Quantum Simulation of the Radical Pair Dynamics of the Avian Compass

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ABSTRACT: The simulation of open quantum dynamics on quantum circuits has attracted wide interests recently with a variety of quantum algorithms developed and demonstrated. Among these, one particular design of a unitary-dilation-based quantum algorithm is capable of simulating general and complex physical systems. In this paper, we apply this quantum algorithm to simulating the dynamics of the radical pair mechanism in the avian compass. This application is demonstrated on the IBM QASM quantum simulator. This work is the first application of any quantum algorithm to simulating the radical pair mechanism in the avian compass, which not only demonstrates the generality of the quantum algorithm but also opens new opportunities for studying the avian compass with quantum computing devices.

A n open quantum system is a quantum system that interacts with an external environment or bath. The interaction between the system and the bath is often too complex to be simulated exactly and thus requires approximations to average out the effects of the bath—this results in the nonunitary dynamics of open quantum systems. Simulating the dynamics of quantum systems has been a main focus of quantum computing research, yet relatively few quantum algorithms have been developed for simulating the dynamics of open quantum systems. To this end, we have developed and demonstrated a general quantum algorithm for open quantum dynamics that is capable of simulating general and complex physical systems. The quantum algorithm leverages the Sz.-Nagy unitary dilation approach to convert nonunitary time evolution operators into corresponding unitary operators, which can then be implemented on a quantum circuit. This quantum algorithm has been applied to a variety of physical systems, including the amplitude damping channel described by the Kraus representation, the Jaynes–Cummings model described by the Kraus representation, the Fenna–Matthews–Olson (FMO) complex described by the Lindblad master equation, and the spin-boson model described by the generalized quantum master equation (GQME).

In this work, we apply the general quantum algorithm to simulating the radical pair mechanism in the avian compass and further demonstrate its generality. The radical pair mechanism (RPM) is a theory proposed to explain the magneto-reception and navigation abilities of certain bird species. Many animals possess extraordinary abilities to sense the direction by perceiving the geomagnetic field. This is probably the result of natural selection over a very long time of evolution, since the ability to sense the direction is crucial for certain animals to find their habitats, such as migratory birds that change habitats from season to season.

In brief, the RPM involves two spatially separated electrons, which are correlated with each other and affected by the external weak magnetic field and internal nuclear spin couplings. The basic scheme of the RPM includes three stages: 1) the photons with certain energies activate a certain type of molecules located in the bird’s eyes, enabling an electron transfer reaction and generating a radical pair in the singlet state; 2) the state of the radical pair converts between the singlet state and the triplet state under the influence of the external magnetic field (the hyperfine coupling effects); 3) the radical pairs in different states will generate different chemical products which can induce a detectable signal for birds to recognize the direction. The RPM is a promising hypothesis that can explain the three unusual properties of the avian compass: 1) The inclination compass: the functional mode of the avian magnetic compass is based on the inclination of the field lines instead of their polarity. 2) The light dependence: light with an energy above a certain threshold is needed for the RPM to work. 3) The narrow range of...
responsive magnetic field intensities: both higher and lower magnetic fields will disable a birds’ ability of navigation.\textsuperscript{36}

To understand the RP mechanism, E. M. Gauger et al. proposed a way to model the dynamics of the RPM system with a Lindblad master equation by adding two “shelving states” for the singlet yield and triple yield.\textsuperscript{37} In the following, we treat the same Lindblad formulation of the RPM dynamics with our general quantum algorithm for open quantum dynamics and simulate the RPM dynamics on the IBM QASM quantum simulator. To our best knowledge, this is the first ever demonstration of any quantum algorithm applied to simulating the RPM dynamics. This work not only shows the generality of the quantum algorithm but also opens new potential ways to study the avian compass with quantum computing devices.

The general quantum algorithm for open quantum dynamics has different versions that can evolve the Kraus representation,\textsuperscript{17} the Lindblad master equation,\textsuperscript{16} and the generalized quantum master equation.\textsuperscript{19} In this work we use the version for the Lindblad master equation.\textsuperscript{16}

We first review how the generalized quantum algorithm converts the nonunitary matrix that encodes the open quantum dynamics into a unitary evaluation based on the Sz.-Nagy unitary dilation procedure.\textsuperscript{17,18} We assume the initial density matrix that describes the physical system is composed of a set of unique pure quantum states $|\phi_i\rangle$ that are weighted by their corresponding probabilities $p_i$:

$$
\rho = \sum_i p_i |\phi_i\rangle \langle \phi_i|
$$

We want to simulate the time evolution of $\rho(t)$ given the initial $\rho$ and the Kraus operators $M_k$’s. This task can be achieved by preparing each input state $|\phi_i\rangle$ in a vector form $v_i$ in a given basis and then building a quantum circuit that generates the quantum state:

$$
|\phi_k(t)\rangle = M_k v_i \xrightarrow{\text{unitary dilation}} U_{M_k}(v_i^T, 0, ..., 0)^T
$$

The $U_{M_k}$ is generated via the 1-dilation of $M_k$:

$$
U_{M_k} = \begin{pmatrix} M_k & D_{M_k} \\ D_{M_k}^\dagger & -M_k \end{pmatrix}
$$

where $D_{M_k} = \sqrt{I - M_k^2M_k}$ and $D_{M_k}^\dagger = \sqrt{I - M_kM_k^\dagger}$.\textsuperscript{38} After obtaining each $|\phi_k(t)\rangle$, we can calculate the population of each basis state in the current basis from the diagonal vector:

$$
\text{diag}(\rho(t)) = \sum_k p_k \text{diag}(|\phi_k(t)\rangle \langle \phi_k(t)|)
$$

where $\text{diag}(|\phi_k(t)\rangle \langle \phi_k(t)|)$ can be efficiently obtained by applying projection measurements on the first half subspace of $U_{M_k}(v_i^T, 0, ..., 0)^T$.

For the dynamics of an open quantum system, the time evolution of the density matrix can be represented as

$$
\rho(s + \delta s) = \sum_k M_{ik} \rho(s) M_k^\dagger
$$

where $\rho(s)$ is the density matrix at time step $s$ and $\delta s$ is considered as the discrete time step, during which the Kraus operators $M_k$ are assumed to be constant. Equation 4 can be used iteratively until reaching the time of interest. Explicitly, the dynamics of the density matrix is described as

$$
\rho(1) = \rho(1) = \sum_k M_{ik} \rho(0) M_k^\dagger
$$

$$
\rho(2) = \rho(2) = \sum_k M_{ik} \rho(1) M_k^\dagger
$$

$$
\rho(3) = \rho(3) = \sum_k M_{ik} \rho(2) M_k^\dagger
$$

Here without losing any generality, the Kraus operators $M_k$ are indexed by the time step $s$, which allows each time step to have a different set of Kraus operators. However, the RPM dynamical model used in this work is a Markovian process described by the Lindblad master equation; therefore, all the time steps have the same set of Kraus operators $M_k$ where the time step index $s$ has been removed.

The basic scheme of the RPM is shown in Figure 1. To simplify the fields, we assume that only the electron near the donor interacts with the nucleus, and the electron away from the donor is not affected by the anisotropic hyperfine coupling.\textsuperscript{57} Therefore, the Hamiltonian of the system is

$$
H = \gamma [\hat{I} \cdot \mathbf{A} \cdot \hat{S}_z + \mathbf{B} \cdot (\hat{S}_x + \hat{S}_z)]
$$

Figure 1. Basic scheme of the radical pair mechanism. After absorbing light, donor (D) and acceptor (A) molecules form radical pairs in its singlet state. Then, under the influence of the magnetic fields, the states of the radical pair converse between the singlet states and triplet states. Finally, the singlet and triplet radical pairs end up with different products.
where $A$ is the anisotropic hyperfine tensor coupling the nucleus and one of two spatially separated electrons, and $A = \text{diag}(A_x, A_y, A_z)$ with $A_x = A_y = A_z/2$; $B = B_0(\cos \varphi \sin \theta, \sin \varphi \sin \theta, \cos \theta)$, and $B_0$ is the magnitude of the geomagnetic field; $\varphi$ is the angle between the $x$-axis of the radical pair and the external magnetic field; $\theta$ is the angle between the $z$-axis of the radical pair and the external magnetic field; and $\gamma = \frac{1}{2}\mu_0 g\mu_B$ and $\mu_B$ is the Bohr magneton and $g = 2$ is the electron-spin $g$-factor.

To model the dynamics of the system with a quantum master equation formulation, two “shelving states” were added to the 8-dimensional Hilbert space of the three spins (two electron spins and one nuclear spin).\textsuperscript{37} We employ operators as shown in eq 9 to represent the spin-selective relaxation into the singlet shelf $|S\rangle$ from the electron singlet state, or the triplet shelf $|T\rangle$ from the electron triplet state. The final populations of $|S\rangle$ and $|T\rangle$ give the singlet and triplet yields.

With the usual definition of singlet $|s\rangle$ and triplet states $|t\rangle$ in the electronic subspace, while $|1\rangle$ and $|1\rangle$ describe the states of the nuclear spin, we define the following decay operators:

\[
P_i = P_{i\uparrow\uparrow} = |S\rangle\langle s\uparrow\uparrow| \quad P_2 = P_{\uparrow\uparrow\downarrow} = |T\rangle\langle t\uparrow\downarrow|
\]

\[
P_3 = P_{\downarrow\uparrow\uparrow} = |T\rangle\langle t\downarrow\uparrow| \quad P_4 = P_{\downarrow\downarrow\downarrow} = |T\rangle\langle t\downarrow\downarrow|
\]

\[
P_5 = P_{\downarrow\downarrow\downarrow} = |S\rangle\langle s\downarrow\downarrow| \quad P_6 = P_{\downarrow\uparrow\uparrow} = |T\rangle\langle t\downarrow\uparrow|
\]

\[
P_7 = P_{\uparrow\downarrow\uparrow} = |T\rangle\langle t\uparrow\downarrow| \quad P_8 = P_{\uparrow\uparrow\downarrow} = |T\rangle\langle t\uparrow\downarrow|
\]

This gives a standard Lindblad master equation:

\[
\dot{\rho} = \frac{i}{\hbar} [H, \rho] + k_d \sum_{i=1}^{8} \left[ P_i \rho P_i^\dagger - \frac{1}{2} (P_i^\dagger P_i \rho + \rho P_i^\dagger P_i) \right]
\]

where $k_d$ is the decay rate of the singlet and triplet states. Note the decay rate $k_d$ is independent of the radical pair, so we have assigned the same decay rate $k_d$ to all eight projectors.

Now to apply the general quantum algorithm to the RPM dynamics, we first consider the nonunitary part on the right side of eq 10, which can be rewritten as

\[
\frac{\delta \rho(t)}{\delta t} = L(\rho) = k_d \sum_{i=1}^{9} \left[ P_i \rho P_i^\dagger - \frac{1}{2} (P_i^\dagger P_i \rho + \rho P_i^\dagger P_i) \right] + O(\delta t^2)
\]

Given a very small $\delta t$, eq 11 becomes

\[
\rho(t + \delta t) - \rho(t) = k_d \delta t \sum_{i=1}^{8} \left[ P_i \rho(t) P_i^\dagger - \frac{1}{2} (P_i^\dagger P_i \rho + \rho P_i^\dagger P_i) \right] + O(\delta t^2)
\]

Assuming $M_0 = \sqrt{I - \frac{i}{\hbar} k_d \delta t \sum_{k=1}^{8} P_k^\dagger P_k}$ and $M_k = \sqrt{k_d \delta t P_k}$ for $k > 0$, and ignoring the second order of $\delta t$ as $\delta t \to 0$, eq 12 can be rewritten as

\[
\rho(t + \delta t) = M_0 \rho(t) M_0^\dagger + \sum_{k=1}^{8} M_k \rho(t) M_k^\dagger
\]

with $M_0 = \sqrt{\text{diag}(I)}$. This “oscillating part” of the dynamics is unitary and thus can be easily realized by multiplying each Kraus operator by a unitary matrix obtained through the diagonalization of the Hamiltonian.\textsuperscript{18}

With the parameters in Table 1, we simulated the RPM dynamics by iteratively applying eq 13 on the IBM QASM quantum simulator and then used the output results to calculate the singlet and triplet yields. Also, we assume the initial state of the two electron spins is $\frac{1}{\sqrt{3}}(|\uparrow\downarrow\rangle + |\downarrow\uparrow\rangle)$, and the initial state of the nuclear spin is $\frac{1}{\sqrt{2}}(|\uparrow\rangle - |\downarrow\rangle)$. In our simulation, the time interval $\delta t$ is set as $0.5/k_d = 5 \times 10^{-5}$ s. We then apply the procedure shown in eqs 5–7. As mentioned above, the populations of $|S\rangle$ and $|T\rangle$ are the singlet and triplet yields, respectively, after the system reaches the steady state. The populations are calculated by the procedure explained in eq 3, where the diagonal elements of the density matrix are obtained by projection measurements into the computational subspace.

The results are compared with those obtained from classical methods in Figure 2, where the quantum algorithm results are highly consistent with the classical method results. Figure 3 shows the dynamical evolution of the singlet and triplet yields (when $\theta = \frac{\pi}{2}$) as simulated by the general quantum algorithm on the IBM quantum simulator. After $2 \times 10^{-4}$ s, the yields almost reach steady state, which is consistent with the chosen decay rate of $k_d = 1 \times 10^4$ s$^{-1}$.

One factor that contributes to the complexity of the quantum algorithm is the system’s size. For a density matrix of the size $n \times n$, the cost to realize the unitary dilation of a most general $n \times n$ Kraus operator $M_k$ is $O(n^2)$.\textsuperscript{17,18} However, in our calculation of the dynamics of the RPM, the Kraus

Table 1. Parameter Values Used in the Calculation

| symbol | description | values |
|--------|-------------|--------|
| $A_x$ | anisotropic hyperfine tensor | $1 \times 10^{-4}$ T |
| $B_0$ | magnitude of the geomagnetic field | $5 \times 10^{-3}$ T |
| $\gamma$ | half of the product of the Bohr magneton and electron-spin $g$-factor | $9.27 \times 10^{-24}$ J/T |
| $h$ | reduced Planck constant | $1.05457 \times 10^{-32}$ J-s |
| $\varphi$ | angle between the $x$-axis of the radical pair and the magnetic field | $0$ |
| $k_d$ | decay rate of the singlet and triplet states | $1 \times 10^4$ s$^{-1}$ |
operators each represent a single elementary physical process and thus the $M_k$ matrices are often sparse with few nonzero elements. This means the practical complexity scaling of implementing each $M_k$ matrix on a quantum circuit can be greatly reduced to $O(\log^2 n)$. Taking into account the total $K$ number of $M_k$ matrices to be simulated on the quantum circuit, the total complexity scaling is $O(K \log^2 n)$ for our given system. It is worth noting that the $K$ is determined on a case-by-case basis by the dynamical model and different $M_k$ matrices can be evolved in parallel; therefore, the scaling in $K$ is a “soft” scaling that does not contribute to either the depth or the width of each individual quantum circuit.\textsuperscript{17,18} Another contributing factor to the quantum algorithm’s complexity is the number of time steps. In the most general case, as can be seen from eq 5 to eq 7, taking $s$ steps requires $K$ matrices to be evolved, which is an exponential scaling in the number of time steps. However, fortunately, the actual number of matrix terms to be simulated can be greatly reduced once again due to the sparsity of the $M_k$ matrices. As mentioned above, the Kraus operators represent elementary physical processes and thus the $M_k$ matrices are often sparse with very few nonzero elements: this means that most matrix product terms in, e.g., eq 7 are zero matrices or matrices with negligible norms. The actual number of matrix products we need to evaluate is determined on a case-by-case basis. In the current simulation, in theory the total number of terms in $n$-th iteration will be $9^n$ with nine Kraus operators \{$M_k | k = 0, 1, 2, ..., 8 \}$. However, since the product of each pair of the decay operators \{$P_i | i = 1, 2, 3, ..., 8 \}$ is 0, the product of each pair of the Kraus operator \{$M_k | k = 1, 2, 3, ..., 8 \}$ is 0. Therefore, there will be only 8 more terms when adding one more iteration. Thus, there will be $8 \times n + 1$ terms in $n$-th iteration, reducing the terms significantly. More simulation details are in the Supporting Information.

Figure 2. Comparison of the singlet (in blue) and triplet (in red) yields between the results obtained from exact calculation and the quantum simulation of the RPM dynamics. The exact curves are generated from the cubic interpolation of the exact calculation of the yields at each data point. The dots represent the results simulated by the general quantum algorithm as implemented on the IBM QASM simulator. The parameters used are shown in Table 1. The yields are calculated around $7.5 \times 10^{-4}$ s after the system has already reached the steady state. The $y$-axis is the final singlet/triplet yields, i.e., the populations of singlet/triplet shelf state, and the $x$-axis is the angle between $z$-axis of the radical pair and the magnetic field.

Figure 3. Comparison of the dynamics of the singlet (in blue) and triplet (in red) yields between the exact and quantum simulated results. The exact curves are generated from the cubic interpolation of the exact calculation of the yields at each data point. The dots represent the results simulated by the general quantum algorithm as implemented on the IBM QASM simulator. After about $2 \times 10^{-4}$ s, both yields reach the steady state. The $y$-axis is the final singlet/triplet yields, i.e., the populations of singlet/triplet shelf state, and the $x$-axis is the time.
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Notes

The authors declare no competing financial interest.

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