**Biexciton Hamiltonian for the correlated triplet pair states in singlet fission: Revealing the contest between triplet-triplet exchange and triplet-triplet energy transfer coupling**

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**Abstract:** The correlated triplet pair state $^1(TT)$ is a key intermediate in the singlet fission process, and understanding the mechanism by which it separates into two independent triplet states is critical for leveraging singlet fission for improving solar cell efficiency. This separation mechanism is dominated by two key interactions: (i) the exchange interaction ($K$) between the triplets which leads to the spin splitting of the biexciton state into $^1(TT)$, $^3(TT)$, and $^5(TT)$ states, and (ii) the triplet-triplet energy transfer integral ($t$) which enables the formation of the spatially separated (but still spin entangled) state $^1(T...T)$. In this work, we use simple ab initio calculations to compute the triplet-triplet exchange ($K$) and triplet-triplet energy transfer coupling ($t$). For a model 1D system, we show how these parameters affect the biexciton energy manifold using a steady state approximation. Our key findings reveal a new condition for successful correlated triplet pair state dissociation. If the $K$ to $t$ ratio is comparable to or less than one, biexciton dissociation is favored for large chromophore assemblies. Additionally, for smaller chromophore assemblies, the biexciton exchange interaction needs to be negligible compared to the triplet energy transfer for favourable dissociation. We also explore the effect of chromophore packing to reveal geometries where the triplet energy transfer integral is significantly larger compared to the triplet-triplet exchange.

**I. INTRODUCTION**

Singlet fission is a spin allowed process where an excited singlet state ($S_1$) is converted to two low energy triplet states ($T_1 + T_1$). Although discovered around 50 years ago, it was recently shown that the process can be used to overcome the Shockley-Queisser limit for efficiency of single junction organic solar cells. The singlet fission mechanism involves a singlet excited state ($S_1$) forming a spin coupled triplet excited state $^1(TT)$ before splitting into two independent triplet states. $^1(TT)$

![Diagram of singlet fission process](image)

**FIG. 1:** Schematic representation of the proposed mechanism of singlet fission dissociation process.

The correlated triplet pair state $^1(TT)$ serves as the key spin conserving intermediate between the final triplet states and the initial singlet state for the singlet fission process. While the separation of the $^1(TT)$ to individual triplets was initially thought to be a fast process in fs region, recent studies have shown it to stay correlated until ns timescales. Being a long lived state, multiple competing decay pathways exist for the $^1(TT)$, beyond simply separating into individual triplets. The formation of the $^1(TT)$ state, which is the quintet coupled triplet pair state is also reported recently. One of the main advantages of forming the $^1(TT)$ state is that it doesn’t have any decay channel other than to separate into individual triplets unlike $^1(TT)$ or $^3(TT)$. These long lived $^1(TT)$ states might also have important applications in quantum information. Being a multichromophoric process, the interaction between molecules is very important for the singlet fission process. To realize the ideal of fast singlet-fission, the electronic interaction between chromophores should be strong enough for the triplet states to be entangled, yet not too strong such that the $^1(TT)$ becomes a trap state.

The triplet-triplet exchange interaction ($K$) provides insight into the energy manifold of the different spin states of biexcitonic (TT) character. This interaction is usually antiferromagnetic leading to the following state energy ordering: $^1(TT) < ^3(TT) < ^5(TT)$. Closely packed (or covalently bonded) chromophores often exhibit large exchange interaction. Usually, this close packing increases the electronic coupling between the initial singlet state and the $^1(TT)$, facilitating unfavorable decoherence via triplet-triplet annihilation. Consequently, it has been shown that lowering this coupling can lead to higher yields of the separated triplets from the $^1(TT)$ in case of intramolecular singlet fission.

Even after establishing the spin-state orderings of (TT), the dissociation of the correlated triplet pair state is still not well understood. Spatially separated yet spin entangled $^1(T...T)$ states have recently been proposed as a possible intermediate and have subsequently been characterized experimentally. The triplet-triplet energy transfer (or Dexter) mechanism has been proposed for the formation of these $^1(T...T)$ states. The triplet-triplet energy transfer rate largely depends on the square of the triplet-triplet energy transfer coupling ($t$) which is an exchange type interaction and decays exponentially with distance between donor and acceptor.
The purpose of this manuscript is two fold: First, we present an efficient ab initio-based technique to characterize the multi-excitonic states in terms of a simple model Hamiltonian consisting of physically meaningful quantities, $t$ and $K$. This model Hamiltonian, in turn, allows us to identify values of $t$ and $K$ which lead to favorable energetics for dissociation of the (TT) state into independent triplet excitons. Second, we perform numerical calculations to resolve the connection between chromophore packing and the $t$ and $K$ parameters to identify which geometries might be preferable for efficient triplet separation.

Theoretical calculations\textsuperscript{22,39,40} play an important role in characterizing the $^1$(TT) state, with its first theoretical report by Zimmerman and coworkers\textsuperscript{10} much before its first experimental detection.\textsuperscript{10} Being a doubly excited state, conventional single excitation methods (e.g., CIS, TDDFT, and RPA) cannot represent this state.\textsuperscript{10} Hence, calculations require very expensive multi-reference methods like CASSCF\textsuperscript{51} or MRPT\textsuperscript{52}. Spin flip\textsuperscript{62} methods are an alternate approach where the multi-configurational systems can be studied by starting from a well-defined high spin single reference. In this work, we use a single spin flip method to obtain the two key electronic couplings ($K$ and $t$ ) needed to describe the multieexciton space. We present numerical evidence that the multieexciton space is well represented using these two couplings and generate a biexciton Hamiltonian for the (TT) state.

II. METHODS

Restricted active space spin-flip\textsuperscript{53,59} (RAS-SF) methods have been successfully used to study the singlet fission process.\textsuperscript{51,57,61} Even though the RAS-SF calculation can produce reliable results for medium sized systems, the calculations get intractable as the number of chromophores increases. The reason, is that the number of spin-flip excitations increases combinatorially with number of chromophores. Recently, our group proposed a hybrid numerical/analytical method where only a simple 1-spin flip calculation is needed to obtain the biexciton manifold instead of factorially scaling $n$ spin flips.\textsuperscript{62} This is carried out by extracting a spin Hamiltonian using the single spin flip method and subsequently using the spin Hamiltonian to form the excited states, hence having the singly, doubly, and higher excited triplet states.\textsuperscript{53} However, having extracted a spin Hamiltonian does not fully alleviate the scaling problem. Even though the spin Hamiltonian is smaller compared to the ab initio Hamiltonian, the scaling is still factorial with system size. In this work, we further simplify this Hamiltonian to form a biexciton model, which only involves the doubly excited space of the spin Hamiltonian. We further demonstrate that this Hamiltonian can be formed solely by using the biexciton exchange and the biexciton hopping term.

A. Setup

While spin models have commonly been used to represent the coupled (TT) state\textsuperscript{17,24,34} these models mapped each chromophore to a spin 1 site. While these approaches were able to describe the dimer systems, the 1-to-1 mapping prevented the descriptions of biexcitons on model systems containing more than two chromophores. In Ref. \textsuperscript{62} a 1-to-2 mapping was introduced, wherein a chromophore system with $n$ chromophores can be cast as a $2n$ site spin lattice, allowing application to systems larger than chromophore dimers. In brief, this process involves computing the high spin ROHF (or DFT) calculation with multiplicity $2n+1$, and then carrying out a single spin flip calculation (CAS-SF, RAS-SF, SF-TDDFT) to obtain the lowest $2n$ eigenvectors in the $2n - 1$ multiplicity space (Figure 2b). The singly occupied orbitals are localized (Figure 2b) which organizes the determinant basis into either neutral or ionic configurations. We finally project the 1-SF eigenstates into the neutral determinant basis and reorthonormalize to obtain a new basis. The ab initio eigenvalues can be represented as a Bloch effective Hamiltonian matrix\textsuperscript{62} and the spin half exchange interaction can be obtained from the off diagonal elements of this effective Hamiltonian.\textsuperscript{62}

For a dimer system as shown in Figure 2a, starting from a quintet guess and using a single spin flip gives the two triplet states, $^3$(TT) and $^5$(TT) in the $m_s = +1$ space. Using these four states, we can form the spin=$\frac{1}{2}$ Hamiltonian using the procedure as mentioned in Ref. \textsuperscript{62}

$$\hat{H}^{\text{spin}} = -2 \sum_{ij} J_{ij} \hat{S}_i \hat{S}_j$$

1

The spin Hamiltonian represents each chromophore as a system with 2 sites. Therefore a system with $n$ chromophores can be approximated with $2n$ spin lattices (Figure 2a). The ability to construct an accurate spin model offers multiple advantages such as deeper conceptual insight, numerical efficiency, and ease of implementation of periodic boundary conditions. Analyzing the spin Hamiltonian produced using the 1-SF calculation, it can be seen that the interactions between the sites which are on the same chromophore are much larger than the interchromophore interactions. Hence we can write the Hamiltonian as

\textsuperscript{57,58}
FIG. 2: a) The final energy levels/biexciton spectrum for the bound triplet state energy manifold can be obtained using CAS-nSF and also by using CAS-1SF+ spin Hamiltonian. In this work we introduce the 1SF-Bloch method to obtain the biexciton spectrum which scales polynomially instead of exponentially. b) Illustration showing the transformation from the molecular lattice into spin lattice and the subsequent biexciton model. c) Singly occupied localized orbitals in the singly occupied active space for the pentancene dimer.

\[ \hat{H} = \sum_I \hat{H}_I + \sum_{IJ} \hat{H}_{IJ} \]

where \( I \) and \( J \) are chromophore indices. The intra chromophore exchange interactions are much larger than inter terms, and hence we can define the zeroth order as the intra terms. These intra chromophore interactions give an estimate of the excitation energy for a triplet state for each chromophore. We can diagonalize \( \sum_I \hat{H}_I \) to form exciton subspaces corresponding to single, double and higher excited triplet states (Figure 3). The energy manifold of each of these excited subspaces is well separated from the manifolds of other subspaces.

Since we are interested in the biexciton space, we can therefore ignore the effect of other states including the reference lowest energy state and other excited manifolds. We can diagonalize the Hamiltonian in the biexciton space to obtain the energies of the (TT) states. We test this model by analyzing the biexciton energy by diagonalizing the spin Hamiltonian and using our model for a tetracene heptamer in Section III A. By removing the ground state from the diagonalization, this approxi-
mation provides size consistent energies\cite{52} and can be extended to larger systems. This diabatic biexciton space has mainly two types of interactions, the exchange interaction between the triplet states and the energy transfer coupling. We will refer to our method to obtain the exchange interaction and the transfer coupling as 1SF-Bloch approach.

B. Biexciton Exchange Integral

The decoherence of the (TT) state for molecular dimers depends largely on the inter triplet exchange interaction ($K$)\cite{62}. For a chromophore dimer system, we can compute the $K$ value from the spin Hamiltonian derived using the 1SF calculation. The biexciton basis for a dimer system in the m$_s$ = 0 space can be formed using the zero order space $|T^+T^-\rangle$, $|T^0T^0\rangle$ and $|T^-T^+\rangle$ where the first index corresponds to chromophore A and the second index corresponds to chromophore B.

The $|T^+\rangle$, $|T^0\rangle$ and $|T^+\rangle$ can be written as

$$
|T^-\rangle = |\beta\beta\rangle \\
|T^0\rangle = \frac{1}{\sqrt{2}}|\alpha\beta + \beta\alpha\rangle \\
|T^+\rangle = |\alpha\alpha\rangle
$$

(3)

The matrix element between any two of these states can be evaluated quite easily for the spin lattice. For example, the matrix element between the $|T^+T^-\rangle$, $|T^0T^0\rangle$ and $|T^-T^+\rangle$ where the first index corresponds to chromophore A and the second index corresponds to chromophore B.

The $|T^+T^-\rangle |T^0T^0\rangle$ and $|T^-T^+\rangle$ in terms of the spin half exchange matrix element comes as,

$$
\langle T^+T^- | H | T^0T^0 \rangle = -\frac{1}{2}(J_{13} + J_{14} + J_{23} + J_{24})
$$

(4)

Here site 1 and 2 are in chromophore A and site 3 and 4 are in chromophore B (Figure 1).

This integral is the triple-triplet exchange interaction ($K$) between the two triplet states in the biexciton space. Hence we can write $K$ in terms of the inter chromophore exchange interactions.

$$
K_{AB} = -\frac{1}{2}(J_{13} + J_{14} + J_{23} + J_{24})
$$

(5)

The $3 \times 3$ Hamiltonian for the dimer using basis $|T^+T^-\rangle$, $|T^0T^0\rangle$ and $|T^-T^+\rangle$ can be written as,

$$
\hat{H} = \begin{bmatrix}
E - K_{AB} & K_{AB} & 0 \\
K_{AB} & E & K_{AB} \\
0 & K_{AB} & E - K_{AB}
\end{bmatrix}
$$

(6)

where $E$ is the reference energy for the biexciton space, and $K_{AB}$ is the exchange interaction between the two triplets in chromophores A and B. Diagonalization of this Hamiltonian leads to the three spin components of the multie exciton, $^1(TT)$, $^3(TT)$ and $^5(TT)$ with energies $E - 2K_{AB}$, $E - K_{AB}$ and $E + K_{AB}$ respectively and eigenvectors:

$$
|^{1}(TT)\rangle = \frac{1}{\sqrt{3}} (|T^+T^-\rangle - |T^0T^0\rangle + |T^-T^+\rangle)
$$

(7)

$$
|^{3}(TT)\rangle = \frac{1}{\sqrt{2}}(|T^+T^-\rangle - |T^-T^+\rangle)
$$

(8)

$$
|^{5}(TT)\rangle = \frac{1}{\sqrt{6}} (|T^+T^-\rangle + 2|T^0T^0\rangle + |T^-T^+\rangle)
$$

(9)

The $K$ parameter also plays an important role that it contributes to the exciton binding energy ($E_b$) defined as the energy difference between $^1(TT)$ and separated triplets. There are studies reporting the biexciton binding energy as low as meV\cite{64} to as high as 100’s of meV\cite{65}$. Connecting singlet fission chromophores through a covalent linker\cite{66} increases the interaction between the chromophores and usually results in large $E_b$ compared to crystals\cite{67}. One of the major limitations with covalently connected chromophores is that the rate of recombination to form the singlet state increases, diminishing the final triplet population\cite{68,69}. Recently, Basel et. al studied singlet fission using a non-conjugated linker between the pentacene monomers thereby decreasing the inter-chromophore exchange interaction\cite{70}. Another approach is by connecting the singlet fission chromophores through meta position of a bridging benzene molecule to change the sign of the exchange interaction\cite{71,72}. The final triplet yield is increased in such cases. Hence it can be said that low/ negative $K$ value can lead to improved triplet yield.

The exchange interaction (Equation 5) derived here is similar to the two orbital representations using the diabatic pseudo canonical orbitals with HOMO and LUMO orbitals in both the chromophore. The exchange interaction comes down to the sum of exchange integral between the orbitals that in representation as well\cite{62}.

C. Biexciton Transfer Integral

Recent studies have proposed that the decoherence of the $^1(TT)$ state occurs by evolving into the $^3(T\ldots T)$ state by triplet-triplet energy transfer\cite{73,74,75}. The magnitude of TT coupling, which is exchange type in nature depends exponentially on the distance between triplet donor and acceptor\cite{76}. Theoretical methods using singly excited methods have been previously used to estimate the triplet-triplet energy transfer integral using the fragment spin difference method\cite{77,78}. Charge transfer states are also important for the triplet-triplet energy transfer process\cite{79,80,81} and are indirectly (via the effective interactions) included in our model\cite{82}.

Given the spin Hamiltonian, the triplet-triplet hopping can be extracted easily by looking at the biexciton or
FIG. 4: The 4 site spin model for a two chromophore system. The biexciton exchange integral splits the three spin components energetically. The biexciton transfer integral plays an important role in formation of (T\textsubscript{T}) from (TT).

single exciton space. For example, consider a three chromophore system, where the biexciton can shift from one pair to another. For the trimer, the size of the biexciton space is \( 3C_2 \times 3 = 9 \) which includes three of each \( ^1\text{TT} \), \( ^3\text{TT} \) and \( ^5\text{TT} \). By closely analyzing the Hamiltonian, it can be seen that the biexciton subspace contains both the exchange interaction \( K \) between the chromophores and a new integral corresponding to triplet transfer, \( t \). This becomes evident when we work with the spin coupled biexciton space. This makes the Hamiltonian block diagonal with dense singlet, triplet and quintet blocks. The resulting Hamiltonian after spin coupling has the \( K \) terms in the diagonal.

The triplet-triplet hopping between two diabatic biexcitons \( ^1\text{TT}\)\textsubscript{AB} and \( ^1\text{TT}\)\textsubscript{AC} can be defined as the matrix element between the two states using the spin Hamiltonian.

\[
t_{AB} = \left\langle ^1\text{TT}\right|_BC \hat{H} \left| ^1\text{TT}\right|_AC \right\rangle \quad (10)
\]

Similar to \( K \), the \( t \) parameter between chromophore A and B can be written as a linear combination of the inter chromophore exchange terms.

\[
t_{AB} = \frac{1}{2}(-J_{13} + J_{14} + J_{23} - J_{24}) \quad (11)
\]

For the trimer, the “singlet-biexciton” block of the Hamiltonian can be represented as using basis of \( ^1\text{TT}\)\textsubscript{AB}, \( ^1\text{TT}\)\textsubscript{AC} and \( ^1\text{TT}\)\textsubscript{BC} can be written as,

\[
\hat{H}\left[ ^1\text{TT} \right] = \begin{bmatrix}
E - 2K_{AB} & t_{BC} & t_{AC}
\end{bmatrix}
\end{bmatrix}
\begin{bmatrix}
t_{BC}
E - 2K_{AC}
t_{AB}
\end{bmatrix}
\begin{bmatrix}
t_{AC}
E - 2K_{BC}
\end{bmatrix}
\quad (12)
\]

The transfer integral for the quintet and triplet multie excitons are the same as the singlet state. The off diagonal elements for the model Hamiltonians for those states look exactly same as Equation 12.

There are other methods based on single excitation quantum chemistry which can be used to obtain the triplet-triplet energy transfer coupling. The fragment spin difference (FSD)\textsuperscript{35,83} is a successful tool used for computing the triplet energy transfer coupling. We compare the coupling using the 1SF-Bloch model with the triplet-triplet energy transfer from the FSD scheme. In Figure 5, we present the comparison between the \( t \) parameter obtained using FSD scheme with our 1SF-Bloch method. For this comparison, we consider the tetracene dimer whose center of mass is 5.0\textdegree away and at 90 degrees with each other. The relative orientation is such that one of the chromophores is perpendicular to the other. Using the 1SF-Bloch model, we can see that the \( J_{13} \) and \( J_{23} \) are equal and so is \( J_{14} \) and \( J_{24} \) for this geometry. Hence using a spin model, this coupling should be zero, and indeed we find this value to be zero.

Next, we calculate the triplet-triplet energy transfer coupling using the FSD method with the triplet states obtained using CIS method. We can see that this coupling becomes zero at 90 degrees from Figure 5 which has been previously reported for the charge transfer couplings as well\textsuperscript{84}.

We compute the energy transfer integral for more
points near by rotating by a small angle and it can be seen that the FSD coupling and 1SF-Bloch coupling are very close for all of the points despite being formally very different approaches and using different references. This consistency between the two methods for computing $t$, helps validate that although our method simultaneously provides $t$ and $K$, we retain the accuracy of the FSD method which specializes on computing $t$.

**D. Biexciton Hamiltonian**

By using the 1SF-Bloch method, we have derived parameters that model the biexciton subspace. The $(TT)$ energy manifold for all three spin states can be obtained using the biexciton space. The final biexciton Hamiltonian for the $^1(TT)$ state using the two parameters can be written as:

$$\hat{H}_{^1(TT)} = \sum_{A,B} -2K_{AB}|A,B\rangle \langle A,B| + t_{AB}|A,C\rangle \langle C,B|.$$  \hfill (13)

The basis $|A,B\rangle$ is the biexciton state where chromophores A and B are in their triply excited states. The full biexciton Hamiltonian with all three spin states for an $n$ chromophore system scales as $3 \times nC_2$. Hence, this Hamiltonian scales quadratically with the number of chromophores compared to the factorial scaling for the parent spin Hamiltonian. In this work we diagonalize this Hamiltonian as a steady state approximation to look at the biexciton energy spectrum and understand the effect of $K$ and $t$ on the biexciton spectrum. Numerically these couplings are small and usually in meV scale which is much smaller than the excitation energies.

With the aide of the 1SF-Bloch model, we can now use a simple 1-spin flip calculation from a high spin reference to simultaneously obtain both the biexciton exchange and transfer integrals. In terms of the impact on singlet-fission triplet yield, the two parameters play opposing roles. Large positive exchange interaction can lead to $^1(TT)$ being a strongly bound state and reduces final triplet yield. Meanwhile having a larger hopping integral compared to the exchange should facilitate the formation of spatially separate ($T...T$) state. Hence, we expect systems with large $t$ and small/negative $K$ to facilitate in the dissociation of the biexciton state.

Even though both of these interactions are on the same scale, their relative magnitudes can change based on the interaction between the chromophores. For example, having two chromophores perpendicular to each other cancels the exciton transfer integral while the biexciton exchange interaction stays non zero. Since, the biexciton exchange interaction is a sum of the spin Hamiltonian terms, it does not really become zero for molecular assemblies when all sites have antiferromagnetic interaction.

While our proposed model is accurate in that it can reproduce more expensive $n$-spin flip calculations, and provides consistent results with the FSD approach, by it’s very nature it is incomplete. Most importantly, our model does not include the initial bright state. This prevents our model from describing both the initial formation of the $^1(TT)$ state, as well as the important triplet-triplet annihilation decay mechanism. Additionally, we have not included spin dipole interactions into our model. However, this is not an essential limitation, and we plan to include this in future work to study the $^1(TT)/^3(TT)$ mixing.

**FIG. 5:** Comparison of the triplet energy transfer integral obtained using FSD on top of a CIS wavefunction with our 1SF-Bloch approach.

**FIG. 6:** Comparison of the biexciton model with the exact result from the spin Hamiltonian for a system with seven tetracene units.
III. RESULTS AND DISCUSSION

We focus on three different aspects of the biexciton model proposed in this work. In Section III.A we compare numerical data for the biexciton model to the data from the actual diagonalization of the spin Hamiltonian for a large chromophore assembly. Secondly, in Section III.B we analyze the effect of different ratios of exchange to transfer parameter and look at the biexciton spectrum for a model 1D system. Finally, in Section III.C we investigate how the exchange and transfer parameters vary as we change the chromophore orientations and suggest geometric orientations where improved triplet dissociation is possible using our model.

A. Tetracene 7-mer

As described above, the mapping of the ab initio Hamiltonian to a quantum spin lattice does not completely remove the combinatorial complexity of the problem. While the dimension of the spin-Hamiltonian Hilbert space is indeed smaller (it is the square root of the fermionic problem), it is still intractable to diagonalize for large systems. However, because we are only concerned with the biexciton states (in contrast to tri-excitons and so forth), we propose to build and diagonalize the spin-Hamiltonian only the the biexcitonic manifold. This not only reduces the computational cost to polynomial scaling, but also restores size consistency by removing the ground state from the diagonalization space.

In this section, we evaluate the magnitude of the errors obtained by diagonalizing in only the biexcitonic subspace instead of the full space. In our previous work\cite{previous_work} we computed the biexciton spectrum using exact diagonalization (ED) of the full spin Hamiltonian for a large cluster with seven tetracene monomers. In Figure 6 we show the comparison of the $^1$(TT) spectrum using ED and the biexciton model Hamiltonian proposed in this paper. There are a total of 21 $^1$(TT) states for this system. As seen in Figure 6, the spectrum obtained from diagonalizing only the biexcitonic manifold is essentially indistinguishable from the exact diagonalization. We have also derived a rigorous downfolding using quasi-degenerate perturbation theory, but since the zeroth-order results presented here are accurate enough, we only provide these slightly more accurate results in the supplementary information.

The broadening of the biexciton spectrum is largely due to static disorder of the tetracene monomers (which would vanish if periodic boundary conditions were used in the ab initio calculations, which will be a focus of future work) and other acenes having two different geometries in the herringbone crystal structure\cite{herringbone}. The energy gap between these two different structures is as large as 50 meV which is larger than the exchange interaction in crystal tetracene molecule\cite{tetracene}.

B. Biexciton energy manifold for 1D system

We generate the biexciton energy manifold for a model 1D system with three chromophores. With three chromophores, there are total $3 \times 3 = 3 \times 3$ biexciton states. We analyze the effect of varying $K$ and $t$ on the biexciton spectrum.

We present the biexciton energy values for both antiferromagnetic and ferromagnetic exchange interactions ($K$) in Figure 8. The sign of $t$ does not change the biexciton spectrum. We fix the $K$ parameter and vary $t$ on the right side of the grey line and on the left we fix $t$ and vary $K$ for both antiferromagnetic (Figure 8a) and ferromagnetic (Figure 8b) $K$. For the antiferromagnetic case with very low $t$, it can be seen that the $^1$(TT) state is much lower in energy than the $^3$(TT) and $^5$(TT) states since $K$ value dominates here.

As we increase the $t$ parameter, we can see that one of each $^1$(TT), $^3$(TT) and $^5$(TT) states becomes more stable until $K = t$. When $K = 0$, all three spin states are degenerate and would facilitate spin mixing. As we increase the $K$ parameter, the $^1$(TT) state becomes lowest energy for the antiferromagnetic case and the $^5$(TT) becomes lowest energy for the ferromagnetic case. The spectrum is flipped vertically for the ferromagnetic $K$ values. Having the $^5$(TT) lower in energy would improve the final triplet yield since the $^5$(TT) state does not have other decay pathways. Hence for ideal singlet fission dissociation, we would expect low/negative $K$ values such that the $^1$(TT) and $^5$(TT) are very close in energy.

In the future, we will include the spin dipole interactions into the model to investigate the effect of spin mix-
FIG. 8: Comparison of the biexciton energy manifold for varying values of \( K \) and \( t \) for a) antiferromagnetic (positive \( K \)) and b) ferromagnetic (negative \( K \)) exchange interactions. We fix \( t = 1 \) meV for the left of the vertical gray line and vary \( K \) uniformly from 0 to 1. On the right side of gray line, we vary the \( t \) parameter from 1 to 0.

C. Biexciton coupling for tetracene dimer

The decoherence of the \(^1(TT)\) largely depends on the \( K \) and \( t \) parameters for the model presented. While large \( K \) leads to longer coherence time, large \( t \) helps in the formation of the \((T...T)\) state. In this section, we investigate how these parameters vary as the molecular packing is changed. The chromophore dimer units are generated using optimized geometries of tetracene molecules at B3LYP/6-31G(d). We present \( t \) and \( K \) parameters as a function of chromophore orientation.

a. Center of mass-coordinate

In this first panel, Figure 9(a), the location of one chromophore is fixed and the other chromophore is translated along the z axis. The \( K \) and \( t \) parameters decay exponentially as the distance between the chromophores \( (d) \) increases with exponential factors of \( e^{-2.76d} \) and \( e^{-2.77d} \), respectively. Beyond 5 Å the couplings fall below 1 meV. Hence having a large distance between the chromophores can increase spin mixing as the only non-exponentially decaying term remaining are spin dipolar interactions. Since both \( t \) and \( K \) decay similarly as a function of center of mass separation (the \( K/t \) ratio stays fairly constant at around 0.73), this therefore is not a useful coordinate for optimizing triplet-triplet separation.

b. \( \theta \)-coordinate

The energetically favorable herringbone structure of simple chromophores stabilize dimers which are rotated with respect to each other. Here we analyze how this rotation angle coordinate between the two chromophores influences the biexciton exchange and transfer integrals. We fix the distance between the centre of mass of the two molecules to be 5.0 Å and rotate one of the chromophores. Since the center of mass is fixed, by rotating we are bringing the two chromophores closer, hence at larger angles the parameters increase in magnitude, although they are still less than 10 meV. From Figure 9(b), it can be seen that the exchange interaction between the two chromophores becomes very low at about 64 degrees. This is the geometry where we have realized the smallest \( K/t \) ratio, which (based on the considerations discussed above), should provide the fastest \(^1(TT)\) \( \rightarrow \) \((T...T)\) separation and decoherence.

Consistent with models based on orbital overlap\(^{33}\), the perpendicular geometry drives the triplet hopping term becomes zero, essentially shutting down any formation of \((T...T)\). The spin lattice in the case of the exact perpendicular geometry is similar to a orthogonal dimer arrangement which is basically a 1D Shastry-Sutherland spin lattice\(^{33}\).

d. \( y \)-coordinate

We also study the parameters as we move the chromophore in the y direction (Figure 9(d)). We fix the distance between the planes of the two chromophores to be 3.75 Å. We can see that there is an interesting variation of \( K \) and \( t \) in this case and the periodicity is almost similar. While the sign of \( t \) oscillates along this coordinate, this does not affect the hopping rate since it depends on \( t^2 \).\(^{30,38}\) In terms of minimizing \( K/t \), one can see a favorable region occurring at around \( d = 2.75 Å \) where \( t \) reaches a local maximum. The exchange interaction between the two triplet states is ferromagnetic for the x shift from 5 Å to 5.6 Å and 7.6 Å to 8.2 Å. For these geometries, the \(^3(TT)\) state is lower in energy compared to the \(^1(TT)\) state.

c. \( x \)-coordinate

Next, we study the effect on biexciton exchange and transfer integrals when one chromophore is moved in the x direction (Figure 9(c)). We fix the distance between the planes of the two chromophores to be 3.75 Å (similar results were obtained at 3.75 Å). As seen
in the previous case, the values of the $t$ and $K$ parameters have similar periodicity. In terms of $K/t$ ratio, this coordinate minimizes this ratio only at small values of $d$, where the two chromophores are directly on top of one another. Hence we do not expect fast dissociation in any of the geometries along this coordinate.

**IV. SUMMARY**

In this work we present a simple biexciton model for the correlated triplet pair state intermediates in singlet fission materials. The biexciton energy manifold is mainly decided by two key parameters, the exciton transfer term ($t$) and the triplet-triplet exchange interaction ($K$). We demonstrate that these parameters can be derived from a simple ab initio calculation by using a single spin flip calculation from a high spin reference using the 1SF-Bloch approach as presented in this work. $K$ and $t$ are competing interactions, for the antiferromagnetic exchange, $K \gg t$ can lead to bound biexcitons whereas $t \gg K$ leads to small energy gap between $^1\text{(TT)}$ and $^5\text{(TT)}$, and favourable formation of ($\text{T...T}$) states. For a ferromagnetic exchange, the quintet biexciton states is more stable than the singlet state even for larger chromophore assemblies and hence can lead to higher triplet yield.

These different ratios of $K$ and $t$ depend on chromophore interactions, hence we investigate the effect of molecular packing. For most of the geometries, the $K$ value is antiferromagnetic. We find ferromagnetic interaction for a small region when the two chromophores are shifted along the $x$ axis around $\sim 5.3\text{A}$ and $\sim 7.9\text{A}$. Very few chromophore arrangements lead to conditions where $t$ is significantly larger than $K$. We find large exciton transfer compared to exchange along the $\theta$ rotation coordinate around 60 degrees. For these molecular packings, the biexciton dissociation should be favoured either by the fast formation of ($\text{T...T}$) state or the $^5\text{(TT)}$ state.

In the future, we will be developing a Marcus theory based model which includes the entanglement of the triplet pair. The biexciton model is simple and adding
spin-dipole effects to this model would be straightforward and will be investigated in future work. The model can be further improved by introducing hole and particle excitations either perturbatively or variationally or using an effective Hamiltonian derived using equation of motion coupled cluster (EOM-CC) Vibrational effects can be introduced for the model either by extracting the exchange parameters at different geometries of the vibrational mode or by introducing the vibrational coupling explicitly using phonon Hamiltonian. The 1SF-Bloch model can also be used for obtaining the coupling in triplet-triplet energy transfer reactions for artificial photosynthesis.

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References


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