Quantum trajectory tests of radical-pair quantum dynamics in CIDNP measurements of photosynthetic reaction centers

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ABSTRACT

Chemically induced dynamic nuclear polarization is a ubiquitous phenomenon in photosynthetic reaction centers. The relevant nuclear spin observables are a direct manifestation of the radical-pair mechanism. We here use quantum trajectories to describe the time evolution of radical-pairs, and compare their prediction of nuclear spin observables to the one derived from the radical-pair master equation. While our approach provides a consistent description, we unravel a major inconsistency within the conventional theory, thus challenging the theoretical interpretation of numerous CIDNP experiments sensitive to radical-pair reaction kinetics.

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1. Introduction

Quantum coherence in the light-harvesting process of photosynthesis [1–15] is a central theme in the growing field of quantum biology [16]. Equally important for understanding photosynthesis and potentially designing biomimetic devices harvesting solar energy is the charge separation following the trapping of the exciton’s energy in the photosynthetic reaction center. Charge separation proceeds via a cascade of electron transfers between radical-ion pairs, as for example bacteriochlorophyll and bacteriopheophytin. In parallel with charge, spin also plays a ubiquitous role [17], with the effect of chemically induced dynamic nuclear polarization (CIDNP) manifested in a wide range of photosynthetic reaction centers [18–31].

Nuclear spin effects in CIDNP are regulated by the radical-pair mechanism, which describes a class of spin-dependent chemical reactions studied by spin chemistry [32–38]. We have recently addressed [39–44] the fundamental quantum dynamics of the radical-pair mechanism, using concepts from quantum information science to describe the intertwined effects of coherent spin motion and spin-dependent charge recombination of radical-pairs, arriving at what we understand as a fundamental master equation describing the time evolution of ρ, the radical-pair’s spin density matrix. The traditional (also called Haberkorn) theory [32] used until now is a limiting case of our theory valid in the regime of strong spin relaxation.

Interestingly, in CIDNP measurements the lifetimes of singlet and triplet radical-pairs (RPs) are small enough to allow observation of quantum effects without them being masked by spin relaxation. Hence CIDNP appears to be an ideal setting to test our understanding of the quantum dynamics of the radical-pair mechanism. To this end, we demonstrated [42] a notable difference between our theory and Haberkorn’s approach in predicting a CIDNP effect at earth’s field. However, measurements at earth’s field are still challenging. Moreover, in [42] we assumed equal singlet and triplet recombination rates, kS = kT, whereas in real reaction centers we encounter asymmetric recombination rates, kT > kS [28].

Instead of attempting a comparison of the two different approaches based on an absolute quantitative prediction, we here use CINDP as a testbed to study the internal consistency of the two theories, Haberkorn’s and ours, and we do so using the realistic (i.e. asymmetric) recombination rates and high magnetic fields pertinent to CIDNP experiments. It is a basic fact of the theory of open quantum systems that the time evolution of an ensemble of systems described by a master equation must exactly reproduce the average calculated from single-system realizations, called quantum trajectories. We here use a central CIDNP observable, the nuclear spin polarization of the radical-pair’s reaction products, and show that Haberkorn’s theory produces vastly different predictions when using Haberkorn’s master equation compared to averaging Haberkorn’s quantum trajectories, while our theory produces largely consistent predictions either way.
2. Quantum measurement approach to radical-pair quantum dynamics

2.1. Master equation approach

In recent years we have addressed the fundamental quantum dynamics of radical-pair reactions, schematically depicted in Figure 1. This biochemical system is both a leaky and an open quantum system, losing population due to the spin-dependent recombination reactions, while simultaneously suffering decoherence. The latter aspect of the dynamics was addressed in [39], and from a slightly different perspective in [43], using quantum measurement theory. The former was phenomenologically considered in [40], while a first-principles approach was developed in [44], where we formally defined a singlet–triplet coherence measure, called $P_{\text{coh}}$. We also utilized the quantum–communications concept of quantum retrodiction to derive the spin-dependent reaction terms of the master equation. We will briefly recapitulate the results of [44] for completeness of this work.

The full master equation we arrived at in [44], the consistency of which we will explore in this work, reads

$$\frac{d\rho}{dt} = -i[\mathcal{H}, \rho]$$

(1)

$$-k_s + k_T \frac{1}{2}(\rho_{QS} + \rho_T - 2Q_S \rho_{QS})$$

(2)

$$-\{1 - P_{\text{coh}}\}(k_S Q_S \rho_{QS} + k_T Q_T \rho_{QT})$$

(3)

$$-P_{\text{coh}} \frac{d\rho}{dt} = \frac{1}{Tr[\rho]} \times \left( Q_S \rho_Q + Q_T \rho_Q + P_{\text{coh}} Q_S \rho_{QT} + P_{\text{coh}} Q_T \rho_{QS} \right)$$

(4)

The density matrix $\rho$ describes the spin state of the radical-ion-pair, consisting of the two unpaired electrons and the nuclear spins residing in the two radicals. The density matrix and all other operators are represented by $d \times d$ matrices, where $d$ is the dimension of the total spin space of the radical-pair. In this work we will consider a radical-pair having just one nuclear spin, hence $d = 8$, and the 8 basis kets are $|S \rangle \otimes |\uparrow \rangle$, $|S \rangle \otimes |\downarrow \rangle$, $|T_+ \rangle \otimes |\uparrow \rangle$, $|T_+ \rangle \otimes |\downarrow \rangle$, $|T_0 \rangle \otimes |\uparrow \rangle$, $|T_0 \rangle \otimes |\downarrow \rangle$, $|T_- \rangle \otimes |\uparrow \rangle$, $|T_- \rangle \otimes |\downarrow \rangle$. The two-electron state is on the left of the tensor product and the nuclear spin state is on the right. The two-electron spin subspace is spanned by the singlet $|S \rangle = (|\uparrow \downarrow \rangle - |\downarrow \uparrow \rangle)/\sqrt{2}$ and the triplets $|T_+ \rangle = |\uparrow \uparrow \rangle$, $|T_0 \rangle = (|\uparrow \downarrow \rangle + |\downarrow \uparrow \rangle)/\sqrt{2}$, $|T_- \rangle = |\downarrow \downarrow \rangle$.

![Figure 1](image_url)

**Figure 1.** A charge transfer from a photoexcited donor–acceptor dyad DA produces a radical-ion pair in the singlet state $1^D(S\uparrow\downarrow)$. Intermolecular magnetic interactions the Hamiltonian $\mathcal{H}$) induce a coherent interconversion between singlet and triplet radical-pair, $1^D(S\uparrow\downarrow) \leftrightarrow 1^D(T\uparrow\downarrow)$, terminated by the recombination event, that results into a singlet, DA, or a triplet neutral product, TDA. Radical-pairs are leaky systems because real transitions to the singlet and triplet vibrational reservoir states lead to population loss at the recombination rate $k_s$ and $k_T$ respectively. They are also open quantum systems, because virtual transitions to the same reservoir states and back to the radical-pair lead to singlet–triplet dephasing at the rate $(k_s + k_T)/2$.

The operators $Q_S$ and $Q_T$ project the radical-pair spin state onto the electron singlet and triplet subspace, respectively. They are orthogonal, complete and idempotent, i.e., $Q_S Q_T = Q_T Q_S = 0$, $Q_S + Q_T = 1$, where $\mathcal{d}$ is the $d \times d$ unit matrix, and $Q_S = Q_S^2 = Q_T = Q_T^2$. Writing $S_D$ and $S_A$ for the spin operator of the donor’s and acceptor’s unpaired electron, respectively, it is $Q_S = \frac{1}{2} - S_D \cdot S_A$.

The rates $k_s$ and $k_T$ are the singlet and triplet recombination rates. If at $t = 0$ we prepare an RP ensemble in the singlet (triplet) electron spin state, and assume that there is no singlet–triplet (S–T) mixing, then the RP population would decay exponentially at a rate $k_s$ ($k_T$). These two rates are properties of the particular RP under consideration. All in principle they can be calculated from electron transfer theory, but in practice they are determined from experiment. There are RPs for which $k_s = k_T$, and there are RPs for which $k_s \neq k_T$.

The term (1) in the previous master equation is the ordinary unitary evolution driven by the intramolecular magnetic interactions contained in the Hamiltonian $\mathcal{H}$ (Zeeman, hyperfine, etc.). The particular Hamiltonian used in this work is relevant to CINDP measurements, and will be outlined in Section 4. Since singlet and triplet states are not eigenstates of $\mathcal{H}$, the term (1) generates S–T coherence.

This is dissipated by the Lindblad term (2), which we derived in [39,43], and which describes a continuous quantum measurement of $Q_S$ performed by the vibrational reservoir $T_0$, the results of which are unobserved. This measurement effects projections to the electron singlet or triplet subspace suffered by individual RPs at random times. The interruptions of the coherent S–T mixing at the single-molecule level lead to S–T decoherence at the ensemble level.

The final two terms (3) and (4) are the so-called reaction terms, reducing the RP population, given by $Tr[\rho]$, in a spin-dependent way. The normalization of the initial density matrix is $Tr[\rho_0] = 1$, if we choose a small enough $dt$ so that $k_s dt, k_T dt \ll 1$, the fraction of the RP population that will recombine into singlet and triplet neutral products within the interval $dt$ around time $t$ is $d\rho = k_s dt Tr[\rho_Q S] = k_T dt Tr[\rho_Q T]$, respectively. Based on how coherent is the RP ensemble at time $t$, quantified by $P_{\text{coh}}$, which is a function of $\rho$ straightforward to calculate [44], we use the theory of quantum retrodiction to probabilistically estimate the pre-recombination state of the observed reaction products and thus arrive at (3) and (4). As expected, it is $dt Tr[\rho] = -d\rho - d\rho_T$. In the single-molecule picture, $d\rho_T (d\rho_T)$ is the probability that a single radical-pair will recombine in the electron singlet (triplet) channel during the time interval $dt$.

The values of $P_{\text{coh}}$ range from 0, describing a maximally incoherent mixture of singlet and triplet RPs, to $0 < P_{\text{coh}} < 1$ describing partially coherent, to $P_{\text{coh}} = 1$ describing maximally coherent RPs. When $P_{\text{coh}} = 0$, the whole term (4) vanishes.

Given (i) the two rates $k_s$ and $k_T$, (ii) the specific Hamiltonian $\mathcal{H}$, and (iii) the initial state $\rho_0$, Eqs. (1)-(4) can be used to propagate $\rho$ in time and thus calculate the time-dependence of any physical observable of interest.

2.2. Quantum trajectory approach

The new physics of RP quantum dynamics that we introduced in [39,40] is that during its lifetime, a radical-pair undergoes random projections to the singlet or triplet electron spin subspace. These projections take place at different and random times for each radical-pair. They run simultaneously with and independently of the second kind of random event, the RP charge recombination, which terminates the reaction. To generate quantum trajectories for an RP being in the state $|\psi(t)\rangle$ at time $t$, we thus have to consider in total 5 possible events that can take place in the following time interval $dt$ (Table 1):

<table>
<thead>
<tr>
<th>Event</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k_s$</td>
<td>Singlet recombination</td>
</tr>
<tr>
<td>$k_T$</td>
<td>Triplet recombination</td>
</tr>
<tr>
<td>$Q_S$</td>
<td>Singlet projection</td>
</tr>
<tr>
<td>$Q_T$</td>
<td>Triplet projection</td>
</tr>
<tr>
<td>$F$</td>
<td>Electron transfer reaction</td>
</tr>
</tbody>
</table>
3. Haberkorn approach to radical-pair quantum dynamics

3.1. Master equation approach

The traditional, or Haberkorn master equation reads

$$\frac{d\rho}{dt} = -i[H, \rho] - \frac{k_S}{2}(Q_S \rho + \rho Q_S) - \frac{k_T}{2}(Q_T \rho + \rho Q_T)$$

(5)

Interestingly, Haberkorn’s master equation follows from our master equation (1)–(4) by forcing $p_{\text{coh}}$ to be zero at all times. In any case, as we have done for our approach in the previous section, Haberkorn’s approach must be able to provide the equivalent picture of single-molecule quantum trajectories. However, the concept of quantum trajectories has not been utilized in spin chemistry so far. Furthermore, although a recent experiment [46] provided evidence for the physical reality of the S–T decoherence process we introduced, a general consensus on what exactly is the quantum state evolution of surviving RPs is still missing. From our perspective, we do not see how Haberkorn’s approach, being phenomenological, can incorporate quantum trajectories without going into the derivations of [39,43], which lead to events K1 and K2, but we leave it as an open question to be addressed by the proponents of the conventional theory. Nevertheless, we will here outline some rather strong guidelines as to how the consistency check of Haberkorn’s approach can in principle unfold.

3.2. Quantum trajectory approach

From Haberkorn’s theory point of view, it is clear that one cannot agree with possibilities K1 and K2, since these lead to our approach. Hence one has to suggest what is the specific state evolution of surviving RPs, i.e. what is, if any, the state change of radical-pairs until the instant of their recombination into a neutral product.

We will now show that there is limited freedom in doing so. This is because in order to secure consistency in the dynamically simple case $k_S = k_T$, one has to accept what has been until recently the intuitive answer to the previous question, namely that nothing (besides Hamiltonian evolution) happens to surviving RPs. As mentioned in Section 2.1, the rates $k_S$ and $k_T$ are parameters entering into the master equation, which obviously must be valid for any choice of those parameters. The case $k_S = k_T$ is rather simple dynamically, since in this case RP population decays exponentially at a rate $k = k_S = k_T$, without the decay affecting the state of the surviving RPs. This means the following. Consider for example a 50/50 mixture of singlet and triplet RPs, having no magnetic interactions ($\mathcal{H} = 0$). If $k_S = k_T$, the same number of singlet and triplet RPs will recombine in the interval $dt$, hence at time $t + dt$ the mixture will still be 50/50, albeit having a smaller total population. On the other hand, if $k_S \neq k_T$, the spin character of this mixture would change, becoming more (less) singlet if $k_S < k_T$ ($k_S > k_T$).

To summarize, (i) assuming that having a different fundamental theory for different radical pairs (i.e. different combinations of $k_S$ and $k_T$) is not an acceptable option, and (ii) being unable to propose what happens in the general case to non-recombining RPs from Haberkorn’s point of view, except in the special case $k_S = k_T$, where consistency forces one to accept that nothing else happens besides unitary evolution (to be proved in the following), we take this to be the general answer, and hence Haberkorn’s quantum trajectories are formed by the three events presented in Table 2.

There are two comments to be made. First, the recombination probabilities $dR$ and $dR'$ are the same in both Haberkorn’s approach and ours, since both theories agree in how the singlet and triplet reaction yields, $Y_i = \int dr_i$, with $j = S, T$, are calculated. Second, we can now easily prove our previous statement about H1–H3 ensuring consistency in the special case $k_S = k_T$. Indeed, taking into account the completeness relation $Q_S + Q_T = 1$, it follows that when $k_S = k_T$, Haberkorn’s master equation (5) becomes $d\rho/dt = -i[H, \rho] - k\rho$. Defining $\rho = e^{k \mathcal{R}}\mathcal{K}$, it follows that $d\rho/dt = -i[H, \mathcal{K}]$. It is thus evident that apart from an exponential decay of $Tr(\rho)$, the only radical-pair state change is due to the Hamiltonian evolution. In terms of the quantum trajectories H1–H3, we can retrieve the master equation $d\rho/dt = -i[H, \rho] - k\rho$ as follows. Since $Q_S + Q_T = 1$, it is $d\rho/dt = k dR/Tr(\rho)$. The single-RP density matrix is $\rho = Tr(\rho)$, hence averaging H1–H3 leads to

$$\rho + d\rho = dR \left( \rho - \frac{\rho}{Tr(\rho)} \right) + dR \left( \rho - \frac{\rho}{Tr(\rho)} \right)$$

(6)

from which it easily follows that indeed $d\rho/dt = -i[H, \rho] - k\rho$. To reiterate, the quantum trajectories formed by H1–H3 exactly reproduce the master equation (5) in the special case $k_S = k_T$. In the following we will show that this consistency check fails in the general and more interesting (in terms of realistic applications) case $k_S \neq k_T$.

4. Testing the consistency of Kominić’s and Haberkorn’s approaches using CIDNP observables

We will now demonstrate that while our approach is largely (but still not perfectly) consistent, Haberkorn’s approach is highly inconsistent. We will use a simple one-nuclear-spin radical-ion-pair with parameters relevant to CIDNP experiments. We stress that how many nuclear spins we consider, or which particular Hamiltonian we pick to exhibit the aforementioned inconsistency is of no concern, since as well known, it takes many supporting cases to establish a theory, but just one counterexample to invalidate it. Nevertheless, we choose a Hamiltonian of the same form considered in CINDP works like [47],

$$\mathcal{H} = \frac{\Delta \omega}{2} S z - \frac{\Delta \omega}{2} S z + e_0 I_z + A S \chi I_z + B S \chi k,$$

(7)

where $\Delta \omega$ is the difference in the Larmor frequencies of donor and acceptor electrons due to $\Delta g, \omega$, the nuclear Larmor frequency, and $A$ and $B$ isotropic and anisotropic hyperfine coupling constants. As in [26], we take a magnetic field of 5T along the z-axis. For the rest of the parameters we use $A = 4 \times 10^{-4}$, $A = \Delta \omega$ and $B = A/2$. 

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Quantum trajectory time evolution according to Kominić’ approach.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>Event</td>
</tr>
<tr>
<td>K1</td>
<td>Projection to the singlet state $\frac{Q_S(\rho)}{\sqrt{Q_S(\rho)}}$</td>
</tr>
<tr>
<td>K2</td>
<td>Projection to the triplet state $\frac{Q_T(\rho)}{\sqrt{Q_T(\rho)}}$</td>
</tr>
<tr>
<td>K3</td>
<td>Singlet recombination $dr_S = k_S dr(T)$</td>
</tr>
<tr>
<td>K4</td>
<td>Triplet recombination $dr_T = k_T dr(T)$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Quantum trajectory time evolution according to Haberkorn’ approach.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>Event</td>
</tr>
<tr>
<td>H1</td>
<td>Singlet recombination $dr_S = k_S dr(T)$</td>
</tr>
<tr>
<td>H2</td>
<td>Triplet recombination $dr_T = k_T dr(T)$</td>
</tr>
<tr>
<td>H3</td>
<td>Hamiltonian evolution $1 - dr_S - dr_T$</td>
</tr>
</tbody>
</table>
Finally, we use the asymmetric recombination rates $k_s = (20 \text{ ns})^{-1}$ and $k_T = (1 \text{ ns})^{-1}$ pertinent to photosynthetic reaction centers [17].

We now calculate the nuclear spin deposited to the reaction products, $dI_z$, as a function of time [30].

### 4.1. Density matrix propagation

If $\rho$ is the RP density matrix at time $t$, during $dt$ there will be $d\rho = k_s dT(\rho Q_s)$ singlet and $d\rho = k_T dT(\rho Q_T)$ triplet neutral products, the properly normalized density matrix of which is $\rho_s = Q_s \rho Q_s / Tr(\rho Q_s)$ and $\rho_t = Q_T \rho Q_T / Tr(\rho Q_T)$, respectively. Hence the total (singlet+triplet) ground-state nuclear spin accumulated during $dt$ is

$$dI_z = d\rho_s I_z(\rho_s) + d\rho_t I_z(\rho_t)$$

$$= \rho_s I_z(\rho_s) + k_T dT[I_z Q_T(\rho Q_T)]$$  \hspace{1cm} (8)

### 4.2. Quantum trajectories

If $|\psi\rangle$ is the state of the radical-pair at the random instant of recombination, then the properly normalized state of the singlet and triplet reaction product is $|\psi_s\rangle = Q_s |\psi\rangle / \sqrt{\langle \psi |Q_s|\psi\rangle}$ and $|\psi_t\rangle = Q_T |\psi\rangle / \sqrt{\langle \psi |Q_T|\psi\rangle}$, respectively. Hence the nuclear spin deposited to the reaction product is $|\psi_s I_z|\psi_s\rangle$ for a trajectory terminating with a singlet recombination, and $|\psi_t I_z|\psi_t\rangle$ for a trajectory terminating with a triplet recombination.

We evolve the quantum trajectories, and we numerically solve the master equation for two different initial conditions, $(I_{11}) |\psi_0\rangle = |S\rangle \otimes |\uparrow\rangle$, and $(I_{12}) |\psi_0\rangle = |S\rangle \otimes |\downarrow\rangle$. We then average the results of $(I_{11})$ and $(I_{12})$ in order to simulate the realistic scenario of starting with an unpolarized nuclear spin. Finally, we integrate the averaged traces in order to find $\int dI_z$, which is directly accessible in CIDNP experiments. This is normalized by the Boltzmann equilibrium nuclear spin, which for the considered parameters is $I_{\text{thermal}} = 10^{-5}$.

In Figure 2 we show the main result of this work. The level of inconsistency of Haberkorn’s approach is evident just by a visual inspection of Figure 2(d)–(f). Even more impressive is the result for $dI_z$, reproduced for convenience in Table 3. The results of Haberkorn’s master equation and Haberkorn’s quantum trajectories are not only different in magnitude by 740%, but are also of different sign. In our approach the two results are of the same sign and different in magnitude by 50%. In Appendix A we describe in detail how the quantum trajectories are produced, while in Appendix B we elaborate on the accuracy of our calculations.

### 5. Discussion

The consistency of a theory is not proof of correctness, but on the contrary, the inconsistency of a theory is proof of its inadequacy. Hence while this work provides a supporting argument that our approach is in the right direction, it unravels a major inconsistency within the traditional approach to radical-pair dynamics.

Until now, all the interesting information about the physical properties of photosynthetic reaction centers were extracted from CIDNP signals based on the traditional understanding of the
radical-pair mechanism. In other words, several mechanisms so far understood to produce enhanced nuclear spin polarizations are mostly based on the combined action of very specific Hamiltonian interactions and radical-pair reactions kinetics. It is clear that the extracted physical information contained in the former will be skewed by the inconsistent description of the latter.

In the conclusions of [48] it was stated that 'Until an experimental instance is found that requires an alternative description of the recombination kinetics, we recommend continued use of the conventional approach'. We believe that a failed consistency check of the conventional theory is quite stronger than an experimental instance challenging the theory. Even more so in light of the previous comment, namely that inconsistent reaction kinetics can contrive with skewed interaction Hamiltonians to produce a deluding agreement with experiments.

To our understanding, singlet and triplet projections are an integral part of the dynamics, the omission of which is responsible for the inconsistent behavior of the conventional theory. The reason is that the Hamiltonian coupling the spin degrees of freedom of the radical-pair to the vibrational reservoir (Figure 1) is responsible for both effects, random projections and recombination. The former within 2nd-order perturbation theory and the latter through 1st-order perturbation theory. As shown in [43], both effects are of order γ².

Clearly, there are still unresolved problems that must be addressed. While the quantum trajectory picture of our approach captures what we think are the underlying physics, our master equation does not do a perfect job in matching Monte Carlo. Hence more work is required to address this issue. Lastly, we recommend that the theoretical interpretation of a large number of CIDNP measurements performed over the last several decades should be meticulously revisited in light of a deeper understanding of radical-pair reaction kinetics.

Acknowledgement

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Appendix A. Generation of quantum trajectories

For each of the two initial states, we average 2 × 10⁵ quantum trajectories. A single quantum trajectory is generated as follows. We split the time from t = 0 to t_max = 20/kT, at which point the reaction is practically over, into n = 5 × 10⁵ steps of duration dt = t_max/n. We start with the initial state |ψ₀⟩ at time t = 0, and in each time step we draw a random number x uniformly distributed between 0 and 1. In our approach we calculate the probabilities dₜₛ, dₜₚ, dₜₛ and dₜₚ, and split the real interval [0, 1] into five intervals. If the random number x falls within the

- 1st interval of length dₜₛ, we realize K1 and move on to the next time step.
- 2nd interval of length dₜₚ, we realize K2 and move on to the next time step.
- 3rd interval of length dₜₛ, we realize K3 and terminate the particular trajectory.
- 4th interval of length dₜₚ, we realize K4 and terminate the particular trajectory.
- 5th interval of length 1 – dₜₛ – dₜₚ – dₜₛ – dₜₚ, we realize K5 and move on to the next time step.

In Haberkorn’s approach we calculate the probabilities dₜₛ and dₜₚ and split [0, 1] into three intervals. If the random number x falls within the
Examples of single quantum trajectories are shown in Figure 3. For visualizing the dynamics, we also plot the evolution of the singlet probability, $|Q_2|$, depicting S–T oscillations. Along those oscillations there are random singlet/triplet projections in Kominis’ trajectories, whereas they are completely missing from Haberkorn’s trajectories. In both theories, the trajectory is terminated by the recombination event.

Appendix B. Accuracy of the calculations

The accuracy of the master equation results (first line of Table 3) is limited just by the number of time steps $n$. By increasing $n$ beyond $5 \times 10^5$, the results converge to numbers within 1% of the ones stated here.

In order for the code to run in a practical amount of time, we limited the Monte Carlo simulation (second line of Table 3) to $2 \times 10^6$ trajectories. The whole simulation (the four Monte Carlo traces of Figure 2) takes about 9 h in a 4-core machine running at 2.5 GHz. The running time of propagating the density matrix discussed previously is negligible compared to one Monte Carlo trace, since the former involves just one time propagation of an $8 \times 8$ matrix, compared to $2 \times 10^5$ time propagations of an $8 \times 1$ vector required for the latter.

The accuracy of the integral $\int dt$, is easily estimated by taking e.g. 100 consecutive points along a relatively flat part of the traces in Figure 2, plotting the $y$-axis values in a histogram, and fitting with a Gaussian. The relative error is at the level of 10%. Thus the positive sign of Haberkorn’s Monte Carlo result is correct to within 10σ.

References