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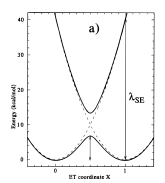
Effects of Bridge Redox State Levels on the Electron Transfer and Optical Properties of Intervalence Compounds with Hydrazine Charge-Bearing Units

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This work concerns zero driving force thermal electron transfer (ET) reactions within charge-localized symmetrical intervalence (SIV) compounds. SIV compounds have different charges on otherwise identical charge-bearing units that are connected by a bridge, and most examples studied have had transition-metal charge-bearing units.¹ SIV compounds show charge-transfer (CT) bands from which two fundamental parameters for ET, the vertical reorganization energy (λ) and the electronic coupling matrix element (V), may be obtained using Marcus-Hush theory.² We designate the CT band corresponding to ET between the chargelocalized minima, as well as the λ and V associated with this band, with subscript SE (for superexchange).³ In the Marcus— Hush model (see Figure 1a), the diabatic SE energy surfaces are parabolas centered at 0 and 1, respectively, on an ET coordinate X. Electronic coupling through the bridge, measured by the offdiagonal matrix element ($V_{\rm SE}$) in a 2 \times 2 secular determinant produces a ground-state double-well adiabatic energy surface and a single minimum excited-state surface. Their energy separation at the ground-state minimum is λ_{SE} , which is the transition energy of the CT band at its maximum ($h\nu_{\text{max}}$). Hush derived a simple equation for evaluation of $V_{\rm SE}$ from the ${\rm CT}_{\rm SE}$ band. ^{2e} The most direct test of ET parameters obtained from a CT_{SE} band is comparing the calculated rate constant for intramolecular ET with that measured experimentally (k_{et}) , but this test has not been applied to metal-centered examples because the ket values calculated are too large to measure.4 Hydrazines have far larger internal vibrational reorganization energies (λ_v) than metal complexes, so their λ_{SE} values are much higher. This allows k_{et} to be in the measurable range even when $V_{\rm SE}$ is rather large, ⁵ making the CT band intense enough to observe easily. The durene-bridged compound $\mathbf{1}^+$ has $k_{\rm et} = 2.6 \times 10^8 \, \rm s^{-1}$ at $-8 \, ^{\circ}\mathrm{C}$ in CH₃CN, determined by dynamic electron spin resonance (ESR) spectroscopy.⁶ It has a large enough $V_{\rm SE}$ to make $k_{\rm et}$ fall in the adiabatic regime, where it is very sensitive only to $\Delta G^{*.2}$ A slight modification of the Marcus-Hush analysis of CT_{SE} bands allows



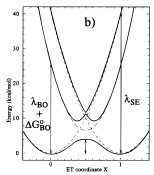


Figure 1. (a) Marcus—Hush plot for 1⁺ showing the diabatic SE surfaces (broken lines) and the adiabatic surfaces (solid lines) obtained using λ_{SE} = 40.3 kcal/mol and $V_{\rm SE}$ = 3.3 kcal/mol derived from the experimental CT band in acetonitrile. (b) Three-state model¹¹ plot for 2⁺ using parameters (kcal/mol) $\lambda_{\rm SE} = 40.3$, $V_{\rm SE}^{\rm eff} = -1.5$, $\lambda_{\rm BO} = 19.2$, $\Delta G^{\circ}_{\rm BO} =$ 6.5, $V'_{\rm BO} = 3.5$, for which $\Delta G^* = 4.2$ kcal/mol. Diabatic surfaces are shown as broken lines and adiabatic ones as solid lines.

accurate calculation of $k_{\rm et}$ from the $\lambda_{\rm SE}$ and $V_{\rm SE}$ obtained for $\mathbf{1}^+$, its analogue with two fewer methyl groups,6b and three bis-(hydrazines) with saturated bridges; therefore, the Hush $V_{\rm SE}$ equation^{2e} is a rather good approximation.⁷

This paper principally concerns 2^+ , which we expected to have a $k_{\rm et}$ value no larger than that of 1^+ because the $N_{\rm Ar}$ lone pair, aryl π system twist angle ϕ , and, hence, λ_{SE} should be nearly the same. The ϕ values of crystalline 1^+ average 50.5° at the neutral

hydrazine unit and 66.2° at the cationic hydrazine unit.6b X-ray data for neutral 3 ($\phi = 53.2^{\circ}$) and 3⁺ ($\phi = 66.7^{\circ}$) provide models for the twist of the hydrazine units of 2^+ (their structures are reported in the Supporting Information). V_{SE} depends on overlap at the bonds connecting the charge-bearing units to the bridge, so it depends on $\cos \phi$ and the orbital coefficients at the atoms involved. The larger π system of the bridge for 2^+ should lower its V_{SE} relative to that for 1^+ , so smaller k_{et} for 2^+ than for 1^+ might be expected. However, $k_{\rm et}$ of 2^+ is far larger than that of 1⁺. The ESR spectrum of 2⁺ is that of a rapidly exchanging hydrazine-centered species at all accessible temperatures in CH₃CN and CH₂Cl₂, but a dynamic alternating line width effect was observed in acetone, allowing determination of $k_{\rm et}$ at -105, -100, and -95 °C as 1.6, 2.1, and 2.9×10^8 s⁻¹, respectively. ESR data for 1^+ in CH₃CN give rate ratios for $2^+:1^+$ of 173, 148, and 135 at these temperatures, corresponding to a ΔG^{\dagger} decrease of 1.7 kcal/mol. The ET barrier ΔG^* for 2^+ in acetonitrile is estimated at 1.7 kcal/mol less than for 1^+ in acetonitrile. 6b or ~ 4.2 kcal/mol.

The principal effect of changing from the durene bridge of 1⁺ to the anthracene bridge of 2^+ is lowering the energy of bridge redox intermediates for 2^+ . The optical spectrum of 2^+ is complex, and we use the spectra of the monohydrazine analogues 3 and 3^+ to help understand those of 2 and 2^+ (these spectra are shown in the Supporting Information). We attribute the visible

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(3) (</sup>a) "The term superexchange in the ET literature is generally meant to imply the mixing of donor and acceptor states by a virtual electronic state of the bridging species. In particular, usage of the term generally implies that p 46). (b) Kosloff, R.; Ratner, M. A. *Isr. J. Chem.* **1990**, *30*, 45.

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absorption for the neutral compounds to bridge reduction charge transfer (CT_{BR}), $Hy^0-B^0 \rightarrow Hy^+-B^-$, where Hy is the hydrazine unit and B is the bridge. The CT_{BR} bands in methylene chloride have the following parameters: 3, $h\nu_{\rm max} = 21\ 500\ {\rm cm}^{-1}\ (\lambda_{\rm max} =$ 465 nm), $\epsilon_{\rm max} = 3800~{\rm M}^{-1}~{\rm cm}^{-1}$ (all ϵ values are reported using these units); 2, $h\nu_{\rm max} = 19\,600 \,{\rm cm}^{-1} \,(\lambda_{\rm max} = 510 \,{\rm nm}), \,\epsilon_{\rm max} =$ 6000.8 Local excitations of the anthracene ring are also observed. That corresponding to the Clar p band shows vibrational fine structure, and the 0,0 band occurs at 1200 cm⁻¹ lower energy for **2** than **3**. The Clar β bands are very broad: **3**, \sim 34 800 cm⁻¹ $(\epsilon_{\text{max}} = 7300); \ \mathbf{2}, \ \sim 34\ 300 \ (\epsilon_{\text{max}} = 11\ 000).$ Species $\mathbf{3}^+$ in acetonitrile shows two visible bands at 12 900 cm⁻¹ (755 nm) $\epsilon_{\rm max} = 450 \text{ and } 19\,000 \text{ cm}^{-1} \text{ (526 nm) } \epsilon_{\rm max} = 2000.$ We assign them as bridge oxidation bands, BO1 and BO2, respectively, corresponding to $Hy^+-B^0 \rightarrow Hy^0-B^+$ CT, from the highest occupied molecular orbital (HOMO) and either HOMO-1 of anthracene or a band of more complex origin, respectively. CT bands from orbitals below the HOMO have been reported for transition-metal-centered compounds.⁴ The BO₁ band of 2⁺ in acetonitrile has $h\nu_{\rm max,BO}=90\bar{0}0~{\rm cm}^{-1}$ (1110 nm), $\epsilon_{\rm max}=1400$, $\Delta v_{1/2} = 4300 \text{ cm}^{-1}$ (estimated from the low-energy side because of band overlap on the high-energy side) and has nearly the same parameters in acetone. Both BO bands are much lower in energy for 2⁺ than those for 3⁺, presumably because the anthracene ring of 2⁺ has an electron-releasing neutral hydrazine substituent. Species 2⁺ shows the optical bands expected for both oxidized and reduced hydrazine units, as required for a localized system. The CT_{SE} band expected for **2**⁺ near 14 100 cm⁻¹ is probably present but is not resolved from the BO₂ band.

The striking feature of the absorption spectrum of 2^+ is that a CT_{BO} band occurs at significantly lower energy than its Hushtype CT_{SE} band, indicating that the anthracene ring oxidized state, 2^+ (BO), cannot lie far above the hydrazine-centered 2^+ ground state. The BO₁ transition energy, $hv_{\text{max,BO}} = 25.7$ kcal/mol = $\lambda_{\text{BO}} + \Delta G^{\circ}_{\text{BO}}$ for 2^+ , making it unlikely that $\Delta G^{\circ}_{\text{BO}}$ is larger than about 6 kcal/mol.⁹

Does such a low-lying $2^+(BO)$ surface mean that it will be an intermediate for ET between the hydrazine units of 2^+ ? Extensive theoretical work on how coupling of charge-bearing units with the orbitals of the bridge affects $V_{\rm SE}$ has been discussed by Newton. The assumption of a small $V_{\rm SE}/\Delta G^\circ$ ratio is clearly not valid for the anthracene HOMO of 2^+ , so a different approach is required. We suggest that it is useful to consider 2^+ as a three-state system having Marcus—Hush parabolic diabatic surfaces corresponding to charge localized on each hydrazine unit and a third parabolic energy surface corresponding to $2^+(BO)$ centered between them (see Figure 1b). Whether $2^+(BO)$ is a minimum on the ground-state surface depends on the relative sizes of the matrix elements $V_{\rm BO}$ and $V_{\rm SE}^{\rm eff.}$ Hush analysis of the $CT_{\rm BO}$ band of 2^+ in acetonitrile gives $V_{\rm BO} = 4726/d_{\rm BO}$ cm $^{-1}$. We believe that $d_{\rm BO}$ should be larger than half the distance between

the nitrogens because charge in $2^+(BO)$ is delocalized over the 14-atom π system of the aryl ring; $V_{\rm BO}=4.5~{\rm kcal/mol}$ (using $d_{\rm BO}$ of 3.0 Å) and 3.9 kcal/mol (using 3.5 Å). V values evaluated from optical spectra which are smaller than Hush ones by a factor of $n_{\rm D}^{-\hat{1}/2} = 0.86$ for acetonitrile ($n_{\rm D}$ is the solvent refractive index) have been suggested by Young and co-workers,12 and these lower V values better fit the observed $k_{\rm et}$ values for five bis(hydrazine) radical cations.⁷ This would lower the $V_{\rm BO}$ estimated from the optical spectrum to 3.9 kcal/mol (using $d_{BO} = 3.0 \text{ Å}$) and 3.3 kcal/mol (using 3.5 Å). At $V_{\rm BO} = 3.5$ kcal/mol, the three-state model fits $\lambda_{\rm BO}$ + $\Delta G_{\rm BO}$ = 25.7 and ΔG^* = 4.2 kcal/mol for ΔG°_{BO} rising from 6.3 (at $V_{SE}^{eff} = 1.0$) to 7.3 (at $V_{SE}^{eff} = 3.0$) kcal/mol, and the ground-state surface is very flat-topped, but $4^{+}(BO)$ is not an intermediate (see Figure 1b). At $V_{BO} = 2.5$ kcal/mol, fit is obtained for $\Delta G^{\circ}_{BO} = 5.1$ kcal/mol (at $V_{SE}^{eff} =$ 1.0) and 5.8 (at $V_{\rm SE}^{\rm eff} = 3.0$) kcal/mol and there is a minimum on the ground-state surface at X = 0.5 that is less than RT kcal/ mol deep (0.13 kcal/mol at $V_{\rm SE}^{\rm eff} = 1.0$, falling to zero at $V_{\rm SE}^{\rm eff}$ = 1.0) (see Supporting Information for more details). It appears from this modeling that 2+ is near the borderline for which 2+-(BO) becomes a very shallow dip on the ground-state energy surface and that the barrier for ET between the hydrazine units is affected little by whether $2^+(BO)$ is an intermediate or not.

This approximate analysis 13 indicates how much information about ET reactions involving the bridge is present in the rich absorption spectrum of 2^+ . Its ground state has charge localized on one hydrazine unit, but the bridge-oxidized state $2^+(BO)$ only lies a few kcal/mol higher in energy. ΔG°_{BO} is small enough to significantly lower the ET barrier, although if $2^+(BO)$ is an intermediate on the ground-state energy surface, the energy gap to the transition state for ET is very small. The bridge-reduced state lies too high in energy to affect the adiabatic ET surface for 2^+ significantly.

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Supporting Information Available: Preparation of **2** and **3** and their radical cations, and crystallographic data for **3** and $3^+\text{Ph}_4\text{B}^-$ and their absorption spectra; plot of ΔG^o_{BO} vs V_{SE}^{eff} for three-state fits to 2^+ for V_{BO} 1.5, 2.5, 3.5, and 4.5 kcal/mol (15 pages). An X-ray crystallographic CIF format file is available on the Web only. See any current masthead page for ordering information and Web access instructions.

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(11) (a) The three-state secular determinate is shown in eq 1:

$$\begin{vmatrix} H_{AA} - E & V_{\text{SE}}^{\text{eff}} & V_{\text{BO}}' \\ V_{\text{SE}}^{\text{eff}} & H_{BB} - E & V_{\text{BO}}' \\ V_{\text{BO}}' & V_{\text{BO}}' & H_{CC} - E \end{vmatrix} = 0$$

 $H_{AA} = \lambda_{\rm SE} X^2, \ H_{BB} = \lambda_{\rm SE} (1-X)^2, \ {\rm and} \ \lambda_{\rm SE} = 40.3 \ {\rm kcal/mol} \ ({\rm that \ for} \ 1^+) \ {\rm are} \ {\rm employed}. \ H_{CC} = \lambda_{\rm BO} (1-2X)^2 + \Delta G^{\circ}_{\rm BO}, \ {\rm giving} \ H_{CC} = \lambda_{\rm BO} + \Delta G^{\circ}_{\rm BO} \ {\rm at} \ X = 0 \ {\rm and} \ 1, \ {\rm as} \ {\rm required} \ {\rm to} \ {\rm giv} \ h \nu_{\rm max,BO}. \ ({\rm b}) \ {\rm Hush} \ V \ {\rm values}^{2e} \ {\rm are} \ {\rm for} \ {\rm two\text{-}state} \ {\rm system} \ {\rm because} \ {\rm system} \ {\rm sean} \ {\rm dev} \ {\rm system} \ {\rm because} \ {\rm system} \ {\rm sean} \ {\rm dev} \ {\rm$

(13) It is not clear how realistic it is to represent BO and SE processes with a single X value, as we have done.

⁽⁸⁾ Because replacing a hydrogen by a hydrazine unit presumably makes the ring harder to reduce, we would expect ΔG°_{BR} for 2 to be larger than for 3. The transition energy $h\nu_{\max,BR} = \lambda_{BR} + \Delta G^{\circ}_{BR}$. It is not obvious to us why λ_{BR} would be smaller for 2 than for 3, which is what the experimental result appears to imply.

Agr. Hosti of the series in the series of the series and identical charge-bearing units, $\mathbf{1}^+$ ($hv_{\text{max,SE}}$) appears to imply. (9) (a) With similar ϕ values and identical charge-bearing units, $\mathbf{1}^+$ ($hv_{\text{max,SE}}$) = 14 100 cm⁻¹ = 40.1 kcal/mol in acetonitrile) should be a good model for λ_{SE} of $\mathbf{2}^+$, but λ_{BO} will be smaller than λ_{SE} because significantly smaller reorganization energy is involved. Estimating $\lambda_s \approx 13-15$ kcal/mol for $\mathbf{1}^+$ in acetonitrile, λ_v , involving two hydrazine units, is $\sim 25-27$ kcal/mol. $\lambda_{v,\text{BO}}$ for $\mathbf{2}^+$ should be the average of that for its hydrazine unit and the bridge. The estimate for λ_v for anthracene itself calculated by the published method⁹⁶ is 6.5 kcal/mol, resulting in a $\lambda_{v,\text{BO}}$ estimate for $\mathbf{2}^+$ of $\sim 16.3 \pm 0.5$ kcal/mol. We estimate $\Delta G^\circ_{\text{BO}}$ as $\sim (9.4 - \lambda_{s,\text{BO}})$ kcal/mol. $\lambda_{s,\text{BO}}$ will be smaller than that for the SE ET, but we have no way of estimating its size accurately. Because $hv_{\text{max,BO}}$ is 3.4 kcal/mol larger in acetonitrile than it is in methylene chloride, it seems likely that $\lambda_{s,\text{BO}}$ in acetonitrile is at least 3.4 kcal/mol although $\Delta G^\circ_{\text{BO}}$ could change too. Using this number, $\Delta G^\circ_{\text{BO}}$ is estimated at no more than 6 kcal/mol. However, substantial mixing between the hydrazine and the bridge might be argued to lower $\lambda_{v,\text{BO}}$ from the value assumed above. (b) Nelsen, S. F.; Blackstock, S. C.; Kim, Y. J. Am. Chem. Soc. 1987, 109, 677.

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