Direct simulations of small multi-fermion systems

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Abstract

I explore computer simulations of the dynamics of small multi-fermion lattice systems. The method is more general, but I concentrate on Hubbard type models where the fermions hop between a small number of connected sites. I use the natural mapping of fermion occupation numbers onto computer bits. Signs from fermion interchange are reduced to bit counting. The technique inherently requires computer resources growing exponentially with the system volume; so, it restricted to modestly small systems. Large volume results would require combining these techniques with further approximations, perhaps in a recursive renormalization group manner.

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Quantum systems involving fermions have proven elusive for computer simulation. A plethora of cancellations in all but a few cases impedes the use of Monte Carlo methods, which have been so successful for bosonic systems. Here I explore the direct application of the relevant Hamiltonians to wave function representations stored in computer memory. I work with large but sparse matrices acting in a finite dimensional Hilbert space. As the methods are inherently exponential in volume, I concentrate on smaller systems. Thus I am admitting defeat in terms of a thermodynamic limit, hoping instead to find interesting physics with finite systems. To obtain results for larger volumes, these techniques could form the basis for approximate techniques, such as solving small blocks of variables to form the starting point for a renormalization group approach. Note that I am discussing directly simulating the behavior of quantum systems on a classical computer. In this way one can also study various models for quantum computers. The storage required does grow exponentially with the number of q-bits under study, but if this is fairly modest, say of order 20, the methods exploited here are straightforward.

Consider a Fock basis \(|n_0, \ldots, n_{N-1}\rangle\) for a many fermion system, where the \(n_i\) are the occupation numbers for some set of orthogonal single particle states. Each \(n_i\) is either 0 or 1. For a lattice model these might represent the occupations on given sites with given spins. This basis naturally maps onto computer words, which are sets of bits also either 0 or 1. A 1 bit is “set” and a 0 bit “unset.” This raw mapping is instinctive for numerical simulations of multi fermion systems. A subroutine representing a creation operator for a fermion would set the corresponding bit in the appropriate word. An annihilation operator resets the bit. Such can all be done with simple bitwise logical operations. This naive observation, however, requires embellishment so that fermion exchanges will give rise to the appropriate relative negative signs. Simple bit counting techniques allow us to track these.

To start, consider \(N\) creation/annihilation pairs \(\{a_i^\dagger, a_i\}\) for \(0 \leq i < N\). These satisfy the usual fermion commutation relations

\[
\begin{align*}
[a_i, a_j^\dagger]_+ &= a_ia_i^\dagger + a_j^\dagger a_i = \delta_{ij} \\
[a_i, a_j]_+ &= 0
\end{align*}
\]

The vacuum state \(|0\rangle\) is annihilated by all \(a_i\); i.e. impose \(a_i|0\rangle = 0\) for every \(i\). A general Fock state is given by applying creation operators to this state

\[|f\rangle = |n_0, \ldots, n_{N-1}\rangle = (a_0^\dagger)^{n_0} \ldots (a_{N-1}^\dagger)^{n_{N-1}} |0\rangle\]
Each of the occupation numbers \( n_i \) is either zero or one. Note that my sign conventions are buried in the ordering convention with increasing index. The concept being explored here identifies each of these basis states with an integer \( n(|f\rangle) \) whose binary representation is given by the corresponding occupation numbers

\[
n(|f\rangle) = \sum_{i=0}^{N-1} n_i 2^i
\]

Given a Fock state, the occupation number of a particular site is then easily determined by testing whether the bitwise logical operation \( n \& (1 \ll i) \) is zero or not. (I assume the reader is familiar with standard C notation for logical operations.)

Of course, in a computer the natural word length is finite, 32 bits in today’s typical personal computer, 64 in most larger machines. If we want to study systems of more fermions, we need to combine several words into a higher precision integer. This technicality is straightforward, and I will not discuss it further here.

A general quantum state is a superposition of states in this Fock basis

\[
|\psi\rangle = \sum_n \psi_n |n\rangle
\]

This involves specifying a complex number \( \psi_n \) for each integer representing one of our Fock states. The computer storage required to hold this information grows exponentially. Indeed, if I need to keep all \( N \) bit states, I need storage for \( 2^N \) complex numbers. In many cases symmetries allow us to reduce this number considerably, although the basic growth with \( N \) remains exponential.

One particularly useful symmetry occurs when fermion number is conserved. Then for a given filling \( N_f \), we only need to keep track of integers up to \( 2^N \) containing exactly \( N_f \) set bits. For half filling this involves a memory saving by a factor of order \( \sqrt{\pi N/2} \), while for other fillings the saving is greater. If we deal with two species of fermions each of which is separately conserved, such as the Hubbard type models discussed later, the resulting savings can be even more. To enumerate states with a given total occupation it is useful to have a function that returns the next integer with the same number of bits as its argument; an implementation is discussed briefly in Appendix A.

Another symmetry in many systems is translational invariance, which manifests itself in momentum conservation. For a given momentum, states which are related by translation have their wave functions related by a phase, and thus only one of them needs to be stored. For my discussion here I will not make use of this symmetry.
So given $N_s \leq 2^N$ states to be stored, we can do this in various ways. One is to use a hash table, keying wave function components to the corresponding Fock states, as discussed in [1]. If the Fock states are ordered in some manner, one can instead use a binary search. This is the method used here, where for each wave function under consideration I keep an array of $N_s$ complex numbers. By having the state table ordered, a given state can be quickly located. Then the corresponding location in the coefficient table contains the desired component of the wave function.

As storage represents the main bottleneck in this type of algorithm, tricks to reduce this are desirable. If we can deal with a real Hamiltonian, the wave function storage drops by a factor of two. With several species each separately conserved, the table of states separates into multiple tables representing the Fock states for the individual species. In this case the bulk of the storage is for the complex numbers representing the wave function. In any case, the type of simulation discussed here spends the dominant amount of its computer time doing the searches through these tables for desired states. Floating point arithmetic operations tend to be insignificant.

I now become more specific and consider an annihilation operator $a_j$. I implement this as subroutine which takes as arguments the specific location $j$ and a pointer to a Fock state labeled as above by an integer $n$. It should return zero unless bit $j$ is set in $n(|f\rangle)$. A simple test for this is whether, the integer $n\&(1 << j)$ is non-vanishing. When this is true the function returns the corresponding sign obtained after the annihilation flips this bit. The resulting Fock state is represented by $n \wedge (1 << j)$. The sign returned is associated with bringing the operator $a_j$ into the canonical ordering above. This is determined by the parity of the number of set bits lower than $j$. This can be quickly found by logical operations, beginning with a masking off of the lower bits by the considering the integer $n\&(1-(1<<j))$. We need to include a negative sign if the population count of this integer is odd. The population count of an integer is the number of set bits it contains. Appendix A discusses one simple way to implement such a count. For a creation operator, one does exactly the same thing except checking that the initial bit is not set.

To apply some combination of creation and annihilation operators to a wave function $|\psi\rangle$ involves looping over all the component Fock states $|f\rangle$. On each of these in succession we apply the above subroutines, multiply the component of the wave function by the returned sign, and then store the result. For a simple hopping Hamiltonian, it is useful to make a
table of the bit locations for the neighbors of any given site. This small array is set up once at the beginning of a simulation. Then the hopping term in the Hamiltonian becomes another subroutine which loops over sites, spins, and neighbors. It successively applies the corresponding creation and annihilation pairs to the components of the source wave function. The results accumulate in a destination wave function.

These bit manipulations do not depend in any deep way on the type of fermion interaction used. Here for convenience I consider a simple Hubbard form \[2\]. Thus I will consider two types of fermions distinguished by “spin” and have them interact by adding an energy \( U \) for each site with both spin states occupied. Such an interaction is also easily implemented by logical operations; on looping over states one multiplies the coefficient of each component of the wave function by \( U \) times the number of doubly occupied sites. The latter is the population count of \( n_{↑} \& n_{↓} \).

To be more specific, consider a set of sites with nearest neighbors connected by bonds. On each site \( i \) the operator \( a_{i,s}^\dagger \) creates a fermion with a spin index \( s \in \{↑, ↓\} \). My Hamiltonian is

\[
H = U \sum_i n_{i,↑} n_{i,↓} - \sum_{\{i,j\},s} (a_{i,s}^\dagger a_{j,s} + a_{j,s}^\dagger a_{i,s})
\]

(6)

Where \( n_{i,s} = a_{i,s}^\dagger a_{i,s} \) and \( \{i, j\} \) denotes the set of neighboring pairs. For simplicity I set the energy scale to make the hopping parameter unity. In one dimension this model is exactly solvable \[3\]. For a more detailed discussion of this solution see \[4\]. Monte Carlo methods for this system are discussed in Ref. \[5\]. In more dimensions the Monte Carlo approach only works effectively for the half filled case \[6\].

I now illustrate several simple numerical “experiments.” I will first play with a six membered ring of sites, thus mimicking a benzene molecule. With a nearest neighbor hopping, we have the standard undergraduate example for using linear combinations of atomic orbitals to illustrate hybridization of the pi electrons. When \( U = 0 \) this has single fermion states with energies \(-2, -1, -1, 1, 1, 2\). Filling the lowest levels with three spin up and three spin down electrons gives a total ground state energy of -8. This is the energy gained from the delocalization of the electron wave functions. This should be compared with the value -6 which would be obtained from three fixed double bonds and no hybridization.

Turning on the Hubbard interaction raises the ground state energy. As \( U \) goes to infinity the ground state energy rises, with the largest components of the wave function alternating
spin up and down around the ring. They prefer to alternate rather than some other pattern since as long as $U$ is not infinity, this state maximizes delocalization. For this case of half filling, i.e. 3 electrons of each spin, there are 400 Fock states. This corresponds to twenty possible arrangements for each set of spins. This is not a particularly large matrix and could presumably be treated by conventional matrix methods, but it enables fast experiments with the table manipulation ideas discussed here. The computer time for these experiments is insignificant, easily practical on a PC.

For most of the following experiments I start with a random initial state. To construct such, I set each of the components of the wave function to a Gaussian random number and then normalize the state. From this I repeatedly apply the Hamiltonian in various ways discussed below. Formally I work with what is known as a Krylov space. Note that when there are degenerate states, a Krylov procedure does not span the full space, but leaves the relative contribution of the degenerate states unchanged. To separate them one must consider multiple starting states. For example, one can first extract a state of interest, construct a random state orthogonal to it, and construct a new Krylov space which will involve different combinations of the degenerate states. If the degeneracy is due to some symmetry, an alternative is to use starting states which are eigenvectors of this symmetry.

A particularly intuitive way to find the ground state is to start with a random state as above and directly apply $e^{-Ht}$. For large $t$ this should project out the ground state. For moderate $t$ one can use the rapidly convergent power series expansion for the exponential. For larger $t$ break the evolution into smaller time intervals and apply the exponentiated Hamiltonian repeatedly. In Fig. 1 I show the behavior of the expectation value of the energy as a function of $t$ for our benzene system where I take the parameter $U = 2$.

There are a variety of ways to obtain information on the first excited state from this experiment. The value of its energy can be extracted from the approach to the ground state via the formula

$$\langle E \rangle = E_0 + \alpha e^{-2(E_1-E_0)t} + \ldots$$  \hspace{1cm} (7)

Solving three successive times for $E_1$ gives the results shown by boxes in Fig. 2. Alternatively, as time evolves we can extract the part of $H|\psi\rangle$ that is orthogonal to $|\psi\rangle$ via the construction

$$|\psi_1\rangle = H|\psi\rangle \langle\psi|\psi\rangle - |\psi\rangle \langle\psi|H|\psi\rangle$$  \hspace{1cm} (8)

While $|\psi\rangle$ evolves, this should be dominated by the first excited state. Measuring the
FIG. 1: The expectation value of the energy in the state $e^{-Ht}|\psi\rangle$ with the initial $|\psi\rangle$ chosen randomly. The system is a six membered ring with Hubbard interaction $U = 2$. The expectation of the Hamiltonian in this state gives the points represented by bursts in Fig. 2. Both the above techniques will fail when $t$ is large enough that we have the ground state to machine precision.

The above evolution is effectively in “imaginary time,” and damps the system to its ground state. Since all signs are being included, one can also work in real time and calculate the evolution of a state under application of $e^{-iHt}$. Since this involves no damping, it will leave the expectation value of the energy unchanged. Fig. 3 shows a simple experiment where the initial state has all six fermions placed on the first three sites and then observes the expected occupation number of either spin on the sites as a function of time. Again I use $U = 2$. Observe the particles spreading towards a uniform distribution. Note that if I did not include the interaction $U$, then all energy levels are spaced by integer amounts and instead of a relaxation we have a sloshing of the fermions back and forth with a periodic return to the original state. Working in real time provides a means to study finite temperature, or more precisely, allows a micro-canonical evolution at energies above the ground state.

While working well for this small system, the calculation of $e^{-Ht}$ is somewhat tedious. A more efficient but still simplistic algorithm for finding the ground state is to repeatedly apply
FIG. 2: Extracting the first excited state. Considering $|\psi(t)\rangle = e^{-Ht}|\psi(0)\rangle$, the boxes represent the expectation of the Hamiltonian in the combination of $|\psi\rangle$ and $H|\psi\rangle$ that is orthogonal to $|\psi\rangle$. The bursts represent the energy of the first exited state extracted from three successive measurements of the expectation value of the energy. For comparison, the crosses replot the data from Fig. 1.

$H$ to the current state and then form the linear combination of $\psi$ and $H\psi$ that minimizes the expectation value of the energy. At each stage this requires calculating $H^2\psi$ as well, but is straightforward to implement. Fig. 3 shows the convergence of this procedure for our half filled benzene system with $U = 0$ and $U = 2$. Using this technique to find the ground state energy, I plot in Fig. 4 the ground state energy as a function $U$.

The methods described here do not have any “sign problems” since all signs are kept track of at all times. While the above experiments were done at half filling, there is nothing that requires this. In Fig 5 I show the ground state energy as a function of filling fraction for the 6 membered ring. I keep the spin up and spin down filling fraction the same. Note how at $U = 4$ the on-site repulsion is sufficiently strong to make the lowest energy state at $1/3$ filling, rather than the $1/2$ of the free case. The $U = 0$ points in this figure show the successive fermions contribute increasing values to the energy, demonstrating the Pauli exclusion principle as the lower levels are filled.

The standard technique for dealing with these large sparse matrices is the Lanczos scheme.
FIG. 3: The relaxation of the occupation number distribution as our wave function evolves in real time. The initial state has all fermions on the first three sites. For this picture $U = 2$.

FIG. 4: The expectation value of the energy as a function of the number of iterations, where for each iteration I take the linear combination the current state with the Hamiltonian on that state that minimizes the resulting energy expectation.
FIG. 5: The ground state energy of the Hubbard model on a 6 member ring as a function of the coupling $U$.

FIG. 6: The ground state energy of the Hubbard model on a 6 member ring as a function of the filling for three values of the coupling $U$. 
This iteratively constructs a sequence of vectors \( |g_i\rangle \) that form a basis under which the Hamiltonian is real and tri-diagonal; it has non-vanishing elements only between diagonal and sequential states in the sequence. The construction is recursive and makes use of an auxiliary sequence of states \( |d_i\rangle \) which satisfy a matrix orthogonality condition \( \langle d_i | H | d_j \rangle = 0 \) whenever \( i \neq j \). The construction starts from an arbitrary initial \( |g_0\rangle = |d_0\rangle \). The higher states are given by

\[
|g_n\rangle \propto H|d_{n-1}\rangle - |g_{n-1}\rangle \langle g_{n-1} | H | d_{n-1}\rangle
\]  
(9)

\[
|d_n\rangle \propto |g_n\rangle \langle d_{n-1} | H | d_{n-1}\rangle - |d_{n-1}\rangle \langle d_{n-1} | H | g_{n-1}\rangle
\]  
(10)

For convenience I choose the proportionality constants so that both vectors are normalized. The orthogonality of the \( |g_i\rangle \) and the matrix orthogonality of the \( |d_i\rangle \) are easily proven by induction. The matrix elements of the tri-diagonalized Hamiltonian can be calculated during the recursion without generating any additional vectors by expanding \( |d\rangle \)

\[
|d_i\rangle = \sum_0^i |g_i\rangle \langle g_i | d_i\rangle
\]  
(11)

and using the orthogonality constraints to obtain

\[
\langle g_{i+1} | H | g_i\rangle = \frac{\langle g_{i+1} | H | d_i\rangle}{\langle g_i | d_i\rangle}
\]  
(12)

\[
\langle g_i | H | g_i\rangle = \frac{\langle g_i | H | d_i\rangle - \langle g_i | H | g_{i-1}\rangle \langle g_{i-1} | d_i\rangle}{\langle g_i | d_i\rangle}
\]  
(13)

The iteration procedure should formally terminate at the dimension of the Krylov space generated by applications of \( H \) to \( |g_0\rangle \). If there are no degenerate eigenvalues and if \( |g_0\rangle \) has non-zero overlap with all states, then this is the dimension of our Hilbert space. However, any degenerate states cannot be separated in this process, and thus the dimension of the generated space is reduced by one for each degenerate state. In a practical simulation on a large system one will usually stop the series at much earlier stage. For a small system, however, the termination is usually signaled by an extremely large normalization factor in the construction of \( |g_n\rangle \), which would be zero were it not for finite machine precision. After this occurs the orthogonality with earlier \( g_i \) is lost.

In Fig. 4 I plot the eigenvalues of the truncated tri-diagonal Hamiltonian from our benzene system as a function of the number of steps taken in this process. The lowest and highest eigenvalues rapidly converge to their respective state energies, while new eigenvalues appear
FIG. 7: The eigenvalues of the Hamiltonian in the Lanczos basis truncated to size $n$, plotted as a function of $n$. Note how the highest and lowest eigenvalues stabilize the most quickly while new states appear between them. The horizontal lines are the ground and first excited state energies from the last points in Fig. 3.

in the middle of the spectrum. The Lanczos procedure converges more rapidly to the true ground state energy than the previous schemes. For long runs, however, it can become unstable as roundoff errors accumulate in the sequence. The approach also gives information on the higher levels, although, as can be seen in the figure, intermediate states taking successively longer to converge. Note that for the filling discussed here both the first and second excited states should be doubly degenerate with non-zero angular momentum around the ring. As discussed earlier, this single Krylov space approach cannot separate such degeneracies.

Note how in only a few tens of iterations we have a reasonably accurate estimate of the ground state energy. Compared to other methods discussed above, this involves a substantially smaller number of applications of the Hamiltonian to our starting state, and thus is generally regarded as the method of choice for larger systems. Using the states generated in this procedure as a basis reduces our 400 state system to, say, a 20 by 20 tridiagonal matrix that reproduces well the lowest few eigenvalues. Truncating to states of this reduced matrix
could provide a useful starting point for an approximate iterative growth to larger systems. Nevertheless, for the small systems considered here, the computer time is insignificant; thus, the earlier more intuitive but less efficient methods still work quite well.

Fig. 7 displays an interesting symmetry between the highest and the lowest energy levels. This is a consequence of half filling on a bipartite lattice. Changing the sign of the fermionic operators on half of our lattice shows that the sign of the kinetic term does not affect the spectrum our Hamiltonian. Doing a particle hole transformation for a fermion \( a_i \leftrightarrow a_i^\dagger \) changes \( n_i \leftrightarrow 1 - n_i \). Doing this on all sites for either value of the spin and then shifting the spectrum by the filling of the other spin changes the sign of the potential term. Thus, whenever either spin state is half filled, we have a symmetry in the spectrum under a combined shift and sign change.

Having the ground state at hand enables one to look at correlations. In Fig. 8 I plot the correlation between a spin up on one site and either spin up or spin down on another site as a function of the separation between them. This graph is for \( U = 2 \). Note the tendency for the system to become anti-ferromagnetic; such a configuration maximizes the stabilization by delocalization. Note also the stronger correlation between same versus opposite spins. When \( U = 0 \) the up and down spins are totally decorrelated, while the Pauli principle leaves a correlation between parallel spins. Of course the correlation between parallel spins on the same site is the filling factor, .5 in this case. For larger systems this figure should match onto the Monte Carlo results shown in Fig. 8 of Ref. [5].

The primary difficulty with these direct approaches is that memory needs grow exponentially with system size. Generalizing the benzene system to an \( N \) site ring, the number of basis states for the half filled case is \( \left( \frac{N^!}{(N/2)!} \right)^2 \). The 400 states needed for the 6 site case rapidly rises to 853,776 for a 12 member ring. The half filled Hubbard model on a two dimensional lattice of size 4 by 4 involves 165,636,900 states. After a few megabytes of storage, one leaves the realm of current personal computers. In Fig. 9 I show the ground state energy density \( E_0/N \) as a function of the ring size \( N \). The filling is one half in all cases, with the number of up and down spins differing by zero (one) for the odd (even) rings. The points for \( U = 0 \) were obtained from the analytic formula. Note the extra stability when the ring size is twice an odd number. In this case Ref. [4] proved that the ground state is unique. For odd ring sizes the ground state should be doubly degenerate. This is the case in the non-interacting case where the final fermion has non-zero angular momentum around
FIG. 8: The correlation $\langle n_{i,s} n_{j,s'} \rangle$ between spins as a function of their spatial separation. This is for $U = 2$ on our six site cyclic system. Note the anti-ferromagnetic tendency.

I note that as with the algorithm discussed in Ref. [1], this approach does parallelize quite well if storage and computation can be done in parallel. If a creation or annihilation operator flips a high bit in a Fock state, this will relate components that are far apart in storage. Thus the algorithm requires long range communications. Nevertheless, the loops over components do not need the results immediately. Thus the results for the new wave function can be sent off to storage while a given processor continues to work on further components that are locally stored. These vectors tend to be quite long, and thus most communication is completed before the new results are needed. Thus we expect good performance from massively parallel MIMD machines, including ones designed primarily for local communication, such as the QCDSP [7] and the QCDOC [8]. As the problem is primarily combinatorial, the performance is not determined or properly measured in terms of floating point operations.
FIG. 9: The ground state energy density $E_0/N$ as a function of the size of an $N$ membered ring. The upper points are for $U = 2$ and the lower ones for $U = 0$. When the interaction is turned off these numbers are from the analytic formula; the interacting points are from the simulation.

**APPENDIX A: COUNTING BITS**

Counting the set bits in a given computer word lies at the heart of the discussions in this paper. There are a variety of ways to accomplish this. Some computers have an assembly instruction that directly returns this “population count.” However in the interest of portability it is useful to implement this counting in a higher level language. A variety of fast schemes exist, but one well known approach increments a counter while $i$ is non-vanishing and repeatedly takes $i$ to $i \& (i - 1)$. This latter operation resets the lowest non-vanishing bit of $i$, and the repetition stops when all bits are cleared. Thus the following implementation in C:

```c
inline int bitcount(int i) {
    /* counts the set bits in a word */
    int result = 0;
    while (i) {
        result++;
        i &= i - 1;
    }
    return result;
}
```
i &= (i - 1);  /* finds and resets rightmost set bit */
}
return result;
}

Another useful operation given an integer $i$ is to find the next integer with the same number of set bits. For this first locate the lowest run of set bits. Move the highest of these up one position and slide the remainder down to start at bit 0. For example, if our integer in binary is, say, (0011001110), we take the run of three set bits from position 1 to 3, move the highest of these to position 4, drop the other two to the beginning of the word, and obtain (0011010011) as the next integer with five set bits. This leads to the following implementation

```
int nextone(int i){
    /* find the next integer with the same bitcount as i */
    int bit=1,count=-1;
    if (i==0) return 1<<nsites;
    /* find first one bit */
    while (!(bit&i)){
        bit<<=1;
    }
    /* find next zero bit */
    while (bit&i){
        count++;
        bit<<=1;
    }
    if (!bit) die("overflow in nextone");
    i &= ~(bit-1);  /* clear lower bits */
    i |= bit | ((1<<count)-1); /* put them in new places */
    return i;
}
```

Defining a variety of routines for manipulating wave functions can be quite useful. For these I define a “wavefunction” type as a pointer to a complex array. Once a generic set of routines
is set up, one can quickly run through a variety of experiments as discussed above. Some such functions whose action should be clear from their names are

```c
    double complex overlap(wavefunction psi1, wavefunction psi2);
    double norm2(wavefunction psi);
    double normalize(wavefunction psi);
    void cmultiply(double complex factor, wavefunction psi);
    void caxpby(double complex a, wavefunction dest,
```

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