The Three-Legged Tree Tensor Networks with $SU(2)$- and molecular point group symmetry

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Abstract

We extend the three-legged tree tensor network state (T3NS) [J. Chem. Theory Comput. 2018, 14, 2026-2033] by including spin and the real abelian point group symmetries. T3NS inter-sperses physical tensors with branching tensors. Physical tensors have one physical index and at most two virtual indices. Branching tensors have up to three virtual indices and no physical index. In this way, T3NS combines the low computational cost of matrix product states and their simplicity for implementing symmetries, with the better entanglement representation of tree tensor networks. By including spin and point group symmetries, more accurate calculations can be obtained with lower computational effort. We illustrate this by presenting calculations on the bis($\mu$-oxo) and $\mu - \eta^2 : \eta^2$ peroxo isomers of $[\text{Cu}_2\text{O}_2]^{2+}$. The used implementation is available on github.

1 Introduction

The discovery of the density matrix renormalization group (DMRG) by S. White in 1992 introduced a new and very successful way of treating strongly correlated quantum systems in both condensed matter physics and theoretical chemistry. Later on, it was discovered that the DMRG wave function corresponds to a variational ansatz over the set of matrix product states (MPS). An MPS is a state that can be represented by a linear chain of tensors. It is the most simple form of a tensor network. The linear form of the MPS explained the high efficiency of DMRG for systems respecting the area law for entanglement in one-dimensional quantum spin systems.

In condensed matter physics, this kick-started the formulation of other (non-linear) tensor networks suitable for systems with another entanglement structure than the one-dimensional area law. The projected entangled pair states (PEPS) and the multiscale entanglement renormalization ansatz (MERA) are some notable examples.

The linear nature of DMRG is also far from ideal for the entanglement structure of most molecules. Hence, other tensor networks have also been studied in quantum chemistry, like the complete-graph tensor network states (CGTNS), the self-adaptive tensor network states (SATNS) and general tree tensor network states (TTNS). However, due to its favorable computational complexity and the ease for exploitation of symmetries, DMRG is still the tensor network method of choice for quantum chemistry.

Recently, we introduced the Three-Legged
Tree Tensor Network (T3NS) is a subclass of the TTNS which allows an efficient optimization of the wave function while still enjoying the better entanglement representation of the TTNS. The tree-shaped network allows a logarithmic growing maximal distance with system size as opposed to the linear maximal distance for MPS. In this paper, we will further extend the study of the T3NS. The implementation of symmetries will be explained and its advantages will be discussed theoretically as well as illustrated by exemplary calculations. More particularly, we will use the real abelian point group symmetries ($C_1, C_i, C_2, C_s, D_2, C_{2v}, C_{2h}$ and $D_{2h}$) and the $SU(2)$-symmetry present in the non relativistic quantum chemical Hamiltonian to obtain more accurate and faster calculations. We impose these global symmetries on the T3NS by using the Wigner-Eckart theorem as introduced by McCulloch for DMRG.

The paper is structured as follows. In section 2, Three-legged Tree Tensor Networks and the advantages of this subset of the general tree tensor network are discussed. In section 3, the handling of symmetries in T3NS is illustrated. We incorporate the parity symmetry $Z_2$, the number conservation $U(1)$, the spin symmetry $SU(2)$ and the real abelian molecular point group symmetries. We heavily rely on graphical depictions for the needed tensor contractions, as it yields a concise and clear notation for otherwise lengthy equations. In section 4, we show some calculations of T3NS with the previously mentioned symmetries. We compare results obtained for $[\text{Cu}_2\text{O}_2]^{2+}$ with the ones presented in the original T3NS paper. Summary and conclusions are provided in section 5.

The used implementation is open source and can be found on Github under the GNU GPLv3 license.

2 Three-Legged Tree Tensor Networks

While the MPS ansatz of DMRG can be represented by a linear chain of tensors, the TTNS ansatz is built by making a tree-shaped network of tensors, making it the most general loop-less tensor network state.

The TTNS ansatz has considerable advantages over the MPS ansatz. It is able to capture algebraically decaying correlation functions as the maximal distance, i.e. the number of bonds, between two sites grows logarithmic in the number of sites. This is contrasted by the linear growth for DMRG. However, due to the branching of the network, the complexity for optimization becomes quickly unfeasible, especially for two-site optimization.

The Three-Legged Tree Tensor Network is a subset of the general TTNS and largely circumvents the high complexity of the algorithm while still maintaining the advantages. In a T3NS, we make use of branching and physical tensors for the wave function ansatz and we intersperse these in the network. Physical tensors are the same as the tensors appearing in an MPS: they have one physical bond and maximally two virtual bonds. Branching tensors have three virtual bonds and no physical bonds. They allow the tensor network to branch. In a T3NS ansatz, branching tensors are never placed next to each other, since this would worsen the complexity for two-site optimization. We also note that a T3NS ansatz
with only physical tensors corresponds with an MPS.

Next to a lower polynomial complexity, the restriction to tensors with three legs simplifies also the implementation of $SU(2)$-symmetry, as will be seen in section 3.

The resource requirements of the T3NS algorithm are presented in table 1. The scaling is explained in appendix A. An example of a T3NS wave function ansatz is given in fig. 1.

Table 1: Resource requirements for DMRG and T3NS for quantum chemistry with renormalized operators. The underlined terms correspond with the complexity of the most intensive part of the algorithm, i.e. the matrix-vector product used in the iterative solver. The number of orbitals (or physical tensors in the network) is denoted by $k$ and the maximal virtual bond dimension is denoted by $D$.

|               | DMRG             | T3NS             |
|---------------|------------------|------------------|
| CPU time:     | $O(k^4 D^2 + k^3 D^3)$ | $O(k^4 D^2 + k^3 D^3)$ |
| Memory:       | $O(k^3 D^2)$     | $O(k^3 D^2 + kD^3)$ |

3 Symmetries in T3NS

When representing the wave function in a tensor network ansatz, the encoding of symmetries into the network can facilitate the calculations. Depending on the particular tensor network ansatz used, the implementation of symmetries can be relatively straightforward or more involved. The usage of symmetries in the T3NS is facilitated by restricting to a maximum of three legs for every tensor in the wave function ansatz. This makes the treatment of both abelian and non-abelian global symmetries very analogous to DMRG, where this is already well studied.

3.1 Labeling the basis states

In the T3NS the index of a virtual (or physical) leg, specifies the different virtual (or physical) basis states traveling through this leg. The three-legged tensors are, due to the Wigner-Eckart theorem, irreducible tensor operators of the total symmetry group. The basis states need to transform according to the rows of the irreducible representations of the symmetry group. Each basis state $|\alpha\rangle$ (or index of a leg) can thus be labeled with the irrep and the row of the irrep according to which it transforms, i.e. $|\alpha\rangle = |\text{irreps and rows, } \alpha\rangle$.

The label $\alpha'$ is needed to discern the different basis states belonging to the same list of irreps and rows of irreps. For a physical bond in non-relativistic quantum chemistry for example we need the local basis states at every orbital. We get

$$
|\uparrow\rangle = |\pi = 1, N_\uparrow = 1, N_\downarrow = 0\rangle \quad (2)
$$

for the parity $\mathbb{Z}_2 (\pi)$, and two $U(1)$-symmetries for both the spin up and down ($N_\uparrow, N_\downarrow$), or

$$
|\uparrow\rangle = |\pi = 1, I = I_0, N = 0, j = 0, m = 0\rangle \quad (5)
$$

$$
|\downarrow\rangle = |\pi = 1, I = I_k, N = 1, j = \frac{1}{2}, m = -\frac{1}{2}\rangle \quad (7)
$$

for $\mathbb{Z}_2 (\pi)$, $U(1) (N)$, $SU(2) (j, m)$ and the real abelian point-group symmetries ($I$). The labels $\pi, N_\uparrow, N_\downarrow, N, j$ and $I$ represent irreps of the different symmetries, while $m$ labels the row of irrep $j$. $I_k$ is the point-group irrep of the orbital. A double occupation results in the trivial point group irrep $I_0$ since $I_k \otimes I_k = I_0$ for real abelian point group symmetries. In this example, $SU(2)$ is the only symmetry that needs an extra label for the row since the other symmetries have one-dimensional representations.

We omitted the label $\alpha'$, since the irreps and rows already uniquely label the local physical basis states. However for the labeling of the virtual basis states the label $\alpha'$ is still needed.
For calculations with fermions we utilize fermionic tensor networks. For every performed permutation or contraction the fermionic signs are calculated by looking at the parities of the different basis states. One could note that labeling the parity $\pi$ is redundant since it already fixed by the total number of particles in the state. However, we chose to keep explicitly track of the parity of the states. In this way, we can separate completely the fermionic sign handling from the particle numbers in the basis states. This allows a more modular implementation of the different symmetries and the fermionic signs.

### 3.2 Reduced tensors

By using the labeling discussed in section 3.1 we can write every three-legged tensor in the wave function ansatz as shown in eq. 9. Due to the Wigner-Eckart theorem, the reducible tensor $A$ can be rewritten as a reduced tensor $\tilde{T}$ multiplied with the Clebsch-Gordan coefficients of the different symmetries as shown in eq. 10. These coefficients are Kronecker deltas for the real abelian symmetries and are the well-known Clebsch-Gordan coefficients for the recoupling of spins for $SU(2)$.

In eq. 11 the latter is replaced by the Wigner-3j symbol using eq. 12. Furthermore, several shorthand notations are introduced. $x_i$ is shorthand notation for the full labeling of irreps and rows of irreps, while $x_i$ is the labeling of only the irreps and not the rows ($m$) of the basis state.

Shorthand notations are also introduced for the Kronecker deltas and for the square root of the multiplicity of $j_i$ ($[j_i] = \sqrt{2j_i + 1}$). A phase is absorbed in the reduced tensor $T$ (hence the transition from $\tilde{T}$ to $T$). We found the calculations particularly simple with this absorbed phase.

Eq. 11 expresses the reducible tensor $A$ in terms of a reduced tensor $T$ independent of the labels of the rows of irreps in $x_i$, and a symmetry tensor. This symmetry tensor consists of the remaining terms in eq. 11. It contains the complete dependency of $A$ on the rows of irreps and is completely independent of the labels $\alpha_i$.

To construct the wave function, the different tensors in the network should be contracted. However, since the symmetry tensors targets all equivalent states in a multiplet at once, the complete wave function should be normalized by dividing by $[j_{\text{final}}]$, with $j_{\text{final}}$ the total spin quantum number of the target state.

The original network of reducible tensors representing the wave function (e.g. as given in fig. 1) now factorizes into two networks with the same shape. One network consists of reduced tensors. It covers the complete dependency of the labels $\alpha_i$ and is also dependent of the labeling of irreps. The other network is built from symmetry tensors. The dependency of the wave function on the labeling of the rows of irreps ($m$) is completely captured by this network. Furthermore, this network is completely independent of the labels $\alpha_i$.

#### 3.2.1 Graphical depiction

![Figure 2: The ansatz for a three-legged tensor with symmetries.](image)

The triangle represents the Wigner-3j symbol and the various Kronecker deltas. The hollow circle represents $(-1)^{j_3 - m_3}$ while the full circle represents $[j_3]$. The direction of the arrows on the legs fixes the sign of $m_i$ in the Wigner sign. An incoming arrow corresponds with $m_i$, while an outgoing arrow is $-m_i$. Such a convention is also needed for a correct treatment of the fermionic signs. An incoming leg for $x_i$ represents $|x_i\rangle$ while an outgoing leg represents $\langle x_i|$.
\[
A = \sum_{\pi_1 I_1 N_1 j_1 m_1 \alpha_1, \pi_2 I_2 N_2 j_2 m_2 \alpha_2, \pi_3 I_3 N_3 j_3 m_3 \alpha_3} A_{\pi_1 I_1 N_1 j_1 m_1 \alpha_1} |\pi_1 I_1 N_1 j_1 m_1 \alpha_1\rangle |\pi_2 I_2 N_2 j_2 m_2 \alpha_2\rangle |\pi_3 I_3 N_3 j_3 m_3 \alpha_3\rangle 
\langle \pi_3 I_3 N_3 j_3 m_3 \alpha_3|\pi_2 I_2 N_2 j_2 m_2 \alpha_2|\pi_1 I_1 N_1 j_1 m_1 \alpha_1| \]  
(9)

\[
= \sum_{x_1 \alpha_1 x_2 \alpha_2 x_3 \alpha_3} \delta_{I_1 I_2 I_3} \delta_{\pi_1 + \pi_2 \mod 2, \pi_3} \delta_{N_1 + N_2, N_3} \langle j_1 m_1 j_2 m_2 j_3 m_3 | \tilde{T}_{x_1 \alpha_1 x_2 \alpha_2 x_3 \alpha_3} | x_1 \alpha_1 \rangle | x_2 \alpha_2 \rangle | x_3 \alpha_3 \rangle. 
\]  
(10)

\[
\langle j_1 m_1 j_2 m_2 | j_3 m_3 \rangle = \left( \begin{array}{ccc} j_1 & j_2 & j_3 \\ m_1 & m_2 & -m_3 \end{array} \right) \sqrt{2j_3 + 1}(-1)^{j_1 - j_2 + m_3}. 
\]  
(11)

The arrow in the triangle is needed to fix the order of the bra and kets in eq. (11). The arrow runs from the first bra or ket to the last bra or ket. Since the tensor is not invariant under permutation of these bras and kets, due to its fermionic character, the fixing of the order in our graphical depiction is needed. The arrow also represents the cyclic invariance of the Wigner-3j symbol. A phase has to be included for the Wigner symbol when changing the direction of the arrow but not when rotating the arrow.

### 3.3 The Chemical Hamiltonian

The non-relativistic quantum chemical Hamiltonian is given by

\[
H = \sum_{ij} t_{ij} \sum_{\sigma} c_{i\sigma}^\dagger c_{j\sigma} + \frac{1}{2} \sum_{ijkl} V_{ijkl} \sum_{\sigma\tau} c_{i\sigma}^\dagger c_{j\tau} c_{l\tau} c_{k\sigma}, 
\]  
(13)

where \(i, j, k\) and \(l\) are indices for the different spatial orbitals and \(\sigma, \tau\) represent the spin degree of freedom (\(\uparrow\) or \(\downarrow\)).

For the four point interactions this separates into the following cases:

\[
V_{ijkl} c_i^\dagger c_j^\dagger c_k c_l, \quad i = j, k = l 
\]  
(14)

\[
V_{ijkl} \left( c_i^\dagger c_j^\dagger c_k c_l + c_j^\dagger c_k^\dagger c_i c_l^\dagger \right), \quad i = j, k < l 
\]  
(15)

\[
V_{ijkl} \left( c_i^\dagger c_j^\dagger c_k c_l + c_j^\dagger c_k^\dagger c_i c_l^\dagger \right), \quad i < j, k = l 
\]  
(16)

\[
V_{ijkl}(c_i^\dagger c_j^\dagger c_k c_l + c_i^\dagger c_j^\dagger c_l c_k^\dagger + c_i^\dagger c_j^\dagger c_k^\dagger c_l^\dagger + c_i^\dagger c_j^\dagger c_k c_l^\dagger) \quad i < j, k < l 
\]  
(17)

where we only used \(V_{ijkl} = V_{jilk}\). The complete Hamiltonian can be constructed by summing these terms for all possible \((i, j, k, l)\)-combinations.

The creation and annihilation operators \(c_{k\sigma}^\dagger\) and \(c_{k\sigma}\) do not yet transform according to the
rows of the $SU(2)$-irreps. As is well known, an additional transformation is given by

\begin{align}
\tilde{c}^\dagger_{k\sigma} &= c^\dagger_{k\sigma} \\
\tilde{c}_{k\sigma} &= (-1)^{\frac{1}{2}+\sigma} c_{k-\sigma}.
\end{align}

In this way, we can again split off the Clebsch-Gordan coefficients into a symmetry tensor. The decomposition of the annihilation operator into a reduced tensor and a symmetry tensor is graphically depicted in fig. 3. One can easily calculate the tensor elements for the reduced tensors in fig. 3. For the creation operators the graphical depiction is completely equivalent.

The most illustrative example is given by eq. 17 since here four different spatial orbitals can be involved. The construction of these terms is graphically shown in fig. 4 where

\begin{equation}
F_J = -[J] (V_{ijkl} + (-1)^J V_{ijk}) .
\end{equation}

The tensor network in fig. 4 should be manipulated to the same geometry of the tensor network of the wave function. For obtaining a T3NS structure out of fig. 4 two types of transformations are needed. First, the order of the operators (e.g. switching $i$ and $k$) can be changed by using properties of the Wigner symbols and taking an appropriate fermionic sign into account. One can also insert identities into the network in fig. 4. These two transformations suffice to change the network in fig. 4 to an arbitrary T3NS network.

Figure 3: Graphical depiction of the annihilation operator on site $k$. The operators are represented by a symmetry tensor and a reduced tensor. $x_k$ and $x_{k'}$ are the local basis states for orbital $k$. The third index $x_{kk'} = (1, I_k, -1, \frac{1}{2}, \sigma)$ serves the purpose of correctly coupling different operators. No bonds here need an extra $\alpha$ label, as stated in section 3.1.

3.4 The optimization

In the previous sections 3.2 and 3.3 we have shown that both the wave function ansatz and the Hamiltonian can be factorized into a reduced tensor network and a symmetry tensor network. In this section, we briefly discuss how this representation can be used during the optimization algorithm of the T3NS.

The optimization of a T3NS occurs in a similar way as for DMRG, i.e. we sweep through the network and optimize only two tensors at a time. During this local optimization of the network, the effect of the Hamiltonian on the other tensors (i.e. the environment) can be efficiently captured by renormalized operators.\cite{32,29,30} The usage of renormalized operators reduces the quartic scaling of the total chemical Hamiltonian as a function of the number of orbitals to a quadratic scaling for the effective Hamiltonian.

These renormalized operators are partial contractions of the Hamiltonian and the T3NS wave function. More specifically, the energy of a wave function $E = \langle \Psi | H | \Psi \rangle$ can be calculated by sandwiching the Hamiltonian terms in their network form between the wave function and its adjoint in the same network form. This triple-layered network of reduced and symmetry tensors can be fully contracted to obtain the energy. However, during the optimization of only two sites of the network, a lot of the triple-layered network can be precontracted and reused since it does not change during this optimization step. The renormalized operators are
Figure 4: Graphical depiction of the terms in eq. 17. The upper gray layer are the reduced tensors and the lower layer are the symmetry tensors. Labels of connected bonds are written fully for clarity. The different operators need to couple to the trivial singlet, since the term in eq. 17 transforms trivially under the used symmetries.

Figure 5: Graphical depiction of a renormalized operator. Both the symmetry tensor and the reduced tensor $O$ are shown. The index $x_1$ ($x_{11}$) originates from the T3NS ansatz of the ket (bra) wave function. The index $x_{11}''$ corresponds with a bond of the network representation of the Hamiltonian. The tensor $O$ is the resulting reduced tensor from partial contraction of the triple-layered network.

After the optimization of the two sites, the sweep algorithm moves on to a neighboring pair of two sites, which has one site in common with the previous pair. These sites are optimized by a new effective Hamiltonian. In order to calculate this effective Hamiltonian through renormalized operators, one can recycle certain of these operators and update others. This is in fact equivalent with the usage of renormalized operators in DMRG for quantum chemistry.

In fig. 6, an example of such an update is given. A site operator (as given in fig. 3) is appended to an already existing renormalized operator. The resulting tensor is then manipulated to another form, for an easier construction of the effective Hamiltonian. In the example, the manipulation of the symmetry tensors gives rise to extra factors. A Wigner-9j symbol arises due to the recoupling of the spin basis states. The phase $(-1)^{\pi_1\pi_2}\pi^\prime_2$ originates from the fermionic character of the tensors.

Wigner symbols are used for the update of renormalized operators and the construction of the effective Hamiltonian when a branching tensor is optimized. However, when only physical tensors are optimized, no Wigner symbols are needed. This is discussed in more depth in appendix B.2. The optimization of two adjacent physical tensors is exactly the same as in DMRG.

As a side remark, the present implementation, omitting branching tensors, can perform a DMRG algorithm which only needs Wigner
symbols during the update of renormalized operators and not during the iterative optimization step.

More details for the construction of the effective Hamiltonian and the updating of the renormalized operators are given in appendix B.

4 Calculations

In this section, we present several calculations with T3NS. The used implementation for the calculations can be found on github. This implementation is able to exploit $\mathbb{Z}_2$, $U(1)$, $SU(2)$ and the real abelian point group symmetries. The symmetries can be included in a modular way. This enables us to compare calculations with and without $SU(2)$ and point group symmetries for the same implementation.

After the optimization of two sites, this two-site object has to be split into two separate sites again. This is done by the Singular Value Decomposition (SVD). The truncation of the virtual bond dimension during this step is done in two different ways.

First, a fixed maximal bond dimension can be imposed. The algorithm will keep as many singular values into consideration as possible. It will first select the largest singular values until no non-zero singular values are left or the maximal bond dimension is reached. The remaining singular values and their corresponding basis states are discarded.

Second, the dynamic block state selection (DBSS) can be used. With this method, the algorithm keeps the largest singular values until a threshold for a cost function is reached. The cost function used in our implementation is given by

$$w_{\text{disc}} = \sum_{i_{\text{disc}}} s_{i_{\text{disc}}}^2$$

i.e. the sum of the squares of all discarded singular values. This corresponds with $\langle \Psi_{\text{disc}} | \Psi_{\text{disc}} \rangle$, with $| \Psi_{\text{disc}} \rangle$ the discarded part of the wave func-
tion during truncation. Other cost functions can be easily implemented.

Next to a targeted threshold, a minimal and maximal bond dimension should be specified. Throwing away too many basis states at a certain stage can impede the optimization at later stages, even though the truncation error is only minimal at that point. Specifying a minimal bond dimension ensures a certain flexibility at all time. The maximal bond dimension is needed to prevent a large increase in both run time and memory usage when the imposed threshold can not be reached.

As noted in section 3.2.2, the usage of symmetries with irreps that are more than one-dimensional, such as $SU(2)$, introduces a compression of the wave function. Different basis states belonging to the same multiplet can be represented by a singular reduced basis state. Analogous to the bond dimension being the number of basis states kept in the bond, the reduced bond dimension is defined as the number of reduced basis states kept. When using $SU(2)$, the reduced bond dimension of the bonds, and not the bond dimension, will reflect the computational complexity.

Just as with DMRG for quantum chemistry, keeping track of the renormalized operators is the most taxing part on memory for T3NS. The amount of renormalized operators needing to be stored for T3NS are of the same order as for DMRG. However, T3NS calculations can be performed on a considerably lower bond dimension for a similar accuracy. Consequently, this lowers the storage requirements for the renormalized operators and allows us to keep all tensors on memory at all time. No checkpoint files need to be written to disk or read from disk during the algorithm for the present system sizes and bond dimensions.

4.1 The Bisoxo and Peroxo Isomer of $[\text{Cu}_2\text{O}_2]^{2+}$

We revisit the bis($\mu$-oxo) and the $\mu = \eta^2 : \eta^2$ peroxy $[\text{Cu}_2\text{O}_2]^{2+}$ isomers as a test case for T3NS with $SU(2)$ and abelian point group symmetries. These transition-metal clusters have been previously studied by other ab initio methods such as the complete active space self-consistent field theory (CASSCF), the complete active space self-consistent field theory with second order perturbation theory (CASPT2), the restricted active space self-consistent field theory with second order perturbation theory (RASPT2) and DMRG. It was the largest system studied in the initial T3NS paper using only $U(1) \otimes U(1)$ symmetry (i.e. conservation of both particle number and spin projection).

We perform calculations for both isomers in an (26e, 44) active space. The same active space is used as in refs. 13,31 Both isomers have a $D_{2h}$ point group symmetry and their ground state is a singlet state in the $A_g$ irrep of $D_{2h}$. When using the $SU(2)$ and/or point group symmetry adapted version of T3NS, states corresponding to these irreps will be targeted. The same T3NS shape and orbital ordering is used as in ref. 13.

4.1.1 The Bisoxo isomer with and without spin symmetry

In order to compare the present spin adapted version of T3NS with its non-adapted predecessor, we perform several calculations for the bisoxo isomer at different bond dimensions and with different symmetries included.

In fig. 7(a), timings for the last sweep are shown for several fixed bond dimensions. Calculations were performed with $U(1) \otimes U(1)$ symmetry and with $SU(2) \otimes U(1)$ symmetry. For both, calculations with and without the $D_{2h}$ point group symmetry are done. For the spin-adapted versions, the bond dimensions shown are the reduced bond dimensions.

As expected, the usage of the point group symmetry introduces a lot of sparsity in the tensors which speeds up calculations considerably. Calculation time improved by a factor of 6 and 13 at $D = 500$ for $U(1) \otimes U(1)$ and $SU(2) \otimes U(1)$ respectively when including $D_{2h}$. For both calculations with or without spin symmetry, the inclusion of $D_{2h}$ yields practically the same energies and maximal truncation errors as when performing the calculation without the point group symmetry.

For tensors of the same size, calculations
including spin symmetry are computationally more intensive than without spin symmetry as can be seen in fig. 7(a). This is expected since the reduced tensors are more dense than the reducible tensors. However, the compressed nature of the reduced tensors (see section 3.2.2) makes spin-adapted calculations at a certain reduced bond dimension more accurate than calculations without spin symmetry at an equal bond dimension. This can be seen in figure 7(b). The maximal unreduced bond dimension during SU(2) calculations is also given in this figure. For the present calculations, the maximal unreduced bond dimension is approximately twice as large as the imposed maximal reduced bond dimension. When comparing wall time with achieved accuracy, the calculations with SU(2) included are considerably faster, as is shown in fig. 7(c).

4.1.2 The bisoxo and peroxo isomers with spin symmetry

Several calculations are performed for both isomers. Both SU(2) and $D_{2h}$ symmetry are used. The inclusion of spin and point group symmetry considerably improves our calculations and allows us to go to much larger bond dimensions. Both a constant maximal bond dimension and DBSS are used. Some obtained results are shown in table 2 alongside previously published results. Calculations at a fixed re-

Figure 7: Results for the bisoxo isomer obtained on an 18-core Intel Xeon Gold 6140 (Skylake at 2.3 GHz). Calculations with a fixed maximal bond dimension of $D = 100, 200, 300, 400$ and $500$ were performed. Results for $U(1) \otimes U(1)$ combined with or without $D_{2h}$ are given by crosses. Results for $U(1) \otimes SU(2)$ combined with or without $D_{2h}$ are given by triangles. The bond dimensions are in this case the reduced bond dimensions, except for (b) where also unreduced bond dimensions are shown. (a) shows the wall time of the last sweep for different bond dimensions and different symmetries. (b) shows the ground state energy for different bond dimensions. For the spin-adapted calculations, both the maximal reduced as unreduced bond dimensions are given. (c) shows wall time of last sweep in function of ground state energy.

Figure 8: Extrapolation of the energy for both the bisoxo and peroxo isomer. Extrapolation is done by using the results for $D = 600, 700, 800, 900$ and $1000$. The DBSS calculations of table 2 targeting a truncation error of $5 \cdot 10^{-6}$ are also shown in the figure. They are however not used for the extrapolation.
Table 2: Energy gaps and ground state energies between the bisoxo and peroxo isomers. The energy gaps are given in kcal/mol. Ground state energies are given in Hartree. For T3NS calculations with fixed bond dimension, the bond dimension is given in the first column. For T3NS calculations using DBSS, the minimal bond dimension and the truncation error is given. For T3NS with SU(2) the given bond dimensions are the reduced ones. Maximum bond dimensions around 2000 were reported for both clusters during the DMRG calculations of ref. 31.

| Method            | $E_{\text{bisoxo}}$ [Hartree] | $E_{\text{peroxo}}$ [Hartree] | $\Delta E$ [kcal/mol] |
|-------------------|-------------------------------|-------------------------------|---------------------|
| DMRG[31]          | -541.53853                   | -541.58114                    | 26.7                |
| T3NS with $U(1) \otimes U(1)$[14] | -541.53820                   | -541.58094                    | 26.8                |
| T3NS with $U(1) \otimes SU(2) \otimes D_{2h}$ |                       |                               |                     |
| 300               | -541.53869                   | -541.58119                    | 26.7                |
| 500               | -541.53954                   | -541.58171                    | 26.5                |
| 500, $5 \cdot 10^{-6}$ | -541.53986                   | -541.58183                    | 26.3                |
| 1000              | -541.53993                   | -541.58197                    | 26.4                |
| 1000, $5 \cdot 10^{-6}$ | -541.53997                   | -541.58198                    | 26.4                |
| Extrapolated     | -541.54012                   | -541.58210                    | 26.3                |

In this paper we extended the T3NS ansatz with SU(2) and real abelian point group symmetries. This ansatz combines the computational efficiency of DMRG and its ease of implementing symmetries with the rich entanglement representation of the TTNS. We show that implementing spin and real abelian point group symmetries is not much more involved than for DMRG.

Several calculations for the bisoxo and peroxo isomers of [Cu$_2$O$_2$]$^{2+}$ are presented. Calculations are performed both with and without spin and point group symmetries. They illustrate the substantial advantages of using the SU(2) and point group symmetries of the chemical Hamiltonian. For a given accuracy, the computational time decreases with every included symmetry.

For these calculations, the same, rather intuitive, orbital ordering as in the preliminary T3NS-paper is used.[13] Advanced techniques for the ordering of orbitals, like the usage of entanglement measures,[31] and its effect on the accuracy will be of interest in subsequent research. Furthermore, results can be improved through the development of post-T3NS methods, in similarity to post-DMRG methods. Some notable examples of post-DMRG methods which can readily be adapted are DMRG-SCF,[37,38] DMRG-CASPT2[39,40] and DMRG-TCCSD (DMRG-tailored coupled cluster with single and double excitations).[41,42]

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A Computational complexity

Two differences can be noted between the computational complexity given in the original T3NS-paper and the one given in this paper.

First, no disk resource requirements are given here, since our current implementation does not store intermediate results on disk. All tensors are kept in memory at all time.

Second, the CPU time of the T3NS has been lowered from \( O(k^5D^2 + k^3D^4) \) to \( O(k^4D^2 + k^3D^4) \). In the previous paper, we stated the complexity for updating renormalized operators with branching tensors to be \( O(k^5D^2) \) for quantum chemistry. It was noted that the recombination of two single operators in both sets of renormalized operators to a complementary double operator was the most intensive part. The worst case scenario for this is indeed \( O(k^4D^2) \) and since there are \( O(k) \) branching tensors in the network where this can occur, we previously obtained \( O(k^5D^2) \). However not all branching tensors will result in this worst case scaling of \( O(k^4D^2) \), and only the most central ones in the network will have this scaling. A more rigorous analysis showed an overall upper bound of \( O(k^4D^2) \) instead.

B Working with symmetry tensors

In this appendix we explain the T3NS with fermionic reduced tensors and their symmetry tensors in more depth. In Appendix B.1 we use the gauge freedom of the tensor network to define a canonical form of the tensors, in correspondence with the canonical form in DMRG. In appendix B.2 we give a few examples of contractions of renormalized operators with two-site tensors. In appendix B.3 some cases for updating the renormalized operators are given.

B.1 The canonical form

The representation of a wave function by a tensor network is not unique. To show this gauge freedom, we contract the tensor \( A[x] \) with an invertible matrix \( Y \) and the neighboring tensor \( A[x + 1] \) with \( Y^{-1} \).

\[
\tilde{A}[x]_{ijk} = \sum_k A[x]_{ijk}Y_{kk'} \tag{22}
\]

\[
\tilde{A}[x + 1]_{k'lm} = \sum_k (Y^{-1})_{k'k}A[x + 1]_{klm}. \tag{23}
\]

When contracting these two new tensors \( \tilde{A}[x] \) and \( \tilde{A}[x + 1] \), we obtain the exact result as contracting the original tensors. This freedom is present at every virtual bond in the tensor network.

Although one can use the gauge freedom in general tensor networks to define a canonical form, it is more straightforward in loopfree finite tensor networks. In this canonical form one tensor is chosen as orthogonality center. The currently optimized tensor is normally chosen for this. Other tensors are orthogonal with respect to contraction over all bonds but the one leading to the orthogonality center. Calculating the overlap of the tensor network with itself now simplifies to a complete contraction of the orthogonality center with its adjoint since the contributions of the other tensors simplify to unit tensors. This allows us to optimize the orthogonality center through an ordinary eigenvalue problem instead of a general one.

All tensors of the T3NS wave function are of the form given in eq. 9. Depending on the indices that are contracted, tensors of this form can be orthogonal in different ways. For such a reducible tensor which is orthogonal to a contraction over index 1 and 2 we propose that the reduced tensor should be orthogonal as well with respect to a contraction over index 1 and 2. Furthermore, the adjoint of the reduced tensor (i.e. the corresponding reduced tensor for the bra wave function) is given by its hermitian.

A graphical depiction for this case is given in fig. 9. In the bottom of fig. 9 it is shown how the symmetry tensors simplify the contraction. Indeed, with help of the symmetry tensors and
If the same reducible tensor has to be orthogonal with respect to a contraction over index 2 and 3, the reduced tensor should be again orthogonal with respect to that contraction. We use the gauge freedom to move a factor of $\sqrt{2j_1 + 1}$ from a neighboring tensor to this tensor and a factor of $\sqrt{2j_3 + 1}$ from this tensor to a neighboring tensor (see the different position of the solid circle in fig. 9 and fig. 10).

The adjoint of the reduced tensor is given again by its hermitian, but now, also an extra phase needs to be introduced. This can be seen in fig. 10. The phase is needed for imposing orthogonality due to the fermionic nature of the tensors. The introduction of this phase does not pose a problem as long the contraction over all adjoint tensors of the network results in the expected bra wave function. This can be ensured by correcting all the introduced phases in the adjoint of the orthogonality center. We get indeed for the orthogonalized tensor given in fig. 10 that

$$C(AA^+) = \sum_{x_1x_1} |x_3\alpha_3\rangle\langle x_3\alpha_3| .$$

(25)
this optimization step, no Wigner symbols are needed. It also means that, in this formulation, one can do an optimization of a DMRG-chain which only needs Wigner symbols when appending site-operators to the renormalized operators.

If one of the two sites is a branching site, a TNS optimization step is needed. During this optimization, three sets of renormalized operators are used to calculate the effect of the effective Hamiltonian on the two site object. An example of this optimization step is shown in fig. 11(b). Depending on where the physical tensor is attached to the branching tensor, variants of fig. 11(b) are needed, giving rise to slightly different prefactors.

**B.3 Updating renormalized operators**

After every optimization step of the algorithm the renormalized operators need to be updated with the newly found optimized tensors. In section 3.4 we already discussed appending a site-operator to a renormalized operator (see fig. 6). Here, we show how a physical tensor or a branching tensor can be used to update the renormalized operators.

In fig. 12, a physical tensor is used to update a renormalized operator with a site-operator appended to it. The physical tensor used in this figure is orthogonal with respect to a contraction over leg 2 and 3 (see fig. 10).

In fig. 13, a branching tensor is used to recombine two renormalized operators to a new one. The branching tensor in this example is orthogonal with respect to a contraction over leg 1 and 2 (see fig. 9).

Similar graphical depictions can be made for both fig. 12 and 13 when using tensors orthogonalized in different ways. Both the appending of a site tensor and the recombination of two renormalized operators through usage of a branching tensor need Wigner-9j symbols. Updating of a renormalized operator through usage of a physical tensor does not need any Wigner symbols.

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Figure 11: (a) The effect of the effective Hamiltonian on a two-site object consisting out of two physical sites. (b) The effective Hamiltonian on a two-site object consisting out of one physical and one branching site. Tensor $M$ is a tensor needed for the merging of the renormalized operators and its value can be found by deforming the network in fig. [to the same form as the T3NS network.]
Figure 12: Update of a renormalized operator with a site-operator attached. This operator can be obtained as shown in fig. 6. The update is done by using a orthogonalized physical tensor and its adjoint. In this example, the tensor is orthogonalized with respect to contraction over bond i and 2.

\[
\delta_{N_i} N_i \delta_{N_i' j_i} N_i' \times \\
\delta_{j_i} j_i', \delta_{j_i'} j_i', \\
\delta_{j_i} j_i', \delta_{j_i'} j_i'.
\]

Figure 13: Recombination of two renormalized operators to a new one. The recombination is done by using an orthogonalized branching tensor and its adjoint. In this example, the tensor is orthogonalized with respect to contraction over bond 1 and 2.

\[
\begin{align*}
x_{i'} x_1^0 & = x_{i'} x_1^0 \\
 x_{i'} x_1^0 & = x_{i'} x_1^0.
\end{align*}
\]

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