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Transferrable property relationships between magnetic exchange coupling and molecular conductance

Computed molecular conductances mediated by organic σ/π-bridges are compared to experimentally-determined magnetic exchange coupling parameters in a series of Donor-Bridge-Acceptor Biradical complexes at parity of the bridge fragment. For monomeric and dimeric bridges, biradical magnetic exchange coupling and computed conductance are related by a power law with an observed scale invariance. We also find that the ratio of distance dependent decay constants for bridge-mediated conductance and magnetic exchange coupling does not equal unity. The results of our observations have been cast in valence bond terms.

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Transferrable property relationships between magnetic exchange coupling and molecular conductance†

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Calculated conductance through $\text{Au}_n$–$\text{S}$–$\text{Bridge}$–$\text{S}$–$\text{Au}_n$ (Bridge = organic $\sigma/\pi$-system) constructs are compared to experimentally-determined magnetic exchange coupling parameters in a series of $\text{Tp}^{\text{Cum,Me}}$Zn$\text{SQ}$–$\text{Bridge}$–$\text{NN}$ complexes, where $\text{Tp}^{\text{Cum,Me}} = \text{hydro-tris(3-cumenyl-1-methylpyrazolyl)}$ borate ancillary ligand, Zn = diamagnetic zinc(II), SQ = semiquinone ($S = 1/2$), and NN = nitronyl-nitroxide radical ($S = 1/2$). We find that there is a nonlinear functional relationship between the biradical magnetic exchange coupling, $J_{\text{D} \rightarrow \text{A}}$, and the computed conductance, $g_{\text{mb}}$. Although different bridge types (monomer vs. dimer) do not lie on the same $J_{\text{D} \rightarrow \text{A}}$ vs. $g_{\text{mb}}$ curve, there is a scale invariance between the monomeric and dimeric bridges which shows that the two data sets are related by a proportionate scaling of $J_{\text{D} \rightarrow \text{A}}$. For exchange and conductance mediated by a given bridge fragment, we find that the ratio of distance dependent decay constants for conductance ($\beta_g$) and magnetic exchange coupling ($\beta_J$) does not equal unity, indicating that inherent differences in the tunneling energy gaps, $\Delta\epsilon$, and the bridge–bridge electronic coupling, $H_{\text{EB}}$, are not directly transferrable properties as they relate to exchange and conductance. The results of these observations are described in valence bond terms, with resonance structure contributions to the ground state bridge wavefunction being different for $\text{SQ}$–$\text{Bridge}$–$\text{NN}$ and $\text{Au}_n$–$\text{S}$–$\text{Bridge}$–$\text{S}$–$\text{Au}_n$ systems.

Introduction

Electron transport in single-molecule devices is typically interrogated via a combination of experimental and computational probes of conductance using Metal–Bridge–Metal (M–B–M) junctions [Fig. 1A].1–4 To account for inherent variations in molecule–electrode binding geometries, the experimental determination of conductance ($g_{\text{mb}}$) typically requires hundreds to thousands of individual conductance measurements in order to construct conductance histograms, which allow researchers to determine both conductance distributions and the most probable conductance value.5 Since conductance calculations often utilize a single M–B–M geometry and the molecular geometry in the actual junction is unknown, this can make direct comparisons between theory and experiment difficult to interpret. As a result, studies that correlate single-molecule conductance with physical observables such as electron transfer rate constants ($k_{\text{D} \rightarrow \text{A}}$)6,7,8 and magnetic exchange couplings

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**Fig. 1** (A) Metal–Bridge–Metal junctions corresponding to biradical complexes (B). (C) Bridges common to M–B–M junctions and Donor–Bridge–Acceptor complexes (B), along with ferromagnetic exchange parameters, $J_{\text{D} \rightarrow \text{A}}$ for biradical complexes (C). $\text{BCO} = \text{bicyclo}[2.2.2]$ octane. $\text{Ph} = \text{para-phenylene}$, $\text{Th} = 2.5$-thiophene, $\text{Py} = \text{pyridine}$, $\text{Py-Th} = 2$-pyridyl-5′-thiophene, $\text{Ph}_2 = \text{para-biphenylene}$, and $\text{Th}_2 = 2.5′$-bithiophene.
\(J_{D \rightarrow A}\)^n continues to be a topic of current research. A primary impact of these efforts will be to provide key insight into orbital pathways for charge transport through molecules, which is difficult to achieve using traditional approaches. These correlations, coupled with the determination of transferable property relationships, are expected to facilitate improved molecular design concepts for molecular electronics and spintronics.

The crux of these transferable property relationships was first described by Nitzan, who derived a relationship between \(g_{mb}\) and \(k_{D \rightarrow A}\) at parity of the molecular bridge. With some approximations, Nitzan was able to show that conductance could be related to the square of the D–A electronic coupling matrix element, \(H_{D \rightarrow A}^2\), for electron transfer (eqn 1–4).

\[
g_{mb} \approx \frac{8e^2}{\pi^2 \Gamma_{D} |\Gamma_{A}|^2} FCWDS k_{D \rightarrow A} \tag{1}
\]

\[
g_{mb} = \frac{e^2}{\pi \hbar} \Gamma_{LB} \Gamma_{RB} [g_B]^2 \tag{2}
\]

\[
k_{D \rightarrow A} = \frac{2\pi n |H_{DB}H_{BA}| |g_B(E)|^2} {\hbar} FCWDS \tag{3}
\]

\[
G_B = \frac{H_{BB} n^{-1}} {\Delta \varepsilon} \tag{4}
\]

Here, the \(H_{ij}\) are pairwise electronic coupling matrix elements that connect D, B, and A, the \(\Gamma_i\) are broadening functions that represent the coupling between the molecule and electrodes, \(\Delta \varepsilon\) is the tunneling energy gap, \(G_B\) is the bridge Green’s function with the overbar indicating that \(G_B\) may differ for electron transfer (or magnetic exchange) compared to conductance, and \(n\) is the number of bridge units. These transferable property relationships indicate that any physical observable that is a function of \(H_{D \rightarrow A}^2\) can be related to \(g_{mb}\) (e.g. eqn 5).

\[J_{D \rightarrow A} \propto H_{D \rightarrow A}^2 \propto k_{D \rightarrow A} \propto g_{mb}\tag{5}\]

This leads to a corresponding relationship between these parameters and the distance decay constants \(\beta_i\) for electron transfer, conductance, and magnetic exchange as indicated in eqn (6),

\[k_{D \rightarrow A} \propto g_{mb} \propto J_{D \rightarrow A} \propto \exp(-\beta_i \times nL)\tag{6}\]

where \(nL\) is the distance spanned by the molecular bridge (e.g. \(R_{DA} - R_0\)).

There is both experimental and computational support for a correlation between \(k_{D \rightarrow A}\), \(g_{mb}\), and \(J_{D \rightarrow A}\). Beratan and Waldeck have built on Nitzan’s work to show that the relationship between \(k_{D \rightarrow A}\) and \(g_{mb}\) is nonlinear for alkane and peptide nucleic acid oligomer bridges, but they are related by a power law dependence. Their work supports an argument that the bridge Green’s function, \(G_B\), is not the same for electron transfer and conductance. As a result, Beratan and Waldeck were able to show that the distance decay constants for \(k_{D \rightarrow A}\) and \(g_{mb}\) are, in general, inequivalent and a linear relationship between \(k_{D \rightarrow A}\) and \(g_{mb}\) will only occur if the individual \(\beta_i\) are identical.

Since the magnetic exchange interaction \(J_{D \rightarrow A}\) is proportional to \(H_{D \rightarrow A}^2\) via eqn (5), we have initiated experiments designed to explore the correlation of \(J_{D \rightarrow A}\) between donor–acceptor biradical centers that are spanned by molecular bridge units (D–B–A biradicals = SQ–Bridge–NN; SQ = S = 1/2 semiquinone, and NN = S = 1/2 nitronylnitroxide radical, Fig. 1B and C). With computed molecular conductance values, \(g_{mb}\). An advantage to understanding the magnitude of \(g_{mb}\) by correlating with experimental \(J_{D \rightarrow A}\) parameters is highlighted by the fact that the variable-temperature magnetic susceptibility measurements used to determine \(J_{D \rightarrow A}\) are performed on solid-state samples with known geometries that have been determined by X-ray crystallography. Thus, there is a single conformation present that contributes to \(J_{D \rightarrow A}\). Unlike experimental conductance measurements in a contact geometry and solution electron transfer rate measurements, the magnitude of \(J_{D \rightarrow A}\) determined by solid state magnetic susceptibility measurements is not affected by structural and solvent bath induced inhomogeneities or by decoherence effects.

Relating molecular orbital pathways for \(J_{D \rightarrow A}\) to orbital transport channels for \(g_{mb}\) has its own set of challenges, and these include (1) differences in the molecular fragments covalently attached to the bridge unit, (2) the molecular identity of surfactant groups, (3) variations in surface contacts, (4) the choice of electrode material and surface morphology, and (5) inhomogeneities in molecular structure (e.g., conformation) and in contact geometries. As alluded to above, an important related question was recently posited by Herrmann, which concerns the transferability of geometric and electronic structure contributions of the bridge moiety to \(k_{D \rightarrow A}\), \(g_{mb}\), and \(J_{D \rightarrow A}\) (Fig. 2). Herein, we address these difficulties and the concept of bridge electronic structure transferability between conductance and exchange. We then use our results to explicitly determine distance decay constants \(\beta_i\) for \(g_{mb}\) from prior studies that have determined the distance dependence of \(J_{D \rightarrow A}\). Our results are described in valence bond terms, with resonance structure contributions to the ground state bridge wavefunction being different for SQ–Bridge–NN and Au–S–Bridge–S–Au systems.

Fig. 2. Illustration of how bridge-mediated electronic coupling is related to conductance and magnetic exchange coupling. Top: Molecular bridge (B) connects metallic electrodes to give a Metal–Bridge–Metal device. Bottom: Molecular bridge (B) connects spin centers to facilitate magnetic exchange coupling.
This provides a path forward for understanding the power law relationship between $g_{mb}$ and $J_{D→A}$ in the absence of structural inhomogeneities and decoherence effects.

**Experimental**

**General**

The synthesis and characterization of SQ–BCO–NN, SQ–Ph–NN, SQ–Th–NN, SQ–Ph2–NN, SQ–Py–Th–NN, and SQ–Th2–NN have been published previously.1,17,26,35,39,41,42 See the ESI† for the preparation and characterization (spectroscopic data, X-ray crystal structure, and magnetic characterization) of the pyridyl-bridged biradical complex, SQ–Py–NN.

**Resonance Raman spectroscopy**

Solution (methylene chloride) resonance Raman spectra were collected in either a 180° backscattering (780 nm) or 90° scattering geometry using 407 nm excitation from a Coherent Innova 70 Kr+ (1 W) ion laser. The scattered radiation was passed through a longpass filter (Semrock RazorEdge) to remove Rayleigh scattered laser light and then dispersed onto a liquid nitrogen-cooled Infrared Associates CCD detector using a Princeton Acton spectrograph. The laser power at the sample was typically kept between 40 and 100 mW in order to prevent possible photo- and thermal degradation of the sample. The sample was sealed in a glass capillary tube and spun with a custom-made sample holder. All data were scan-averaged, and any individual data set with vibrational bands compromised by cosmic events were discarded.

**Molecular electronic structure calculations**

Thiophene and benzene calculations were performed at the density functional level of theory using the Gaussian 09W software package.44 Input files were prepared using the molecular builder function in GaussView. Calculations employed the B3LYP hybrid exchange–correlation functional and a 6-31g(d,p) split valence basis set with polarizability functions were used for all atoms. Frontier molecular orbitals (MOs) were generated from fully optimized ground states.

**Transport calculations**

Transport calculations were performed using ATK 2016.0 v16.0.44–46 as detailed in our earlier work.1 The device configuration consists of the left Au electrode, the molecular bridge (scattering region), and the right Au electrode. Both the left and right gold electrodes consist of nine layers with each layer comprised of a 3 × 3 array of Au(111) atoms, for a total of 81 gold atoms per electrode. For the electrodes, the C or z direction is periodic in the system. This is also the direction of electron transport. Initial geometry optimizations of the bridge molecules BCO, Py, Ph, Th, Ph2, Py–Th, Th2 were performed using Gaussian 09 (ref. 43) with a 6-31g(d,p) basis set and the B3LYP functional. Electrode surfaces were constructed by cleaving the bulk crystal and using a copy of this to form the second electrode. The molecule was then placed on the surface of the left electrode and subsequently connected to the right electrode through terminal sulfur atoms to create the final device geometry. Prior to creating the final device geometry, the bulk configuration was optimized using a single-zeta basis set for the gold atoms (to save computational time) and a double zeta basis set for all other atoms. The optimization was performed using ATK-DFT with a Perdew–Zunger local density approximation (LDA-PZ) exchange–correlation. After this bulk optimization, the configuration was converted into a final device geometry for all subsequent calculations. A $5 \times 5 \times 51$ K-point sampling has been used in x, y, z-directions, respectively. The boundary conditions were Dirichlet (fixed boundary condition) in the z direction, which is the direction of transport. The boundary conditions in the x and y directions were disabled so they could not be changed but the default parameterization is periodic for the x and y directions. The NEGF formalism was used to calculate the non-equilibrium electron density of the central region of the device.

The Landauer–Büttiker formula77 relates transmission probabilities to conductance, $g_{mb}$ ($g_{mb} = I/V$). This formula is used to calculate the voltage dependent current, $I(V)$, across a molecular junction, which is determined by integrating the transmission function, $T(E,V)$, according to eqn (7).

$$I(V) = \frac{e}{h} \sum_{\sigma} \int \frac{f(E - \mu_q)}{k_B T_R} \left( \frac{f(E - \mu_q)}{k_B T_R} - \frac{f(E - \mu_i)}{k_B T_L} \right) dE \quad (7)$$

here, the Fermi–Dirac distribution functions for the right and left electrodes are given by $f(E - \mu_q)/k_B T_R$ and $f(E - \mu_i)/k_B T_L$, with $\mu_q$ and $\mu_i$ being the chemical potentials for the right and left electrodes, respectively, $e$ is the charge of the electron, $h$ is Planck’s constant, $eV_{bias}$ is the bias window equivalent to $\mu_q - \mu_i$, (where $V_{bias} = V_L - V_R$), $T_{\sigma}(E)$ is the transmission coefficient for the spin component, $\sigma$, describing the junction at an energy $E$ and a bias voltage $V_{bias}$. All electron transport properties were computed using the ATK software package that includes virtual nanolab associated analysis modules. The molecular projected self-consistent Hamiltonian (MPSH) technique has been used to understand the molecular orbital origin of the resonant peaks in the transmission spectra.48,49

**Results and discussion**

**Exchange coupling in SQ–Bridge–NN complexes**

Magnetic exchange coupling constants ($J_{D→A}$) for the monomeric SQ–Bridge–NN biradical complexes (Fig. 1B and C) SQ–BCO–NN, SQ–Ph–NN, SQ–Th–NN,17,26,44 and for the dimeric bridge molecules SQ–Py–Th–NN,5 SQ–Ph2–NN,37 and SQ–Th2–NN7 have previously been determined by magnetic susceptibility measurements. The dimer exchange spin Hamiltonian (eqn (8)):

$$\mathcal{H}_{ex} = -2J_{D→A}S_{SQ}S_{NN} \quad (8)$$

has been used in the analysis of these data, including the analysis of new magnetic susceptibility data for the pyridyl-bridged complex (SQ–Py–NN), which yields $J_{D→A} = +72$ cm$^{-1}$ (see ESI†). Collectively, these SQ–Bridge–NN biradical systems possess a large range of $J_{D→A}$ values, bridge-dependent electronic structures, and both $\sigma$- and $\pi$-coupling units. Notably, all
of these molecules are ferromagnetically exchange coupled with $J_{D \rightarrow A}$ values that vary from a low of $J_{D \rightarrow A} = +1 \text{ cm}^{-1}$ for $\sigma$-mediated exchange in SQ–BCO–NN, to a markedly larger $J_{D \rightarrow A} = +220 \text{ cm}^{-1}$ for $\pi$-mediated exchange in SQ–Th–NN.

Exchange coupling within the context of a valence bond configuration interaction model

Given the geometric structures of these SQ–Bridge–NN biradicals, we can conveniently divide them into two distinct bridge categories: the first possesses single-ring molecular bridges, while the second contains two-ring molecular bridges where each bridge ring is connected to the other via a $\sigma$-bond. The electronic origins of $\pi$-mediated exchange coupling in SQ–Ph–NN and $\sigma$-mediated exchange in SQ–BCO–NN have been rigorously evaluated using a combination of spectroscopy, magnetic susceptibility, and theory.19,20,26,27 For $\pi$-mediated exchange, a valence-bond configuration interaction (VBCI) model has been used.19,20,26 Within the context of more traditional orbital pathways for understanding exchange coupling, the VBCI method conveniently illustrates how specific excited state configurations admix with the ground configuration to affect the magnitude of $J_{D \rightarrow A}$. The NN(SOMO), SQ(SOMO), NN–Bridge(HOMO), and the NN–Bridge(LUMO) (Fig. 3) orbitals form a convenient minimal active space to generate the ground and excited state configurations that contribute to $J_{D \rightarrow A}$ for conjugated SQ–Bridge–NN systems. Previous work has demonstrated the crucial role of the LUMO(B–NN) in determining the magnitude of $J_{D \rightarrow A}$.17,19,20,26,27 Resonance structures that illustrate both the spin and the charge distributions of ground state SQ–Ph$_2$–NN configurations and excited state configurations that arise from one-electron promotions within this active space are shown in Fig. 4. Here, the ground SQ–Bridge–NN configuration is represented by GC, EC1 and EC2 represent the configuration that results from an intraligand SQ(SOMO) $\rightarrow$ NN–Bridge(LUMO) electron promotion, and the NN–Bridge(HOMO) $\rightarrow$ SQ(SOMO) one-electron promotion is represented by the EC3 and EC4 resonance structures. Similar resonance structures can be drawn for other SQ–Bridge–NN biradical complexes. For the conjugated SQ–Bridge–NN systems discussed here, the charge transfer configuration generated by the intraligand SQ(SOMO) $\rightarrow$ NN–Bridge(LUMO) one-electron promotion (EC1 and EC2) has been shown to be the dominant contributor to the ground state ferromagnetic exchange coupling in SQ–Ph–NN, with the NN–Bridge(HOMO) $\rightarrow$ SQ(SOMO) (EC3 and EC4) playing a less important role.17,19,20,26,27 The relative contributions of these two intraligand charge transfer (ILCT) configurations will be dependent on the nature of the bridge. The use of more electron donating bridge molecules (e.g., thiophene) is expected to increase the relative NN–Bridge(HOMO) $\rightarrow$ SQ(SOMO) charge transfer contribution to the magnetic exchange by raising the energy of the bridge HOMO and LUMO orbitals. In addition, we have shown that bond torsions which disrupt Ph–NN $\pi$-coupling result in the ILCT transition being shifted to higher energy, with a concomitant reduction in the ILCT intensity relative to configurations that possess more planar Ph–NN conformations.41 As a result, an ILCT transition is not observed in SQ–BCO–NN and the lack of a CT feature in the visible region of the electronic absorption spectrum correlates with the weak $\sigma$-mediated exchange coupling.26,41 Our understanding of how the bridge HOMO and LUMO orbitals affect the magnitude of both $J_{D \rightarrow A}$ and $H_{D \rightarrow A}$ allow us to begin an assessment of how
these orbitals contribute to the transferability of $J_{D\rightarrow A}$, $k_{D\rightarrow A}$, and $H_{D\rightarrow A}$ to conductance in a biased electron transport configuration (Fig. 2).

**Correlation of exchange and conductance**

Electron transport calculations have been performed on M–B–M constructs (Au–S–Bridge–S–Au) in order to compare the magnitude of bridge-mediated $J_{D\rightarrow A}$ with $g_{mb}$ (i.e., current). In the transport computations, the bridge molecules in Fig. 1C are connected to voltage-biased Au electrodes via thiol anchoring groups. Thus, the bridges are the molecular fragments depicted in Fig. 1C with sulfur atoms attached to the bridge carbons. These calculations treat the M–B–M ensemble at the SCF level with the computational results being interpreted in terms of the Landauer–Büttiker formula, which relates transmission probabilities to conductance, $g_{mb}$, according to eqn (7). In marked contrast to the dominant role of the bridge LUMO in SQ–Bridge–NN biradical magnetic superexchange coupling, the dominant orbital contributions to our computed conductance values are the bridge HOMOs. This is clearly illustrated in Fig. 5, where the bias-dependent transmission coefficient for M–Th–M (note that Th is a dithiol in the M–B–M construct) is largest for occupied energy levels that are comprised of thiophene bridge HOMOs. This derives from the bridge HOMOs being effectively pinned to the electrode Fermi level. The molecular HOMO character is clearly present in the molecular projected self-consistent Hamiltonian (MPSH) state contributions to the transmission at energies just below the Fermi level for M–Th–M and M–Ph–M (Fig. 5 and S3†). Closer inspection of the MPSH HOMO reveals the presence of anchor sulfur p-orbital contributions, which represent a key difference in the nature of the orbitals that promote M–B–M conductance vs. SQ–Bridge–NN magnetic exchange.

Following the work of Beratan, Yoshizawa and coworkers have conveniently related the zeroth order Green’s function for the bridge $(\tilde{G}_B^{0}(E_F) - \tilde{G}_B)$ to the MO coefficients of the bridge orbitals, the Fermi energy, and the individual MO energy levels according to:

$$G_B^0(E_F) = \sum_C \frac{C_{iC}^* C_{iC}}{E_F - \epsilon_i \pm i\eta}$$

where $C_{iC}$ and $C_{iC}^*$ correspond to the MO coefficients for the sulfur atoms located on the left- and right-hand side of the bridge molecule that directly connects to the electrodes. We note that these sulfur atoms are connected to the same carbon atoms of the organic bridge as the SQ-donor and NN-acceptor moieties in our SQ–Bridge–NN biradical systems (Fig. 5B, right). The sulfur orbital contributions to the MPSH states are important in modulating the conductance, since they directly connect the electrodes to the molecule and allow for delocalization of the bridge wavefunction in the scattering region.

When considering only the HOMO contributions to the conductance, eqn (9) reduces to:

$$G_B^0(E_F) = \frac{C_{\text{HOMO}}^* C_{\text{HOMO}}}{E_F - \epsilon_{\text{HOMO}} \pm i\eta}$$

Thus, our *a priori* expectation is a difference in bridge-mediated coupling for conductance and magnetic exchange that results from the importance of sulfur atom character in the M–B–M MPSH HOMOs and the importance of the bridge LUMO that facilitates magnetic exchange in SQ–Bridge–NN biradicals. The computed conductance as a function of the experimentally determined SQ–Bridge–NN magnetic exchange coupling constants using monomeric and dimeric bridge molecules is presented in Fig. 6. We have fit these data to the power law expression:

$$g_{mb} = a(c^\gamma J_{D\rightarrow A})^\gamma$$

where $J_{D\rightarrow A}$ is the exchange parameter for SQ–NN with no bridge ($J_{D\rightarrow A} = 550$ cm$^{-1}$). The best fit to the data shown in Fig. 6 yields $a = 2012$ and $\gamma = 0.675$. One immediately notices two important aspects of this $J_{D\rightarrow A}$ vs. $g_{mb}$ correlation. The first is that although there is a functional relationship between $J_{D\rightarrow A}$ and $g_{mb}$, it is not a linear relationship (i.e., $\gamma \neq 1$). The second observation is that the data for different bridge types (monomer vs. dimer) do not lie on the same curve. However, the fits to the data do indicate that there is a scale invariance that relates $J_{D\rightarrow A}$ and $g_{mb}$, for monomeric and dimeric bridges ($c = 1$ and $c = 0.28$ for monomeric and dimeric bridges, respectively), showing that

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Fig. 5  (A) Illustration of the HOMO conductance mechanism. The biased electrodes (blue) are filled to the chemical potential of the electrodes and the variable bias window $|\Delta E_{\text{cap}} = V_{\text{bias}}|$ is highlighted in yellow. Only the HOMO is dominant in the bias window and therefore it maximally contributes to $g_{mb}$. The LUMO is at higher energy and lies outside the bias window. As such, it does not contribute to $g_{mb}$. (B) Left: Computed zero-bias energy vs. transmission plot for S–Th–S connected to Au electrodes. A 2 V bias window is depicted in yellow and the Fermi energy is shown as a dashed line. Inset: Molecular projected self-consistent Hamiltonian (MPSH) state that dominantly contributes to $g_{mb}$, which is primarily comprised of the Th HOMO. Right: HOMO and LUMO of Th and Ph. Bridge carbon atoms of contact are indicated by blue arrows.
in order to understand the relative ratios of distance dependent decay constants for conductance ($\beta_d$) and magnetic exchange coupling ($\beta_j$). The best fits of eqn (12) to our $g_{mb}$ vs. $J_{D-A}$ data yield $\beta_d/\beta_j = 1.15$ for the phenylene series and $\beta_d/\beta_j = 1.58$ for the thiophene series (Fig. 7). Beratan and Waldeck also observed a nonlinear relationship between $g_{mb}$ and heterogeneous electron transfer rate constants, $k_{ET}$.\textsuperscript{4,5} However, in their study they showed that $\beta_d/\beta_j < 1$ for a series of alkanethiol and peptide nucleic acid oligomer bridges, with electron transfer decaying more rapidly with increasing bridge distance than conductance. Here, we observe the opposite correlation, with conductance falling off more rapidly with distance than the magnetic exchange coupling ($\beta_d/\beta_j > 1$), albeit using very different molecular bridges.

The relationship between $g_{mb}$ and $J_{D-A}$ for thiophenes is markedly more nonlinear than what we observe in the phenylene data. Eqn (13) describes how differences in these exponential decay constants ($\beta_i$) directly relate to inherent differences in the tunneling energy gaps, $\Delta_e$, the bridge–bridge electronic coupling, $H_{BB}$, and $L$, the length of the bridge unit.\textsuperscript{56,57} The tunneling energy gap ($\Delta_e$) is defined as the energy required to promote an electron from the donor (anode) to the bridge LUMO, or to promote a hole from the acceptor (cathode) to the bridge HOMO, and the distance between the donor and acceptor (or the electrodes) is given by $R_{DA} = R_0 + nL$. Thus, a combination of large $H_{BB}$ and a small $\Delta_e$ lead to low $\beta$-values and more shallow distance decays that promote long range magnetic superexchange coupling and conductance. Eqn (14) relates these distance decay constants to effective barrier heights, $\Delta E_{eff}$, with $m_e$ being the mass of the tunneling electron.\textsuperscript{56,57}

$$\Delta E_{eff} = \left(\frac{\hbar}{2m_e}\right) \beta^2 = \left(0.952 \text{ eV } A^{-2}\right) \beta^2$$

Beratan and Waldeck elegantly attributed their observed deviations in $\beta_d/\beta_j$ from unity to a combination of charge transfer energy barrier differences between electron transfer in D–B–A ensembles and electron transport in M–B–M configurations, bath-induced decoherences on the bridge molecule, and bridge–bridge electronic coupling ($H_{BB}$).\textsuperscript{4,5} Their work comparing conductance values and electron transfer rate constants indicated that differences in effective barrier heights derive from differences between the electrode work function in a transport geometry and donor/acceptor redox potentials in D–B–A systems.\textsuperscript{4,5} With respect to $\beta$ values, differences in the work function and redox potentials derive from different values for $\Delta_e$. In general, due to the interdependence of $\Delta_e$ and $H_{BB}$ on $\beta$, values it is difficult to separate these individual contributions to $\Delta E_{eff}$.\textsuperscript{59}

Since decoherence effects are not important for the $J_{D-A}$ vs. $g_{mb}$ correlation presented here, the dominant contributors to the nonlinearity between $J_{D-A}$ and $g_{mb}$ must relate to differences in bridge–bridge electronic coupling ($H_{BB}$) and the tunneling energy.
gaps (ΔE). For our SQ–Bridge–NN biradical complexes, we have used the ILCT energy as an approximation to the tunneling energy gap. This ILCT transition energy increases going from SQ–Ph–NN to SQ–Ph2–NN and decreases when going from SQ–Th–NN to SQ–Th2–NN.37 This contributes to the observed differences in βi between SQ–(Ph)1–NN and SQ–(Th)1–NN, but also contributes to the magnitude of βR/βi and differences in ΔEeff for magnetic exchange coupling and conductance.

Previously, we used magnetic susceptibility measurements to determine ΔEeff for superexchange-mediated magnetic coupling in SQ–Bridge–NN biradical systems, and this yielded ΔEeff = 1173 cm⁻¹ and 373 cm⁻¹ for phenylene and thiophene bridges, respectively.37 Given the conductance to magnetic exchange distance decay ratios (βR/βi) for phenylene and thiophenes determined here, we compute conductance barrier heights (ΔEeff) of 1555 cm⁻¹ and 941 cm⁻¹ for phenylene and thiophene bridges that span electrodes in a transport geometry. This translates to a decrease in ΔEeff for magnetic exchange coupling of ~25% (phenyles) and ~60% (thiophenes) when compared to conductance values using these same organic bridge fragments. Our βi values determined from magnetic susceptibility experiments (βi = 0.39 Å⁻¹ (Ph)b; βi = 0.22 Å⁻¹ (Th)b)17 and the magnitude of βR/βi determined in this work reveal the corresponding βR values for conductance using these molecular bridges (Table 1). The βR = 0.45 Å⁻¹ value we determine for para-phenylene bridges is very close to the βR = 0.42 Å⁻¹ determined by conducting probe atomic force microscopy on oligo(phenylene)-monothiols suspended between metal contacts.29 Additionally, the βR = 0.35 Å⁻¹ value that we have determined for oligothiophene bridges is also in good agreement with conductance studies performed on thiophene bridges with repeat units (n) equal to 1, 2, 3 and 5 (βR = 0.29 Å⁻¹).40

**Table 1** Comparison of decay constants, barrier heights, tunneling energy gaps, and bridge–bridge coupling ratios

| Compound/device | Parameter | Value |
|-----------------|-----------|-------|
| M–Ph–Mb         | βR        | 0.45 Å⁻¹ |
|                 | ΔEeff     | 1555 cm⁻¹ |
|                 | ΔR/ℏbb,kg| 2.63 (~ΔR/ℏbb,kg) |
|                 | βi        | 0.39 Å⁻¹ |
| SQ–Ph–NNb,17    | βR        | 0.35 Å⁻¹ |
|                 | ΔEeff     | 1168 cm⁻¹ |
|                 | ΔR/ℏbb,j | 2.31 |
| M–Th–Mb         | βR        | 0.22 Å⁻¹ |
|                 | ΔEeff     | 941 cm⁻¹ |
|                 | ΔE/ℏbb,kg| 1.79 (~ΔE/ℏbb,kg) |
| SQ–Th–Nnb,17    | βR        | 1.53 |
|                 | ΔEeff     | 372 cm⁻¹ |

* Data calculated using distances Rb from X-ray crystal structures of SQ–Bridge–NN with one single bond included = 4.30 Å (Ph) and 3.87 Å (Th)b n = 1 and 2. * Consistent with Hbb,j > Hbb,kg and/or ΔR > ΔE/j.

Resonance contribution to bridge mediated conductance and exchange

Since βi and ΔEeff are also a function of Hbb, the bridge–bridge electronic coupling, differences in the nature of the frontier bridge MOs that promote both gnb and JD→A will also affect the magnitude of the distance decay constants. It is generally assumed that the bridge–bridge electronic couplings encountered in electron transfer, conductance, and magnetic exchange are equivalent, and the donor, acceptor, and electrode contacts do not affect the magnitude of Hbb. However, this assumption may not be true. A convenient vehicle for understanding electronic structure contributions to βi values is the use of contributing resonance structures to highlight excited state configurations that mix with the ground state configuration within a valence bond framework.41,61–64

As introduced above, the resonance structures shown in Fig. 4 describe the charge and spin distributions of the ground state configuration (GC) and four excited state configurations (ECs) for the SQ–Ph2–NN biradical complex. Quinoidal EC1 and EC3 possess enhanced Ph–Ph π-coupling, which provides a mechanism for increasing the magnitude of Hbb. Contributions of resonance structures EC1–EC4 to the ground state all lead to enhanced magnetic exchange coupling in SQ–Bridge–NN biradicals. This occurs via a combination of B–B, D–B, and B–A electronic couplings (Hbb, Hdb, Hba), which are different than those provided by the coupling of these same bridge molecules with the electrodes ([IΓb], Γlb, Γrb). This is partly due to the fact that only high energy zwitterionic and biradical resonance structures can be drawn for the S–Ph–Ph–S electrode linkage. In contrast, these CT configurations lead to a low-energy ILCT state at ~24 000 cm⁻¹ in SQ–Bridge–NN biradicals, which has been shown to admix with the electronic ground state by resonance Raman spectroscopy.28,41 Specifically, the effect of this enhanced electronic coupling has been observed in SQ–Bridge–NN biradical compounds by optical pumping into the excited state ILCT and observing a large resonance enhancement of the phenylene quinoidal stretching

**Fig. 8** Top: Resonance Raman spectrum collected on resonance with the SQ → Ph–NN ILCT band showing large resonance enhancement of the 1600 cm⁻¹ quinoidal stretch (inset). Bottom: Computed electron density difference map (EDDM) that highlights the nature of the SQ → Ph–NN ILCT. Electron density loss in the transition is shown in red, while the electron density gain is shown in blue.
vibration, which decreases in intensity with increased SQ–Bridge and Bridge–NN bond torsions.\(^5\) For di-bridged SQ–Ph\(_2\)–NN, a large resonance enhancement of the 1600 cm\(^{-1}\) quinoidal stretch is also observed when pumping into the SQ → Ph–NN ILCT band\(^1\) (Fig. 8). The nature of the charge transfer is indicated by the computed electron density difference map (EDDM) in Fig. 8, which shows net charge transfer from the SQ donor (red) to the Ph–NN acceptor fragment (blue). Thus, the nature of the charge transfer in di-bridged SQ–Ph\(_2\)–NN is analogous to what has been observed previously in the mono-bridged SQ–Ph–NN biradical,\(^{20,39}\) highlighting quinoidal resonance structure contributions to the ground state wavefunction.

Conclusions

This work provides a detailed account of the transferability of electronic coupling with regard to exchange coupling and conductance. We have developed an electronic structure description of the functional relationship between \(J_{FD\to A}\) and \(g_{mb}\), which we observed to be nonlinear (\(\beta_g/\beta_f \neq 1\)). Plots of \(J_{FD\to A}\) vs. \(g_{mb}\) for monomeric and dimeric bridges lie on different curves due to a scale invariance between these bridge types, which indicates a proportionate scaling of the exchange interaction, \(J_{FD\to A}\). Our \(g_{mb}\) vs. \(J_{FD\to A}\) data yield \(\beta_g/\beta_f = 1.15\) for the phenylene series and \(\beta_g/\beta_f = 1.58\) for the thiophene series, and these ratios accurately replicate recently reported \(\beta\) values for electron transport through oligothiophene and oligophenylene bridges.\(^{29,46}\) In addition, these \(\beta_g/\beta_f\) ratios have been used to determine conductance tunneling barrier heights for oligophenlenes (\(\Delta E_{\text{eff}} = 1555\) cm\(^{-1}\)) and oligo-thiophenes (\(\Delta E_{\text{eff}} = 941\) cm\(^{-1}\)) that span electrodes in a transport geometry. This translates to a decrease in effective barrier heights \(\Delta E_{\text{eff}}\) for magnetic exchange coupling of \(~25\%\) for phenyles and \(~60\%) for thiophenes when compared to conductance values using these same organic bridge fragments.

Tunneling gaps and bridge–bridge electronic coupling, \(H_{mb}\), calculated from experimental data for the phenylene- and thiophene series provide a convincing explanation for the more pronounced \(g_{mb}\) vs. \(J_{FD\to A}\) nonlinearity observed in the thiophene series compared to the phenylene series. Additionally, resonance Raman spectroscopy indicates that \(H_{mb}\) may be enhanced by quinoidal resonance structure contributions in SQ–Bridge–NN biradicals relative to M–B–M devices for conductance. Namely, contributions from ECN resonance structures (Fig. 4) are expected to be more pronounced for SQ–Th\(_n\)–NN than for SQ–Ph\(_n\)–NN or Au\(_n\)–S–Bridge–S–Au\(_n\) constructs.

Resonance enhancement of the bridge quinoidal stretching mode illustrates the key bridge distortion that is the hallmark of a large \(H_{mb}\), and the magnitude of this excited state distortion is proportional to the degree of SQ → B–NN charge transfer, which contributes to the magnitude of the magnetic exchange interaction in SQ–Bridge–NN biradicals. From a valence bond/ resonance perspective, contributing ECN resonance structures accurately reflect both the spin- and charge distribution of low-lying SQ–Bridge–NN CT states, for which the corresponding quinoidal bridge-containing excited states in M–Bridge–M constructs lie far higher in energy. With respect to the importance of quinoidal resonance contributions, aromaticity has been shown to reduce conductance values in single molecule junctions.\(^6\) Critically, the quinoidal resonance structure contributions in Fig. 4 result in diminished aromaticity and contribute more to exchange than to conductance, leading to \(\gamma \neq 1\) and \(\beta_g/\beta_f > 1\) for these molecular bridges.

Conflicts of interest

There are no conflicts to declare.

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