Mapping Quantum Chemical Dynamics Problems to Spin-Lattice Simulators

Debadrita Saha, Srinivasan S. Iyengar, Philip Richerme,*, Jeremy M. Smith, and Amr Sabry

ABSTRACT: The accurate computational determination of chemical, materials, biological, and atmospheric properties has a critical impact on a wide range of health and environmental problems, but is deeply limited by the computational scaling of quantum mechanical methods. The complexity of quantum chemical studies arises from the steep algebraic scaling of electron correlation methods and the exponential scaling in studying nuclear dynamics and molecular flexibility. To date, efforts to apply quantum hardware to such quantum chemistry problems have focused primarily on electron correlation. Here, we provide a framework that allows for the solution of quantum chemical nuclear dynamics by mapping these to quantum spin-lattice simulators. Using the example case of a short-strong hydrogen-bonded system, we construct the Hamiltonian for the nuclear degrees of freedom on a single Born–Oppenheimer surface and show how it can be transformed to a generalized Ising model Hamiltonian. We then demonstrate a method to determine the local fields and spin–spin couplings needed to identically match the molecular and spin-lattice Hamiltonians. We describe a protocol to determine the on-site and intersite coupling parameters of this Ising Hamiltonian from the Born–Oppenheimer potential and nuclear kinetic energy operator. Our approach represents a paradigm shift in the methods used to study quantum nuclear dynamics, opening the possibility to solve both electronic structure and nuclear dynamics problems using quantum computing systems.

I. INTRODUCTION

The quantum mechanical treatment of electrons and nuclei is critical for a wide range of problems that are of significance to biological, materials, and atmospheric studies. For example, hydrogen transfer processes are ubiquitous in reactions critical to human health, alternative energy sources, food security, and environmental remediation. Yet, the detailed treatment of such problems is confounded by the presence of nontrivial quantum nuclear effects, such as hydrogen tunneling, coupled with electron correlation. For the study of electron correlation in most molecular systems, several powerful approximations have been developed for classical computing platforms, and these are known to provide significant speedups compared to exponentially scaling full-configuration interaction calculations. Indeed, chemical accuracy may be obtained for many systems using the well-known CCSD(T) method that has an associated scaling cost of $O(N^6)$, where $N$ represents the number of electrons.

More recently, algorithms to solve electron correlation problems in small molecular systems have been implemented on quantum hardware devices using trapped atomic ions, photons, nuclear spins, quantum dots, Rydberg atoms, and superconducting circuits. The mapping of most electron correlation problems to quantum hardware is facilitated by the Jordan–Wigner, parity, or Bravyi–Kitaev transformations, where a product of Fermionic creation and annihilation operators are transformed to a chain of Pauli spin operators. In contrast, the intrinsic spin statistics of quantum nuclear dynamics problems, arising from the permutation symmetries of the wavefunctions that describe the constituent nuclear degrees of freedom, do not play a role under conditions prevalent in biological, materials, and atmospheric systems, such as hydrogen transfer reactions under ambient conditions. As a result, most such quantum dynamics studies are currently constructed on classical computing platforms using basis sets and on grids. Furthermore, many of these problems are known to display anomalous nuclear quantum effects that are challenging to study on classical hardware due to the exponentially scaling computational cost of quantum dynamics with increasing degrees of freedom. Unlike several recent
The primary goal of this paper is to develop a set of mapping protocols to allow the study of quantum nuclear dynamics problems on quantum hardware that do not require considering spin statistics. We provide and analyze an approximate algorithm to map exponentially scaling quantum nuclear dynamics problems on a single Born–Oppenheimer surface, onto a general class of Ising model Hamiltonians. Such Ising-type Hamiltonians may be implemented on a range of quantum computing platforms, such as ion traps, superconducting coils, Bosonic processors with photons, solid-state devices and quantum dots inside cavities, and Rydberg atoms. Since quantum nuclear dynamics problems under ambient conditions do not need to be encoded using a set of Fermionic or Bosonic operators, we do not write the Ising model and molecular Hamiltonians in their respective second quantized forms. Instead, we first probe the structure of the Ising Hamiltonian matrix in its exponential scaling space of spin basis vectors. This exponential space is admittedly intractable. Yet, our analysis of the Ising Hamiltonian matrix reveals an intrinsic structure where specific blocks appear within the Ising Hamiltonian matrix, and the corresponding matrix elements are only controlled by a subset of the externally controlled field parameters that dictate the dynamics of the model. To the best of our knowledge, such a structure has never been noted, or exploited, before in the literature. This structure allows us to characterize the general class of problems that may be “computable” using such hardware systems, and in this paper, we further inspect the extent to which quantum chemical dynamics studies may be conducted on such systems, when the statistics of particle permutation need not be included.

The most significant features of the mapping algorithm are summarized in Figure 1, with a more detailed illustration provided in Figure 2. An example of a quantum nuclear problem is shown in Figure 2a and also in Figure 1 (left). Here, we depict a system containing a short-strong hydrogen bond with anharmonic vibrational behavior along the donor–acceptor axis. This problem is prototypical and is representative of a broad range of systems that occur during hydrogen transfer reactions and in hydrogen-bonded systems. We precompute the Born–Oppenheimer potential using electronic structure calculations and obtain a discrete version of the quantum nuclear Hamiltonian. To map this Hamiltonian onto a spin-lattice Ising-type model, the key insights in this paper are as follows: (i) A projected subspace of a specific unitary transformation of the diagonal elements of the quantum nuclear Hamiltonian (related to the Born–Oppenheimer potential) maps to and defines the local magnetic fields applied on each lattice site of an Ising model Hamiltonian. (ii) A similarly projected subspace of a related unitary transformation of the off-diagonal elements of the quantum nuclear Hamiltonian (related to the nuclear kinetic energy operator) defines is mapped onto the intersite coupling terms in the Ising model. These features of our map are highlighted in Figure 1. Importantly, we do not use a quantum circuit model. Instead, the matrix elements of the nuclear Hamiltonian that describe the molecular dynamics, inform the choice of local magnetic fields applied on each lattice site and laser pulse intensities that dictate the intersite coupling, and govern the dynamics of the ion-trap quantum computing platform. In this manner, we provide a direct map of the two quantum systems. Thus, we take a critical step toward solving quantum nuclear dynamics problems, and more generally problems that may not obey Fermi statistics, by mapping them to Ising-type Hamiltonians realizable on ion-trap quantum hardware.

This paper is organized as follows: In Section II, we inspect the block structure of the Ising Hamiltonian, which informs the general class of problems that may be computable on hardware architectures used to realize such Ising-type Hamiltonians. Following this, we then introduce the quantum nuclear Hamiltonian matrix on a single Born–Oppenheimer surface in Section III and a class of Givens rotation-based matrix transformations in Section III.I to represent the quantum nuclear Hamiltonian matrix in a form that is commensurate with the transformed form of the Ising model Hamiltonian in Section II. This transformation leads to our approximate mapping protocol that is outlined in Section IV. Numerical results for the anharmonic molecular vibrations of the shared proton in a symmetric short-strong hydrogen-bonded system are provided in Section V. These include explicit numerical propagation of both the molecular dynamics problem and the spin-lattice dynamics governed by Ising-type Hamiltonian, where the Ising Hamiltonian parameters are chosen based on the mapping protocol in Section IV. The results match exactly for the case of three qubits, and error estimates beyond three qubits are given in Section IV. Conclusions are given in Section VI. Technical aspects of the discussion are further supported through a set of appendices.

II. BLOCK STRUCTURE OF ISING-TYPE HAMILTONIAN MATRICES OBTAINED FROM APPROPRIATE CLASSIFICATION OF THE COMPUTATIONAL BASIS

Ising-type Hamiltonians can be implemented on a range of available quantum computing platforms, which makes these one of the most commonly used quantum computing models today. However, for specificity, we will illustrate our mapping protocols for ion-trap-based quantum architectures, where ions form defect-free arrangements and can support quantum coherence times longer than 10 min. Interactions between ions map to interactions between effective quantum spin states and quantum-harmonic-oscillator bath states—each of which can be precisely controlled and programmed using laser light. Site-resolved detection of
each ion’s spin state can be achieved with near-unit fidelity.\(^{57}\) These features have made trapped ions the leading platform for establishing atomic frequency standards\(^{38-39}\) and one of the leading candidates for performing quantum simulations and quantum computations on such interacting spin systems.\(^{39,65,66}\)

For ion-trap quantum hardware, the generalized Ising Hamiltonian is represented by a spin-lattice of qubits, where (a) the energy gap between the states at each qubit, \(\gamma\), and their relative orientations, are controlled by local effective magnetic fields, \(\{B_i^x, B_i^y, B_i^z\}\), and (b) the spin–spin coupling between different lattice sites, \(i\) and \(j\), is controlled using laser pulses, also spatially nonisotropic, and represented as \(\{f^x_{ij}, f^y_{ij}, f^z_{ij}\}\). Thus, the most general Hamiltonian achievable within the ion trap quantum hardware at low temperatures is

\[
\mathcal{H}_{IT} = \sum_{\gamma} \sum_{i=1}^{N-1} \sum_{j=i+1}^{N} J_{\gamma}^{ij} \sigma_i^\gamma \sigma_j^\gamma + \sum_{\gamma} \sum_{i=1}^{N} B_i^\gamma \sigma_i^\gamma
\]

where \(\gamma \in \{x, y, z\}\) and \(N\) is the number of qubits (or ion sites). The quantities \(\{\sigma_i^\gamma\}\) are the Pauli spin operators acting on the \(i\)th lattice site along the \(\gamma\)-direction of the Bloch sphere. It is critical to note that the expression above is more general than that commonly used in condensed matter physics and quantum chemistry, where only nearest-neighbor interactions are considered. In fact, the set of programmable Ising-type Hamiltonians on an ion trap quantum computer depicts a complete graph that connects all qubits in a spin-lattice system with programmable interactions.\(^{55}\) This aspect is completely captured in eq 1. Furthermore, a critical aspect that differentiates a quantum computer evolving according to eq 1 from a quantum simulator is one where the quantum computer can precisely address and control all \(\{f^\gamma_{ij}\}\) values, whereas a quantum simulator programs a single functional dependence involving a \(\{f^\gamma_{ij}\}\), thus resulting in a reduced set of controllable parameters.\(^{39,65,66}\)

In this paper, we map the Born–Oppenheimer nuclear Hamiltonian to eq 1, thus allowing the two quantum systems to undergo analogous quantum dynamics. Toward this, the parameters \(\{B_i^\gamma, f^\gamma_{ij}\}\) are “programmed” as per the elements of the classically determined Born–Oppenheimer Hamiltonian matrix. To arrive at such a map, we first examine the intrinsic symmetries that are present within such generalized Ising Hamiltonians.

The ion-trap Hamiltonian, \(\mathcal{H}_{IT}\), is naturally represented in a basis of \(2^N\) spin states, where, for example, \(\{|\uparrow\uparrow\rangle, |\uparrow\downarrow\rangle, |\downarrow\uparrow\rangle, |\downarrow\downarrow\rangle\}\) form a basis for a two-qubit system. These now provide us with a “computational basis” with programmable handles, \(\{B_i^\gamma, f^\gamma_{ij}\}\). To gauge the set of mappable problems, we introduce a general set of permutations on the computational basis vectors to reveal a novel block structure of the Ising Hamiltonian matrix. Specifically, the \(2^N\) spin states are partitioned into two sets that are created by the span of even and odd total spin raising operators. Toward this, the basis vectors created using an even number of lattice-site spin raising operators \(\{S_i^+\}\) acting on the full-down spin state, \(|2^N-1\rangle = |\downarrow\downarrow\cdots\rangle\) yield the set \(\{|\downarrow\cdots\rangle; S_i^+ S_j^+ |\downarrow\cdots\rangle; S_i^+ S_j^+ S_k^+ |\downarrow\cdots\rangle; \cdots\}\) that is grouped as part of one block of the ion-trap Hamiltonian. See the set of vectors in Figure 3b, and the bottom left row of Figure 4, where this idea is illustrated for a three-qubit system. For the notation in this paper, we have used the binary representation \(|11\cdots\rangle\) for spin state \(|\downarrow\cdots\rangle\) and the corresponding integer representation \(|2^N-1\rangle\) obtained from the bit-sequence encoded in \(|11\cdots\rangle\).

Similarly, the states obtained using an odd number of raising operators \(\{S_i^+ |\downarrow\cdots\rangle; S_i^+ S_j^+ S_k^+ |\downarrow\cdots\rangle; \cdots\}\) are grouped into a second block and are shown in Figure 3a and on the bottom right row of Figure 4.

---

**Figure 2.** Algorithm that converts the Born–Oppenheimer potential surface and kinetic energy terms in a quantum nuclear problem to a set of controllable parameters and facilitates the dynamical evolution of quantum states in an ion trap. Box (a) shows the Born–Oppenheimer potential and kinetic energies for a short-strong hydrogen-bonded system. This system Hamiltonian is mapped onto an ion trap quantum simulator shown in box (b). Discrete representation of the nuclear Hamiltonian and appropriate rotations yield ion-trap parameters \(\{\{B_i; f^x_{ij}, f^y_{ij}, f^z_{ij}\}\}\) to determine the Ising model used to control the dynamics of lattice spin states. Also see Figure 1.
Thus, the two sets independently span \( S^{n}_{\uparrow \downarrow \cdots} \) and \( S^{n}_{\downarrow \downarrow \cdots} \), where \( S^{n} \) is the total spin raising operator. When the spin basis vectors are partitioned in this manner, the Ising Hamiltonian in eq 1 separates into the block structure that is illustrated in Figure 4 for a three-qubit system. Specifically, the matrix that determines the time evolution of the hardware system separates into two diagonal blocks that can only be coupled by turning on \( \{ B_{i}^{x}; B_{j}^{x} \} \), and this is shown in Figure 4 as part of the gray square. Thus, eliminating these \( \{ B_{i}^{x}; B_{j}^{x} \} \) fields would yield two separate diagonal blocks allowing the treatment of systems that may have a similar block structure. Similarly, the off-diagonal matrix elements within each diagonal block are determined by the laser field parameters, \( \{ f_{i}^{x}; f_{j}^{x} \} \). While the structure derived here is completely general, it is illustrated in Figure 4 for a three-qubit system. The diagonal elements of the matrix, not shown in Figure 4 to maintain clarity, contain linear combinations of \( \{ B_{i}^{x}; f_{j}^{x} \} \).

For a larger number of qubits, the block structure has a recursive form, and this aspect is further elaborated in Appendix A. This block form of the Ising-type Hamiltonian and the associated structure in Figure 4 are a significant general result in this paper, and as we find below, this analysis is critical toward mapping arbitrary problems.

### III. GRID-BASED QUANTUM NUCLEAR HAMILTONIAN COMPUTED ON CLASSICAL HARDWARE

The quantum nuclear Hamiltonian for the molecular system, \( \mathcal{H}^{\text{Mol}} \), is constructed on classical hardware, for the purpose of this paper. In the coordinate representation with basis elements, \( \{ |x\rangle \} \), the Hamiltonian matrix elements are given by

\[
\langle x | \mathcal{H}^{\text{Mol}} | x' \rangle = K(x, x') + V(x)\delta(x - x')
\]  

For local potentials, the potential energy operator, \( \hat{V} \), is diagonal in the coordinate representation. The expression in

Figure 3. 2\(^N\) spin (computational) basis states partitioned into (a) the span of odd powers of the total spin raising operator acting on the down-spin state \( \{ S^{2n-1} \downarrow \downarrow \cdots \} \) and (b) the span of even powers of the total spin raising operator acting on the down-spin state \( \{ S^{2n} \downarrow \downarrow \cdots \} \). This partitioning is illustrated here for the case of three qubits and leads to a block form of \( H_{IT} \) as illustrated in Figure 4.

Figure 4. Upper triangular part (excluding the diagonal) of the Ising Hamiltonian, \( H_{IT} \) (eq 1), in the permuted computational basis illustrated for a three-qubit system. Spin (computational) basis state kets, along with their corresponding binary and integer representations, are presented at the base of the figure. These states are partitioned into odd, \( \{ S^{2n-1} \downarrow \downarrow \cdots \} \), and even, \( \{ S^{2n} \downarrow \downarrow \cdots \} \), spans of the total spin raising operators. The interaction between any two states, \( |i\rangle \) and \( |j\rangle \), is the \( ij^{th} \) matrix element of the ion trap Hamiltonian. For example, \( \langle 11 \uparrow | H_{IT} | 11 \downarrow \rangle \equiv | J_{1}^{+} - J_{2}^{-} |. \) The off-diagonal block that couples the vectors obtained from the odd, \( \{ S^{2n-1} \downarrow \downarrow \cdots \} \), and even, \( \{ S^{2n} \downarrow \downarrow \cdots \} \), spans of the total spin raising operators are marked by the gray square.
eq 2 yields a continuous representation, and in practice, the matrix and the corresponding vectors that the matrix acts on are represented on a discretized Cartesian grid. In this case, the Dirac delta function above is replaced by the Kronecker delta. The potential energy in the above equation is obtained from electronic structure calculations that may also be performed on quantum hardware, independently, in future.

The kinetic energy operator may be approximated in a number of ways. One approach is to recognize that this operator is diagonal in the momentum representation, and hence fast Fourier transforms are commonly employed. In this paper, we employ an analytic banded Toeplitz distributed approximating functional (DAF)68,69 representation for the grid representation of the kinetic energy operator in eq 2

\[ K(x, x') = K(|x - x'|) = \frac{-\hbar^2}{4m\sigma^2\sqrt{2\pi}} \exp\left(-\frac{(x - x')^2}{2\sigma^2}\right) \sum_{n=0}^{M_{\text{DAF}}/2} \left(-\frac{1}{4}\right) \frac{1}{n!} H_{2n+1}\left(x - x'\sqrt{2\sigma}\right) \]

(3)

The banded Toeplitz representation of the DAF approximation for the kinetic energy operator, where the property of its matrix elements, \( K_{ij} \equiv K(|i - j|) \), has a critical role in reducing the nuclear Hamiltonian to the form of \( \mathcal{H}_T \), depicted in Figure 4. This is further elaborated in the following section. In eq 3, \( H_{2n+1}\left|x - x'\sqrt{2\sigma}\right> \) are the even-order Hermite polynomials that only depend on the spread separating the grid basis vectors, \( |x\rangle \) and \( |x'\rangle \), and \( M_{\text{DAF}} \) and \( \sigma \) are parameters that together determine the accuracy and efficiency of the resultant approximate kinetic energy operator. Appendix D provides a brief summary of the DAF approach for approximating a function in general.

III.I. Unitary Transformations That Yield the Block Structure of the Nuclear Hamiltonian, for Symmetric Potentials, to Make These Commensurate with and Mappable to the Spin-Lattice Hamiltonian, \( \mathcal{H}_T \). The nuclear Hamiltonian, \( \mathcal{H}^{\text{Nuc}} \) from eq 2, has a banded Toeplitz structure due to the kinetic energy being expressed in terms of DAFs. In general, the Hamiltonian in eq 2 represents a multidimensional quantum dynamics problem, where the number of dimensions directly corresponds to the number of nuclear degrees of freedom. In this paper, we examine the map between the Hamiltonian in eq 2 for symmetric one-dimensional potentials and the Ising model Hamiltonian discussed in Section II. Routes from here to unsymmetric potentials and to problems in higher dimensions will be considered as part of future publications. In the proton transfer problem considered here, the one-dimensional potential energy surface along the hydrogen transfer axis, \( V(x) \) in eq 2, is a symmetric double well owing to the isoeNERgetic donor and acceptor sites arising from the symmetry of the system (Figure 2a). We exploit the symmetric structure of the potential and the Toeplitz structure of the kinetic energy operator to construct a unitary transformation that block-diagonalizes the nuclear Hamiltonian.

The unitary transform that leads to the block diagonalization of the nuclear Hamiltonian, similar to the structure of the Ising Hamiltonian, can be expressed as a product of Givens rotations. The effect of the Givens rotations on the grid basis states is to create superposition states of the symmetric grid basis states. To explain this, we introduce a uniform one-dimensional set of \( 2^N \) grid points, \( \{|\vec{k}\rangle\} \), such that the Givens-transformed grid basis, \( \{|\vec{\lambda}\rangle\} \), may be represented as

\[ |\vec{k}\rangle \equiv \frac{1}{\sqrt{2}} \left(|\vec{k}_+\rangle + |\vec{k}_-\rangle\right), \quad 0 \leq i < (n + 1)/2 \]

(4)

\[ \equiv \frac{1}{\sqrt{2}} \left(|\vec{k}_+\rangle - |\vec{k}_-\rangle\right), \quad (n + 1)/2 \leq i \leq n \]

(5)

where \( n = 2^N - 1 \). The grid basis and the Givens-transformed grid basis, for a three-qubit system, are represented on the left columns of Figure 5. Equations 4 and 5 form two mutually orthogonal subspaces and are represented in the top and bottom portions of Figure 5, separated by the dashed line. These subspaces diagonalize the nuclear Hamiltonian for symmetric potentials. This process is illustrated for a three-qubit system (2³-grid points) in Figure 6.

The \( i^{th} \) matrix element of the resultant molecular Hamiltonian in the Givens-transformed grid basis is explicitly written as

\[ \mathcal{H}_d^{\text{Mol}} = \frac{1}{2} (\mathcal{H}_{i,i}^{\text{Mol}} + \alpha_i \mathcal{H}_{i,i-1}^{\text{Mol}} + \alpha_i \mathcal{H}_{i,i+1}^{\text{Mol}} + \alpha_i^2 \mathcal{H}_{i,i+1}^{\text{Mol}}) \]

(6)

where \( \alpha_i = \text{sgn}[i - (n/2)] \). The elements of the diagonal blocks of \( \mathcal{H}_d^{\text{Mol}} \) (matrix on the right in Figure 6) are obtained from eq 6 as
The elements of the unitary transform, $\alpha_i$, are, in fact, the characters of the $C_i$ point group. The right-hand side of the above equation, therefore, represents a symmetry-adapted transformation of the nuclear Hamiltonian, and the term $(1/2)[V(x_i) + V(x_{n-i})]$ symmetrizes the potential energy surface in one dimension. By extension, for the elements of the off-diagonal blocks of $\hat{H}_d^{\text{Mol}}$ in Figure 6, $\alpha_i = -\alpha_i$, and

$$\hat{H}_d^{\text{Mol}} = \frac{1}{2}(\hat{H}_l^{\text{Mol}} + \alpha_l\hat{H}_{l,n-l}^{\text{Mol}} + \alpha_{l,n-l}\hat{H}_{n,l}^{\text{Mol}} + \hat{H}_{n,n-l}^{\text{Mol}})$$

$$= [K(x_i, x_i) + \alpha K(x_i, x_{n-i})] + \frac{1}{2}[V(x_i) + V(x_{n-i})]\delta_{i,l}$$

(7)

The elements of the unitary transform, $\alpha_i$, are, in fact, the characters of the $C_i$ point group. The right-hand side of the above equation, therefore, represents a symmetry-adapted transformation of the nuclear Hamiltonian, and the term $(1/2)[V(x_i) + V(x_{n-i})]$ symmetrizes the potential energy surface in one dimension. By extension, for the elements of the off-diagonal blocks of $\hat{H}_d^{\text{Mol}}$ in Figure 6, $\alpha_i = -\alpha_i$, and

$$\hat{H}_d^{\text{Mol}} = \frac{1}{2}(\hat{H}_l^{\text{Mol}} - \alpha_l\hat{H}_{l,n-l}^{\text{Mol}} + \alpha_{l,n-l}\hat{H}_{n,l}^{\text{Mol}} - \hat{H}_{n,n-l}^{\text{Mol}})$$

$$= \frac{1}{2}[V(x_i) - V(x_{n-i})]\delta_{i,l}$$

(8)

where the kinetic energy contribution is identically zero purely due to the Toeplitz nature of eq 3, and only the antisymmetric portion of the potential, $(1/2)[V(x_i) - V(x_{n-i})]$, contributes to the antidiagonal part of $\hat{H}_d^{\text{Mol}}$. Thus, for symmetric potentials such as those considered here, eq 8 is identically zero. This observation will become useful when we generalize the approach presented here, first to general potentials and then to problems of higher dimensionality in future publications.

IV. MAPPING PROTOCOL FOR QUANTUM CHEMICAL DYNAMICS

The structure of the ion-trap Hamiltonian constrains the class of mappable problems. These constraints dictate the accuracy with which quantum chemical dynamics simulations can be performed on an ion-trap system given by eq 1. To summarize our discussion thus far (see Figure 5), we began with a computational basis $|\lambda\rangle$ used to describe the Ising Hamiltonian, $\hat{H}_{IT}$ in eq 1, and the grid basis $|\xi\rangle$ used to represent the quantum nuclear Hamiltonian, $\hat{H}^{\text{Mol}}$ in eq 2. In the interest of matching the structures of the two Hamiltonians, we first obtained a permuted computational basis: $|\lambda\rangle \rightarrow |\tilde{\lambda}\rangle$ (Section II, and also summarized on the right side of Figure 5) and a unitary (Givens) transformed quantum nuclear basis: $|\xi\rangle \rightarrow |\tilde{\xi}\rangle$ (Section III.I). In doing so, our goal becomes

$$\langle \tilde{\lambda}|\hat{H}^{\text{Mol}}|\tilde{\xi}\rangle \leftrightarrow \langle \tilde{\lambda}|\hat{H}_{IT}|\tilde{\xi}\rangle$$

(9)

where we first introduce a map between the transformed quantum nuclear wavefunction bases and the permuted computational bases that represent the Ising spin-lattice system as

$$|\tilde{\xi}\rangle \leftrightarrow |\tilde{\lambda}\rangle$$

(10)

This map is illustrated within the central box in Figure 5, where the left side of the central box represents the Givens-transformed grid basis and the right side represents the permuted computational basis. Furthermore, the mapped basis states are separated into blocks by a dashed horizontal line. For the molecular Hamiltonian, the coupling across these blocks is identically zero for symmetric potentials, while for the Ising Hamiltonian, the coupling across these blocks is identically zero when the terms $B_j^z$ and $B_j^x$ are eliminated from eq 1 (see Figure 4). The effectiveness of the maps in eqs 9 and 10 will essentially dictate the accuracy to which the dynamics captured within the ion-trap quantum simulator controlled by an Ising Hamiltonian accurately predicts the quantum nuclear dynamics.

In this section, we will show that, due to the structure of the Hamiltonians discussed in the previous sections, the diagonal and off-diagonal elements of each individual diagonal block of mappable Hamiltonians, such as eq 2, are Hadamard-transformed to provide $\{B_j^z, f_j^z\}$ and $\{f_j^x, f_j^x\}$, respectively. As a consequence of the discussion in Section III.I, both Hamiltonians, eqs 1 and 2, by construction, take the form depicted in Figure 2c and the right side of Figure 6, respectively. Our quantum nuclear dynamics test case that will be mapped to the aforementioned Ising Hamiltonian (Section...
V) exploits the block structure discussed above, and we illustrate the map by studying a symmetric hydrogen-bonded system displayed in Figure 2a, where a symmetric double-well potential is also shown. For these cases, as seen from eqs 8 and 7 and the discussion in Section II and eq A4 in Appendix A, the block structure of both the Hamiltonian matrices allows the two blocks of each Hamiltonian matrix (Ising and molecular) to be propagated independently, and, potentially on different quantum simulators, for the Ising Hamiltonian. We exploit this feature to evaluate a separate set of \( \{B_i^z; f_j^x\} \) values, below, for each of the two diagonal blocks of the molecular Hamiltonian, while maintaining \( \{B_i^z; f_j^y\} \) to be identically zero.

### IV.I. Obtaining Ion-Trap Parameters \( \{B_i^z; f_j^x\} \) from the Diagonal Elements of the Molecular Hamiltonian

The diagonal elements of the molecular Hamiltonian are directly mapped to those of the spin-lattice Hamiltonian after invoking the map of the unitary-transformed grid basis \( \{\hat{x}\} \) to the permuted computational basis \( \{\hat{\lambda}\} \). Each diagonal element of the molecular Hamiltonian in the transformed grid representation \( \langle \hat{\lambda} | \hat{H}_{\text{Mol}} | \hat{x} \rangle \) is equivalent to the corresponding element of the ion-trap Hamiltonian \( \langle \hat{\lambda} | \hat{H}_{\text{IT}} | \hat{\lambda} \rangle \) in the permuted computational basis representation. In doing so, the set of on-site and intersite coupling parameters \( \{B_i^z; f_j^x\} \) of the ion-trap that occur along the diagonal of \( \hat{H}_{\text{IT}} \) can be evaluated. The mapping expression between the diagonal elements of the molecular Hamiltonian and the corresponding elements of the ion-trap Hamiltonian may be written as

\[
\langle \hat{\lambda} | \hat{H}_{\text{Mol}} | \hat{x} \rangle \equiv \langle \hat{\lambda} | \hat{H}_{\text{IT}} | \hat{\lambda} \rangle
\]  

(11)

Using eq 7 and eq 1, for the left and right sides of eq 11, we obtain

\[
[K(x_i, x_i) - K(x_i, x_{i-n})] + \frac{1}{2} [V(x_i) + V(x_{i-n})]
\]

\[
= \sum_{j=1}^{N} (-1)^j B_j^z + \sum_{j=1}^{N-1} \sum_{k>j} (-1)^{j+k} f_j^x f_k^x \quad \text{for } i < n/2
\]  

(12)

\[
[K(x_i, x_i) + K(x_i, x_{i-n})] + \frac{1}{2} [V(x_i) + V(x_{i-n})]
\]

\[
= \sum_{j=1}^{N} (-1)^j B_j^z + \sum_{j=1}^{N-1} \sum_{k>j} (-1)^{j+k} f_j^x f_k^x \quad \text{for } i > n/2
\]  

(13)

where \( \oplus \) on the right side denotes the addition modulo 2 and \( \hat{\lambda} \) is the \( j \)th bit of the bit representation of \( |\hat{x}\rangle \) with values 0 or 1 for up- or down-spin, respectively, as shown in Figure 3 and Figure 4.

Our goal is to use the diagonal elements of \( \hat{H}_{\text{Mol}} \) to obtain ion-trap parameters \( \{B_i^z; f_j^x\} \). The expressions needed for this purpose are derived from eqs 12 and 13, and a detailed discussion on this map is provided in Appendix B. In summary, we show in Appendix B that the ion-trap control parameters \( \{B_i^z; f_j^x\} \) are specific Hadamard transforms of \( \langle \hat{\lambda} | \hat{H}_{\text{Mol}} | \hat{x} \rangle \), that is

\[
\langle \hat{\lambda} | \hat{H}_{\text{Mol}} | \hat{x} \rangle \xrightarrow{\text{Hadamard}} \{B_i^z; f_j^x\}
\]  

(14)

This is a particularly key result in this paper, since the Hadamard transforms, like Fourier transforms,\(^6\) are unitary and hence the ion-trap parameters, \( \{B_i^z; f_j^x\} \), are Walsh–Hadamard-transform\(^7\) components of \( \langle \hat{\lambda} | \hat{H}_{\text{Mol}} | \hat{x} \rangle \). However, while this map is general for arbitrary number of qubits, we will also describe in Appendix B that the linear transformations between \( \langle \hat{\lambda} | \hat{H}_{\text{Mol}} | \hat{x} \rangle \) and \( \{B_i^z; f_j^x\} \), in eqs 11, 12 and 13, are rank-deficient and hence error estimates are also presented in Appendix B that apply for arbitrary number of qubits.

Before we conclude this section, we note that the map in eqs 12 and 13 may be used for the potential-free particle in a box case by eliminating the potential leading to

\[
[K(x_i, x_i) - K(x_i, x_{i-n})] = \sum_{j=1}^{N} (-1)^j B_j^z
\]

\[
+ \sum_{j=1}^{N-1} \sum_{k>j} (-1)^{j+k} f_j^x f_k^x \quad \text{for } i < n/2
\]  

(15)

\[
[K(x_i, x_i) + K(x_i, x_{i-n})] = \sum_{j=1}^{N} (-1)^j B_j^z
\]

\[
+ \sum_{j=1}^{N-1} \sum_{k>j} (-1)^{j+k} f_j^x f_k^x \quad \text{for } i > n/2
\]  

(16)

### V. PERFORMANCE OF THE MAPPING PROTOCOL FOR A SYMMETRIC HYDROGEN-BONDED SYSTEM

We examine the map by simulating the quantum dynamics of the molecular system and the ion-trap dynamics, on classical hardware, independently. In doing so, we study the time evolution of the initial wavepacket states prepared in the respective permuted basis representations for the molecular and Ising model Hamiltonians. As stated, the parameters in the Ising Hamiltonian are determined, and thus controlled, by the precomputed matrix elements of the molecular Hamiltonian. The specific intramolecular proton transfer problem considered here is that in the protonated 1,8-bis(dimethylamino) naphthalene (DMANH\(^+\)) system shown in Figure 7a. The DMAN molecule has an extremely large hydrogen bond in proton sponges is attractive from the point of view of both the nature of the short potentially symmetric hydrogen-bond bridges,\(^74–77\) their infrared spectroscopic behavior, and their propensity to occur in common nitrogen activation catalysts.\(^78–79\) Thus, the DMANH\(^+\) system has been frequently studied as a model for short, low-barrier hydrogen bonds that have a role in certain enzyme-catalyzed reactions. In solution, the shared proton delocalization in DMANH\(^+\) is controlled by a low-barrier symmetric double-well potential, with barrier height being influenced by solvent and temperature.\(^80,81\) In fact, the environment and variables such as solvent and temperature influence the donor–acceptor distance fluctuations, thus having a critical role in the quantum mechanical nature of the shared proton. The effect of these donor–acceptor variables is seen in Figure 7b, where we present the shared proton one-dimensional symmetric potentials (red curves on the right side of Figure 7b) for a
V.I. Precomputing the Molecular Hamiltonian ($H_{\text{Mol}}$ in Equation 2) on Classical Hardware. To compute the potential energy surface for the intramolecular proton transfer in the molecular system DMANH$^+$ (Figures 2a and 7a), we locate a symmetric stationary point with the shared proton at the center of the donor–acceptor axis. For the case of DMANH$^+$, this stationary point turns out to be a transition state with one imaginary frequency that is obtained from the eigenstates of the electronic structure Hessian matrix, with the vibrational mode corresponding to the intramolecular proton transfer direction. At this geometry, the shared proton is symmetrically located between the donor and acceptor nitrogen atoms. These calculations are performed using standard electronic structure methods. The level of electronic structure theory used is density functional theory with hybrid functional, B3LYP, and an atom-centered Gaussian basis set containing polarization and diffuse functions on all atoms, that is, 6-311++G(dp). Future work will also include mapping of this Hamiltonian precomputation step onto quantum hardware. A reduced dimensional potential energy surface calculation for one-dimensional proton motion along the donor–acceptor axis is performed at the aforementioned stationary point geometry. This is also done for a set of donor–acceptor distances with significant classical Boltzmann populations as seen in Figure 7b. The potential energy surfaces are obtained on a grid defined along the donor–acceptor axis. We choose $2^N$ number of equally spaced grid points, symmetrically located about the grid center, and perform electronic structure calculations at these points, on a classical computing platform, at the level of theory mentioned above. The molecular Hamiltonian is computed (eq 2) and unitary-transformed to achieve a block structure according to Section IIII.

V.II. Quantum Simulation of Proton Transfer Dynamics. Given the block structure of both molecular and Ising Hamiltonians in the permuted and Givens-transformed basis representations, the initial wavepacket for the ion-trap system is chosen as a coherent linear combination of the spin basis states: $\{01\} + \{10\} / \sqrt{2}$ on a three-qubit system.

Given the block structure of the Ising Hamiltonian with $\{B_0^+, B_0^-\}$ turned off, the components of this initial state, $\{11\}$ and $\{\downarrow\downarrow\}$, are not coupled. Additionally, these states will not couple as might be the case in the presence of $B_3^+ - iB_3^-$ in the off-diagonal blocks: for example, pathways such as $\{11\} \rightarrow |1\downarrow\downarrow\rangle \rightarrow |\downarrow\down\downarrow\rangle \rightarrow |\down\down\down\down\rangle$

will remain unpopulated. Hence, in essence, $|\down\down\down\rangle$ gets propagated as per the unitary evolution corresponding to the top diagonal block of the Ising Hamiltonian and $|\down\down\down\rangle$ as per the bottom block. This critical feature allows us to treat the two separated blocks as arising from two different ion traps with two different sets of $\{B_0^+, B_0^-\}$ parameters. Given the direct map in eq 10 between the permuted computational basis and the Givens-transformed molecular grid basis, the initial wavepacket for the molecular system is to be chosen in an analogous manner to the initial wavepacket of the ion-trap,
which is \( \frac{|\psi_0\rangle + |\psi_1\rangle}{\sqrt{2}} \). This essentially leads to the initial wavepacket for the quantum nuclear dynamics problem as being chosen on one end of the grid, that is, a state localized closer to one of the nitrogen atoms in Figures 2a and 7a. This choice results in the initial nuclear wavepacket being symmetrically located at either end of the Givens-transformed basis (eq 5). The spin-lattice and molecular wavepackets are then independently propagated for each potential obtained for different donor–acceptor separations and compared to gauge accuracy of the quantum simulation.

Given the recursive form of the matrix representation of the Ising Hamiltonian in eq 1, as discussed in Appendix A (see eq A4), the ion-trap hardware initial wavepacket state is directly propagated by the choice of \( \{ B'_i; F_i^a \} \) for arbitrary time segments. In this study, we do not seek experimental validation using a real ion-trap simulator but emulate the time evolution of the ion-trap system according to the Hamiltonian in eq 1 on classical hardware, using the eigenstates of the Ising Hamiltonian in Appendix A. The time-dependent probabilities resulting from the projection of the resultant time-dependent wavepacket on the computational basis, at each interval of time, is shown using dashed lines in Figures 8 and 9 for a donor–acceptor distance of 2.53 Å and for the full set of donor–acceptor distance values in Figure 10. (The donor–acceptor distance of 2.53 Å corresponds to the most stable structure, but as seen from Figure 10, there are several other geometries that are also populated (at 300 K) even from a purely classical Boltzmann estimation.) Similarly, we determine the time evolution of the initial wavepacket for the molecular system using the eigenstates of the transformed Hamiltonian in eq 6, and the resulting probabilities from the projection of the time-dependent wavepacket on the Givens-transformed grid basis \( \{ F'_i \} \) are shown using solid lines in Figures 8–10. The probabilities match exactly, apart from numerical round-off error \((10^{-15})\), for the quantum simulation of the dynamics of the two systems. Clearly, this is also true for much longer time intervals as can be seen in Figure 9. Given the exact match between the spin-lattice dynamics and the quantum chemical dynamics, the features present in ion-trap dynamics must also exist in the chemical dynamics problem. Thus, through the isomorphism constructed above, our algorithm shows the ability to probe any entanglement that may be present in chemical systems.

### VI. CONCLUSIONS AND OUTLOOK

The successful simulation of quantum nuclear dynamics on quantum hardware promises a new paradigm for studying a
broader class of coupled electron nuclear transfer problems. In this study, we provide a general but approximate mapping procedure between a quantum chemical dynamics problem, constructed on a single Born–Oppenheimer surface, and an ion-trap quantum simulator where the dynamics is dictated by a generalized form of the Ising model Hamiltonian. The key step involved in facilitating our map is the partitioning of the coupled qubit space into two zones using only odd or even powers of the total spin raising operators that are used to generate such a coupled qubit space. Once the coupled qubit computational basis set is partitioned in such a way, the Ising model Hamiltonian reduces into a block form, thus allowing the possibility to map all problems that may be written in a similar block form. In some sense, we have also taken here the necessary steps to detail the kinds of general problems that can be solved exactly on a quantum system whose dynamics is dictated by a generalized form of the Ising model Hamiltonian. In this particular paper, though, we consider a symmetric proton transfer problem and then go on to show how such a problem can be mapped to an ion-trap system and

Figure 10. Similar to Figure 8 but for multiple donor–acceptor distances ($d_{DA}$) between the nitrogen atoms for the molecule in Figure 2a. Boltzmann populations ($\rho_B$) are computed at 300 K relative to the population of the configuration used in Figure 8 that has a $d_{DA}$ value of 2.53 Å. The correlated changes in the $|0\rangle$ and $|7\rangle$ projections are clearly facilitated by components along other basis vectors, and these may have a critical role in the reactive process as a function of temperature. The fact that the ion-lattice dynamics displays the same dynamical trends provides an additional probe to complex chemical systems.
also show that the dynamics of the two systems is identical provided the parameters of the ion trap are chosen in concert with that of the molecular system obtained from classical precomputation. We also provide error bounds for this approximate algorithm for arbitrary number of qubits.

General quantum nuclear dynamics problems, however, have unsymmetric potential energy surfaces and are generally performed in higher dimensions. This work will become critical in extending our mapping protocol to general potentials in higher dimensions, as will be considered in Figure 11.

Figure 11. Complements Figure 4. At the base of each figure are the computational basis state kets. The interaction between any two states \( |\lambda_i\rangle \) and \( |\lambda_j\rangle \) can be read off from the graph, by starting at the two states and following the lines to their intersection. The node at the intersection gives the interaction between the two. For example, \( |2\rangle \) and \( |7\rangle \) in (c) have an off-diagonal matrix element of \( [J_{13}, J_{13}] \). The blank nodes are zero and show the block diagonal form of the Ising Hamiltonian when \( \{B_x; B_y\} \) are set to zero.
future publications. In addition, the next set of steps also includes inspection of nuclear wavepacket basis set dependence on the accuracy of the proposed map. Using appropriate basis sets, it may be possible to reduce the number of independent descriptors within the molecular Hamiltonian, thus tailoring the accuracy of the map according to the constraints provided in this paper.

### APPENDIX A: RECURSIVE, BLOCK STRUCTURE OF THE ISING HAMILTONIAN MATRIX

The Ising Hamiltonian matrix, $H_N$, for a spin-lattice system with $N$ qubit sites, when written in the computational basis

$$H_N = \begin{bmatrix} H_N^{D1} & B_N \\ B_N^{T} & H_N^{D2} \end{bmatrix}$$

where $I_N$ denotes an identity matrix of size $2^N$. The quantities $J_{xy,N}$ and $J_{yz,N}$ are matrices that appear in the recursive definition of the diagonal blocks, labeled with superscripts $D_1$ and $D_2$, respectively, and contain intersite coupling of the $N$th spin site with the remaining $N-1$ sites. To arrive at the matrix elements belonging to $|\lambda\rangle J_{xy,N} \langle \lambda'|$ and $|\mu\rangle J_{yz,N} \langle \mu'|$ in the equation above, a bitwise XOR operation is constructed between the corresponding computational bases, $|\lambda\rangle$ and $|\lambda'|$. The XOR operation provides the identity of the spin sites where the computational basis vectors $|\lambda\rangle$ and $|\lambda'|$ differ, that is, when the spin states are flipped between $|\lambda\rangle$ and $|\lambda'|$. When the bases differ at two spin-lattice site locations, $i$ and $j$, the corresponding matrix element of $J_{xy,N}$ or $J_{yz,N}$ is given by $J_{xy,N}^{i,j}$. The phase preceding the $J_{xy,N}^{i,j}$ values results from an XNOR operation on the $i, j$ lattice sites discovered through the XOR operation above.

The terms, $J_{x,N}$ and $J_{y,N}$, in eq A2 are also defined in a similar manner. Both $J_{x,N}$ and $J_{y,N}$ matrices are diagonal in form. Thus, the diagonal elements of $H_N^{D1}$ and $H_N^{D2}$ are incremented by a linear combination of all possible intersite couplings of the $N$th spin site with the remaining $N-1$ sites given by $\sum_{i=1}^{N-1} (-1)^{\hat{\lambda} \cdot \hat{\mu}} J_{xy,N}^{i}$. (Also see eqs 12 and 13.)

As noted in the paper, setting all of the transverse local qubit magnetic fields, $B_N^{x}$, and $B_N^{y}$ values to zero in eq 1 block-diagonalizes eq A2 and may be recursively written as

\[
H_N = \begin{bmatrix} H_N^{D1} & B_N \\ B_N^{T} & H_N^{D2} \end{bmatrix} \begin{bmatrix} 0 & 0 \\ 0 & 0 \end{bmatrix} = \begin{bmatrix} H_N^{D1} & B_N \\ B_N^{T} & H_N^{D2} \end{bmatrix}
\]

At this stage, it is critical to realize that the two blocks in the equation above are completely decoupled and basis vector components that undergo unitary evolution due to the top block are never influenced by elements from the bottom block and vice versa. This is explicitly elaborated in Figure 11 for the case of two and three qubits. This presents us with an additional degree of flexibility for our quantum simulation. We exercise this flexibility here and map separately the top and bottom blocks of the equation above, to two different $N$-qubit ion trap systems controlled by parameters $\{B_N^{x}, J_N^{x}\}$ and $\{B_N^{x}, J_N^{y}\}$ respectively. It is important to note here that while the underlying structure of each block in the Ising Hamiltonian matrix remains the same, two different sets of ion-trap control parameters are used to simulate the top and bottom blocks, respectively, thus providing greater flexibility in simulating real systems. We, therefore, introduce a subtle change in denoting the corresponding Ising model Hamiltonian as $H_N$ and allow the diagonal blocks to be independently determined in the following manner.
where the top block is controlled by parameters \( \{ B_i^z; f_{ij}^y \} \) and the bottom block is controlled by a different set of ion-trap parameters \( \{ B_i^z; f_{ij}^y \} \). The molecular Hamiltonian is mapped to the above form of the Ising model Hamiltonian matrix.

We now illustrate the above form of Ising Hamiltonian for the two- and three-qubit systems. But we note that the aforementioned basis set partitioning and Hamiltonian structure is completely general and applies to all cases. Explicitly written, for the case of two qubits, eq A4 takes the form

\[
\mathcal{H}_2 = \begin{pmatrix}
B_1^z + B_2^z + J_{12}^z & J_{12}^x - J_{12}^y & 0 \\
J_{12}^x - J_{12}^y & -B_1^z - B_2^z & 0 \\
0 & 0 & 0
\end{pmatrix}
\]

\[
\mathcal{H}_3 = \begin{pmatrix}
H_{N-1}^{D1} + B_N^z I_{N-1} + J_{N,N}^z & J_{N,N}^y & 0 & 0 \\
J_{N,N}^y & H_{N-1}^{D2} - B_N^z I_{N-1} + J_{N,N}^y & 0 & 0 \\
0 & 0 & \tilde{H}_{N-1}^{D1} + \tilde{B}_N^z I_{N-1} + \tilde{J}_{N,N}^z & \tilde{J}_{N,N}^y \\
0 & 0 & \tilde{J}_{N,N}^y & \tilde{H}_{N-1}^{D2} - \tilde{B}_N^z I_{N-1} + \tilde{J}_{N,N}^z
\end{pmatrix}
\]

(A4)

where again we have highlighted the distinction between ion-trap simulators that represent the top block, \( \{ B_i^z; f_{ij}^y \} \), and those that control the bottom block, \( \{ \tilde{B}_i^z; \tilde{f}_{ij}^y \} \). The three-qubit Hamiltonian is then recursively obtained from the two-qubit Hamiltonian as prescribed by eq A4 and may be written in compact form as follows

\[
\mathcal{H}_3 = \mathcal{H}_3^{D1} \oplus \mathcal{H}_3^{D2}
\]

(A7a)

Here, \( \mathcal{H}_3^{D1} \) and \( \mathcal{H}_3^{D2} \), as defined for the general case in eq A4, refer to the top and bottom diagonal blocks of the two-qubit Ising Hamiltonian (eq A5) simulated using the ion-trap parameters \( \{ B_i^z; f_{ij}^y \} \) while \( \mathcal{H}_2^{D1} \) and \( \mathcal{H}_2^{D2} \) refer to the top and bottom blocks of the two-qubit Ising Hamiltonian (eq A5) controlled by \( \{ \tilde{B}_i^z; \tilde{f}_{ij}^y \} \). While most of \( \mathcal{H}_3^{D1} \) and \( \mathcal{H}_3^{D2} \) is preserved and appear as diagonal blocks of the two-qubit Hamiltonian, the Nth qubit on-site term \( B_i^z \) and intersite coupling terms with all \( N - 1 \) qubits \( f_{ij}^y \), \( f_{ij}^z \) with appropriate phases are added to each diagonal element. The quantities \( J_{N,N}^y \) and \( J_{N,N}^z \) in the top block and \( \tilde{J}_{N,N}^y \) and \( \tilde{J}_{N,N}^z \) in the bottom block capture the interaction of qubits 1 and 2 with qubit 3 in the form of the intersite coupling terms for the two ion traps, respectively. Explicitly, the three-qubit system Hamiltonian becomes

\[
\mathcal{H}_3 = \mathcal{H}_3^{D1} \oplus \mathcal{H}_3^{D2}
\]

(A7b)
We further clarify that the top block, $\mathbf{H}_{13}^{D_1}$, is controlled by parameters $\{B_i^z, f_i^z\}$, whereas the bottom block, $\mathbf{H}_{33}^{D_1}$, is controlled by $\{B_i^z, f_i^z\}$.

**APPENDIX B: OBTAINING THE ION-TRAP CONTROL PARAMETERS $\{B_i^z, f_i^z\}$ FROM EQUATIONS 12 AND 13**

The goal of eqs 12 and 13 is to compute the values for $\{B_i^z, f_i^z\}$, given the values on the left side of eqs 12 and 13. It must first be noted that both equations are a linear set of equations, but the number of free parameters available on the right side, that is, the number of ion-trap control variables within the set $\{B_i^z, f_i^z\}$, is $N+N(N-1)/2$ or $N(N+1)/2$. In Appendix C, the number of parameters that control the ion-trap is analyzed. On the contrary, given that the two blocks of the molecular Hamiltonian are to be propagated independently, as outlined in Appendix A, eq A4, and also in Section IV, the number of free parameters on the left sides of eqs 12 and 13 are $2^{N-1}-1 = \frac{2^{N-1}-1}{2}$. This quantity is obtained by realizing that each block contains $2^{N-1}$ diagonal elements that arise from the Givens transform of the potential energy surface, and of the nuclear kinetic energy, as seen from eqs 12 and 13. But the dynamics is invariant to any shift to the potential energy surface, and thus, eliminating the average value of the diagonal elements of the molecular Hamiltonian yields a reduction of the number of free parameters in the diagonal elements of the molecular Hamiltonian leading to the quantity $\frac{2^{N-1}-1}{2}$.

Thus, in general, the number of ion-trap control parameters $\{B_i^z, f_i^z\}$, in eqs 12 and 13, is not always equal to the number of independent control variables that arise from the molecular Hamiltonian. In this section, we present the best case solution to the set $\{B_i^z, f_i^z\}$ for an arbitrary $N$, given the diagonal elements of the Givens-transformed molecular Hamiltonian. Furthermore, we are also in a position to provide error bounds for the case of an arbitrary number of qubits as we do later in this section.

To appropriately invert eqs 12 and 13 to determine a suitable set of $\{B_i^z, f_i^z\}$ values, we first rewrite these equations to expose the fact that the ion-trap control parameters $\{B_i^z, f_i^z\}$ are specific Hadamard transforms of $\{\alpha \mathcal{H}^{\text{Mol}} \}$ as stated in eq 14. To achieve this, we treat all parameters in $\{B_i^z, f_i^z\}$ on an equal footing and assign these to a new set of variables $\{B_i^z, f_i^z\} \equiv D_i$, where the index “$T$” runs over all integer values less than or equal to $B_i^z_{\text{min}} = \min\left(N(N+1)/2, \frac{N-1}{2}\right)$.

as is clear from the above discussion. We, thus, write the mapping expression for the diagonal elements, from eqs 12 and 13 as

$$
\begin{align}
K(x_j, x_y) &= K(x_j, x_{y-j}) + \frac{1}{2} [V(x_y) + V(x_{y-j})] \\
&= \sum_{j=1}^{N} (-1)^j B_j^z + \sum_{k>j}^{N-1} (-1)^{k+j} \frac{\lambda_{jk}}{\gamma^{k+j}} \equiv \sum_{i=1}^{N} T_{ij}^z D_i^z
\end{align}
$$

for $i < n/2$ (B2)

and

$$
\begin{align}
K(x_j, x_y) &= K(x_j, x_{y-j}) + \frac{1}{2} [V(x_y) + V(x_{y-j})] \\
&= \sum_{j=1}^{N} (-1)^j B_j^z + \sum_{k>j}^{N-1} (-1)^{k+j} \frac{\lambda_{jk}}{\gamma^{k+j}} \equiv \sum_{i=1}^{N} T_{ij}^z D_i^z
\end{align}
$$

for $i > n/2$ (B3)

where $\{\lambda\}$ corresponds to either of the two sets of permuted computational basis states (see Figure 3) that independently span $\{S_{z}^{\lambda} \mid \lambda \rangle \cdots \rangle$ or $\{S_{-z}^{\lambda} \mid \lambda \rangle \cdots \rangle$ and $T_{ij}$ represents a coefficient matrix for the phase preceding the corresponding $D_i^z$ in the equations above. The upper limit, $B_{\text{max}}$, to the summation in eqs B2 and B3 thus denotes the maximum number of independent parameters in $\{B_i^z, f_i^z\}$ that will be used to encode the diagonal part of $\hat{\mathcal{H}}^{\text{Mol}}$ for a given number of qubits. Thus, the diagonal elements of the transformed Ising Hamiltonian $\langle \alpha \mathcal{H}^{\text{Mol}} \rangle$ encode both the Givens-transformed Born–Oppenheimer potential energy surface, $V(x)$, and the nuclear kinetic energy that appear due to the block diagonalization process, needed to make the two Hamiltonians have the same structure (see eqs 7, 12, and 13). The columns of the transformation matrix on the right side of eqs B2 and B3, with elements, $T_{ij} = \pm 1$, resemble a subset of columns that span an $N - 1$-dimensional Hadamard matrix, which is an $(N - 1)$-th-order tensor product of the standard 2x2 (or one-qubit) Hadamard transform. Thus, the columns of $T$ in eqs B2 and B3 form an orthonormal set and thus the $\{B_i^z, f_i^z\}$ values represent a rotation of the elements in $\langle \alpha \mathcal{H}^{\text{Mol}} \rangle$.

$$
D_i^z = \frac{1}{2^{N-1}} \sum_{(\lambda)} T_{ji} \langle \alpha \mathcal{H}^{\text{Mol}} \rangle
$$

(B4)

Owing to the equivalence of $\langle \alpha \mathcal{H}^{\text{Mol}} \rangle$ and $\langle \alpha \mathcal{H}^{\text{Mol}} \rangle$, as seen in eqs 9 and B4, we use the precomputed diagonal elements of the unitary-transformed molecular Hamiltonian in eq B4 to obtain the on-site parameters of the ion trap. As per
eq 11, \( \langle \tilde{\lambda} | \mathcal{H}_{\text{mol}} | \tilde{\lambda} \rangle \) may be replaced by \( \langle \tilde{x} | \mathcal{H}_{\text{mol}} | \tilde{x} \rangle \) in eq B4 leading to

\[
D^z = \frac{1}{2^{N-1}} \sum_{\{\tilde{x}\} \in \{\tilde{\lambda}\}} \mathbf{T}_{\tilde{\lambda}}(\tilde{x}) \mathcal{H}_{\text{mol}}(\tilde{x})
\]

(B5)

where we tersely assume the summation over \( \{\tilde{\lambda}\} \) to also correspond to the summation over \( \{\tilde{x}\} \) as allowed by the correspondence in eq 10. Furthermore, since the N-qubit-Hadamard transform is unitary, we have simply transposed the elements of the transformation matrix, \( \mathbf{T} \), in writing eq B5. For example, for the three-qubit case studied here, the tensor product of two Hadamard matrices yields

\[
H_z = \frac{1}{2} \begin{pmatrix}
1 & 1 & 1 & 1 \\
1 & -1 & 1 & -1 \\
1 & 1 & -1 & -1 \\
1 & -1 & -1 & 1
\end{pmatrix}
\]

(B6)

Hence, the transformation matrix in eq B5 is obtained from eq B6, by removing the zero-frequency component (first row) leading to

\[
[T]^T = \begin{pmatrix}
1 & -1 & -1 & 1 \\
1 & -1 & 1 & -1 \\
1 & 1 & -1 & -1
\end{pmatrix}
\]

The first row of eq B6 is the average of the diagonal elements of the molecular Hamiltonian and, as noted at the top of this section, the diagonal elements, \( \langle \tilde{x} | \mathcal{H}_{\text{mol}} | \tilde{x} \rangle \), are shifted so that this average value is zero. This uniform shift of the diagonal elements results in a constant shift between the eigenvalues of the molecular Hamiltonian and the Ising Hamiltonian, which results in no change to the dynamics, as is clear from Figures 8–10.

The transformation matrix in eq B5, that is, \( \mathbf{T}^T \), is also illustrated in Figure 12b,c for the 2^4-dimensional sub-blocks of a five-qubit Ising and for the 2^3-dimensional sub-blocks of a three-qubit Ising Hamiltonian in Figure 12a. The dimension \( 2^{N-1} \times \frac{N(N+1)}{2} \) of the \( \mathbf{T} \) matrix is apparent from this figure. The latter dimension of the \( \mathbf{T} \) matrix that depends on the number of independent \( D^z \) values is at most \( \frac{N(N+1)}{2} \) (or \( B^z_{\text{min}} \)) and is found to be 3 for the three-qubit system and 15 for the five-qubit system. While the figure is only presented for three- and five-qubit systems, the transformation is completely general.
ERROR BOUNDS ON MAPPING
\langle \tilde{\sigma} | \mathcal{H}^{\text{Mol}} | \tilde{\epsilon} \rangle \leftrightarrow \langle \tilde{\lambda} | \mathcal{H}_{1T} | \tilde{\lambda} \rangle \) FOR A LARGER NUMBER OF QUBITS

Arising from the above discussion, the error, \( \epsilon \), associated with such a partial Hadamard transform of the diagonal elements of the molecular Hamiltonian, can be obtained from the orthogonal complements of the transformation matrix in the corresponding Hadamard matrix. This can be expressed in a closed form as

\[
e = \frac{1}{2^{N-1}} \sqrt{\left(\langle \mathcal{H}^{\text{Mol}} \rangle_{\text{Diag}} P_T [\mathcal{H}^{\text{Mol}}_{\text{Diag}}]^T \right)}
\]

where \( \mathcal{H}^{\text{Mol}}_{\text{Diag}} \) contains the diagonal elements of \( \mathcal{H}^{\text{Mol}} \), that is, \( \langle \tilde{\sigma} | \mathcal{H}^{\text{Mol}} | \tilde{\epsilon} \rangle \), in the equations above and \( P_T \) is a projector on to the orthogonal complement of transformation matrix \( T \) (depicted in Figure 12 for three and five qubits) as obtained from the Hadamard matrix.

\[
P_T \equiv 2^{N-1} \mathcal{H}^{\otimes (N-1)} \mathcal{H}^{\otimes (N-1)}^T - T T^T
\]

where the quantity \( \mathcal{H}^{\otimes (N-1)} \) is the \((N-1)\)-th order tensor product of the standard \(2 \times 2\) (or one-qubit) Hadamard transform. Thus, cases where the diagonal part of the molecular Hamiltonian is exactly captured within the subspace represented by eq B5 may be exactly modeled using the ion-trap simulator/computer. In all of these cases, the orthogonal complement in eq B8 is identically zero. In the general case, the problem becomes that of suitably representing the diagonal elements of the molecular Hamiltonian within the subspace of Hadamard transforms represented by \( T^T \).

APPENDIX C: NUMBER OF DEGREES OF CONTROL IN THE ISING HAMILTONIAN, EQUATION 1

For a given number of qubits, \( N \), the number of ion-trap handles in eq 1 that control various sectors of the Hamiltonian matrix scale as

\[
\{N + N(N - 1)/2 \} + \{N(N - 1)\} + \{2N\} \rightarrow O(N^2)
\]

(C1)

Here, the first quantity, \( \{N + N(N - 1)/2 \} \), refers to the parameters, \( \{B^j, f_0^j\} \), that control the diagonal elements of the matrix (actively discussed in Appendix B), the second quantity on the left, \( \{N(N - 1)\} \), refers to the parameters, \( \{f_b^j \pm f_0^j\} \), that control the coupling between the basis vectors inside each block, and \( \{2N\} \) refers to the parameters, \( \{B^j \pm B^j\} \), that control the coupling across the sets of basis vectors created using the odd and even raising operators described above. This characterization not only elucidates the degrees of freedom of the Ising model Hamiltonian in eq 1 but also provides the sectored availability of these control parameters.

At this stage, there are two cases that become interesting insofar as mapping to realistic systems is concerned. In the first case, the structure of the Ising Hamiltonian is used as is, including the \( \{B^j \pm B^j\} \) terms, and the number of degrees of freedom is as given above and must match the same for the problem at hand to produce an exact map. For the second case, if the \( \{B^j \pm B^j\} \) handles are eliminated, the system reduces to two separate blocks that may be propagated independently, perhaps even on two different sets of ion-trap architectures arranged in parallel, or Trotterized on one single ion-trap architecture. It is this second case that we consider in this paper as it allows the ability to have different Ising model parameters for the two diagonal blocks, and in this case, the number of ion-trap handles become

\[
2\{N + N(N - 1)/2 + N(N - 1)\}
\]

(C2)

which is, in fact, greater than the number of Ising model handles available in an \((N - 1)\)-qubit system when \(N > 17\). The above discussion also implies that for Hamiltonians containing \(2^N\) independent terms, only approximate computation is possible. In this sense, the current paper takes a first step toward providing the necessary accuracy bounds in Appendix B.

APPENDIX D: THE “DISTRIBUTED APPROXIMATING FUNCTIONAL” (DAF) REPRESENTATION FOR THE KINETIC ENERGY OPERATOR (EQUATION 3)

The DAF approach has been well reviewed in the literature.\(^{68,69}\) But given the significance of the banded Toeplitz form toward the mapping algorithm, where as seen in eq 8, the contributions to the off-diagonal blocks from the kinetic energy go to zero due to the banded Toeplitz representation, we present a brief summary of the DAF representation here. The starting point is to expand the wavepacket at time \(t = 0\) using a local set of symmetric fitting functions, \(a(x - x_i)\), such that

\[
\chi(x; t = 0) = \sum_i \Delta x_i a(x - x_i) \chi(x_i; t = 0)
\]

(D1)

where \(\Delta x_i\) is the grid spacing (not in general uniform). The functions \(a(x - x_i)\) are local fitting functions, the choice for which may, in general, depend upon the point \(x_i\). One of the most common directions at this point is to assume that \(a(x - x_i) \equiv \delta(x - x_i) \equiv (x|x_i)\) is a suitable approximation to the Dirac delta function. Subsequent resolution of the identity in terms of some complete set of basis functions leads to a representation of the wavepacket in that basis. The DAF approximation differs from these approaches by assuming that a suitable local representation\(^{69,84}\) can be directly constructed for \(a(x - x_i)\), that is

\[
a(x - x_i) \equiv a_n(x - x_i; \sigma)
\]

\[
= \sum_n b_n H_n\left(\frac{x - x_i}{\sqrt{2}\sigma}\right) \exp\left[-\frac{(x - x_i)^2}{2\sigma^2}\right]
\]

(D2)

where \(H_n(x)\) are the Hermite polynomials generated from Gaussians according to

\[
H_n(y) \exp[-y] = (-1)^n \frac{d^n}{dy^n} \exp[-y]
\]

(D3)

Note that eq D2 is different from the expression obtained using a standard basis set approximation for \(a(x - x_i)\), wherein the appropriate expression would be
with \( c_n = (\sqrt{\pi} 2^n n!)^{-1} \). Note that eq D4 is separable in \( x \) and \( x_i \) whereas eq D2 only depends on \((x - x_i)\). The local spectral form in eq D2 has many computational advantages not the least of which is the fact that eq D2 yields a banded Toeplitz matrix at any level of approximation. The choice of Hermite functions here is by no means a requirement; it is however a convenient choice. Using the orthogonality of the Hermite functions and the fact that \( a(x - x_i) \) must be symmetric with respect to interchange of \( x \) and \( x_i \) (since it approximates the Dirac delta function), one obtains

\[
\begin{align*}
 b_{2n+1} &= 0 \\
 b_{2n} &= \frac{1}{\sigma \sqrt{2n}} \left( \frac{-1}{4} \right)^{n} n! 
\end{align*}
\]

where we have used the identity\(^{86}\)

\[
\int dy \exp\left(-y^2\right)H_n(y) H_m(y) = \delta_{n,m} 2^n n! \sqrt{\pi} 
\]

thus resulting in eq 3. The variables \( M \) and \( \sigma \) determine the accuracy and width (or computational efficiency), respectively, of the DAF. It has been shown\(^{85,87,88}\) that these parameters are not independent, and for a given value of \( M \), there exists a \( \sigma \) that provides optimal accuracy for the propagation. The accuracy of this method in conjunction with \textit{ab initio} dynamics has been benchmarked in ref 88. For an approximation controlled by choice of parameters \( M \) and \( \sigma \), eq 3 only depends on the quantity \((x - x_i)\), that is, distance between points in the coordinate representation, and goes to zero as this quantity becomes numerically large due to the Gaussian prefactor. This yields a banded matrix approximation to eq 3, for any \( M \) and \( \sigma \). Furthermore, on account of its dependence on \((x - x_i)\), a matrix representation of eq 3 has the property that all diagonal elements of this matrix are equal; similarly, all n-th super (and sub)diagonal elements are the same. Such a matrix is called a Toeplitz matrix. The dependence on \((x - x_i)\) also implies a translational symmetry reminiscent of wavelet theories.

We finally note that the DAF approach differs from other approaches that use Hermite functions to represent the wavepacket\(^{93,94}\) based on Heller’s Gaussian wavepacket formalism.\(^{95}\) Within these formalisms,\(^{93,94}\) a locally harmonic approximation to the potential\(^{95}\) allows the reduction of the time-dependent Schrödinger equation to classical-like equations to propagate the width and center of the Gaussian wavepackets. In our case, no assumption is made on the nature of the potential.

### ACKNOWLEDGMENTS

This research was supported by the National Science Foundation grant OMA-1936353 to S.S.I., P.R., J.M.S., and A.S. The authors are grateful to Dr. Miguel Ángel López Ruiz for valuable comments.

### REFERENCES

1. Weinberg, D. R.; Gagliardi, C. J.; Hull, J. F.; Fecenko Murphy, C.; Kent, C. A.; Westlake, B. C.; Paul, A.; Ess, D. H.; McCafferty, D. G.; Meyer, T. J. Proton-Coupled Electron Transfer. *Chem. Rev.* **2012**, *112*, 4016–4093.

2. Nagel, Z.; Klinman, J. Tunneling and Dynamics in Enzymatic Hydride Transfer. *Chem. Rev.* **2006**, *106*, 3095.

3. Soudackov, A. V.; Hammes-Schiffer, S. Probing Nonadiabaticity in the Proton-Coupled Electron Transfer Reaction Catalyzed by Soybean Lipoygenase. *J. Phys. Chem. Lett.* **2014**, *5*, 3274.

4. Hydrogen-Transfer Reactions, Hynes, J. T.; Klinman, J. P.; Limbach, H.-H.; Schowen, R. L., Eds.; Wiley-VCH: Weinheim, Germany, 2007.

5. Cukier, R. I.; Nocera, D. G. Proton-coupled electron transfer. *Annu. Rev. Phys. Chem.* **1998**, *49*, 337–369.

6. Mayer, J. M. Proton-coupled electron transfer: A Reaction Chemists View. *Annu. Rev. Phys. Chem.* **2004**, *55*, 363–390.

7. Hammes-Schiffer, S. Catalysts by Design: The Power of Theory. *Acc. Chem. Res.* **2017**, *50*, 561–566.

8. Raghavachari, K.; Trucks, G. W.; Pope, J. A.; Head-Gordon, M. A fifth-order perturbation comparison of electron correlation theories. *Chem. Phys. Lett.* **1989**, *157*, 479–483.

9. O’Malley, P. J. J.; Babbush, R.; Kivlichan, I. D.; Romero, J.; McClean, J. R.; Barends, R.; Kelly, J.; Roushan, P.; Tranter, A.; Ding, N.; Campbell, B.; Chen, Y.; Chen, Z.; Chiaro, B.; Dunsworth, A.; Fowler, A. G.; Jeffrey, E.; Lucero, E.; Megrant, A.; Mutus, J. Y.; Neeley, M.; Neill, C.; Quintana, C.; Sank, D.; Vainsencher, A.; Wenner, J.; White, T. C.; Coveney, P. V.; Love, P. J.; Neven, H.; Aspuru-Guzik, A.; Martinis, J. M. Scalable quantum simulation of molecular energies. *Phys. Rev. X* **2016**, *6*, No. 031007.  

10. Kandala, A.; Mezzacapo, A.; Temme, K.; Takita, M.; Brink, M.; Chow, J. M.; Gambetta, J. M. Hardware-efficient variational quantum eigensolver for small molecules and quantum magnets. *Nature 2017*, 549, 242.

11. Xia, R.; Kais, S. Quantum machine learning for electronic structure calculations. *Nat. Commun.* **2018**, *9*, No. 4195.
(89) Grossmann, A.; Morlet, J. Decomposition of Hardy Functions into Square Integrable Wavelets of Constant Shape. *SIAM J. Math. Anal.* 1984, 15, 723−736.

(90) Strang, G. Wavelets and Dilation Equations: A Brief Introduction. *SIAM Rev.* 1989, 31, 614−627.

(91) Daubechies, I.; Heil, C. Ten Lectures on Wavelets. *Comput. Phys.* 1992, 6, 697.

(92) Strang, G.; Nguyen, T. Q. Wavelets and filter banks; SIAM, 1996.

(93) Billing, G. D. Quantum dressed classical mechanics. *J. Chem. Phys.* 2001, 114, 6641−6653.

(94) Ben-Nun, M.; Quenneville, J.; Martinez, T. J. Ab Initio Multiple Spawning: Photochemistry from First Principles Quantum Molecular Dynamics. *J. Phys. Chem. A* 2000, 104, 5161−5175.

(95) Heller, E. J. Time-dependent approach to semi-classical dynamics. *J. Chem. Phys.* 1975, 62, 1544−1555.