Lazy skip lists, a new algorithm for fast hybridization-expansion quantum Monte Carlo

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The solution of a generalized impurity model lies at the heart of electronic structure calculations with dynamical mean-field theory (DMFT). In the strongly-correlated regime, the method of choice for solving the impurity model is the hybridization expansion continuous time quantum Monte Carlo (CT-HYB). Enhancements to the CT-HYB algorithm are critical for bringing new physical regimes within reach of current computational power. Taking advantage of the fact that the bottleneck in the algorithm is a product of hundreds of matrices, we present optimizations based on the introduction and combination of two concepts of more general applicability: a) skip lists and b) fast rejection of proposed configurations based on matrix bounds. Considering two very different test cases with \( d \) electrons, we find speedups of \( \sim 25 \) up to \( \sim 500 \) compared to the direct evaluation of the matrix product. Even larger speedups are likely with \( f \) electron systems and with clusters of correlated atoms.

I. INTRODUCTION

One of the frontiers in condensed matter systems is the realistic modeling of strongly-correlated materials. The combination of density functional theory (DFT), a workhorse for electronic structure calculations of weakly-correlated materials, with dynamical mean-field theory (DMFT)\(^1\), originally designed to handle strong correlations in simple models, has allowed insights into strongly-correlated compounds at a level of realism previously unobtainable. Comparisons of momentum-resolved spectral functions, densities of states, and optics between theory and experiment are routine.

Lying at the core of this combined theory, named DFT+DMFT\(^2\)–\(^8\), is the solution of a generalized Anderson impurity model. In the strongly-correlated regime, the method of choice is the hybridization expansion continuous time quantum Monte Carlo (CT-HYB)\(^9\)–\(^12\), a numerically exact algorithm capable of handling arbitrary local interactions on the impurity site, in particular, the full atomic Coulomb potential needed to capture the \( d \) and \( f \) electron physics present in strongly-correlated materials. Enhancements to the CT-HYB algorithm are important for bringing new physical regimes within the reach of current computational resources.

In the context of model Hamiltonians, CT-HYB is also commonly used as an impurity solver for cluster generalizations of DMFT.\(^13\)–\(^25\) CT-HYB is particularly useful in the strongly correlated case.\(^26\) Here, we present optimizations based on skip lists\(^28\) and matrix bounds which result in a speedup of \( \sim 25 \) up to \( \sim 500 \) as compared to the straightforward implementation of CT-HYB (see Fig. 1). These speedups are obtained for two very different test cases where the materials contain correlated \( d \) electrons. In the low-temperature and strongly-correlated regimes of interest, the most computationally expensive step is the evaluation of the expectation value of a time-ordered sequence of (possibly thousands of) creation and annihilation operators acting on the impurity degrees of freedom, schematically notated as \( \langle d_1^\dagger d_2 d_3 d_4^\dagger d_5 d_6 \cdots \rangle \). When the complete basis of impurity states are inserted between each operator, the problem is transformed into (the trace of) a product of hundreds of matrices, called the impurity trace, which must be evaluated at each Monte Carlo step.

Our algorithm, which we dub “lazy skip lists”, optimizes the matrix product by combining the following two ideas. First, we take advantage of the fact that between subsequent Monte Carlo steps, the matrix product only changes by the insertion or removal of two operators, for example, \( \langle d_1^\dagger d_2 d_3 d_4^\dagger d_5 d_6 \cdots \rangle \rightarrow \langle d_1^\dagger d_2 d_3^\dagger d_4 d_5 d_6 \cdots \rangle \) in the case of insertion. We observe that the intermediate products \( d_1^\dagger d_2 d_3 d_4^\dagger \) and \( d_1^\dagger d_5 d_6 \cdots \) are unchanged. Using skip lists, we efficiently store these intermediate products to minimize recomputation. A similar idea based on binary search trees is presented in Ref. 12. However, skip lists are easier to implement and statistically just as
Second, we often can avoid performing the matrix product altogether by quickly rejecting proposed Monte Carlo moves via a “lazy” evaluation of the impurity trace. This implementation was first carried out in Ref. 29 and already successfully used in Ref. 30. In normal Monte Carlo sampling, we compute an acceptance probability
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In this section, we briefly summarize the key steps which generate the hybridization expansion formulation of impurity models. The goal is to quickly arrive at a description of the structure of the impurity trace imposed by the physics and to discuss what it implies for the Monte Carlo algorithm.

A general impurity model consists of a local interacting system $H_{\text{loc}}$ describing the impurity degrees of freedom, immersed in a non-interacting electronic bath:

$$H = H_{\text{loc}}(d^\dagger_i, d_i) + \sum_\mu \epsilon_\mu a^\dagger_\mu a_\mu + \sum_\mu (V_\mu a^\dagger_\mu d_i + \text{h.c.}), \quad (1)$$

where $\epsilon_\mu$ is the bath dispersion and $V_\mu$ the amplitude for particles to hop from the impurity orbital $i$ to the bath orbital $\mu$. The spin index is absorbed into the index $i$.

### A. Partition Function Sampling

In CT-HYB, we transform the partition function $Z = \text{Tr} e^{-\beta H}$ of the impurity model into a form amenable for Monte Carlo sampling (described in detail in Ref. [12]). One uses the interaction representation with the unperturbed Hamiltonian the sum of the local and bath Hamiltonians. The hybridization is the interaction term. Then, we expand the resulting expression in powers of this hybridization term, giving

$$Z = Z_{\text{bath}} \sum_{k=0}^{\infty} \int_0^\beta d\tau_1 \cdots \int_0^\beta d\tau_k \int_0^\beta d\tau'_1 \cdots \int_0^\beta d\tau'_{k-1} \times \sum_{i_1 \cdots i_k} \sum_{i'_1 \cdots i'_{k-1}} w\{(i_1, \tau_1) \cdots (i_k, \tau_k)\}, \quad (2)$$

where the integrand is

$$w\{(i_1, \tau_1) \cdots (i_k, \tau_k)\} = \text{Det} \Delta \times \text{Tr}_{\text{loc}}[T e^{-\beta H_{\text{loc}}} d_{i_k}(\tau_k)d^\dagger_{i_k}(\tau'_k) \cdots d_{i_1}(\tau_1)d^\dagger_{i_1}(\tau'_1)]. \quad (3)$$

Since the impurity and bath degrees of freedom are decoupled, the trace over the bath has been performed. The bath is contained in the determinant of a $k \times k$ matrix $\Delta$ with elements evaluated from the hybridization function $(\Delta)_{mn} = \Delta_{i_m i_n}(\tau_m - \tau_n)$ whose Matsubara definition is

$$\Delta_{ij}(i\omega_n) = \sum_\mu V_{\mu i}^* V_{\mu j}/i\omega_n - \epsilon_\mu. \quad (4)$$

The average over the impurity $\text{Tr}_{\text{loc}}$ in general cannot be further decomposed. Its evaluation requires converting the sequence of operators (and intervening time-evolution
operators) into matrices in the basis of the impurity Hilbert space $\mathcal{H}$.

The Monte Carlo sampling of Eq. 2 proceeds as follows: the integrands $w$ of the partition function sum define the weights of a distribution over the configuration space $\{(i_1, \tau_1) \ldots (i_k, \tau_k)\}$ which is sampled with the Metropolis-Hastings algorithm. At each step, a new configuration is proposed with probability $A$ and accepted with probability

$$p = \min \left(1, \frac{A'|w'|}{A|w|} \right),$$

where $w$ and $w'$ are the weights of the new and the old configuration respectively, and $A'$ is the proposal probability of the inverse update.

The bottleneck is that the weights $w$, and the expensive impurity trace contained within, must be computed in order to decide whether to accept each new proposed configuration. In terms of computational effort, if $N = |\mathcal{H}|$ is the size of the local Hilbert space, and we are sitting at perturbation order $k$, the impurity trace costs $O(N^3k)$ while the hybridization determinant costs $O(k^3)$ (which can be reduced to $O(k^2)$ for local updates). The average expansion order $\langle k \rangle$, which is typically in the hundreds, is proportional to the inverse temperature $\beta$, whereas the $N$ grows exponentially with the number of impurity orbitals ($N = 1024$ for the $d$-shell). Thus, except at very low temperatures, the calculation of the impurity trace is the bottleneck in these Monte Carlo simulations.

Alluded to in the above discussion, the impurity trace contains a time-evolution operator between each creation and annihilation operator, which we denote by $P_r = e^{-\tau H_{\text{loc}}}$. We also write $(F_i)_{mn} = \langle m | d_i | n \rangle$ for the matrix representation of the creation and annihilation operator, where $m$ and $n$ index the states in $\mathcal{H}$. In this notation, the impurity trace explicitly becomes an alternating matrix product:

$$\text{Tr}_{\text{loc}} P_{\beta - \tau_k} F_{a_k} P_{\tau_k - \tau_i} F_{i_k}^\dagger \cdots F_{i_1} P_{\tau_1 - \tau_i} F_{i_1}^\dagger P_{\tau_i}. \quad (6)$$

For simplicity, we have assumed that the imaginary times in Eq. 3 are time-ordered as they appear.

**B. Symmetries, Sectors and Block Matrices**

We can make a key simplification to the impurity trace using symmetries prior to developing computational algorithms. The local Hamiltonian $H_{\text{loc}}$ generally possesses abelian symmetries (e.g. particle number, spin, momentum), which allow us to decompose the impurity Hilbert space as a direct sum $\mathcal{H} = \bigoplus_{q \in \mathbb{Z}/2} \mathcal{H}(q)$. Here, $q$ enumerates the sectors of the Hilbert space, each of which is characterized by a definite set of quantum numbers (e.g. particle number, spin, momentum).

Using these symmetries one defines a new basis for the creation-annihilation operators. A creation or annihilation operator, which we denote by a generalized index $\alpha$ formed by combining its quantum numbers with the type of operator (creation or annihilation), maps each sector $q$ either to 0 or uniquely to one other sector $q'$. This leads to block matrices $F_{\alpha_1}(q)$ which can be combined with a sector mapping function $s_{\alpha_1}$ defined by $s_{\alpha}(q) = q'$. The time-evolution operator maps each sector onto itself.

In the sector basis, the operator product in Eq. 6 becomes $PF_{\alpha_2}PF_{\alpha_3-1} \cdots F_{\alpha_2}PF_{\alpha_1}$ $P$ that maps a sector $q_0$ onto $q_k$ defined by the string $q_0 \rightarrow q_1 := s_{\alpha_1}(q_0) \rightarrow \cdots \rightarrow q_{2k} := s_{\alpha_2}(q_{2k-1})$. The impurity trace decomposes into a sum over sector traces:

$$\text{Tr}\sum_{q_0} Tr P(q_{2k})F_{\alpha_2}(q_{2k-1}) \cdots F_{\alpha_1}(q_0) P(q_0), \quad (7)$$

and only sectors $q_0$ which are not mapped on 0 contribute. Such mapping on 0 generally occurs because of the Pauli principle. In a typical 3$d$ impurity model with the full atomic Coulomb interaction, the number of sectors is $\sim 100$ and the number of surviving strings ranges from 1 to $\sim 20$.

**III. SKIP LISTS**

We first begin with a motivation for skip lists. Then the skip list and the way it is used to store matrix subproducts is described. The final subsection explains how matrix multiplications can then be performed efficiently when operators are inserted or removed.

**A. Motivation for Skip Lists**

At each Metropolis-Hastings step, a matrix product needs to be computed to decide whether the proposed configuration is accepted or rejected. One possibility is to always calculate all the products from scratch. However, only two matrices are typically inserted or removed, so this strategy is not only expensive, but also highly redundant.

To avoid multiplying almost all the time the same matrices, we may pair them off and store their product. This way almost every second multiplication is skipped when calculating the product of a proposed configuration. However, this is not yet optimal. One can store products of four, eight matrices etc. leading to a collection of subproducts that will allow us to minimize the number of redundant multiplications. This storage strategy may be represented as shown in Fig. 2, where we omit the propagators for simplicity. The arrows store the sub-products of operators they span, including the operator they start from and excluding the operator they point to.

Inserting now a matrix $F$, some of the stored subproducts expire, as shown on the lower panel of Fig. 2. These are the sub-products of arrows that span over the
There is an arrow starting and ending at the top end of the inserted matrix. To calculate the product of the products with a bold red multiplication sign need to be calculated in order to obtain the total product.

Insertion means that a sub-product is ideally always the product of two moving matrices at random places. Equilibrated means that this storage scheme works only if the expansion order in the number of operators in the product. However, this configuration is proposed, the product is calculated by filling the sampling more efficient. Hence, we need a flexible multiplication algorithm, which is discussed in the next section.

C. Skip Lists and Matrix Multiplication

To calculate the new product after an arbitrary sequence of insertions and/or removals with a minimal number of matrix multiplications, we proceed in two steps. First the matrices are inserted and/or removed, one after the other. At each time, this invalidates some sub-products $M = PF_1PF_2\ldots PF_n$ stored in the blue arrows. These sub-products are thus emptied. Once the new configuration is proposed, the product is calculated by filling up the emptied sub-products.

When inserting an operator in the skip list, a sub-product expires if the operator lies between the head and the tail of the corresponding arrow, see Fig. 3.
ify all such arrows, we follow the skip list insertion algorithm and begin at the tail of the top arrow. This arrow necessarily spans over the operator to insert, and its sub-product is emptied. Moving down the red arrow on the right in Fig. 3 to the next lower blue arrow, we test if the operator to insert lies between the head and tail of this arrow. If yes, the sub-product is emptied, and the next lower blue arrow is tested. If not, the arrow is traversed and the process is repeated until we end up by emptying the sub-product at the blue arrow just above the place where the operator will be inserted. Proceeding likewise for removal, all expired sub-products are emptied once the new configuration is proposed.

To fill up the emptied sub-products \( M \) once the insertions and/or removals are completed, we proceed recursively. The sub-product at an arrow \( A \) can be calculated from the sub-products \( M_a, M_{a+1}, \ldots, M_b \) stored at the arrows \( A_a, A_{a+1}, \ldots, A_b \) just below. If all of these sub-products have not been emptied, they are multiplied while traversing the arrows \( A_a \rightarrow A_{a+1} \rightarrow \cdots \) and the result is stored at the arrow \( A \). If however one of the sub-products \( M_i \) at an arrow \( A_i \) is missing, we recursively calculate this sub-product from the sub-products below the arrow \( A_i \). This recursion stops at the latest at the bottom of the skip list, where the operators are multiplied with the propagators. The total product is obtained by starting the recursion at the top arrow.

Once the new product is calculated, we decide whether to accept or reject the proposed configuration. To recover the skip list in case of rejection, a backup is taken at the beginning of a trial step.

IV. LAZY TRACE EVALUATION

In the regimes of interest (moderate to low temperatures \( T \lesssim 100 \) K, strong Coulomb interaction \( U \gtrsim 5 \) eV), the probability of accepting a proposed move is low, generally lying below 10% and often below 1%. The Pauli principle and time-evolution operators \( e^{-\Delta t H_{\text{loc}}} \) place strong constraints on the insertion/deletion of operators, causing the low acceptance probabilities. Developing techniques to reject improbable moves with minimal computational effort is crucial.

The Pauli constraint is computationally negligible, as it can quickly be determined by following the string of sector mappings \( q_0 \rightarrow q_1 \rightarrow q_2 \cdots \) and checking that not all strings are annihilated (i.e. mapped to 0). In contrast, the time-evolution operators are interspersed within the matrix product. Proposed moves often drive transitions to high-energy sectors, where the exponentials \( e^{-\Delta t H_{\text{loc}}} \) strongly suppress the acceptance probability. Here, we describe a “lazy trace” algorithm which leverages these exponentials to efficiently reject moves with low acceptance probability, largely avoiding a full evaluation of the impurity trace.

The first component of the lazy trace algorithm is fast bounding of the impurity trace in each symmetry sector. Writing in shorthand Eq. 7 as \( \text{Tr} = \sum_q \text{Tr}_q \), assume we can quickly compute bounds \( B_q \geq |\text{Tr}_q| \) for each sector trace. This provides a maximum bound on the trace via the triangle inequality:

\[
|\text{Tr}| \leq \sum_q |\text{Tr}_q| \leq \sum_q B_q.
\] (8)

Using the expression for the acceptance probability \( p \) (Eq. 5), and writing the weight of the old configuration as \( w' = \text{Det}' \cdot T \gamma \), we obtain an upper bound

\[
p_{\text{max}} = \frac{A'}{A} |\text{Det}| \sum_q B_q \end{equation}

This bound can be refined as follows: take the sector \( q_{\text{max}} \) with the largest \( B_q \) and compute the exact sector trace \( \text{Tr}_{q_{\text{max}}} \). Applying the reverse triangle inequality gives

\[
|\text{Tr} - |\text{Tr}_{q_{\text{max}}}| \leq \sum_{q \neq q_{\text{max}}} B_q,
\] (10)

producing refined bounds

\[
\frac{p_{\text{max}}}{p_{\text{min}}} = \frac{A'}{A} |\text{Det}| \sum_{q \neq q_{\text{max}}} \left( |\text{Tr}_{q_{\text{max}}}| \pm \sum_{q \neq q_{\text{max}}} B_q \right).
\] (11)

This procedure can be continued, generating successively tighter bounds, until we obtain the exact trace. The sequence of bounds is likely to tighten most rapidly if we choose the sectors in decreasing order of \( B_q \).

The second key idea is to flip the Monte Carlo coin first to obtain the acceptance threshold \( u \), before computing the above approximation to the acceptance probability. If \( p_{\text{max}} < u \), and it often is, we can reject the move outright. If \( p_{\text{min}} > u \) we accept the move. If neither of these possibilities occur, we successively refine the bounds on
p until we can either accept or reject the move, as illustrated in Fig. 4. In the following, we describe the construction of the bounds $B_q$.

The basic equation is the formula

$$|\text{Tr} A_1 A_2 \cdots A_n| \leq C \cdot \|A_1\| \|A_2\| \cdots \|A_n\|,$$  \hspace{1cm} (12)

proven in Appendix A. Here $A_k$ are matrices (not necessarily square, although the entire product must be), $\|\cdot\|$ is a sub-multiplicative matrix norm, and $C$ is a constant which depends on the specific matrix norm chosen and the dimension of the matrices. In the lazy trace algorithm, the spectral norm (see Appendix A) is used. For rectangular matrices $A_k \in \mathbb{R}^{N_k \times M_k}$, the constant $C$ becomes the dimension of the smallest matrix within the product, $C = \min\{N_k\}$. The spectral norm is unity for a creation or annihilation operator, and $e^{-\Delta \tau_i E_0(q_i)}$ for time-evolution operator, where $E_0$ is the ground state energy of the sector $q_i$ and $\Delta \tau_i$ is the time spent in this sector.

Application to the trace of a single sector in Eq. 7 gives

$$|\text{Tr} P(q_{aK}) F_{a1} F_{a2} \cdots F_{aK} P(q_0)|$$

$$\leq \text{dim} \{H(q_i)\} \cdot \exp \left( - \sum_{i=0}^{2K} \Delta \tau_i E_0(q_i) \right),$$  \hspace{1cm} (13)

While extremely cheap to calculate, this bound precisely captures the vast variations in magnitude caused by exponentials in the time-evolution operators. The bounds for each sector $B_q$ decrease extremely rapidly; in many cases, the initial $p_{\text{max}}$ is sufficient to reject a proposed move.

When a move is accepted, the trace needs to be evaluated exactly, up to numerical accuracy, to be able to compute the acceptance probability of the next move.

V. LAZY SKIP LISTS

In this section, we begin by combining the algorithms presented in Sec. III and Sec. IV. In a second step, we show how the bounds on the sector traces in Sec. IV may be improved using this combined algorithm.

A. Skip Lists and Lazy Trace Evaluation

When iteratively refining the bounds in the lazy trace evaluation, we only need the contribution to the trace of one sector $q_0$ at a time in Eq. 7. To achieve this with the skip lists in Sec. III B, we begin by taking into account the block structure of the matrices.

The operators $F$ and the sub-products $M$ are stored in their block form as pairs $s(q), F(q)$ and $s(q), M(q)$ of mapped sectors and corresponding matrix blocks. Similar to the total product which splits into strings in Sec. II B, this splits a sub-product $PF_b \cdots PF_a$ into sub-strings $P(q_{a+1}) F_b(q_a) \cdots P(q_{b+1}) F_a(q_b)$. Such a sub-string is stored in the matrix block $M(q_a)$ together with the mapped sector $s(q_a) := q_{a+1}$.

To calculate one string in the total product, we only need one of the sub-strings of a given sub-product. When recursively updating the sub-products in the skip list as in Sec. III C, we thus have to specify at each arrow $A$ the requested sub-string by a start sector $q_a$. To select the entries in the block matrices $M_i$ (stored in $A_i$ below $A$) which need to be multiplied to obtain the requested sub-string $M_b(q_b) \cdots M_{a+1}(q_{a+1}) M_a(q_a)$, one maps the start sector $q_a$ into $q_{a-1}$ using the sector mappings $s_i$ at the arrows $A_i$, namely $q_a \rightarrow q_a+1 := s_a(q_a) \rightarrow \cdots \rightarrow q_b := s_{b-1}(q_{b-1})$. The product is then stored in the matrix block $M(q_a)$ at the arrow $A_i$, together with the mapped sector $s(q_a) := q_{a+1}$. Again, if a matrix block $M_i(q_i)$ at an arrow $A_i$ is empty, we proceed recursively.

The combination of the skip lists and the lazy trace evaluation is now straightforward. First, expiring sub-strings are emptied when inserting and/or removing operators in the skip list, similar to Sec. III C. Once the new configuration has been proposed, we start the recursion at the top arrow of the skip list separately for each sector needed by the lazy trace evaluation.

B. Sub-products and Trace Bounds

The bounds on the sector traces in Eq. 13 are calculated from the product of the norms of each propagator and operator individually. Tighter bounds may be obtained by using the norms of stored sub-products. In Fig. 2 for example, the trace is bounded by

$$|\text{Tr}| \leq C \cdot \|F_0 F_2 \| \|F_6\| \|F\| \|F_5\| \|F_4 F_3 F_2 F_1\|$$  \hspace{1cm} (14)

after insertion of the matrix $F$. Such bounds for a given sector trace Tr$_q$ are obtained recursively, in a manner analog to the block-matrix product of the corresponding string.

Calculating the spectral norm of a stored matrix block is expensive, so the Frobenius norm is used here instead. While this norm is larger than the spectral norm, its numerical cost is small compared to a matrix multiplication. However, this means that this bound is not necessarily smaller than the one in Sec. IV. Other choices for the norms are discussed in Appendix A.

VI. TWO EXAMPLES

In this section we benchmark the skip lists (Sec. III A taking into account the block structure described in Sec. VA), the lazy trace evaluation (Sec. IV) and the lazy skip lists (Sec. VA and Sec. VB). To this end, we consider Anderson impurity problems that appear in DFT+DMFT electronic structure calculation for thin
film of LaNiO$_3$ (LNO)$^{27,33}$ and FeTe bulk compound$^1$, using experimental structure of Ref. 34 and Ref. 35, respectively.

In both cases, the impurity is a d-shell system, and the associated Hilbert Space splits into 132 sectors. The expansion orders are $\langle k \rangle \approx 225$ for LNO and $\langle k \rangle \approx 515$ for FeTe. The benchmarks are performed using two kinds of Metropolis-Hastings updates: i) standard ones$^{36}$ with low acceptance ratio and ii) efficient updates with high acceptance ratio higher by a factor 10 to 25.

Fig. 1 shows the speedups of the different optimizations presented in this paper compared with, as a baseline, a straightforward implementation (Sec. II B) that takes the block structure into account. Note the logarithmic scale. The skip lists alone accelerate the simulations for both test cases by a factor of about 20. While the lazy trace evaluation gives a substantial speedup for LNO, essentially no speedup is obtained for FeTe. This also shows in the performance of the combined algorithms, the lazy skip lists, which, with speedups of order 500, perform much better for LNO. The reasons for this difference between LNO and FeTe will become clear below.

Fig. 5 shows, in addition to the speedup, the reduction in matrix multiplications and the reduction in floating point operations. While combining different optimizations does not always result in an additional speedup, in our case the lazy trace evaluation and the skip lists work well together. The reduction in matrix multiplications for the lazy skip lists (Sec. VA) is essentially the product of the reductions for the lazy trace evaluation and the skip lists separately. While the reduction in matrix multiplications for the lazy skip lists in Sec. VB is less evident to anticipate, there is always an additional speedup that comes from calculating the bounds using the norms of the stored sub-products in the skip list.

Note that speedups are smaller than expected from the reduction in matrix multiplications and floating point operations, in particular for the lazy skip lists of Sec. VB. This is due to the optimization overhead and to the fact that other parts than the local trace evaluation in the CT-HYB expansion, such as the evaluation of the determinants, are beginning to take a significant proportion of the total time.

To understand why most of the speedup comes from the lazy trace evaluation for LNO while it comes from the skip list for FeTe, it is useful to consider the sector weights. We use standard updates. In Fig. 6a) we show results for LNO and in Figs. 6b) results for FeTe. Note the logarithmic vertical scales. The top panels display the average weights $\langle Tr_q / Tr \rangle$ of the various sectors in the partition function expansion. The lower panels of Figs. 6a) and b) show for each sector $q$ the frequency of $Tr_q$ evaluation.

Consider first the case of LNO. In contrast to the baseline, it is clear in Fig. 6a) that the sector frequencies for the lazy trace evaluation are largely proportional to the sector weights. Only a few sectors with $N = 7$ to 8 collect most of the weight, and this not only shows where the large reduction in matrix multiplications in Fig. 5a) comes from, but also why the reduction in floating point operations is even bigger. Indeed, the sectors with $N = 7$ to 8 have generally smaller dimension than the ones with $N = 4$ to 6 which are not calculated most of time in the lazy trace evaluation.

Given their negligible sector weights, it would also be possible in principle to just drop the sectors with $N = 0$ to 3. However, the gain from this is small since these sectors have rather small dimension. Dropping the sectors with $N = 4$ to 6 involves more important approximations so one would need careful checks that the truncated sectors do not affect the results. The lazy trace evaluation avoids the calculation of these sectors most of time and there is no approximation involved.
Quantum Monte Carlo algorithms generally involve multiplications of large matrices. In the case of the strong-coupling based CT-HYB algorithm, this is a limiting factor. When updates generate new configurations that have a large probability of being rejected, we have shown that an efficient way of speeding up the algorithm is to first choose the random number and then use matrix norms to bound the Metropolis rejection/acceptation probability. This is called lazy trace evaluation. Skip lists on the other-hand provide a way to store intermediate matrix products and avoid in all circumstances the recomputation of some of the matrix-products. The combination of both algorithms, lazy skip lists, provides a robust algorithm that guarantees large speedups when the trace evaluation takes a large fraction of the computing time.

The speedup of the trace evaluation achieved with the lazy skip lists algorithm is such that parts of CT-HYB that usually take negligible time compared with the evaluation of the trace, for example measurements, calculation of determinants etc., can now become the limiting factor.

Skip lists allow control of memory requirements by changing the probability $p$ to add a level to an inserted bar after an update. We have not discussed further improvements in speed that can be obtained by using the associative property of matrix multiplication to speedup the calculation of products of rectangular matrices, or many other possible optimizations that are dependent on computer architecture, such as caches, parallelism etc.

Some of the ideas developed here can be directly applied to other problems treated by Monte Carlo methods. For example the rejection method based on bounds (see Fig. 4) can be applied to classical Monte-Carlo simulations for spins with long-range interactions.\(^\text{38}\) Take an Ising spin system and consider a single spin-flip Monte Carlo update. The energy associated with this spin can be bounded by

$$E_{i,[\min,\max]} = S_i \sum_{j \leq R} J_{i,j} S_j \pm S_i \sum_{j > R} |J_{i,j}|. \quad (15)$$

The bounds can be refined by successively increasing the range $R$. The sums over absolute values of exchange constants need to be calculated only once. Similar problems are encountered in spin-ice models with dipolar interactions, ordered and/or random spins with both dipolar and RKKY interactions.

Speedups by factors in the hundreds that can be achieved with the lazy skip lists algorithm will bring new physical regimes in correlated electronic-structure calculations and cluster generalizations of dynamical mean-field theories within reach of computational power. Applications of such methods extend as far as molecular biology.\(^\text{40}\)

### VII. DISCUSSION AND CONCLUSION

Moving to the case of FeTe in Fig. 6b), one notices that the sector weights are more uniformly distributed. There are fewer sectors with extremely small weights. Hence the lazy trace evaluation does not give a substantial speedup. The skip lists on the other hand still reduce the number of matrix multiplications.

FIG. 6. On the basis of a LNO thin film simulation (a) and of a FeTe simulation (b) with standard updates: average weight $\langle Tr_q / Tr \rangle$ of a sector $q$ in the partition function expansion (top panel) and frequency with which $Tr_q$ is calculated for a sector (lower panel).

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Appendix A: Trace Bounds via Matrix Norms

Different matrix norms give different bounds for the magnitude of the trace of a matrix product. We consider here induced norms

$$\|A\|_p := \max_{\|x\|_p=1} \|Ax\|_p,$$

where $A \in \mathbb{R}^{N \times N}$, $x \in \mathbb{R}^N$ and $\|x\|_p := (\sum_i |x_i|^p)^{1/p}$ with $p \geq 1$, and the Frobenius norm

$$\|A\|_F := \left( \sum_{ij} A_{ij}^2 \right)^{1/2}.$$

1. Induced Norms

For the induced norms, one obtains $|A_{ij}| \leq \|Ae_{ij}\|_p \leq \|A\|_p$, where $e_{ij}$ is the standard basis of $\mathbb{R}^N$, and hence

$$|\text{Tr} A| \leq N \cdot \|A\|_p.$$

This immediately generalizes to a product

$$\left| \text{Tr} \prod_{l=1}^n A_l \right| \leq \min \{N_l\} \cdot \prod_{l=1}^n \|A_l\|_p \quad \text{(A1)}$$

of rectangular matrices $A_l \in \mathbb{R}^{N_l \times M_l}$, since induced norms are sub-multiplicative. From the cyclicity of the trace, the pre-factor in Eq. 12 becomes $C = \min \{N_l\} = \min \{M_l\}$, the minimal row or column dimension of all the matrices within the product.

For a propagator $P_\tau$, written in the eigenbasis, one obtains $\|P_\tau\|_p = \exp(-\tau E_0)$, where $E_0$ is the smallest eigenvalue. These norms are hence well suited for the lazy trace evaluation in Sec. IV. Especially convenient is the spectral norm ($p = 2$). This norm is one for annihilation or creation operators since

$$\|d\|_2 = \max_{\langle \psi | \psi \rangle = 1} \sqrt{\langle \psi | d^\dagger d | \psi \rangle} = 1$$

by the Pauli principle, and only the exponentials of the propagators enter into the bound given in equation (A1).

2. Frobenius Norm

For the Frobenius norm, Cauchy-Schwarz states

$$|\text{Tr} AB| \leq \|A\|_F \cdot \|B\|_F,$$

and as the Frobenius norm is sub-multiplicative

$$\left| \text{Tr} \prod_{l=1}^n A_l \right| \leq \prod_{l=1}^n \|A_l\|_F,$$

where $n \geq 2$. The Frobenius norm is numerically cheap, so equation (A2) can be used for the lazy skip lists in Sec. V B. Other numerically cheap choices are the induced norms with $p = 1$ and $p = \infty$.

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