approach of all planar π-acceptors. Indeed, QUANTA molecular modeling calculations predict the steric envelope around **HEB** to have an average van der Waals "thickness" of 2r \simeq 6.4 Å, which is significantly larger than that (4.0 Å) for the lower homologue **HMB**. As such, we conclude, that the difference of Δ*r* = 1.2 Å represents an upper limit for the molecular association of any of the π-acceptors with the benzenoid chromophore.

X-ray crystallography establishes the hexamethylbenzene association with chloranil to be optimized at an interplanar distance of $d = 3.51$ Å in the cofacial (sandwich) structure **A**. It is noteworthy that such a donor−acceptor separation corresponds to the sum of van der Waals contact of hexamethylbenzene and chloranil, $39a$ and this conclusion is supported by the **HMB**/**CA** separation of *r* = 3.57 Å calculated with the aid of QUANTA molecular modeling. It is tempting to conclude from this computational analysis that molecular associations leading to charge-transfer absorptions derive from *inner-sphere* interactions involving the intimate van der Waals contact of the donor and acceptor chromophores.^{39b} According to this formulation, any steric encumbrance of either the donor or acceptor (or both) that extends much beyond the sum of van der Waals radii of the chromophores will lead to sharply diminished chargetransfer absorptions. However, shape selectivity can be an ameliorating factor in at least 2 ways. *First*, unsymmetrical steric encumbrances in π-donors [such as those in the mono-and di-*tert*-butyl derivatives **TXY** and **DTT**, as well as in semi-annulated **TMA** and **MEA** (see Chart 1)] allow the close cofacial (inner-sphere) approach of π-acceptors by a small parallel shift or

"slippage" along the aromatic planes, as illustrated by structures **H**, **L**, and **W**. *Second*, small electron-poor molecules such as σ-acceptors [in which the electron deficiency is either largely localized at a single atom as in NO^+ , CBr_4 , Br_2 , etc. or on a small group of atoms as in RNO_2 , O₃, ArN₂⁺, SO₂, etc.] can approach the benzenoid chromophore in sterically encumbered donors [such as hexaethylbenzene **HEB** and the multiply annulated analogues **TET**, **TMT**, **OMA**, **DMA**, and **MEA**2] with van der Waals cavities of sufficient size to allow the nestling of a single (acceptor) center as in the NO+ complexes with structures **N** and **P** or close approach of multiple centers as in the **TNM** complexes with structures **T** and **U**.

The modulation of steric effects also allows the selective complexation of unsymmetrically substituted biphenyls as in the slipped structure **W** and in the tethered bichromophoric donor as in structure **Y**. The chemical consequences of such shape-selective complexations of (poly)chromic donors and acceptors will be presented separately.

Experimental Section

Materials. Hexaethylbenzene (Acros), hexamethylbenzene, durene, 4,4'-dimethylbiphenyl, 1,3,5-tri-*tert*-butylbenzene, and 3,5-di-*tert*-butyltoluene (Aldrich) were purified by repeated crystallization from ethanol and heptane. Mesitylene (Aldrich) and 1-*tert*-butyl-3,5 dimethylbenzene (Wiley) were purified by fractional distillation. Tetrachloro-*p-*benzoquinone and tetracyanoethylene (Aldrich) were sublimed *in vacuo* and recrystallized from benzene. 1,3,5-Trinitrobenzene was purified by crystallization from ethanol. Tropylium tetrafluoroborate (Aldrich) and nitrosonium tetrafluoroborate (Pfaltz and Bauer) were purified by recrystallization from an acetonitrile/dichloromethane mixture. Dimethoxydurene, 40 9,10-dimethoxy-1,2,3,4,5,6,7,8-octahydro-1,4:5,8-dimethanoanthracene,⁴¹ 1,2,3,4,5,6,7,8-octahydro-1,4:5,8dimethanoanthracene, 42 1, 4-dimethoxy-2, 3-dimethyl-5, 6, 7, 8-tetrahydro-5, 8methanonaphthalene:41 [oil, 1H NMR (CDCl3) δ 1.32 (d, *J* = 7.2 Hz, 2H), 1.54 (d, *J* = 8.7 Hz, 1H), 1.76 (d, *J* = 8.7 Hz, 1H), 2.00 (d, *J* = 7.2 Hz, 2H), 2.20 (s, 6H), 3.64 (s, 2H), 3.82 (s, 6H); 13C NMR (CDCl3) 12.64, 27.39, 40.89, 49.00, 61.04, 127.53, 137.64, 148.08], 1,1,4,4,5,5,8,8 octamethyl-1,2,3,4,5,6,7,8-octahydroanthracene⁴³ [mp 222−223 °C (lit.⁴³ mp 220−222 °C); ¹H NMR (CDCl3) δ 1.52 (s, 24H), 1.90 (s, 8H), 7.44 (s, 2H); 13C NMR (CDCl3) 32.25, 34.18, 35.54], 1,1,4,4-tetramethyl-1,2,3,4,5,6,7,8-octahydroanthracene⁴³ [mp 121-122 °C (lit. mp⁴³ 121−123 °C); 1H NMR (CDCl3) δ 1.38 (s, 8H), 1.71 (s, 4H), 1.88 (sym m, 4H), 2.84 (sym m, 4H), 7.11 (s, 2H); 13C NMR (CDCl3) 26.61, 29.30, 32.22, 34.05, 35.44, 127.15, 134.47, 142.31], 2,3,4,5,6-pentaethyl-4'-methylbipheny[l44](javascript:void(0);) [mp 89−91 °C; 1H NMR (CDCl3) δ 1.05 (t, *J* = 7.4 Hz, 6H), 1.35 (t, *J* = 7.5 Hz, 6H), 1.38 (t, *J* = 7.5 Hz, 3H), 2.44 (q, *J* = 7.4 Hz, 4H), 2.52 (s, 3H), 2.82 (q, *J* = 7.5 Hz, 4H), 2.85 (q, *J* = 7.5 Hz, 2H), 7.24 (d, *J* = 7.8 Hz, 2H), 7.31 (d, *J* = 7.8 Hz, 2H); 13C NMR (CDCl3) 16.00, 16.10, 21.46, 22.41, 23.79, 128.53, 130.04, 135.72, 137.62, 138.45, 139.36, 139.58, 140.29], and 2,3,4,5,6,4′-hexamethylbiphenyl⁴⁵ [mp 90−91 °C; ¹H NMR (CDCl3) δ 2.09 (s, 6H), 2.40 (s, 6H), 2.44 (s, 3H), 2.54 (s, 3H), 7.14 (d, *J* = 7.8 Hz, 2H), 7.35 (d, *J* = 7.8 Hz, 2H); 13C NMR (CDCl3) 16.82, 16.98, 18.59, 21.41, 129.14, 129.57, 131.00, 132.45, 134.00, 135.81, 139.99, 140.18], 1,2,3,4,5,6,7,8,9,10,11,12-dodecahydro-1,4:5,8:9,12 triethanotriphenylen[e46](javascript:void(0);) (**TET**), and 1,2,3,4,5,6,7,8,9,10,11,12-dodecahydro-1,4:5,8:9,12 trimethanotriphenylen[e47](javascript:void(0);) (**TMT**) were available from literature procedures. Synthesis of the

tethered 9-[3-(pentamethyl-phenyl)-1-propyloxy]-10-methoxy-1,2,3,4,5,6,7,8-octahydro-1,4:5,8 dimethanoanthracene (**HEB**-**MEA**2) will be described elsewhere. Pentaethyltoluene (**PET**) was prepared from hexaethylbenzene by refluxing a mixture of hexaethylbenzene (6.15 g, 25 mmol) and acetyl chloride (2.0 g, 25.5 mmol) in carbon disulfide (25 mL) in the presence of anhydrous aluminum chloride for 8 h. The resulting deep brown solution was cooled to room temperature, poured over a mixture of ice (200 g), concentrated hydrochloric acid (25 mL), and extracted with ether $(4 \times 50 \text{ mL})$. The combined ether extracts were washed with water and dried over anhydrous magnesium sulfate. Removal of the solvent and recrystallization form ethanol afforded white needles of pentaethylacetophenone⁴⁸ (6.2 g, 95%); mp 136−137 °C (lit.48 mp 136−137 °C). A solution of pentaethylacetophenone (6.0 g, 23 mmol) in trifluoroacetic acid (25.5 mL) and water (4.5 mL) was refluxed for 16 h and cooled to room temperature. The dark brown reaction mixture was poured over ice and extracted with ether $(4 \times 50 \text{ mL})$. The ether layers were washed with water and dried over anhydrous magnesium sulfate. Evaporation of the solvent *in vacuo* furnished pentaethylbenzene as an oil [GC-MS *m/z* 218 $(M⁺)$, calcd for C₁₆H₂₆]. The resulting crude pentaethylbenzene was dissolved in acetic acid (15 mL) and treated with paraformaldehyde (0.75 g, 25 mmol) and 5 mL of a 31 wt % HBr in acetic acid.49 The mixture was heated for 12 h at 80 °C and then poured into 100 mL water. The crystalline precipitate thus formed was filtered off, washed with water, and dried *in vacuo* to afford (bromomethyl)pentaethylbenzene (7.0 g, 22.5 mmol); GC-MS *m/z* 310, 312 (M+), calcd for C17H27Br. To a solution of crude (bromomethyl)pentaethylbenzene (6.22 g, 20 mmol) in anhydrous diethyl ether (100 mL) was added lithium aluminum hydride (0.76 g, 20 mmol), and the mixture was refluxed for 2 h. The usual aqueous workup resulted an oily residue which upon crystallization form ethanol afforded pure pentaethyltoluene as colorless prisms (3.7 g, 80%):  mp 45−46 °C; 1H NMR (CDCl3) δ 1.44 (sym m, 15 H), 2.54 (s, 3H), 2.93 (quintet, 10H); 13C NMR (CDCl3) 14.86, 15.38, 15.91, 16.07, 22.24, 22.51, 23.16, 132.18, 137.71, 138.26, 138.62. GC-MS *m/z* (M+), 232, calcd for C17H28. Anal. Calcd for C17H28:  C, 87.86; H, 12.14. Found:  C, 88.06; H, 12.00. Triethylmesitylene (**TEM**) was prepared by dropwise addition of a 30% solution of HBr in acetic acid (100 mL) to a stirred mixture of paraformaldehyde (12 g, 0.4 mol) and 1,3,5-triethylbenzene (16.2 g, 0.1 mol) in glacial acetic acid (25 mL) at 25 °C.⁴⁹ The resulting mixture was heated at 100 °C for 72 h, cooled to room temperature, and poured over ice−water (250 mL) mixture. The precipitate thus obtained was filtered, washed with water, and dried *in vacuo*. The crude pale brown solid (38 g) was dissolved in diethyl ether (200 mL) and added dropwise to a suspension of lithium aluminum hydride (10 g) in diethyl ether during a 1 h period. The resulting mixture was refluxed for 4 h and cooled to room temperature. The usual workup as above and crystallization from ethanol afforded pure triethylmesitylene (74%):  mp 55−56 °C; 1H NMR (CDCl3) δ 1.22 (t, *J* = 7.5 Hz, 9H), 2.37 (s, 9H), 2.79 (q, *J* = 7.5 Hz, 6H); 13C NMR (CDCl3) 13.97, 15.50, 23.79, 131.43, 138.88. GC-MS *m/z* (M+), 204, calcd for C15H24. Anal. Calcd for C15H24:  C, 88.16; H, 11.84. Found:  C, 88.37; H, 11.68.

Dichloromethane (Mallinckrodt analytical reagent) was repeatedly stirred with fresh aliquots of concentrated sulfuric acid (∼20% by volume) until the acid layer remained clear. After separation, it was washed successively with water, aqueous sodium bicarbonate, water, and aqueous sodium chloride and dried over anhydrous calcium chloride. The dichloromethane was distilled twice from P₂O₅ under an argon atmosphere and stored in a Schlenk tube

equipped with a Teflon valve fitted with Viton O-rings. Acetonitrile (Fischer) was stirred with KMnO₄ for 24 h, and the mixture was refluxed until the liquid was colorless. The MnO₂ was removed by filtration. The acetonitrile was distilled from P_2O_5 under an argon atmosphere and then refluxed over CaH2 for 6 h. After distillation from the CaH2, the solvent was stored in a Schlenk flask under an argon atmosphere. The UV−vis absorption spectra were recorded on a Hewlett-Packard 8450A diode-array spectrometer. The ¹H and ¹³C NMR spectra were recorded on a General Electric QE-300 spectrometer, and chemical shifts are reported in ppm units downfield from tetramethylsilane.

The Formation Constants of Donor−Acceptor Complexes. General Procedure. Typically, in a 1-cm square quartz cuvette (UV cell) equipped with a side arm and Schlenk adapter was placed 1−4 mM solution of chloranil (with the aid of a hypodermic syringe) under an argon atmosphere. A known amount of arene donor was added in increments, and the absorbance changes were measured at the absorption maxima as well as two other wavelength close to the absorption maxima (see Figure 1). The absorbance data were then evaluated with the aid of the Benesi-Hildebrand correlation in eq 2.¹⁶ From the linear plot of [CA]/*A*_{CT} against [arene]⁻ ¹, consisting of at least eight data points, the slope was estimated as $(K_{DA} \epsilon cT)^{-1}$ and the intercept as $(\epsilon c_{\text{T}})^{-1}$. Linear fits obtained by least square method had a correlation coefficient of >0.999. The runs were made in duplicate to ensure the reproducibility of the spectra. Detailed procedures for the quantification of DA complex formation with various electron acceptors such as tetracyanoethylene,¹⁸ 1,3,5-trinitrobenzene,¹¹ tropylium tetrafluoroborate, $\frac{50}{2}$ tetranitromethane, 35 and nitrosonium tetrafluroborate 30 have been described previously. **Estimation of the Formation Constants for DA Complexes from Hindered Donors.** In cases of the hindered arene donors (in which no charge-transfer absorption band was detected), the upper limits of the formation constants (K_{DA}) were estimated as follows. The detection limit of the spectrophotometer was taken as 0.01 absorbance unit for an average charge-transfer extinction coefficient of ϵ c τ = 1000 M⁻¹ cm⁻¹. We calculated the lowest concentration of the donor−acceptor complex detectable by UV−vis spectroscopy to be [D,A] = $0.01/\epsilon$ CT = 1 x 10⁻⁵ M. If no DA complex was detected at a donor concentration of [D] = 0.1–0.5 M and an acceptor concentration of [A] = 4−10 mM, the upper limit for the formation constant was taken as *K*_{DA} = [DA]/[D] [A] < 0.025-0.002 M⁻¹ (see Tables 1 and 2). The DA complexes with K_{DA} < 0.025 M⁻¹ were generally considered to be contact charge-transfer complexes. 8.51

Simulation of the Intermolecular Separations in Donor−Acceptor Complexes by Molecular Modeling Calculations. The most energetically favorable separation of the donor and acceptor in isolated [D,A] complexes was searched with the aid of molecular modeling calculations (CHARMm program,⁵⁴ vers. 22, Molecular Simulation Inc., 1994) which included both intramolecular force field (such as covalent bond, bond angle, and torsion angle potentials) and intermolecular interactions (including Columbic and Lennard-Jones potential terms). The initial molecular models of the interacting arene donors and the various electron acceptors (see Charts 1 and 2) were constructed with the aid of the graphics package QUANTA15 running on a Silicon Graphics ONYX Reality workstation. The optimization of the geometry of the individual molecules was carried out by initial energy minimization using the Steepest Descents (SD) algorithm $\frac{53}{2}$ to remove any obvious improper conformations, and this was followed by 1000 steps of ABNR⁵⁴ (Adopted Basis of Newton-Raphson) algorithm in "CHARMm" to reach the minimum of molecular potential energy. The charges of the individual molecules were balanced using Gasteiger's method,⁵² thereby resulting in neutral molecules. The energy minimization package CHARMm allowed any number of molecules within a single structure file to be simulated. Thus, the optimized donor and acceptor molecules were put together in random orientation and the intermolecular separation distance between them was varied from 3.0 to 5.0 Å, and the relative orientation of the interacting molecules was also varied prior to the energy minimization. The energy minimization of the various resulting aggregates was carried out in a similar way as described for the individual molecules, i.e., using the Steepest Descents algorithm⁵³ to remove unfavorable steric contacts followed by 1000 steps of ABNR54 algorithm. The geometry and the separation distances between donor and acceptors molecules of the minimized structures were analyzed using the XP graphical package⁵⁶ after the energy tolerance was satisfied.

Isolation and X-ray Crystallography of Donor−Acceptor Complexes. The chloranil complexes were crystallized from an equimolar solution of **CA** and **HMB**, **TEM**, or **PM-T** by very slow evaporation of dichloromethane. Since the similar treatment of the tethered **HEB-MEA**₂ yielded an amorphous solid, a 1:1 CH₂Cl₂/CCl₄ mixture was used to prepare dark purple crystals of [**HMB-MEA**2,**CA**]. For the nitrosonium complexes, hexaethylbenzene (**HEB**) was added to a flask that contained nitrosonium hexachloroantimonate, and a minimum amount of dichloromethane was added under an argon atmosphere at 0 °C. The undissolved solid was removed by filtration, and the resulting dark colored solution was carefully layered with anhydrous toluene (30 mL) and stored in the refrigerator (−10 °C). After 48−72 h, dark red crystals of **HEB**/NO+ complex were deposited at the dichloromethane/toluene interface. A similar procedure yielded dark red crystals of the **TET**/NO+ complex.

The X-ray crystallography of the donor−acceptor complexes was carried out with a Siemens SMART diffractometer equipped with a CCD detector at −150 °C. The structures were solved by direct methods⁵⁵ and refined with IBM Pentium and SGI Indigo computers. In the various chloranil complexes, the intramolecular dimensions of **CA** and the arene donors were the same as those observed in uncomplexed molecules. However, the arene complexed to nitrosonium cation in [**HEB**,NO+] and [**TET**,NO+] showed slight lengthening of all aromatic C C double bonds when compared with crystal structures of the uncomplexed donors. The pertinent intermolecular orientations as PLUTO plots (generated from the XP graphical package)56 are shown in structures **A**, **C**, **N**, **P**, **W**, and **Y**, and the critical interplanar separations (d) are included. The crystallographic data are on deposit and can be obtained from the Director, Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge, CB2 1EZ, U.K.

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Tables of crystal structure data for [**TEM,CA**], [**PM-T,CA**], [**HMB-MEA**2,**CA**], [**HEB**,NO+SbCl6 -], and [**TET**,NO+SbCl6-] including atomic coordinates, anisotropic thermal parameter, bond lengths, and bond angles and ORTEP diagrams showing thermal ellipsoids (36 pages). See any current masthead page for ordering and Internet access instructions.

PDF [ja9393.pdf \(1.1 MB\)](https://pubs.acs.org/doi/suppl/10.1021/ja9720319/suppl_file/ja9393.pdf)

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- **[23](javascript:void(0);)**For the relationship that connects the separation (*d*) with λ _{CT} and ϵ _{CT}, see: footnote 5.
- **[24](javascript:void(0);)** (a) The barrier to the rotation of the aryl−ethyl bond in **HEB** has been estimated to be 11.8 kcal mol^{-1,21a} (b) The reduced overlap of the donor−acceptor π-orbitals as a result of such a parallel shift is reflected in a corresponding decrease in the ϵ _{CT} values with increasing number of *tert*-butyl groups in Table 3 (see entries 1−4). (c) According to Mulliken,^{4a} the charge-transfer intensity (Κ_{DΑ}ε_{CT}) derives from the transition moment which relates to the overlap integral of the donor and acceptor orbitals (see:  Orgel, L. E.; Mulliken, R. S. *J. Am. Chem. Soc*. **1957**, *79*, 4839.
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- **[27](javascript:void(0);)** See: Foster, R. in ref 4 for the relationship between the donor strength and E_{ox}° , as measured electrochemically (see Experimental Section).
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- **[34](javascript:void(0);)** The van der Waals diameters of the cavities in **HEB** and **TET** were estimated to be ∼3.8 and 3.6 Å, respectively, based on the published structures. $9,21a$
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