Synthesis of Hidden Subgroup Quantum Algorithms and Quantum Chemical Dynamics

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ABSTRACT: We describe a general formalism for quantum dynamics and show how this formalism subsumes several quantum algorithms, including the Deutsch, Deutsch–Jozsa, Bernstein–Vazirani, Simon, and Shor algorithms as well as the conventional approach to quantum dynamics based on tensor networks. The common framework exposes similarities among quantum algorithms and natural quantum phenomena: we illustrate this connection by showing how the correlated behavior of protons in water wire systems that are common in many biological and materials systems parallels the structure of Shor’s algorithm.

1. INTRODUCTION

The promise of solving complex problems efficiently using quantum computing hardware and associated software is a rapidly evolving research frontier. While we are in the very early stages of this upcoming quantum revolution, there are a diverse set of important scientific and technological areas that may greatly benefit from such developments. One key quantum algorithm that started this entire debate approximately 25 years ago is Shor’s algorithm. Here, a quantum system can, in principle, factorize large integers into prime factors using \( N (\log \log \log N) \) fast multiplications. Since this is exponentially faster than the traditional classical approach, which requires \( O\left(e^{\frac{1}{289}(\log N)\frac{1}{2} (\log \log N)^{2+\epsilon}}\right) \) operations, the promise of a second quantum revolution was born.

Orthogonally, the sister fields of atomic and molecular physics and quantum chemistry have learned to wonder if atoms and molecules store and propagate quantum information. While it has been known that such “information” indeed evolves in time as per the laws of quantum theory, one may also ask if chemical reactions and chemical transformations are indeed algebraic transformations that “compute” new information not dissimilar from quantum algorithms. That is, is the time evolution of a molecular process to be interpreted as a computational protocol that is “programmed” by nature or through clever use of synthetic techniques? However, the study of molecular dynamics is complicated by the fact that molecules contain many correlated degrees of freedom. For example, with \( D \) degrees of freedom and \( N \) basis representation per degree of freedom, the complexity of information grows approximately as \( N^D \). As a result, quantum chemical dynamics is thought to be exponentially hard.

To alleviate this rather catastrophic situation, tensor networks (TNs)\(^7\)–\(^8\) have recently become popular. Tensor networks have roots in the tensor decomposition field of multilinear algebra,\(^9\)–\(^10\) and have proven to be effective for efficient representation of many-body quantum states in strongly correlated systems.\(^7\)–\(^25\) While a tensor network treatment adaptively truncates the Hilbert space based on the intrinsic entanglement within the problem, given the advent of novel quantum computing algorithms, tensor networks have also proved to be a natural resource for developing new quantum algorithms.\(^26\)–\(^29\) The approach has been shown to have applications for low-energy states of local, gapped Hamiltonians, which are characterized by satisfying a so-called area law of entanglement.\(^7\)–\(^30,31\) The introduction of the

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density matrix renormalization group (DMRG)\textsuperscript{32–35} was perhaps the catalyst for the excitement in the TN methodology, proving to be very useful for the simulation of one-dimensional quantum lattices,\textsuperscript{36–39} electronic structure calculations,\textsuperscript{40–48} approximations to vibrational states,\textsuperscript{49–56} open-quantum systems\textsuperscript{57,58} and image processing,\textsuperscript{11–14,59} and even machine learning applications.\textsuperscript{60–62}

In this paper, we cast the basic structure present in a family of quantum algorithms that includes Shor’s algorithm in an abstract fashion using the language of tensor networks.\textsuperscript{7} This presentation exposes parallels to more general quantum processes that occur in multidimensional quantum dynamics\textsuperscript{63–71} and in quantum dynamics of open systems\textsuperscript{72–83} that are of significance to many chemical, biological, and materials problems. Hence, we will then show how such an abstraction applies directly to many natural and synthetic chemical processes, thus drawing a connection between existing quantum algorithms and chemical and natural processes.\textsuperscript{83–87} At the end of this exposition, we are forced to ask if natural processes exist that may represent mathematically constructed, number-theoretic algorithms.

Given this overarching theme, this paper is organized as follows. In Section 2, we review the textbook presentation of Shor’s algorithm and generalize it in Section 3, using tensor networks to arbitrary multipartite systems. The formalism used in that generalization leads to our central result in Section 4, which derives the correspondence to generalized Shor-like circuits applicable to general quantum chemical dynamics problems. In Section 5, we exploit it to show that multidimensional quantum dynamics in protonated water clusters such as wire systems that are present in many biological channels and enzyme active sites and are also the subject of several state-of-the-art experimental\textsuperscript{88–91} and multidimensional correlated quantum dynamics studies\textsuperscript{92–94} can be mapped to the circuit model exhibiting the same structure as the family of quantum algorithms under study. Section 6 concludes.

2. QUANTUM ALGORITHMS

Most quantum algorithms that are thought to be exponentially faster than their best-known classical counterparts are algorithms for solving instances of hidden subgroup problems. This family of algorithms includes the textbook quantum algorithms of Deutsch, Deutsch–Jozsa, Bernstein–Vazirani, Simon, and Shor.\textsuperscript{95–102} and are all solved using the same approach illustrated in Figure 1. All of the algorithms start by creating an equal superposition of all relevant possibilities, applying the \( U_f \) block to the superposition, and analyzing the result using the quantum Fourier transform (QFT). The \( U_f \) block, often called the “oracle,” is uniformly defined as

\[
U_f(|x\rangle|y\rangle) = |x\rangle|f(x)\oplus y\rangle
\]

for the specific function \( f \) of interest. The circuit template uses the QFT uniformly as the last step, although—with the notable exception of Shor’s algorithm—the low precision approximation of QFT (which is the Hadamard gate\textsuperscript{103}) is often sufficient.

2.1. Shor’s Algorithm. To be concrete, Figure 2 instantiates the general template to the quantum circuit for an instance of Shor’s algorithm for factoring the number \( N \). In Stage (1), two registers are prepared: the top (input) register of \( q \) qubits is initialized to an equal superposition \( \frac{1}{\sqrt{q}} \sum_{i=0}^{q-1} |i\rangle \). In the bottom (output) register of \( m \) qubits, each qubit is initialized to \( |0\rangle \). In Stage (2), the initial state,

\[
\left\{ \frac{1}{\sqrt{2^{m-1}}} \sum_{i=0}^{2^{m-1}-1} |i\rangle \right\} \otimes |0\rangle
\]

is evolved through a reversible circuit that computes \( [a^r \pmod{N}] \). The resulting state is

\[
\frac{1}{\sqrt{2^m}} \sum_{i=0}^{2^{m-1}-1} |i\rangle \otimes |a^r \pmod{N}\rangle
\]

Equation 3 represents a highly entangled state. The degree to which the two sets of registers above is entangled is probed by computing the Schmidt number from a tensor network decomposition of the unitary evolution operations that lead to eq 3, as detailed in ref 104.

At Stage (3), a measurement of the output register produces some value \( w \); this measurement collapses the input register to a superposition of these states \( |i\rangle \) where \( a^r \pmod{N} = w \). Let the number of those states be \( W \); the input register state is then \( \frac{1}{\sqrt{W}} \sum_{i=0}^{W-1} |i\rangle \) for those states \( i \)'s whose mapping by the function \( a^r \pmod{N} \) produces the same value \( w \). Since the function \( a^r \pmod{N} \) is periodic, all of these states are guaranteed to be of the form \( |a + k\rangle \) for some starting offset \( a \) and some multiple \( k \) of the period \( s \). Put differently, the state of the input register is

\[
\frac{1}{\sqrt{W}} \sum_{k=0}^{(W/s)-1} |a + ks\rangle
\]

It is important to note that a different measurement \( w' \) of the output register would only change the starting offset \( a \) and the total number of states \( W \) in the superposition, but it would not change the period \( s \). Critically, the QFT is largely insensitive to the starting offset and the total number of states in the superposition. Its main effect is to transform a superposition of periodic states \( |a + ks\rangle \) to states in the Fourier basis \( |b\rangle \) such that \( b \) is close to a multiple of \( W/s \). When the period \( s \) is a power of 2, the Fourier states are perfectly aligned with the multiple of \( W/s \), as shown by the formula below:

\[
\text{QFT}\left(\frac{1}{\sqrt{W}} \sum_{k=0}^{(W/s)-1} |a + ks\rangle\right) = \frac{1}{\sqrt{s}} \sum_{m=0}^{(s-1)} e^{(2\pi i/m) a m W/s} |mW/s\rangle
\]

When the period is not a power of 2, the Fourier states with the largest probabilities are the ones close to a multiple of \( W/s \). From such a measurement, some classical postprocessing succeeds, with high probability, in determining the period \( s \) and hence the factors of the number \( N \).

2.2. Factoring Examples. We illustrate the algorithm for \( N = 15 \) and 21. In the first simpler example of factoring \( N = 15 \), we proceed as follows. In a classical preprocessing step, we choose a
value for $a$ that is coprime with 15 (say 2), calculate the needed number of qubits $q = 4$ and $m = 4$, and generate the modular exponentiation circuit for $f(x) = 2^x \mod 15$ using adders and multipliers. The execution of the quantum circuit proceeds as follows. The input register is initialized to the (unnormalized) equal superposition of $|0\rangle + |1\rangle + \cdots + |15\rangle$. At barrier (2), the two registers are entangled, producing the (unnormalized) state $|0\rangle |1\rangle + |1\rangle |2\rangle + |2\rangle |4\rangle + |3\rangle |8\rangle + |4\rangle |1\rangle + \cdots + |15\rangle |8\rangle$. A measurement of the output register may produce 1, 2, 4, or 8 with equal probability. Say we measure 4. The input register then collapses to the (unnormalized) state $|0\rangle + |4\rangle + |8\rangle + |12\rangle$. Say we measure $|12\rangle$. By properties of the QFT, we know that 12 is a multiple of $m = 16/s$, where $s$ is the period we seek, i.e., $12m = 16/s$ or $12/16 = m/s$. The idea is that $m/s$ is guaranteed to be a small irreducible fraction that approximates $12/16$. In this case, we get the exact approximation $3/4$, from which we infer that the period is 4.

From the period, we calculate the two factors of 15 using $\gcd(15, s) = 3$ and $\gcd(15, 5) = 5$. We follow the previous development with $N = 21$, $a = 10$, $q = 5$, and $m = 5$. At barrier (2), the (unnormalized) state is $|0\rangle |1\rangle + |1\rangle |10\rangle + |2\rangle |16\rangle + |3\rangle |13\rangle + |4\rangle |4\rangle + |5\rangle |19\rangle + |6\rangle |1\rangle + \cdots + |31\rangle |10\rangle$. Say we measure 13 at the output register. The input register collapses to the (unnormalized) state $|3\rangle + |9\rangle + \cdots + |27\rangle$. The QFT is not as perfect this time. We get the following approximation $63=4$, from which we infer that the period is 4.

The coefficients $C_{ij}$ capture the degree to which the parts, A and B, are correlated to each other. For example, when $C_{ij}$ is zero for all but one value of $i$ and $j$, then A and B are completely decoupled, and a product approximation suffices. However, when this is not the case, the degree to which A and B influence each other is often important in physical systems. Another popular example of eq 6 are the well-known Bell states $^2$ for two-qubit systems, which are a sum of product states,

$$\psi^1_{\text{Bell}} = \frac{1}{\sqrt{2}} (|0\rangle |1\rangle \pm |1\rangle |0\rangle)$$

and

$$\psi^2_{\text{Bell}} = \frac{1}{\sqrt{2}} (|0\rangle |0\rangle \pm |1\rangle |1\rangle)$$

Equation 6 may also be rewritten using the Schmidt decomposition, as

$$\psi = \sum_i \alpha_i |\psi^A_i\rangle |\psi^B_i\rangle$$

which is essentially a bipartite matrix product state (MPS) type network decomposition and is usually obtained by applying a sequence of singular value decomposition steps on the Tucker-form of the entangled states in eq 6. Such states are common in quantum dynamics, as well as in quantum computing, where the degree of correlation or entanglement between parts A and B are gauged using the $\alpha$-values. For the so-called maximally entangled states (such as Bell states and the Greenberger–Horne–Zeilinger (GHZ) states), $\alpha$ is a constant value for all $i$. (Compare eq 9 with eqs 7 and 8.) In quantum chemical dynamics, such highly correlated states are not common, and in general, the $\alpha$-values may decay in some fashion when the sets of states $\{|\psi^A_i\rangle\}$ and $\{|\psi^B_i\rangle\}$ are appropriately ordered.

For a system containing $D$ separate parts labeled as $A_r \gamma = 1 \cdots D$, one may write the overall wave function in a form similar to eq 6:

$$\psi = \sum_{r \in \gamma} C_{r \in \gamma} |\psi^{A_1}_{\gamma} \cdots |\psi^{A_{D}}_{\gamma}\rangle$$

3. MULTIPARTITE QUANTUM DYNAMICS AS TENSOR NETWORKS

The quantum algorithms of Section 2 and the tensor networks approach share some apparent similarities. In both cases, the systems are composed of multiple correlated parts that evolve quantum mechanically in a system-dependent manner and are interrogated using scenarios that measure one part of the system. This measurement affects the remaining parts of the system, whose spectral properties can then be inferred using the QFT. In this section, we make this intuitive correspondence precise, opening the door for richer connections between mathematically constructed algorithms and systems occurring in nature.

3.1. Multipartite Quantum Systems. We begin by considering a multidimensional quantum system represented as $A \otimes B$ composed of two correlated parts, A and B. Here, A and B can be multiple parts of a complex correlated quantum system, or alternately, A $\otimes$ B can together represent a complex condensed-phase quantum dynamics problem, where it may be more appropriate to refer to these as system and bath variables. The development here includes both descriptions. Such a system can be modeled in the tensor network formalism using a family of orthogonal states that represent subsystem $A$, referred to as $\{\psi^A_i\}$, that may be correlated to (or entangled with) a family of mutually orthogonal states representing $B$ and referred to as $\{\psi^B_i\}$. The overall wave function is then written as a tensor product, or correlated sum, of the two components, namely,

$$\psi = \sum_{i,j} C_{i,j} |\psi^A_i\rangle |\psi^B_j\rangle$$

Figure 2. Quantum circuit for Shor’s algorithm.
where $C_{\alpha_i\ldots}\mathcal{D}$ is a rank-$\mathcal{D}$ tensor and encodes the correlations between the constituents, $\{A_i\}$. In fact, eq 10 is the starting point for the well-known multiconfigurational time-dependent Hartree (MCTDH) approach commonly used in multidimensional quantum dynamics\(^{7,70,107}\) and vibrational spectroscopy. The matrix product state representation of eq 10, obtained from a sequence of bipartite singular value decomposition steps, yields the MPS state

$$|\psi\rangle = \sum_{\tau} |\psi_{\tau}^{A}\rangle |\psi_{\tau}^{B}\rangle |\psi_{\tau+1}^{A}\rangle |\psi_{\tau+1}^{B}\rangle \ldots |\psi_{\tau_{n-1}}^{A}\rangle |\psi_{\tau_{n-1}}^{B}\rangle$$

$$(11)$$

where the coefficients $\{\beta_{\tau}\}$ take on a generalization of the $\alpha_i$ in eq 9 and capture entanglement in a system with $\mathcal{D}$ parts.

### 3.2. Time Evolution of Multiparticle Quantum Systems.

In $A \otimes B$ systems occurring in quantum dynamics and electron–nuclear dynamics, we are often interested in learning about the influence of each part on the other. Toward this goal, without loss of generality, we begin by introducing an initial state of the $A \otimes B$ system that is an uncorrelated bipartite simplification of eq 9, that is,

$$|\psi_0\rangle = |\psi_0^{A}\rangle |\psi_0^{B}\rangle$$

The time evolution of the state $|\psi_0\rangle$ is given using a unitary evolution operator,

$$\mathcal{U}|\psi_0\rangle = |\psi_0^{A}\rangle |\psi_0^{B}\rangle$$

$$(13)$$

which may be further explicated by writing the time-evolution operator $\mathcal{U} \equiv \exp -i\mathcal{H}/\hbar$ as a correlated matrix product operator\(^{41,109,110}\) or a tensor product operator,

$$\mathcal{U} = \sum_{\alpha} \mathcal{U}_\alpha^a \mathcal{U}_\alpha^b$$

$$(14)$$

where, again, the multiple parts of the system are coupled by the overall Hamiltonian (and the time-evolution operator). Thus,

$$\mathcal{U}|\psi_0\rangle = \sum_{\alpha} \mathcal{U}_\alpha^a |\psi_0^{A}\rangle |\psi_0^{B}\rangle = \sum_{\alpha} |\psi_\alpha^{A}\rangle |\psi_\alpha^{B}\rangle$$

$$(15)$$

The structure of the Hamiltonian and the associated time-evolution operator result in the system correlations that are captured within the time-evolution process. This is represented by the sum of product states on the right side of eq 15. It must be noted here that while $\mathcal{U}$ is required to be unitary, in general, no such restrictions are present on $\{\mathcal{U}_\alpha^a, \mathcal{U}_\alpha^b\}$. This implies that while $|\psi_\alpha^{A}\rangle$ and $|\psi_\alpha^{B}\rangle$ may not, in general, be normalized, the overall propagated state is always normalized. For most physical systems, $\mathcal{U}$ has an explicit time dependence and is given by the exponential of a Hermitian operator as noted above, and hence $\langle |\psi_\alpha^{A}\rangle |\psi_\alpha^B\rangle \rightarrow \langle |\psi_\alpha^{A}\rangle |\psi_\alpha^B\rangle \rangle$. In such cases, the nonunitary nature of $\{\mathcal{U}_\alpha^a, \mathcal{U}_\alpha^b\}$, combined with the unitary nature of $\mathcal{U}$, yields a flow of probability between parts $A$ and $B$. When only one term is present on the right side of eq 14, the two parts $A$ and $B$ are uncorrelated, and in such cases, $|\psi_\alpha^{A}\rangle \rangle$ and $|\psi_\alpha^B\rangle \rangle$ remain individually normalized, and there is no flow of information between the two parts.

Since the input states for each part of the system may be chosen from a complete set of states, $\{\{\psi_0^A\}\}$ and $\{\{\psi_0^B\}\}$, we may expand the final states, $|\psi_\alpha^{A}\rangle$ and $|\psi_\alpha^{B}\rangle$, using these as basis functions to obtain the general form of the composite state after time evolution as

$$|\psi_\alpha\rangle = \sum_{j} \sum_{j} c_{j,\alpha}^{A} |\psi_j^{A}\rangle |\psi_j^{B}\rangle = \sum_{j} \sum_{j} c_{j,\alpha}^{A} |\psi_j^{A}\rangle |\psi_j^{B}\rangle$$

$$(16)$$

where $c_{j,\alpha}^{A} = \langle \psi_j^{A}\rangle \langle \psi_j^{B}\rangle$, and similarly for $c_{j,\alpha}^{B}$. Thus, it is the coefficient tensor, $\sum_{\alpha} c_{\alpha}^{A} c_{\alpha}^{B}$ that builds in the correlations in eq 6 and is obtained here through time evolution by $\mathcal{U}$. At this stage, the two parts of the system are completely correlated to the extent allowed by the propagator $\mathcal{U}$. In fact, the extent of such a correlation may be precisely defined by the number of elements in the summation in eqs 14 and 16.

### 3.3. Final-State Analysis.

At this stage in quantum chemical dynamics,\(^{111}\) there are several analysis techniques available to gauge correlations within eq 16. There are two basic types of questions asked of the propagated state in eq 16. In one case, it is of interest to directly Fourier transform eq 16 in the time domain since these now provide the spectroscopic signatures of the Hamiltonian and, hence, the eigenspectrum of the Hamiltonian that governs the dynamics. This is common when only a few degrees of freedom are involved and tends to become prohibitive when the number of dimensions grow. Second, along the lines of the topic here, one is often interested in how the subsystem $A$ evolves and is coupled to the properties of subsystem $B$. In a system-bath context, subsystem $B$ may be considered as the bath degrees of freedom, whereas in a multidimensional quantum system or a reactive problem, subsystem $B$ may contain modes that facilitate a chemical process or transition. This is a general problem and includes both condensed-phase quantum dynamics as well as electron–nuclear dynamics. It may also include chemical and biological sensing phenomena as signified by molecular binding processes.

Correspondingly, we project eq 16 onto a specific bath state $|\psi_\beta^B\rangle$, which is akin to performing a measurement on the bath state, to yield

$$\langle \psi_\beta^B|\mathcal{U}|\psi_0\rangle = \sum_{j} \sum_{j} c_{j,\alpha}^{A} c_{j,\alpha}^{B} \langle \psi_j^{A}|\psi_j^{B}\rangle = \sum_{j} c_{j,\alpha}^{B} |\phi_\alpha^A\rangle$$

$$(17)$$

which represents the state after measurement on the bath state, with the measurement outcome

$$Tr[|\psi_\beta^B\rangle \langle \psi_\beta^B|\mathcal{U}|\psi_0\rangle \langle \psi_0|\mathcal{U}^\dagger|\psi_\beta^B\rangle] = Tr[|\psi_\beta^B\rangle \langle \psi_\beta^B|\mathcal{U}^\dagger\mathcal{U}|\psi_0\rangle \langle \psi_0|\mathcal{U}^\dagger\mathcal{U}^\dagger]$$

$$= Tr\left[\sum_{j} \sum_{j} c_{j,\alpha}^{A} c_{j,\alpha}^{A} |\psi_j^{A}\rangle \langle \psi_j^{A}|\psi_j^{B}\rangle \langle \psi_j^{B}|\psi_j^{B}\rangle\right]$$

$$= Tr\left[\sum_{\alpha} c_{\alpha}^{B} |\phi_\alpha^A\rangle \langle \phi_\alpha^A|\phi_\alpha^A\rangle \langle \phi_\alpha^A|\phi_\alpha^A\rangle\right]$$

$$= \sum_{\alpha} \langle \phi_\alpha^A|\phi_\alpha^A\rangle$$

$$(18)$$

where $c_{\alpha}^{B} = \langle \psi_\alpha^B|\psi_\beta^B\rangle$, and similarly for $c_{\alpha}^{A}$. Thus, it is the coefficient tensor, $\sum_{\alpha} c_{\alpha}^{A} c_{\alpha}^{B}$ that builds in the correlations in eq 6 and is obtained here through time evolution by $\mathcal{U}$. At this stage, the two parts of the system are completely correlated to the extent allowed by the propagator $\mathcal{U}$. In fact, the extent of such a correlation may be precisely defined by the number of elements in the summation in eqs 14 and 16.
Thus, while the result of measurement is the net probability of eq 16 along $|\psi_j|^2$, that is eq 18, the remaining state is as in eq 17. A large number of such measurements on the bath state will yield various outcomes

$$\{ Tr(|\psi_j^B(U|\psi_j^B)|^2) ; |\psi_j^B|^2 \}$$

(19)

where each outcome, $(|\psi_j^B(U|\psi_j^B)|^2$, for bath state, $|\psi_j^B|^2$, is accompanied by the system remaining in the state given in eq 17. In this manner, multiple measurements of $B$ yield multiple states for $A$. Fourier transform of each of these yield,

$$\text{FT}\left\{ \sum_j \sum_\alpha c^{A_j}_\alpha \chi^{B,j}_\alpha \left| \psi_j^A \right\rangle \right\} = \text{FT}\left\{ \sum_\alpha c^{B,j}_\alpha \phi^{A,j}_\alpha \right\}$$

(20)

effectively the state of $A$, and the degree of coupling, or entanglement, between the system ($A$) and bath ($B$) state as originally captured by eq 9. Equations 19 and 20 are critical to multiple areas of metrology in physical and biological sciences. In each case, the interpretation of systems $A$ and $B$ may be different. In sensing, an analyte might bind to system $B$, which collapses the system, and its Fourier transform (or a linear transform) may provide information about the analyte binding to $B$. Similar aspects exist in condensed-phase quantum dynamics and chemical catalysis as well.

3.4. Generalized Phase Kickback. In eq 18, part $B$ is measured by using the same basis, $(|\psi_j^B|^2)$, as that used for the original propagation. Suppose this was not the case and the measurement was done using a specific ket, $|\chi_j^B|^2$ from within a different basis $(|\psi_j^B|^2)$ where

$$|\chi_j^B|^2 = \sum_i d^{B,j}_i |\psi_j^B|^2$$

(21)

and $d^{B,j}_i = \langle \psi_j^B | \chi_j^B \rangle$. In that case, eq 17, takes on a more general form:

$$\langle \chi_j^B | U | \psi_j^B \rangle = \sum_{i,i'} \sum_\alpha c^{i,j}_\alpha c_{i'}^{B,j} \left| \psi_j^A \right\rangle \langle \psi_i^B | \chi_j^B \rangle$$

$$= \sum_{i,i'} d^{B,j}_i |\chi_j^B|^2 \left| \psi_j^A \right\rangle \langle \psi_i^B |$$

(22)

$$= \sum_{i,i',\alpha} d^{B,j}_i \chi^{B,j}_\alpha \left| \phi^{A,j}_\alpha \right\rangle$$

(23)

Equation 23, as we show below, represents a generalized form of the phase kickback property, which is commonly seen in quantum information. This can be illustrated by considering eqs 7 and 8 as our propagated states, $U |\psi_j^B\rangle$. That is, to make a connection between the abstract tensor network formalism and qubits:

$$\{|\psi_j^A\rangle\} \rightarrow \{|0\rangle^A ; |1\rangle^A \}$$

(24)

and similarly for $B$. Furthermore,

$$\{U |\psi_j^B\rangle\} \rightarrow \{|\psi_{\text{Bell}}^1\rangle ; |\psi_{\text{Bell}}^2\rangle \}$$

(25)

The measurement basis for phase kickback is chosen as

$$\{|\chi_j^B\rangle\} \rightarrow \{|+\rangle^B ; |-\rangle^B \}$$

(26)

and therefore $d^{B,j}_i = \pm 1/\sqrt{2}$ for all $i,j$. (See eq 21.) In that case, as per eq 23, when measurement is constructed using $|\pm\rangle^B$, one finds

$$\langle + | \psi_{\text{Bell}}^1 \rangle = \frac{1}{\sqrt{2}} |1\rangle^A \pm \frac{1}{\sqrt{2}} |0\rangle^A$$

(27)

and

$$\langle - | \psi_{\text{Bell}}^2 \rangle = \frac{1}{\sqrt{2}} |1\rangle^A \pm \frac{1}{\sqrt{2}} |0\rangle^A$$

(28)

That is, system $A$ is rotated onto the $X$ basis as a result of this measurement, or the phase angle in $\langle + | \psi_{\text{Bell}}^1 \rangle$ is “kicked” into state $A$, upon measurement. Likewise,

$$\langle + | \psi_{\text{Bell}}^2 \rangle = \frac{1}{\sqrt{2}} |0\rangle^A \pm \frac{1}{\sqrt{2}} |1\rangle^A$$

(29)

and similarly $\langle - | \psi_{\text{Bell}}^2 \rangle$.

These features are captured in a general way within eqs 22 and 23, where the appropriate generalization of the phase kickback is in the terms $d^{B,j}_i \chi^B_j \psi^A_j$ that represent the components of the measurement basis of $B$ with respect to the initial basis and also the additional basis components that are “kicked-back” into system $A$, as per eq 22, after measurement on $B$.

The broader implications of eqs 22 and 23 are as follows: if we consider a general system containing two entangled parts, with the degree of entanglement dictated by a unitary evolution operator and hence an underlying Hamiltonian, a measurement or projection on one part, chosen as $A$ here, is also noted in $B$. Thus, in some sense, $B$ can “sense” the projection in $A$, but the extent of such a sensing process is dictated by the extent of entanglement present within $U |\psi_j^B\rangle$ in eq 16. For the Bell state, the information is directly transferred, whereas for the state in eq 22, the measurement information is convoluted with the extent of entanglement.

4. CIRCUIT MODEL FOR Multipartite QUANTUM DYNAMICS

The development in the previous section can be recast in the circuit model to make the parallels with Shor’s algorithm more explicit. We begin with Figure 3, which provides an instance of the general formalism as a quantum circuit closely relating to the description of Shor’s algorithm in Section 2. Specifically, the initial state in eq 12 is chosen to be a direct product state and represents Stage (1) in Figure 3. (Compare eqs 2 and 12.) It must be noted that this initial state, depicted at the end of Stage (1) in Figure 2, was obtained from a set of Hadamard transforms that essentially provide equal weights to all components of the computational basis embedded within the first wire stream of Figure 2.
Thus, equivalently, $A$ in Figure 3 may be entangled at Stage (1), but importantly, $A$ and $B$ are uncorrelated at this initial stage, and in this sense, eq 12 resembles the initial state for all of the quantum algorithms shown above. This state is then time-evolved as dictated by eq 14, leading to eqs 15 and 16, and is represented as Stage (2) in Figure 3. Similarly, the $U_y$ operator in Figure 2 plays the same role as the propagator in Figure 3 and presents a correlated (or entangled) state, given by eq 3 and represented at Stage (2) in Figure 2. Following this time evolution leading to eqs 15 and 16, a measurement is constructed in all scenarios. We have presented only the analogue to Shor in Figure 3, complemented by the discussion in Section 3.3. The resultant state in Figure 3, given by eq 17, now represents a general and abstract interpretation of the resultant state of Shor’s algorithm at Stage (3), prior to the QFT step.

The next step, as per Figures 2 and 3, is a Fourier transform of the state of the system, $A$, as described by eqs 19 and 20, which yields the momentum representation of the resultant state in subsystem $A$. In a sense, this also presents a more abstract form of the output from the top wire in Figure 2, and one may be induced to ask if we indeed obtain a similar “momentum representation” for the states captured in the top wire (natural numbers) in Figure 2. Thus, at the end of this process, if the Hamiltonian represented within the propagator in eq 14 (or in Shor’s algorithm) entangles the $A$ and $B$ dimensions, then a measurement of $B$ projects it onto a specific state. Following this, a Fourier transform of $A$ yields the momentum representation and, in fact, the power spectrum of $A$ for the specific projection of $B$.

Such a Fourier transform captures the entanglement within the composite $A \otimes B$ supersystem by probing the Fourier space structure of one part of the supersystem, namely, system $A$, for all possible measurement outcomes of system $B$ (assuming that multiple measurements are done on $B$). For the specific choice of unitary in Shor’s algorithm, this Fourier spectrum of $A$ is always the same for any measurement of $B$. This may not, of course, be the case for naturally occurring or physicochemical systems.

Finally, we note a set of problems that may benefit from the analysis above. We will explore these connections in detail in future publications. Equation 6 resembles the total nonadiabatic electron–nuclear wave function for molecular systems and may be written as an expansion in the complete set of electronic wave functions with the coefficients being functions of nuclear coordinates. In multidimensional correlation spectroscopy, vibrational mode coupling may be studied using similar partitioning schemes as in eq 6, where bath variables influence the dynamics of a chosen system. In fact, the example chosen in the next section is related to problems in multidimensional correlation spectroscopy. In chemical catalysis, ligands that surround an active site may influence the reactive process. Such ideas are commonly used in catalyst design.

5. APPLICATIONS: PROTONATED WATER WIRE SYSTEMS

We now exploit the formalism presented above to explore parallels between quantum algorithms and natural phenomena in physical, chemical, and biological systems.

Protonated water wires such as those in Figure 4a,b are encountered in a large variety of biological ion channels, catalytic sites, light-harvesting systems, and fuel cells and form the central part of many condensed-phase chemical processes. Such systems are found in confined media such as ion channels. Quantum effects play a critical role in contributing to the rate of proton transport and in determining vibrational properties.

5.1. Coupled Stretch Modes in Protonated Water Dimer. In this section, we provide a tensor network description of such protonated systems, beginning with the simple case of two protons and extending to a longer water wire chain of multiple shared protons.

We begin with an analysis of the two shared proton dimensions marked as $R_1$ and $R_2$ in Figure 4b. Using the Schmidt decomposition, the wave function for this two-dimensional system may be written as

$$\psi(x_1, x_2) = \sum_i \alpha_i \psi_i^\alpha(x_1) \psi_i^\beta(x_2)$$

(30)

Here, $\{\psi_i(x)\}$ represents a family of functions that depicts the distribution of dimension $R_1$; this family is coupled to the family of functions, $\{\psi_j(x)\}$ that depicts the distribution of dimension $R_2$. Additionally, $\langle \psi_i | \psi_j \rangle = \delta_{ij}$, and thus these functions form an independent orthonormal basis for the two separate dimensions. For a $(\text{H}_2\text{O})^2$ water wire subsystem with degrees of freedom $R_1$ and $R_2$, the wave function components are calculated as shown in Figure 5a,b.

Several techniques can be used to investigate such systems. A basic one is to perform a measurement on one of the dimensions, say $R_1$. This interrogation could be done by projecting the system onto a specific state of $R_1$, say, $\psi_k(x)$. We then obtain

$$\langle \psi_k(x_1) | \psi(x_1, x_2) \rangle = \alpha_k \psi_k^\alpha(x_2)$$

(31)

based on the orthogonality conditions stated above. The associated measurement outcome is

$$|\alpha_k|^2 |\psi_k^\alpha|^2$$

(32)

and for $k = 1$, this quantity is close to 1, as indicated by the values of $|\alpha_i|$ provided in the caption for Figure 5.

More generally, we could also envision a more sophisticated measurement where the interrogation could be performed using the state $\chi(x_1) \equiv 1/\sqrt{2} [\psi_1(x_1) + \psi_2(x_1)]$ leading to a final state:

$$\langle \chi(x_1) | \psi(x_1, x_2) \rangle = \frac{1}{\sqrt{2}} [\alpha_1 \psi_1^\alpha(x_2) + \alpha_2 \psi_2^\alpha(x_2)]$$

(33)
In this case, the superposition in the interrogation state is transferred to the outcome

$$\frac{1}{2} |\alpha|^2 |\psi^2_1| + |\alpha|^2 |\psi^2_2| \approx 0.5 \quad (34)$$

In both interrogation scenarios, the measurement of one dimension influenced the other. Thus, the form of the state in eq 30 has a significant impact on the result of the observation. If one of the states that is already within the family included in eq 30 is used as the measurement basis, the outcome is the corresponding state within that family. However, when a combination of states is used as the measurement basis, the phase kickback mechanism causes the complex phase included in this combination to make its appearance in the measured outcome.

5.2. Coupled Stretch Modes in Protonated Water Pentamer. In biological systems, the water wire described above is often partially confined within an active site or inside an ion channel, as illustrated in Figure 6. In this section, we analyze an instance of such a system consisting of four degrees of freedom.

Deferring the confinement modeling for a moment and focusing on the four degrees of freedom, the matrix product state for the coupled wave function has the form,

$$\psi(x_1, x_2, x_3, x_4) = \sum_{i_1i_2i_3i_4} \psi^1_{i_1}(x_1) \alpha_{i_1i_2} \psi^2_{i_2}(x_2) \alpha_{i_2i_3} \psi^3_{i_3}(x_3) \alpha_{i_3i_4} \psi^4_{i_4}(x_4) \quad (35)$$

The quantities $\alpha_i^j$ in the equation above represent weights for the bond dimensions.

As we did in the previous section, we can model various interrogation scenarios. We show the result of performing a measurement on dimension $R_1$. This reduces eq 35 to produce

$$\langle \psi^1_{i_1}(x_1) \psi(x_1, x_2, x_3, x_4) \rangle = \sum_{i_2i_3i_4} \alpha_{i_1i_2} \alpha_{i_2i_3} \alpha_{i_3i_4} \psi^1_{i_1}(x_1) \psi^2_{i_2}(x_2) \psi^3_{i_3}(x_3) \psi^4_{i_4}(x_4) \quad (36)$$

and the corresponding measurement outcome is simply the magnitude of the vector in eq 36:

$$\langle \psi(x_1, x_2, x_3, x_4) \psi^1_{i_1}(x_1) \psi^2_{i_2}(x_2) \psi^3_{i_3}(x_3) \psi^4_{i_4}(x_4) \rangle \quad (37)$$

This simple analysis ignored the fact that in most biological systems (especially ion channels as well as enzyme active sites), the water molecules that encapsulate dimensions $R_2$ and $R_3$ have a limited degree of flexibility. This results in a limited degree of projection of dimensions $R_2$ and $R_3$ into, for example, a subspace given by

$$\chi(x_2, x_3) = \sum_{i_2i_3} \beta_{i_2i_3} \psi^2_{i_2}(x_2) \psi^3_{i_3}(x_3) \quad (38)$$

Note that such a state is not dissimilar to the measurement basis state used in the phase kickback scenario at the end of Section 3 and reduces the possible outcomes for $x_4$. Thus, after projection of eq 38 onto eq 36, we obtain

$$\sum_{i_2i_3} (\beta_{i_2i_3} \alpha_{i_2i_3} \psi^2_{i_2}(x_2) \psi^3_{i_3}(x_3) \psi^4_{i_4}(x_4) \quad (39)$$

where the bracketed set acts as a combined coefficient that curtails the set of possibilities for $x_4$. Based on the extent of flexibility provided by the restrictions to $\beta_{i_2i_3}$, there are a range of possible outcomes at the far end depicted by $x_4$.

This idea can be generalized for an arbitrary number of degrees of freedom, where, using eq 39, we may write our final result as

$$\sum_{i_2 \cdots i_{N-1}i_N} (\beta_{i_2 \cdots i_{N-1}i_N} \alpha_{i_2 \cdots i_{N-1}} \psi^N_{i_N}(x_N) \quad (40)$$

5.3. Circuit Model for Protonated Water Wire. We now analyze the results from the sections above using the quantum circuit model. The quantum circuits thus derived are based on Figure 3 and are presented in Figures 7 and 8. As discussed in Section 4, all parts of both systems are correlated at the end of Stage (2) with wave functions given by eqs 30 and 35. In the case of the protonated water dimer, a measurement along dimension $R_1$ follows, resulting in the projected state given by eq...
31 at Stage (3), with the measurement probability given by eq 32.

The protonated pentamer problem is complicated due to the phase kickback step resulting from basis rotations (eq 39 on the \(R_2, R_3\) degrees of freedom, as shown in Figure 8). Specifically, the correlated state at Stage (2) given by eq 35 undergoes measurements at \(R_1\) with the outcome, eq 37, where the corresponding resultant state is then projected onto a rotated basis within dimensions \(R_2, R_3\), given by eq 38 to arrive at Stage (3) at the state given by eq 39. Thus, Figures 7 and 8, through the discussion accompanying Figure 3, provide a detailed analogue to the Shor algorithm in Figure 2 by way of an abstract formalism presented in Section 3.

6. CONCLUSIONS

In this article, we have developed an abstract formalism of tensor network-based quantum dynamics applicable broadly for all quantum systems, and we discuss how this general idea captures the key signatures within the well-known Shor’s algorithm as well as other well-known quantum algorithms. In essence, for a general bipartite system, unitary evolution encodes within the evolved state a characteristic correlation or entanglement that bears the signature of the Hamiltonian that defines the evolution operator. Thus, once we interrogate a specific part of a quantum system, the remaining parts of that system automatically get projected based on the extent of correlation that resides within the system, and this is borne out of measurement.

This close resemblance of the general formalism of quantum propagation of multipartite systems and the symmetry of integers as captured by Shor’s algorithm begs the rather profound question of whether natural systems exist or can be designed that may perform the operations that we might interpret as prime-factoring. We do not dwell on this general question in this article, but we ask the opposite question of whether we can exploit this connection to analyze quantum chemical dynamics. Indeed, we find that when we apply this concept along with the generalized definition of phase kickback as obtained from the tensor network description of Shor’s algorithm and quantum dynamics described here, the coupled dynamics of protons in a protonated water wire naturally lends itself to a projected transport interpretation. We find that projection of states at one end of a water wire, along with phase kickback-like constrained operations along the length of the water chain, provide a general description for proton transport that is commensurate with our description of Shor’s algorithm. Future publications are currently planned to exploit this analogy to probe electron–nuclear dynamics in the nonadiabatic setting. Furthermore, given that the surrounding vibrational degrees of freedom may be projected through phase kickback, as noted here, to tailor the dynamics within one mode (\(R_4\) for the protonated pentamer), perhaps chemical catalysis is another area where these broad ideas may find application. These aspects will be probed in future work.

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Notes

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