Rényi entropies of interacting fermions from determinantal quantum Monte Carlo simulations

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Abstract. Entanglement measures such as the entanglement entropy have become an indispensable tool to identify the fundamental character of ground states of interacting quantum many-body systems. For systems of interacting spin or bosonic degrees of freedom much recent progress has been made not only in the analytical description of their respective entanglement entropies but also in their numerical classification. Systems of interacting fermionic degrees of freedom, however, have proved to be more difficult to control, in particular with regard to the numerical understanding of their entanglement properties. Here we report a generalization of the replica technique for the calculation of Rényi entropies to the framework of determinantal Quantum Monte Carlo simulations—the numerical method of choice for unbiased, large-scale simulations of interacting fermionic systems. We demonstrate the strength of this approach over a recent alternative proposal based on a decomposition in free fermion Green’s functions by studying the entanglement entropy of one-dimensional Hubbard systems both at zero and finite temperatures.

Keywords: Hubbard model (experiments), quantum Monte Carlo simulations, analysis of algorithms, entanglement in extended quantum systems (theory)

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1. Introduction

Entanglement is one of the fundamental concepts of quantum mechanics that describes how quantum mechanical objects—e.g. photons, electrons, or spins—are interwoven into a collective state [1]. If such a state can no longer be described as a simple product state of single-particle wavefunctions, one says that the quantum mechanical objects are entangled. Beyond its conceptual relevance quantum mechanical entanglement has turned into a key resource in various fields of modern physics [2]. In quantum information theory it is exploited in storing and manipulating information in so-called qubits [3–5]. In condensed matter physics entanglement has become increasingly appreciated as a measure to classify different states of quantum matter which cannot be distinguished by any local observable such as topologically ordered states [6–9]. The probably stunning realization that oftentimes ground states of quantum many-body systems are far from being highly entangled states has led to the development of a novel class of
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While the notion of entanglement was originally associated with typically a handful of qubits its application to quantum many-body systems requires entanglement measures that allow one to deal with an almost arbitrarily large number of interwoven quantum mechanical degrees of freedom. One such powerful measure is the so-called entanglement entropy [2], which can be calculated from a bipartition of a quantum many-body system into two complementary parts A and B as illustrated in figure 1. Tracing out the degrees of freedom in one subsystem one can calculate a reduced density matrix for the other, e.g. $\rho_A = \text{Tr}_B(\langle \psi | \psi \rangle)$. The information in the density matrix is then condensed into a single number, e.g. the von Neumann entropy [4]

$$S(A) = - \text{Tr}[\rho_A \log \rho_A].$$ (1)

The von Neumann entropy is the most prominent member of a more general family of entanglement entropies, the so-called Rényi entropies [13] which are calculated from the density matrix as

$$S_n(A) = \frac{1}{1-n} \log (\text{Tr}(\rho_A^n)), \quad (2)$$

where the limit $n \to 1$ recovers the above von Neumann entropy.

The strength of these entropic entanglement measures becomes apparent when considering the scaling of the entanglement entropy for varying sizes of the subsystem $A$. In contrast to conventional thermodynamic entropies the entanglement entropy is not extensive, but rather scales with the length of the boundary between the two partitions—the so-called boundary law [6] (which is often also referred to as area-law). Corrections to this prevalent boundary law have received widespread attention for their ability to classify different states of quantum matter [2]. For instance, it has been shown that the topological character of non-local order present in a quantum ground-state wavefunction can be revealed by a universal $O(1)$ correction [8, 9] to the boundary law. Numerical simulations of quantum spin systems have subsequently shown that such an identification is indeed feasible and unambiguously revealed the long conjectured topological order present in ground states of certain frustrated quantum magnets [14–16]. Systems of interacting fermionic degrees of freedom are the conceptually most...
interesting class of interacting quantum many-body systems due to the even more complex variety of possible ground states and corresponding entropic signatures arising from the intricate nodal structure of fermionic wavefunctions. While a generalization of the numerical schemes to calculate entanglement entropies also for these many-fermion systems is highly desirable, progress so far has been limited to variational Monte Carlo techniques \cite{17–20}. The first step to develop an approach based on determinantal Monte Carlo—an unbiased, auxiliary field technique that has become the method of choice for large-scale simulations of interacting fermion systems—has recently been reported by Grover \cite{21}, whose approach is based on a decomposition of the entanglement entropy in terms of free-fermion Green’s functions.

The purpose of this manuscript is to describe an efficient numerical method to calculate the entanglement entropy for systems of interacting fermions in any spatial dimension. Our approach generalizes a replica scheme used to calculate Rényi entropies in world line quantum Monte Carlo approaches for interacting spin or bosonic systems to the framework of determinantal Monte Carlo simulations as detailed in the following section.

We demonstrate the applicability of this approach by simulating one-dimensional Hubbard systems and discuss the strength of our technique in a detailed comparison with Grover’s recent alternative proposal to decompose the entanglement entropy in terms of free-fermion Green’s functions \cite{21} in section 3. We close with an outlook in section 4.

2. Determinantal QMC and the replica trick

We consider a setup where the interactions between spinful fermionic degrees of freedom are captured by a lattice Hamiltonian such as the Hubbard model

\[
\mathcal{H} = -t \sum_{\langle i,j \rangle, \sigma} (c_{i,\sigma}^\dagger c_{j,\sigma} + \text{h.c.}) + U \sum_i n_{i,\uparrow} n_{i,\downarrow} - \mu \sum_i (n_{i,\uparrow} + n_{i,\downarrow}),
\]

(3)

whose physics thrives from the competition of the on-site interaction \(U\) and the hopping \(t\) for fixed chemical potential \(\mu\). Our method is, however, not limited to the specifics of the Hubbard model but can in fact be applied to any fermionic Hamiltonian amenable to a quantum Monte Carlo simulation. The quantum mechanical state of such an interacting many-fermion system can be described via its density matrix \(\rho\). When considering the ground state of the system at zero temperature this density matrix is given by

\[
\rho = \frac{\langle \psi | \psi \rangle}{\langle \psi | \psi \rangle},
\]

(4)

while at finite temperatures it takes the form

\[
\rho = \frac{\exp(-\beta \mathcal{H})}{\text{Tr}(\exp(-\beta \mathcal{H}))}.
\]

(5)

In both cases, we have introduced an explicit normalization constant \(\mathcal{N}\) in the denominator, which not only ensures that the trace of the so-defined density matrix is 1, but
will play an important conceptual role in the following. To be even more explicit, we can rewrite both expressions as

$$\rho = \frac{1}{N'} \rho',$$

(6)

which is the form we will be using in the following.

### 2.1. Rényi entropies and the replica trick

The first step in calculating the Rényi entropy is to determine the reduced density matrix $\rho_A$ by tracing out the degrees of freedom in subsystem $B$

$$\rho_A = \text{Tr}_B \rho.$$

In our numerical calculations we will concentrate on the second Rényi entropy, i.e. the case of $n = 2$, which can be calculated from the reduced density matrix $\rho_A$ as

$$S_2(A) = - \log (\text{Tr}_A (\rho_A^2)).$$

(7)

Note that $\rho_A$ is squared before the remaining degrees of freedom of subsystem $A$ are traced out. Using the notation of equation (6), we thus find the following general expression for the Rényi entropy

$$S_2(A) = - \log \left( \frac{\text{Tr}_A (\rho_A^2)}{N^2} \right),$$

(8)

which is valid for both finite temperature and ground state considerations. In the following we will closely examine how this definition of the Rényi entropy (8) can be translated into an algorithm for its numerical computation. For concreteness we will initially focus on the finite-temperature scenario and expand our discussion to ground-state calculations in a later step. Considering first the denominator in equation (8), we note that

$$N^2 = (\text{Tr} \rho)^2 = Z^2,$$

(9)

i.e. the normalization $N^2$ is equal to the square of the usual partition sum considered in thermodynamics. The numerator of definition (8) is a bit more involved

$$\text{Tr}_A (\rho_A^2) = \sum_{A,A',B,B'} \langle AB'|\rho'|A'B' \rangle \langle A'B|\rho'|AB \rangle \equiv Z[A, 2, T],$$

(10)

where we have defined the partition function $Z[A, 2, T]$. We are now left with the question of how to numerically calculate these two partition functions. To this end, we will first consider their calculation in the framework of world line quantum Monte Carlo techniques, as they are typically used for systems of interacting spin or bosonic degrees of freedom. We will then turn to the framework of determinantal quantum Monte Carlo (DQMC) techniques, typically used to simulate many-fermion systems.

Turning first to the case of world line QMC techniques, it is helpful to translate the two partition sums $Z^2$ and $Z[A, 2, T]$ into their respective world line line representations
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\[ Z^2 \]

\[ Z[A, 2, T] \]

\[ W(\sigma, \psi) = \det(\sigma, \psi). \]

\[ \text{Figure 2.} \] Ensemble switching in a world line picture. The left side shows the square of the regular partition sum \( Z^2 \) where all world lines have to be \( \beta \)-periodic. The right side shows a configuration of allowed world lines in the \( Z[A, 2, T] \) partition sum, where \( 2\beta \)-periodicity is enforced in part A of the system but part B remains \( \beta \)-periodic.

as illustrated in figure 2. On the left side, the world line representation of \( Z^2 \) from the denominator in equation (8) is depicted with two sets of \( \beta \)-periodic world lines extending from 0 to \( \beta \) and from \( \beta \) to \( 2\beta \), respectively. On the right hand site a world line representation is depicted for the partition function \( Z[A, 2, T] \) of the numerator of equation (8), where we consider two replicas of the system connected in imaginary time. Carefully implementing the imaginary time boundary conditions defined in equation (10) results in a \( \beta \) periodicity for part B and a \( 2\beta \) periodicity for part A—a scheme often referred to as the replica trick [22, 23]. To sample world line configurations according to these two partition sums, one simply needs to implement their respective imaginary time boundary conditions—a task, which is relatively straightforward for any existing world line Monte Carlo implementation [23, 24].

2.2. Determinantal QMC

Let us now turn to the framework of DQMC simulations and try to adapt the evaluation of the two partition functions \( Z^2 \) and \( Z[A, 2, T] \) needed to calculate the Rényi entropy of equation (8) for a fermionic system. Describing the DQMC framework in a nutshell, one first decouples quartic terms in the Hamiltonian by a Hubbard–Stratonovich transformation introducing an auxiliary field, which then allows to integrate out the fermions analytically. A Monte Carlo scheme is then used to sample configurations of the auxiliary field. One of the key differences to world line techniques described above is that we do not sample world lines directly. In fact, by integrating out the fermionic degrees of freedom, we sample all possible world line configurations simultaneously for a given configuration of the auxiliary field.

This raises the question of how to properly implement the replica scheme in this framework. We will concentrate our discussion on the key conceptual steps in the following and refer the inclined reader interested in a more concise technical description to appendix A. Considering at first an arbitrary pair of a fermion state \( \psi \) and an auxiliary field state \( \sigma \) its statistical weight \( W(\sigma, \psi) \) is given by a Slater determinant

\[ W(\sigma, \psi) = \det(\sigma, \psi). \]
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A crucial step is to realize that the grand-canonical trace over these Slater determinants can be recast as a single determinant

$$\text{Tr} \det (\sigma, \psi) = \det (\sigma),$$

which implies that one can integrate out the fermionic degrees of freedom and instead consider only configurations of the auxiliary field. However, it should be noted that this identity works only if the weights of the partition sum can be written in terms of single Slater determinants. When considering the replica scheme for the calculation of Rényi entropies, however, one typically encounters statistical weights in the expression for the partition function $Z[A, 2, T]$ that are the product of two determinants

$$Z[A, 2, T] = \sum_{\sigma} \sum_{A', B', B''} \langle AB' | \rho' | A'B' \rangle \langle A'B | \rho' | AB \rangle$$

$$= \sum_{\sigma} \sum_{A', A''} \det (\sigma, AB' | A'B') \det (\sigma, A'B | AB),$$

where the two determinants are denoted as $\det (\sigma, AB' | A'B')$ and $\det (\sigma, A'B | AB)$. They again depend on the auxiliary field configuration $\sigma$ and the arguments $A, B, A', B'$ indicate the imaginary time boundary conditions for a particular pair of auxiliary field and fermionic configurations. This structure seems to suggest that one can no longer integrate out the fermions by taking a grand-canonical trace as in equation (12)—the essential step at the heart of the DQMC framework, which if missing does not allow numerical simulations of the fermionic system at feasible numerical cost.

The key idea in our approach is to represent the replica scheme in a setting where the statistical weights can again be simplified to a form relying on a single Slater determinant thus allowing to take a grand-canonical trace of the form (12). This is achieved by artificially enlarging the system by considering an additional copy $B'$ of subsystem $B$, which we will use to selectively evolve subsystem $B$ in imaginary time. In particular we will show that an imaginary-time Hamiltonian of the form

$$\tilde{H}(\tau) = \mathcal{H}_{AB} \Theta(\tau) \Theta(\beta - \tau) + \mathcal{H}_{AB'} \Theta(\tau - \beta) \Theta(2\beta - \tau)$$

will give direct access to the partition sum $Z[A, 2, T]$. A world line representation of this Hamiltonian is given in figure 3. To see this, suppose that we are given (14) as a

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model Hamiltonian and our task is to determine its physics at an arbitrary temperature, suggestively written as \( 2\beta \). We denote a given basis state by \( |\psi\rangle = |A, B, B'\rangle \), which leads to a partition sum of

\[
\tilde{Z} = \sum_{\{A, B, B'\}} \langle A, B, B'| \exp(-\beta H_{AB}) \exp(-\beta H_{AB})|A, B, B'\rangle.
\]

Although there are two propagation operators, the weight of the system would still be given as a single Slater determinant because only one expectation value has to be evaluated. We proceed to insert a resolution of unity in between the two exponential operators to obtain

\[
\tilde{Z} = \sum_{\{A, B, B', C, D, D'\}} \langle A, B, B'| \exp(-\beta H_{AB})|C, D, D'|\langle C, D, D'| \exp(-\beta H_{AB})|A, B, B'\rangle.
\] (15)

Notice that in the right term, states of \( B' \) do not appear in the Hamiltonian. Thus, independent of the specific form of \( A \) and \( B \), we need to have \( B' = D' \) for any non-vanishing term contributing to this partition function. Similarly, one obtains \( B = D \) from inspecting the left term. Further, if subsystems \( B \) and \( B' \) do not appear in the Hamiltonian for a given imaginary time interval they not only do not evolve, but also remain completely decoupled from the rest of the system over this interval. As a result, they will also not affect the statistical weight

\[
\langle C, B, B'| \exp(-\beta H_{AB})|A, B, B'\rangle
\]

\[
= \langle [C, B] \otimes \langle B'| \rangle \exp(-\beta H_{AB})([A, B] \otimes |B')\rangle
\]

\[
= \langle C, B| \exp(-\beta H_{AB})|A, B\rangle
\]

and thus can safely be ignored. Finally, renaming \( C \) to \( A' \) to match our earlier notation we obtain the following simplified expression

\[
\tilde{Z} = \sum_{A, A', B, B'} \langle AA'| \exp(-\beta H_{AB})|A', B'\rangle \langle A'B| \exp(-\beta H_{AB})|AB\rangle = Z[A, 2, T],
\] (16)

which is precisely the expression for the sought-after partition function \( Z[A, 2, T] \). We have thus shown that one can indeed recast the partition function \( Z[A, 2, T] \) in a way that relies only on single determinants thus allowing us to take the grand-canonical trace (12).

2.2.1. Ground-state formulation. When considering the ground-state DQMC algorithm (see appendix A.2) only minor modifications to the above scheme have to be implemented. The normalization constant \( \mathcal{N} \) introduced in (6) is now given as

\[
\mathcal{N} = \langle \psi | \psi \rangle = \sum_{|A\rangle} \langle \psi | A \rangle \langle A | \psi \rangle = \text{Tr} (|\psi \rangle \langle \psi |).
\]

The ground-state wavefunction \( |\psi\rangle \) is obtained by a projective scheme

\[
|\psi\rangle = \lim_{\theta \to -\infty} e^{-\theta H} |\psi_T\rangle,
\] (17)

applied to a test wavefunction \( |\psi_T\rangle \). If the test wavefunction has a non-zero overlap with the actual ground-state wavefunction, this projective scheme should eliminate all contributions from excited states and converge to the ground-state wavefunction.
Inserting this projection into the definition of the Rényi entropy in equation (8), we find an expression for the canonical trace $\text{Tr} \rho^2_A$ very similar to the finite temperature expression of the grand-canonical trace for $\hat{Z}$ in equation (16) discussed above

$$\text{Tr} \rho^2_A = \lim_{\theta \to \infty} \sum_{A,A',B,B'} \langle A'B' | \exp(-\theta H) | \psi_T \rangle \langle \psi_T | \exp(-\theta H) | A'B' \rangle$$

$$\times \langle A' B | \exp(-\theta H) | \psi_T \rangle \langle \psi_T | \exp(-\theta H) | AB \rangle,$$

where the only difference is the appearance of the density matrices $|\psi_T \rangle \langle \psi_T |$, which however come in handy when taking the grand-canonical trace to make use of equation (12). Precisely because of the occurrence of these density matrices only states with an occupation number identical to the one of the test wavefunction will contribute.

2.2.2. Higher Rényi entropies. While we have concentrated our discussion on Rényi entropies of order 2, it should be noted that our algorithm can be extended in a straightforward way to also compute higher Rényi entropies. For the calculation of the $n$-th Rényi entropy via the replica trick, imaginary time has to be split into $n$ segments which would contribute one determinant each in equation (13). It would thus be necessary to introduce $n$ replicas of the subsystem $B$ and work in an overall system of size $N_A + n \cdot N_B$. We have not implemented this more general case and therefore cannot comment on limiting system sizes or potential numerical instabilities arising in such an extended scheme.

2.2.3. Ensemble switching. The replica scheme outlined above allows us to directly sample partition functions of the form $\hat{Z}[A, 2, T]$. For the calculation of the Rényi entropy, however, we are really interested in determining the ratio of the partition functions $\hat{Z}[A, 2, T]$ and $\hat{Z}^2$ as given in equation (8). This ratio can be directly accessed [25] without explicitly calculating the individual partition functions but by cleverly switching between the two ensembles.

Imagine a two state simulation, where the weight of two states is given by $w_1$ and $w_2$. For any simulation fulfilling detailed balance the random walk would spend $N_1 = w_1/(w_1 + w_2)$ steps in state 1 and $N_2 = w_2/(w_1 + w_2)$ steps in state 2. Thus, the ratio of the weights $w_1/w_2$ corresponds precisely to the relative time spent in the two respective states. We can readily generalize [25] this statement to a situation where we sample a random walk switching back and forth between two ensembles whose partition functions equal the weights $w_1 = \hat{Z}[A, 2, T]$ and $w_2 = \hat{Z}^2$, respectively. Thus the ratio of relative time spent sampling each of the two ensembles can then be used to calculate the entanglement entropy

$$S_2(A) = - \log \left( \frac{\hat{Z}[A, 2, T]}{\hat{Z}^2} \right) = - \log \left( \frac{N_1}{N_2} \right).$$

In practical terms, we start our simulation in one of the two ensembles and after a fixed number of Monte Carlo steps, we calculate the weight of the current configuration in both ensembles and switch ensembles according to Metropolis rules, see figure 4.

When implementing this ensemble switching method, one benefits from an additional advantage of the determinantal QMC framework. The configuration space of

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\begin{align}
C(Z[A, 2, T]) & \quad C(Z^2) \\
C(Z[A, 2, T]) & \quad C(Z^2)
\end{align}

Figure 4. Schematic illustration of the ensemble switching method to calculate the Rényi entropy of equation (8). We start a random walk in one of the configuration spaces corresponding to the ensembles appearing in the numerator and the denominator of equation (8), denoted by \(C(Z[A, 2, T])\) and \(C(Z^2)\), respectively. Whenever we encounter a configuration that is admissible in both ensembles, we compare their relative weights and decide in which ensemble we continue to sample configurations based on a Metropolis scheme.

\(Z[A, 2, T]\) and \(Z^2\) is exactly equal, and the transition probabilities \(p_{1 \to 2}\) and \(p_{2 \to 1}\) are typically spread over the entire range \((0, 1]\). In appendix A.4 we show that the Monte Carlo estimate for the Rényi entropy is actually given as the ratio

\[
\frac{N_1}{N_2} = \frac{\langle p_{2 \to 1} \rangle}{\langle p_{1 \to 2} \rangle},
\]

which can be obtained by two separate simulations. This allows for much quicker convergence than we would obtain by explicitly switching between ensembles and counting how much time we spent in each one of the two ensembles.

It should further be noted that our approach does not necessarily require the iterative build up of the subsystem \(A\) from smaller blocks to achieve convergence [23], as it has been observed in the context of world line Monte Carlo approaches where the overlap between the ensembles might become rather small. We note that such an iterative build-up is also possible in the context of our DQMC approach if needed.

3. The Hubbard chain

To demonstrate the applicability and numerical efficiency of our replica switching method to calculate Rényi entropies within the DQMC framework we study the entanglement of a one-dimensional Hubbard chain, which at half-filling does not suffer from the infamous sign problem. We first concentrate on the zero-temperature properties of the entanglement entropy. We compare our numerical results to the quasi-exact analytical form derived from the conformal field theory description of the gapless theory governing the zero-temperature physics of the Hubbard chain in the presence of a finite on-site interaction \(U\). We then turn to finite-temperature properties and show how the Rényi entropy crosses over from a low-temperature entanglement entropy to a high-temperature thermal entropy. Finally, we discuss the strength of our approach in comparison to a recent proposal to calculate Rényi entropies for interacting fermion systems from a decomposition in free fermion Green’s functions [21]. We demonstrate
that our approach is significantly more efficient in capturing the entanglement properties in the interaction dominated regime of the Hubbard model.

3.1. Zero-temperature physics

In the presence of a repulsive on-site interaction $U > 0$ the ground state of the half-filled Hubbard chain is well known to be a Mott insulator exhibiting quasi-long range antiferromagnetic order. This means that at zero temperature charge fluctuations are frozen out entirely for any $U > 0$, while the localized spin degrees of freedom interact via an effective Heisenberg exchange of order $t^2/U$ thereby building up quasi-long range antiferromagnetic order. The system thus remains gapless and can be described in terms of a conformal field theory with a central charge $c = 1$ corresponding to the number of gapless modes.

The entanglement entropy of such a gapless one-dimensional system is known to exhibit a logarithmic correction to the boundary law, which for a one-dimensional system simply states that the entanglement entropy is a constant for any bipartition. The logarithmic correction, however, does reflect the relative size of the two subsystems in the bipartition and for all Rényi entropies follows the general form

$$S_\eta(l_A) = \frac{c}{6\eta} \left( 1 + \frac{1}{n} \right) \ln \left[ \frac{\eta L}{\pi} \sin \left( \frac{\pi l_A}{L} \right) \right] + O(1),$$

where $c$ is the central charge of the conformal field theory, $L$ is the overall system length and $l_A \leq L$ is the length of subsystem $A$. Open and periodic boundary conditions correspond to $\eta = 2$ and $\eta = 1$, respectively, and further subleading corrections of order $O(1)$ in the system size are neglected. Numerical results obtained with the zero-temperature DQMC algorithm (for details see the appendix) for an open chain of length $L = 64$ are shown in figure 5. We find that the DQMC data is generally in good agreement with quasi-exact results obtained using density matrix renormalization group (DMRG) simulations. We do observe, however, a slight trend of the DQMC data to underestimate (within the statistical error bars) the entanglement entropy for some of the intermediate block sizes—an effect which we find to be absent for smaller system sizes (not shown) and which previously has also been observed in conceptually similar simulations for spin systems using the replica trick in combination with a stochastic series expansion (SSE)\(^1\).

We thus conclude this section with the observation that our replica switching DQMC method is indeed well equipped to efficiently determine the zero-temperature entanglement properties of the half-filled Hubbard chain.

3.2. Thermal crossover of the entanglement

When considering a quantum system at finite temperatures, both quantum and thermal fluctuations contribute to all entropies including the Rényi entropy of interest here.

\(^1\) In the context of DQMC simulations, one additional potential origin for the systematic bias might be the discretization of imaginary time. In our simulations it was generally set to a value of $\Delta \tau = 0.125$, which for the studied onsite interaction of $U = 2.0$ is smaller than typically needed [31].
Increasing the temperature the relative contributions of these two types of fluctuations of course change. As a result the Rényi entropy shows a crossover from a boundary law (with logarithmic corrections) at zero temperature to a more conventional extensive behavior (i.e. a volume law) at high temperature of the form

\[ S(l_A) = l_A \log 4, \]

simply counting the number of possible states in the subsystem.
This thermal crossover of the Rényi entropy from a zero-temperature entanglement entropy to a thermal entropy at high temperatures can easily be observed in our numerical DQMC simulations. This is illustrated in figure 6 for a half-filled Hubbard chain of length $L = 32$ with intermediate on-site interaction $U/t = 2$ in a temperature range $0.025 \leq T \leq 5$ (for $t = 1$). With increasing temperature the arc-like structure of the low-temperature entanglement entropy disappears and gives way to the simple linear form of an extensive thermal entropy. This thermal crossover is also reflected in figure 7 where we plot the Rényi entropy $S_2(L/2)$ of an equal-size bipartition of the chain for different system sizes versus temperature. In particular, we observe the expected data collapse at high temperatures when rescaling the calculated Rényi entropies by the respective system size, see the right panel of figure 7.

To quantitatively determine the crossover temperature $T^*$, below which a finite-sized system is effectively in its ground state, we turn to the so-called purity

$$P = \exp (-S_2(L)),$$

which becomes 1 for a quantum mechanical ground state, since the entropy $S_2(L)$ needs to equal its complement $S_2(\emptyset)$ and thus must vanish for any quantum mechanical ground state—an observation which is ultimately also responsible for the arc-like structure of the entanglement entropy in equation (8). Indeed we find that the purity sharply rises towards 1 as the temperature is lowered in our simulations, see figure 8 where we plot the purity as a function of temperature for different system sizes and a sequence of on-site interactions. On the one hand, we find that for a fixed value of the on-site interaction the crossover temperature decreases with system size in accordance with the fact that the finite-size gap of the system also decreases with increasing system size. On the other hand, we observe that for fixed system size the transition temperature $T^*$ decreases as the on-site interactions $U$ are reduced reflecting the enhancement of charge fluctuations in this weakly coupled regime.
Finally, we note that we generally find somewhat smaller transition temperatures than recent stochastic series expansion (SSE) simulations [29] of the half-filled Hubbard chain, which can be tracked back to the fact that our DQMC simulations employ a grand-canonical ensemble, while the SSE simulation in [29] employed a canonical ensemble.

3.3. Comparison to free fermion decomposition method

We round off our discussion of our replica switching DQMC method by comparing it to a recent proposal [21] to calculate Rényi entropies from a decomposition in free fermion Green’s functions. Such a decomposition might seem natural within the DQMC approach, since the Hubbard–Stratonovich transformation at its heart results in an effective description of free fermions moving in an external potential.

In the spirit of a fair comparison we have implemented the free fermion decomposition method [21] using the same algorithmic optimizations as for our replica switching technique whenever possible. We then ran both codes on identical parameter sets logging the exact same CPU time for both codes. Results from this comparison are
summarized in figure 9 where we show results of both approaches for the entanglement entropy of a half-filled 10-site Hubbard chain at fixed temperature $T = 0.025$ ($\beta = 40$) for different values of the on-site interaction $U/t \in \{1, 2, 4\}$. While the free fermion decomposition method reproduces the arc-like structure of the entanglement entropy for small on-site interaction $U/t = 1$, it shows deviations from this behavior already for moderate values of the on-site interaction $U/t = 2$. By contrast, our replica switching method nicely reproduces the exact diagonalization data up to strong on-site interaction $U/t = 4$. We thus conclude that our approach is significantly more efficient in capturing the entanglement properties in the interaction dominated regime of the Hubbard model.
4. Conclusions

In summary, we have introduced a novel replica switching scheme to efficiently calculate Rényi entropies for interacting fermionic systems in determinantal quantum Monte Carlo simulations. Our approach is capable of efficiently determining not only finite-temperature thermal entropies but also zero-temperature entanglement entropies as demonstrated for the half-filled Hubbard chain over a range of interactions. In comparison to a recent proposal to calculate Rényi entropies from a decomposition in terms of free fermion Green’s functions [21], our approach yields much faster convergence and significantly higher numerical efficiency in the regime of strong correlations $U/t > 1$.

While we have concentrated our discussion on the second Rényi entropy $S_n = 2$, it should be noted that the replica technique described here can be expanded in a straightforward way to also access higher Rényi entropies. Calculating Rényi entropies with higher indices $n > 2$, however, requires one to simulate a system of size $N_A + n \cdot N_B$ at temperature $n\beta$ (where $N_A$ and $N_B$ correspond to the size of subsystems $A$ and $B$, respectively). Thus, the computational cost to access these higher Rényi entropies in our approach sharply increases as the DQMC simulations generally scale as $\beta N^3$ (where $N$ is the total number of sites). This should be contrasted to the free-fermion decomposition method of Grover [21], where the $n$-th Rényi entropy can readily be accessed by simply simulating $n$ replicas of the system at temperature $\beta$—a much more moderate increase in computational expense. Indeed recent numerical simulations [30] for the Kane–Mele model have demonstrated that with the free fermion decomposition method it is feasible to calculate a partial entanglement spectrum, i.e. the spectrum of (lowest) eigenvalues of the density matrix, from a sequence of Rényi entropies. It is likely that such a calculation of the entanglement spectrum is out of reach for our technique.

Although we have focused our discussion on one-dimensional fermion systems in this manuscript, two dimensional systems can be treated equally well. In particular, we point out that the numerical overhead of our method arising from the simulations of the two copies $B$ and $B'$ (see figure 3) for the second Rényi entropy reduces with increasing spatial dimensionality: In one dimension an equal-size bipartition results in an overhead of $L/2$ additional sites simulated at temperature $2\beta$. In two spatial dimensions, where we might consider a subsystem $A$ of size $L/2 \times L/2$ and its somewhat larger complement, we can choose to simulate the smaller subsystem (in this case $A$) twice. Such a simulation would thus only need an overhead of 1/4 of the sites in comparison to a conventional simulation and thus make it possible to calculate Rényi entropies for systems of nearly the same size as conventionally investigated in DQMC simulations.

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Appendix A. DQMC primer

We will give a short introduction to the basics of determinantal Quantum Monte Carlo that should suffice to understand the modifications to it presented in the main text. There are many more extensive, excellent reviews of the DQMC method available, such as [31, 32] among others, on which our exposition is based. We will start with the finite temperature algorithm and later mention what modifications are necessary to perform ground state simulations. For concreteness, we will stay within the context of the Hubbard model.

A.1. Finite-temperature algorithm

The first step in setting up the algorithm is to Trotter decompose imaginary time:

\[
\mathcal{Z} = \text{Tr} \prod_i \exp \left( -\Delta \tau \mathcal{H} \right)
\]

\[
= \sum \langle \psi | \exp \left( -\Delta \tau \mathcal{H} \right) \ldots \exp \left( -\Delta \tau \mathcal{H} \right) | \psi \rangle.
\]  

The occupation number operator \( \mathcal{N} \), needed for the chemical potential, is from now on included in the potential operator \( \mathcal{V} \). The exponential is now separated and the appearing commutator ignored which in turn results in a systematic error of the order \( O(\Delta \tau) \):

\[
e^{A+B} \approx e^A e^B \Rightarrow e^{-\Delta \tau (\mathcal{K}+\mathcal{V})} \approx e^{-\Delta \tau \mathcal{K}} e^{-\Delta \tau \mathcal{V}} + O(\Delta \tau).
\]

The kinetic part \( e^{-\Delta \tau \mathcal{K}} \) has only two operators and can be evaluated directly. The potential part \( e^{-\Delta \tau \mathcal{V}} \) on the other hand is made up of four operators and can therefore not be integrated out analytically. We thus apply the Hubbard–Stratonovich transformation to decouple the interaction. It is based on an identity for Gaussian integrals

\[
e^{\frac{1}{2} s^2} = \sqrt{2\pi} \int_{-\infty}^{\infty} ds \, e^{-\frac{1}{2} s^2 - sA},
\]  

with \( A \) being the original operator, in our case the on-site interaction \( \mathcal{V}_i = U n_{i,\uparrow} n_{i,\downarrow} \). The price we have to pay is the introduction of the eponymous auxiliary field \( s \) that couples to the fermions. The decoupling itself is not unique: we can decouple with respect to the local charge or to the local magnetization. For the repulsive case, we usually couple to the local magnetization to avoid a complex transformation which is nevertheless possible and can be advantageous [33]. We do, however, break \( SU(2) \) symmetry which is only restored when performing enough updates.

It turns out that in the case of the Hubbard model, it is not necessary to work with a continuous auxiliary field, but that we can choose to work with discrete Ising spins \( s \) taking values \( \{-1, 1\} \) [34]. The decoupling has to be performed for each site and at each time slice. Doing so, we obtain

\[
e^{-U \Delta \tau n_{i,\uparrow} n_{i,\downarrow}} = \frac{1}{2} e^{-\frac{1}{2} \Delta \tau U n_i} \sum_{s=\pm 1} e^{-\lambda s m_i} = \frac{1}{2} \sum_{s=\pm 1} \prod_{\sigma=\uparrow,\downarrow} e^{-\left( s \sigma + \frac{1}{2} U \Delta \tau \right) n_{i,\sigma}}
\]
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![General Matrix](image1) × ![B](image2) = ![Result](image3)

**Figure A1.** The computational cost of the algorithm due to the additional sites is lowered by exploiting the matrix structure induced by the imaginary time dependent Hamiltonian (14). The slice matrices comprise blocks of zeros which should explicitly be ignored in steps involving matrix multiplications.

The parameter $\lambda$ can be determined by inserting all possible values for the auxiliary field $s = \pm 1$ and the spins $\sigma = \{\uparrow, \downarrow\}$. We find

$$cosh \ \lambda = e^{2|t|\Delta \tau}.$$  

Inserting this into (A.2) and switching to an explicit vector notation for the operators, we obtain the following form for the partition function:

$$Z = \left(\frac{1}{2}\right)^{N' L} \text{Tr}_s \text{Tr} \prod_{l=L}^{1} \prod_{\sigma = \uparrow, \downarrow} e^{\Delta \tau_c K c e^{-\Delta \tau_c V} e^i c},$$

where $K$ and $V$ are the matrix representations of the $K$ and $V$ operator, respectively.

The partition function now consists of one trace over all auxiliary field configurations and another one over the fermionic states which act on a product over all time slices and spins of the discretized and separated exponential.

We continue to rewrite the operators in matrix form $K$ and $V_s$ where the index $s$ reminds us that the potential part depends on the auxiliary field. A product $e^{\Delta \tau_c K c e^{-\Delta \tau_c V} e^i c}$ will be denoted by $B^s(l)$ and an ordered sequence of all $L \ B$-matrices with arbitrary starting point $\tau$ by

$$B^s(\tau) = B^s(\tau - 1) ... B^s(0) B^s(\beta) ... B^s(\tau).$$  

(A.4)

The time $\tau$ is a multiple of the discretization $\Delta \tau$. The weight of one configuration of fermions and auxiliary field is then given by

$$\langle \psi | B^s(\tau) | \psi \rangle = \text{det} (B^s(\tau)).$$  

(A.5)

We could use this weight to set up the Monte Carlo simulation, sampling both fermion and auxiliary field configurations. But this would be very inefficient because the calculation of determinants is numerically very expensive. Instead, we will integrate out the fermions analytically and sample only auxiliary field configurations.

One can prove that the *grand-canonical* trace over all fermionic states of the $B$-matrix is given by

$$\text{Tr} B = \text{det} (1 + B),$$  

(A.6)

allowing us to explicitly perform the trace over all fermionic states. One can hardly underestimate the importance of this identity for determinantal QMC simulations. By
applying the identity (A.6), we are left only with the problem of sampling the auxiliary field where the weight of each configuration is given by one determinant. The connection to the field is made by the $B$ matrices via the potential. Using these notations, we can rewrite the partition function as

$$Z = \left(\frac{1}{2}\right)^N \sum_{\sigma} \prod_{\sigma} \det (1 + B^\sigma(0)).$$

In principle, we are now able to set up our Monte Carlo simulations. We know the form of the weights and we can sample configurations of auxiliary fields using, for example, the Metropolis scheme mentioned before. For the calculation of Green’s functions and more practical aspects we refer the interested reader to the aforementioned references.

### A.2. Ground-state algorithm

To access ground state properties, we start with a carefully chosen trial wavefunction $|\psi_T\rangle$ and project out the excited states by applying a large power of the Hamiltonian:

$$\lim_{\Theta \to \infty} e^{-\Theta H} |\psi_T\rangle = |\psi\rangle.$$  \hspace{1cm} (A.7)

The trial wavefunction must be non-orthogonal to the true ground state wavefunction $|\psi\rangle$ for the procedure to work. For the Hubbard model at half filling we chose a spin singlet as a trial wavefunction $|\psi_T\rangle$. Setting up the simulation is very similar to the finite temperature algorithm. The projection parameter $\Theta$ plays the role of the temperature $\beta$. The exponential in (A.7) is then decomposed and the interaction term decoupled by a Hubbard–Stratonovich transformation. We will not go into further detail because the calculation of the Rényi entropies for ground state problems was shown to reduce to a modified finite temperature problem.

### A.3. Numerical optimizations

In DQMC simulations we are limited by the $N^2\beta$ scaling and have to pay special attention to numerical instabilities. Simulating an artificially enlarged system may thus appear to be very inefficient. However, enlarging the system and implementing the Hamiltonian (14) goes along with a special matrix structure that we can make use of to lower the numerical cost. In the matrix representation, the hopping terms are of the following form:

$$\mathbf{K}_{AB} = \begin{pmatrix} K_{AA} & K_{AB} & 0 \\ K_{BA} & K_{BB} & 0 \\ 0 & 0 & 0 \end{pmatrix}, \quad \mathbf{K}^*_{AB'} = \begin{pmatrix} K_{AA} & 0 & K_{AB'} \\ 0 & 1 & 0 \\ K_{B'A} & 0 & K_{B'B'} \end{pmatrix},$$

where $K$ is the matrix representation of the hopping operator and the index signifies the part of the lattice it connects. One of the parts $B$ and $B'$, respectively, remains unchanged and thus has a 1 on the diagonal and zeros for the off-diagonal terms. Interaction terms will be of the same form.

One can now readily convince oneself that multiplying two matrices in the same imaginary time interval $(0, \beta)$ or $(\beta, 2\beta)$ will not alter this structure because they are block-diagonal, see figure A1. Mixing matrices of different time intervals, on the other
hand, will typically result in dense matrices. This is inevitably the case when calculating the sequence of $B$ matrices (A.4), except for $\tau \in \{0, \beta\}$. Even when multiplying a dense matrix with one of the slice matrices, it is not necessary to perform a full matrix multiplication, because of the blocks of zeros present in the slice matrix.

### A.4. The ensemble switching method

We will show that to estimate the ratio \( (8) \) it suffices to measure the expectation values of \( \langle p_1 \rightarrow 2 \rangle \) and \( \langle p_2 \rightarrow 1 \rangle \), which are the probabilities to switch ensembles from ensemble 1 to 2 and from ensemble 2 to 1, respectively. Using a Metropolis scheme, the probability to switch ensembles for a given configuration $c$ is given as

\[
p_{1 \rightarrow 2} = \min \left( 1, \frac{W_2(c)}{W_1(c)} \right),
\]

where $W_1(c)$ and $W_2(c)$ are the weights in the respective ensembles. The configurations $c$ are configurations of the auxiliary field and both ensembles share the same configuration space $C$. Writing out the ratio of the expectation values for the switching operation gives

\[
\frac{\langle p_{1 \rightarrow 2} \rangle}{\langle p_{2 \rightarrow 1} \rangle} = \frac{Z_2}{Z_1} \sum_{c \in C(Z_2)} \min \left( 1, \frac{W_2(c)}{W_1(c)} \right) \frac{W_1(c)}{W_2(c)}
\]

\[
= \frac{Z_2}{Z_1} \sum_{W_2(c) < W_1(c)} W_2(c) + \sum_{W_1(c) < W_2(c)} W_1(c)
\]

\[
= \frac{Z_2}{Z_1} \sum_{W_2(c) < W_1(c)} W_2(c) + \sum_{W_1(c) < W_2(c)} W_1(c)
\]

### Appendix B. Comparison to free fermion decomposition

In this appendix we provide further details on our analysis of the comparison between the replica switching and free fermion decomposition DQMC algorithms as presented in figure 9 of the main text.

Our C++ simulation codes of the two algorithms employ a common code base implementing the same optimizations for many underlying core features (such as linear algebra subroutines, sampling improvements or parallelization schemes) for both approaches and further build on the ALPS libraries [35].

We separately ran 16 independent simulations per data point in figure 9 of the main text each with a different random seed and later merged the results via a jackknife analysis using the ALPS tools. The spread of the raw results of all 16 runs before merging are shown in figure A2.

For an allocated computing time of 105 min (per data point and seed) the number of measurements in the free fermion case was around 3500 per data point and seed.
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For the replica switching method, we obtained a considerably smaller number of measurements for the switching probabilities in (20) in the same allocated computing time. For the largest cut, i.e. \( l_A = 10 \), we recorded some 1300 measurements, while for the smallest cut, i.e. \( l_A = 1 \) we recorded only around 100 measurements. The lower number of measurements for the ensemble switching method in a given time frame is due to two effects: First, for the replica switching method we have to perform one simulation for an ensemble of two separate systems and one simulation in the connected ensemble. Both of these simulations have to be thermalized in contrast to only one simulation in the free fermion case. Second, the simulation cell of the connected system in the replica switching method is enlarged. For the case of \( l_A = 1 \) for example, we effectively simulate a system of size \( L = 19 \) which in combination with the \( N^3 \beta \) scaling of the algorithm further reduces the number of possible sweeps in a given time.

Figure A2. Comparison of the spread of the raw data of DQMC runs for 16 independent runs using the replica switching (squares) and free fermion decomposition (circles) algorithms. Shown is the Rényi entropy of a half-filled Hubbard chain with varying on-site interactions \( U/t \) at temperature \( T = 0.025 (\beta = 40) \).
Comparing the results obtained with the two algorithms the spread of the raw data shows two overall trends. First, looking at the dependence of the spread of data points with subsystem size $l_A$, the free fermion decomposition method shows a considerable increase of this spread with increasing subsystem size, while for the replica switching method there is no measurable dependence. Second, with increasing onsite interaction $U$ the data spread clearly increases much more strongly for the free fermion decomposition method in comparison with the replica switching technique (which becomes most poignant for large subsystem sizes $l_A$).

References