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Bethe ansatz approach to quench dynamics in the Richardson model

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By instantaneously changing a global parameter in an extended quantum system, an initially equilibrated state will afterwards undergo a complex nonequilibrium unitary evolution whose description is extremely challenging. A nonperturbative method giving a controlled error in the long time limit remained highly desirable to understand general features of the quench induced quantum dynamics. In this paper we show how integrability (via the algebraic Bethe ansatz) gives one numerical access, in a nearly exact manner, to the dynamics resulting from a global interaction quench of an ensemble of fermions with pairing interactions (Richardson’s model). This possibility is deeply linked to the specific structure of this particular integrable model which gives simple expressions for the scalar product of eigenstates of two different Hamiltonians. We show how, despite the fact that a sudden quench can create excitations at any frequency, a drastic truncation of the Hilbert space can be carried out therefore allowing access to large systems. The small truncation error which results does not change with time and consequently the method grants access to a controlled description of the long time behavior which is a hard to reach limit with other numerical approaches. © 2009 American Institute of Physics.

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I. STATEMENT OF THE PROBLEM

Due to fast dissipation rates, the unitary evolution of out-of-equilibrium quantum systems remained for a long time a question which remained mostly of academic interest. However, recent developments in the field of cold atomic systems,1 in which tunability of the Hamiltonian combined to very low dissipation made this problem experimentally accessible and created a strong interest2–5 in trying to draw a general theoretical understanding of far from equilibrium quantum dynamics. The interaction quench is a simple way to create a prepared out-of-equilibrium quantum state of the system. It is realized by supposing that a quantum system is initially in an eigenstate (the ground state) of Hamiltonian $H_{g_0}$. Here $g_0$ is some global tunable parameter of the Hamiltonian. At time $t=0$, the parameter $g_0$ is instantaneously changed to $g$, and as a consequence the initial state $|\psi_{g_0}^0\rangle$ is no longer an eigenstate of $H_g$. This process puts the system in a far from equilibrium state, whose subsequent evolution is fully determined by the new Hamiltonian $H_g$ and the initial conditions.

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The resulting dynamics of the wave function are easily formulated using elementary quantum mechanics since they simply are given by solving the time dependent Schrödinger equation,

\[ |\psi(t)\rangle = e^{-iHt/\hbar}|\psi(0)\rangle. \]  \hspace{1cm} (1.1)

Naturally, the evolution is greatly simplified by projecting the system onto the eigenbasis of \( H \), defined by a set of eigenvectors \( |\psi^\mu_S\rangle \) with eigenvalues \( \omega^\mu \).

\[ |\psi(t)\rangle = \sum_\mu e^{-i\omega^\mu t} (|\psi^\mu_S\rangle |\psi^\mu_S\rangle). \]  \hspace{1cm} (1.2)

The resulting time-evolved average of any given operator \( \mathcal{O} \) can also be written as a sum over form factors weighted by the projections of the initial ground state onto the eigenbasis of the final Hamiltonian,

\[ \langle \psi(t)|\mathcal{O}|\psi(t)\rangle = \sum_\nu,\nu' e^{-i(\omega^\nu - \omega'^\nu)t} \langle \langle \psi^\nu_S|\psi^\nu_S \rangle \langle \psi^\nu_S|\psi^\nu_S \rangle \rangle \langle \psi^\nu_S|\mathcal{O}|\psi^\nu_S \rangle \langle \psi^\nu_S|\psi^\nu_S \rangle \rangle \hspace{1cm} (1.3)

Not only does one need to be able to compute eigenstates and eigenenergies of both Hamiltonians, the exact treatment also requires the capacity to compute overlaps between the eigenstates of two different Hamiltonians. We will show how this apparent hurdle is simply dealt with, when using the results from the algebraic Bethe ansatz (ABA) within the fermionic pairing model treated here.

Moreover, as is readily seen, obtaining exact results necessitates a double sum over the full Hilbert space. Since its dimension generally grows factorially with system size, this is apparently heavily limiting as far as reachable system sizes is concerned. In the coming sections, we also show how the quench populates predominantly a small subset of the Hilbert space, therefore allowing a drastic truncation with minimal loss of information ultimately granting access to systems much larger than one could naively expect.

Most of the results presented here were previously published in Ref. 6. This paper, however, supplements this reference with additional details on the truncation scheme and also presents novel ideas on the relationship between weak and strong coupling eigenstates of this particular system.

II. SYSTEM

We treat a model of spin 1/2 fermions in a set of single particle energy levels which can accommodate up to two electrons (of opposite spin). Electrons interact through a typical Bardeen-Cooper-Schrieffer (BCS) uniform s-wave pairing term which retains coupling only between time-reversed states,

\[ H = \sum_a \sum_\sigma e_a^\sigma c_a^\dagger \sigma c_{-a}^\sigma - g \sum_{\alpha,\beta} c_{\alpha}^\dagger \sigma c_{\beta}^\dagger \sigma c_{-\alpha}^\sigma c_{-\beta}^\sigma. \]  \hspace{1cm} (2.1)

The model is basically a discrete version of the celebrated BCS Hamiltonian to which it reduces in the thermodynamic limit.\(^7\) An Anderson pseudospin representation\(^8\) also exists by defining \( S_a^\pm = c_{a,\uparrow}^\dagger c_{a,\downarrow} \), which leads to

\[ H_{BCS}(\{\epsilon_j\},g) = \sum_{j=1}^N \epsilon_j S_{j+}^\dagger S_{j+} - g \sum_{a,\beta=1}^N S_a^+ S_{-\beta}^-. \]  \hspace{1cm} (2.2)

Here the \( N \) levels are the unblocked levels, i.e., the ones which are not occupied by a single electron and therefore participate to the dynamics.\(^9\) This model, although initially developed by Richardson\(^8\) to describe pairing in nuclei, has also been used more recently to describe the physics of ultrasmall superconducting metallic grains.\(^10\)
Irrespective of the distribution of the energy levels and of their degeneracies, the model is integrable\textsuperscript{11} and can be diagonalized fully using the ABA.

A. The eigenstates of the systems

Through the use of the ABA,\textsuperscript{10,12,13} one can access a very compact representation of the eigenstates of Hamiltonian (2.2). For any given set of \(M\) complex parameters \(w_j\), we shall call rapidities, the repeated action of the \(C(w_j)\) operator of the monodromy matrix on a fully down polarized state, build a “general” state,

\[
(w_j) = \prod_{k=1}^{M} C(w_k) \downarrow \downarrow \cdots \downarrow = \prod_{k=1}^{M} \left( \sum_{a=1}^{N} \frac{S_a^+}{w_k - \epsilon_a} \right) \downarrow \downarrow \cdots \downarrow .
\]  

(2.3)

All of these \(M\) rapidities states have a definite \(z\)-projection of the total spin operator which is simply given by \(M - N/2\). For any value of \(g\), the un-normalized eigenstates of the BCS Hamiltonian are all built in this fashion. For a given coupling constant \(g\), the complete set of eigenstates correspond to the general states constructed out every set of rapidities which are solution to the Richardson (Bethe) nonlinear algebraic system of equations,

\[
-\frac{1}{g} = \sum_{a=1}^{N} \frac{1}{w_j - \epsilon_a} - \sum_{k \neq j}^{M} \frac{2}{w_j - w_k}, \quad j = 1, \ldots, M .
\]  

(2.4)

The eigenenergy of one of these states is given (up to a constant) by the sum,

\[
E((w_j)) = \sum_{j=1}^{M} w_j.
\]  

(2.5)

At zero coupling, the full set of fixed magnetization (fixed \(M\)) eigenstates of Hamiltonian \(H_0(\{\epsilon_j\}) = \sum_{j=1}^{N} \epsilon_j S_j^z\) is known to be given by the various colinear spin states with \(M\) spins aligned in the +\(\bar{\epsilon}\) direction and the remaining ones in the −\(\bar{\epsilon}\) direction. Each and every one of these states can be expressed in the form (2.3) by using a set of \(M\) real rapidities \(w_j\). By giving the \(M\) rapidities the values of the \(M\) energy levels at which a +\(\frac{1}{2}\) spin is found \((\{w_j\} = \{\epsilon_j\})\), this leads to

\[
(w_j) \propto \left( \prod_{\{\epsilon_j\}}^{M} S_j^+ \right) \downarrow \downarrow \cdots \downarrow ,
\]  

(2.6)

which is just the state with \(M\) flipped spins at the chosen levels \(\{\epsilon_j\}\).

In the opposite regime of infinitely large coupling, the Hamiltonian reduces to

\[
H_{\omega}(\{\epsilon_j\}) = -g \sum_{\alpha, \beta=1}^{N} S_{\alpha}^+ S_{\beta} = -g S_{tot}^+ S_{tot}^- = -g[J^2 - (J_z)^2 + J^2],
\]  

(2.7)

where \(J\) is the total angular momentum operator of the \(N\) spins system and \(J_z\) is its \(z\) component. The eigenvectors of the Hamiltonian are therefore known through the procedure of angular momentum coupling of \(N\) spins \(\frac{1}{2}\) leading to sets of eigenstates of \(J^2\) and \(J_z\). As mentioned previously, the \(z\)-projection of the total angular momentum is equal to the number of rapidities \(M\). For a fixed value of \(M\), one can absorb the \(M\)-dependent terms in a global shift of the energies and consequently the eigenspectrum consists of \(N!/(N-M)!/(M)!\) states subdivided into degenerate subbands of total angular momentum \(J\) with eigenenergies and degeneracies given by

\[
E(j) = -g[j(j+1)],
\]  

(2.8)
In the strong coupling limit, the solutions to Richardson’s equations naturally define an alternative representation for these eigenstates. It is known that for a state defined by $M$ rapidities, the various solutions will be defined by a number $r$ of divergent rapidities. These rapidities can be expressed in the large $g$ limit as $w_i = L_i g$, where the $L_i$’s are the zeros of the Laguerre polynomial,\(^\text{14}\)

\[
L_r^{-(N+2r-2M)}(L_i) = 0 \quad \forall L_i.
\]  

There is a direct correspondence between the number $r$ of diverging rapidities and the total spin eigenvalue $j$. Looking simply at the energies, it was shown\(^\text{14}\) that:

\[
r = j + M - \frac{N}{2}.
\]  

The remaining roots (which remain finite) defining the various eigenstates can then be found by solving the following set of simplified Richardson’s equations which involve only the nondivergent $M-r$ rapidities:

\[
\sum_{a=1}^{N} \frac{1}{w_j - \epsilon_a} - \sum_{k=j}^{M-r} \frac{2}{w_j - w_k} = 0, \quad j = 1, \ldots, M-r.
\]  

This particular set of equations still requires a numerical approach in order to find every possible solution.

One must realize that within a given degenerate subset, both sets of eigenstates (Bethe ansatz and total spin eigenbasis) are chosen differently. This is of course not an issue in the $g = \infty$ case, but the Bethe ansatz representation compares advantageously to the total spin eigenbasis when trying to expand in powers of $1/g$. Whereas in the former, one can directly expand the Richardson equations in powers of $1/g$ in order to describe the system,\(^\text{14}\) in the latter representation we would need to apply perturbation theory on a set of highly degenerate levels which would therefore involve diagonalization of large matrices.

### B. Numerical solution

Except in the previously discussed extreme cases of zero or infinite coupling, no analytical solution is known to the Richardson equations, and one therefore needs to resort to numerical techniques in order to study this particular system. Many efforts have been made to construct efficient algorithms for solving Richardson’s equations\(^\text{15,16}\) but it still remains a difficult task to achieve stable and fast computation of the various solutions. As is always the case when dealing with nonlinear systems of algebraic equations, the convergence of any algorithm toward a given solution highly depends on the capacity one has to generate an appropriate initial approximation of it.

In the absence of any known general features of the solutions in the intermediate coupling regime, it makes it quite natural to use a scanning procedure starting from the known solutions at $g = 0$ and increasing the coupling in small steps in order to reach solutions at a given value of $g$. This guarantees an adequate trial solution from which an iterative procedure can be carried out successfully.

The solutions are such that rapidities are either real or form a complex conjugate pair (CCP) with another one. At the critical values of $g$ at which two real rapidities collapse into a CCP (or vice versa), Richardson’s equations expressed in the form (2.4) have cancelling divergent terms which require appropriate modifications in order to be dealt with numerically. These forming and
splitting apart of CCPs happen exclusively when two rapidities, say \( w_1 \) and \( w_2 \) are equal to one another while at the same time being equal to one of the energy levels \( \epsilon_i \in \{ \epsilon_1, \epsilon_2, \cdots, \epsilon_N \} \). This allows a rewriting of the equations using the two real variables,

\[
\lambda_+ = 2\epsilon_1 - w_1 - w_2,
\]

\[
\lambda_- = (w_1 - w_2)^2.
\]

Contrarily to \( w_1 \) and \( w_2 \), these variables maintain a finite derivative with respect to \( g \) while going from real to complex. One can additionally transform the Richardson equations for \( w_1 \) and \( w_2 \) in the following two equations:

\[
4\lambda_+ + ((\lambda_+)^2 - \lambda_-)G_1 = 0,
\]

\[
((\lambda_+)^2 - \lambda_-)G_2 - 4\lambda_+ = 0,
\]

where

\[
G_1 = -\frac{2}{g} - \frac{1}{g} \sum_{a \neq c} \left[ \frac{1}{w_1 - \epsilon_a} + \frac{1}{w_2 - \epsilon_a} \right] + \frac{2}{M} \sum_{k \neq 1,2} \left[ \frac{1}{w_1 - w_k} + \frac{1}{w_2 - w_k} \right],
\]

\[
G_2 = -\frac{1}{g} \sum_{a \neq c} \left[ \frac{1}{(w_1 - \epsilon_a)(w_2 - \epsilon_a)} \right] + \frac{2}{M} \sum_{k \neq 1,2} \left[ \frac{1}{(w_1 - w_k)(w_2 - w_k)} \right].
\]

Rapidities which form a CCP at a given \( g \) can split apart into two real rapidities at a larger coupling and can also reform a CCP with a different rapidity. This means that one might need to redefine a new set of variables (and equations), at every step in \( g \). This is simply achieved by “pairing” any two real rapidities \( w_i, w_j \) which have the same closest single particle energy level \( \epsilon_i \) and using for them the variables (2.13). For these rapidities we also use the modified version of the Richardson equations written above. This always results in a system of \( M \) equations depending on \( M \) variables, but it can be modified at every step in \( g \) in order to adapt to the local structure of the solutions.

Independently of the pairing used at a given point, the Jacobian is always obtainable analytically in terms of these variables and it therefore allows a straightforward use of Newton’s algorithm for nonlinear algebraic equations. Despite the lack of guaranteed stability (unavoidable in a general set of algebraic equations), one can still manage to compute rapidly any solution in this way, provided we adjust the steps in \( g \) in such a way that at every \( g \) we correctly evaluate the rapidities which need to be paired. In a case where numerous pairings-unpairings happen on a small \( g \) interval, one simply needs to reduce the steps in \( g \) in order to make sure every rapidity crosses only one critical point per step. Since the scanning process gives access to a local evaluation of the derivatives of rapidities, we can always get a good evaluation of the desired \( \Delta g \) for the coming step.

Figure 1 shows the real part of rapidities for a few solutions of the Richardson equations (examples showing also the imaginary part can be seen in other papers,\(^{17}\) for example). These examples show how one can conjecture, for a given \( g=0 \) solution, the number of diverging rapidities that this state will have when deformed from 0 to infinite coupling.

One can understand the passage of a given state from its uncoupled form to the resulting large coupling state by looking at the contiguous blocks of occupied and unoccupied states at \( g=0 \). Any contiguous block of \( m \) unoccupied states will refrain up to \( m \) rapidities above it from diverging to infinity. Starting from high energies, one looks at the first block of occupied states and the first following block of empty states. Supposing they, respectively, contain \( p_1 \) states and \( h_1 \) states, if \( h_1 \geq p_1 \), the \( p_1 \) rapidities will remain finite. If \( h_1 < p_1 \) then \( h_1 \) of these rapidities will remain finite.
and the remaining \( p_1 - h_1 \) can be added to the following block of occupied states. Supposing the next blocks of occupied (empty) states contain \( p_2 \) (\( h_2 \)) states, it would be treated in the same way, but using

\[
p'_2 = \begin{cases} 
  p_2 \\
  p_2 + p_1 - h_1 
\end{cases}
\]

depending on what happened in the previous block. In the three examples presented in Fig. 2, we, respectively, find for the leftmost case: \((p_1=1, h_1=2), (p_2=2, h_2=1), (p_3=2, h_3=1), \) and \((p_4 =3, h_4=3)\) leading to two diverging rapidities (here they form a CCP), in the middle panel we have \((p_1=3, h_1=1), (p_2=2, h_2=1), (p_3=1, h_3=1), \) and \((p_4=2, h_4=2)\), giving three diverging rapidities (two of them forming a CCP) and the last case has \((p_1=3, h_1=3)\) and \((p_2=5, h_2=0)\) giving five divergent rapidities (including two CCPs). Although giving results equivalent to the algorithm proposed in Ref. 16, this approach can also provide some insight on approximative values of the finite rapidities in the large \( g \) limit. This could give access, in this limit, to initial guesses close enough to the real solutions to allow direct computation of the various solutions to Eq. (2.12).
This procedure seems to work well for any state which is half-filled or less, since in that case the diverging rapidities all have real parts going toward $-\infty$, the Laguerre polynomials $L_n$ involved having all their zeros in the left half plane. At higher filling factors this is no longer true, but the alternative choice of representing the eigenstates as the repeated application of the operator $B(w_i)$ on the fully up-polarized pseudovacuum, $\Pi_j^{N}(\sum_a^{N}w_a/(\epsilon_a-\epsilon_b))|\uparrow \uparrow \cdots \uparrow\rangle$, could probably allow similar arguments to be used in this case.

III. QUENCH MATRIX

Having shown how one can access individually any eigenstate of the system, we now need to be able to realize explicit calculations of the projection of the initial state of the system onto the eigenbasis of the final Hamiltonian. For a general Hamiltonian and even for a general integrable Hamiltonian, this problem requires full diagonalization, in a given basis, of the initial and final Hamiltonians. The factorial growth of the Hilbert space dimension (and therefore of the Hamiltonian matrix) limits one to very small system sizes.

For ABA integrable systems there exists a formula, due to Slavnov,\textsuperscript{16} which gives an expression of the overlap between an eigenstate of the system and a state built identically but the alternative choice of representing the eigenstates as the repeated application of the operator $B(w_i)$ in the model discussed here reads explicitly\textsuperscript{13}

\[
\langle \{w_j\}|\{v_j\}\rangle = \frac{1}{\sqrt{\det G'[\{w_j\}]\det G'[\{v_j\}]}} \Pi_j^{M}(w_j-v_j)^{-1} \det J, \quad (3.1)
\]

with the matrix elements of $J$ given by

\[
J_{ab} = \frac{v_b-w_b}{v_b-w_a} \left[ \sum_{k=1}^{N} \frac{1}{(w_a-\epsilon_k)(v_b-\epsilon_k)} - 2 \sum_{c\neq a} \frac{1}{(w_a-\epsilon_c)(v_a-w_c)} \right]. \quad (3.2)
\]

The $M \times M$ Gaudin matrices $G$, whose presence is needed to normalize both eigenstates, are themselves obtained through the limit $\{w_j\} \rightarrow \{v_j\}$ of Eq. (3.1), which gives

\[
G_{ab}\{v_j\} = \left\{ \begin{array}{ll} 
\sum_{k=1}^{N} \frac{1}{(v_a-\epsilon_k)^2} - 2 \sum_{c\neq a} \frac{1}{(v_a-\epsilon_c)^2}, & a = b \\
2 \frac{1}{(v_a-v_b)^2}, & a \neq b.
\end{array} \right. \quad (3.3)
\]

In the precise case treated here, the $C(w)$ operators have no explicit dependency on $g$ [see Eq. (2.3)]. Consequently eigenstates of the Hamiltonian at any value of $g$ are all general states built like Eq. (2.3). This allows one to use formula (3.1) directly to compute the relevant overlaps between two sets of eigenstates $|\psi_g^e\rangle |\psi_g^o\rangle$. One must understand that a different model might not share this property, and if a determinant representation exists for these overlaps in a general integrable model, it remains to be found.

A. Weak coupling to strong coupling

For $M=8$ rapidities and $N=16$ energy level, the dimension of the Hilbert space is 12,870 which is sufficiently small to allow us to compute the full set of eigenstates. Using Slavnov’s formula (3.1) we calculate the overlaps $\langle \psi_g^e |\psi_g^o\rangle$ of the $g=0$ ground state with the full eigenbasis at $g$. For a few quenches, with equally spaced energy levels, the squared norms of these overlaps are plotted on Fig. 2 as a function of the energy of the state $|\psi_g^o\rangle$.

In the limit of a small quench from $g=0$ to $g=\Delta g$, first order perturbation theory theory can be used. It gives nonzero overlaps with the $g=\Delta g$ coupled ground state and the set of states which differed (at $g=0$) from the ground state by a single particle-hole excitation (one up spin moved to a different level). The following order would allow nonzero overlaps with two particle-hole exci-
tation states and so on. This fact is clearly seen in the smallest quench presented in the top panel of Fig. 2 showing a set of decreasingly overlapping subbands, each associated with a different number of particle hole excitations.

When the quench is carried from an initial weak-coupling ground state to a stronger coupling (the two top panels of Fig. 2), one finds that a surprisingly small fraction of the final Hilbert space is importantly populated by the quench. In fact, we see, in the distribution of energies, the large coupling band structure forming and notice that in each of the bands the projection on a single state of the initial ground state heavily dominates. We could easily verify that these states are the finite $g$ deformations of a given set of $g=0$ configurations pictorially shown on Fig. 3. This set of states is obtained (at $g=0$) by promoting contiguous blocks of $N_p\leq M$ rapidities from right below to right above the Fermi level (FL). According to the procedure outlined in Sec. II B, they all lead, at $g\to\infty$, to states whose number of rapidities remaining finite is equal to $N_p$. In fact, they are the lowest energy states having a given number of particle-hole excitations (at $g=0$) equal to the number of finite rapidities (at $g\to\infty$).

For any quench from $g_0=0$ to $g\in[0,1]$, this set of $M+1$ states always account for more than 60% of the total amplitude $\Sigma_g|\langle\psi'_g|\psi\rangle|^2$ of the wave function expressed in the final eigenbasis. Figure 4 shows this total contribution when quenching from an uncoupled initial state. Despite the
presence of a dip in their contribution when quenching to the intermediate coupling regime, they clearly remain strongly dominant for finite-size systems.

It is then natural, when looking for additional large contributions, to assume that states built by slightly deforming the preceding set would be dominant. We therefore add to the reduced eigenbasis the set of states built at $g=0$ by adding a single particle-hole excitation to the block states. Figure 5 shows, for one of the block states discussed previously, the $g=0$ configurations created by adding this single excitation (either above or below the FL).

One can then add a set of states built with two particle-hole excitations over the block states. We include every state obtained with one particle excitation above the FL and one hole excitation below it. These would be constructed by the tensor product of the possible states above the FL, as seen in the first set of states in Fig. 5, with the possible states below the FL in the second set. Naturally, states with two excitations above or below the FL are also possible. A fairly small set of these is actually included, since the study of the quench matrix made for a smaller system indicates that they should be dominant. The treated cases are built by promoting any two rapidities above the FL to any contiguous set of two free (occupied) level above the FL as shown in Fig. 6.

In the case of $M=16$, $N=32$, the inclusion of all the aforementioned states proves sufficient to obtain more than 97% of the weight of the wave function after the quench, i.e., $\Sigma_i \langle \Psi_i | \Psi_f \rangle^2 \geq 0.97$. This figure holds for any final value of $g$ between 0 and 1. One must understand that for a given value of the final coupling only a part of these states will have an important contribution to the wave function. As one can readily see in Fig. 2, the weight is shifted to different parts of the Hilbert space as $g$ changes. For example, when doing a large quench the overlap of the final ground state with the initial one will become quite small, but for a small quench, it would be the dominantly populated state. It is therefore possible to get an accurate projection of the uncoupled ground state onto the coupled eigenbasis by keeping less than a 1000 states out of the 601 080 390-dimensional Hilbert space for $M=16$, $N=32$ for a given quench. However, in order to be able to treat the spectrum of quenches discussed here, we need to compute only a total of 7675 states. This drastic reduction in the effective Hilbert space is the key element which allows us to treat relatively larger system sizes with a minimal error.

Since this truncation scheme is carried out in the eigenbasis of the Hamiltonian driving the time evolution of the system, the error will not change with time. In fact, the time evolution only

![Figure 5](image5.png)

**FIG. 5.** Pictorial representation of the single excitations for a given block state. One should understand that the same process is applied to every one of the block states (see Fig. 3).

![Figure 6](image6.png)

**FIG. 6.** Pictorial representation of the doubly excited states which are kept for a given block state. The same process is applied to every one of the block states (see Fig. 3). Moreover, one should remember the inclusion of states featuring one excitation above and one below the FL (not represented here but discussed in the text).
affects the phase of various components of the wave function and not their amplitude. Apart from
the initial condition, the wave function (and by extension expectation value of operators) at any
time does not depend on the previous history. For very large times, it can therefore be calculated
directly and with a precision set by the small truncation error.

B. Strong coupling to weak coupling

Quenching from a strongly coupled initial ground state down to a weakly coupled system
brings a radically different structure. Looking at Fig. 2, we see that although a small quench still
brings a quench matrix naturally strongly peaked at low energies (with a roughly exponential
decay in energy), when the difference between initial and final \( g \) is made larger, the distribution
flattens significantly.

One can understand this tendency by noting that the infinite coupling ground state is charac-
terized by having every rapidity present diverging [see Eq. (2.5)]. In this case we have
\( \lim_{g\to\infty} C(t) \approx \sum_i S_i^z \). The repeated action of the operator for a set of diverging rapidities therefore
makes the \( M \) rapidity ground state a uniform superposition of every \( g=0\) \( M \)-flipped spins state, i.e.,
\[
|\psi^0_{g\to\infty}\rangle \approx \left( \sum_i S_i^z \right)^M |\downarrow\downarrow\cdots\downarrow\rangle.
\]
The \( g \to \infty \) ground state therefore has a uniform projection onto the \( g=0 \) eigenbasis. In this
extreme scenario, any truncation of the final Hilbert space is impossible. However, when starting
from a finite albeit large initial coupling, it appears that the nearly monotonic energy behavior
could allow one to reduce the effective dimension of the problem by removing the largest energy
states. The slow decrease in the overlaps in energy would still make the problem of very large to
very weak \( g \) quench hard to treat using this technique.

IV. FORM FACTORS

Being in a position to compute the eigenstates, eigenvalues and the quench matrix elements is
sufficient to access the time-evolved wave function of the system. However, to compute the
average value of an observable \( O \), one still needs tractable expressions for the form factors,
\[
\langle \psi^0_0 | O | \psi^0_\mu \rangle,
\]
of the operator we are interested in. In this work we will look at time evolution of the off-diagonal
order parameter defined as
\[
\Psi(t) = \frac{1}{M} \sum_{\alpha\beta} \langle \psi(t) | S_\alpha^z S_\beta^z | \psi(t) \rangle.
\]

In equilibrium and in the thermodynamic limit, this quantity is closely related to the BCS gap,
making it a suitable parameter to quantify the superconducting tendency for finite systems even
out of equilibrium. One could also use in a similar fashion the canonical order parameter
\( \sum_\alpha \frac{1}{2} - \langle S_\alpha^z \rangle^2 \) (as in Refs. 19 and 20), but since both quantities showed the same qualitative
behavior we only present the former here.

Although in the infinite coupling case we have \( \sum_{\alpha\beta} \langle \psi^0_0 | S_\alpha^z S_\beta^z | \psi^0_\mu \rangle \approx \delta_{\mu,0} \) since \( \sum_{\alpha\beta} S_\alpha^z S_\beta^z \approx H_{\text{exc}} \), at
any finite coupling it requires computation, for every couple \( \alpha, \beta \) of the form factors \( \langle \psi^0_0 | S_\alpha^z S_\beta^z | \psi^0_\mu \rangle \).

Using Slavnov’s formula, the solution to the inverse problem, i.e.,
\[
S_i^z = \lim_{u \to \epsilon_i} (u - \epsilon_i) C(u),
\]
\[ S_i^t = \lim_{u \to \epsilon_i} (u - \epsilon_i) B(u), \] (4.3)

and determinant properties, it was shown in Ref. 6 that these form factors for \( S^\alpha, S^\beta \) (and consequently \( S^\alpha, S^\beta \) too) can be written as a sum of \( M \) determinants. This was done by generalizing to the case \( \{v\} \neq \{w\} \) the method of Ref. 17, starting from the double sums given in Ref. 13. For \( \alpha \neq \beta \) we have

\[
\langle \{v\} | S^\alpha_{\alpha^\prime} S^\beta_{\beta^\prime}(w) \rangle = \frac{\Pi_k \left( \frac{v_c - \epsilon_a}{w_c - \epsilon_a} \right) \prod_{k \neq q} (w_k - w_q) \sum_{q=1}^{M} \frac{w_q - \epsilon_a}{w_q - \epsilon_\beta} \det \mathcal{J}_{\alpha, \beta} \rangle.
\] (4.4)

where, defining \( A_{ab} = J_{ab} \Pi_{c \neq b} (v_c - w_b) \) \( [J_{ab} \text{ being defined by Eq. (3.2)]} \), the matrix elements are given by

\[
\mathcal{J}_{ab}^b = A_{ab} - \frac{\Pi_{k \neq b, a} (w_k - w_b)}{\Pi_{k \neq b, a} (w_k - w_{b+1})} \frac{w_b - \epsilon_a}{w_{b+1} - \epsilon_a} A_{ab+1}, \quad b < q - 1,
\]

\[
\mathcal{J}_{aq}^b = A_{aq-1} + \frac{2(w_q - \epsilon_a)(w_{q-1} - \epsilon_a)}{w_{q-1} - w_q} \prod_e \left( \frac{v_e - \epsilon_a}{w_e - \epsilon_\beta} \right) \times \prod_{k \neq q-1} (w_k - w_{q-1}) \left( \frac{2v_a - \epsilon_a - \epsilon_\beta}{v_a - \epsilon_a} \right)^2 \left( v_a - \epsilon_\beta \right)^2,
\]

\[
\mathcal{J}_{aq}^q = 1/(v_a - \epsilon_a)^2,
\]

\[
\mathcal{J}_{aq}^b = A_{ab}, \quad b > q.
\] (4.5)

For \( \alpha = \beta \) it can be easily related to the form factors for \( S^\alpha \) which were already published elsewhere.\textsuperscript{13}

V. ORDER PARAMETER EVOLUTION

Having a numerically tractable expression for every necessary element, it is straightforward to compute the time evolution of the off-diagonal order parameter [Eq. (4.2)]. The upper left corner of Fig. 7 compares the time-averaged value i.e.,
weak coupling, the superconducting correlations between electron pairs are suppressed when the mean-field result obtained in the limit going from large coupling to weak coupling, the results show stronger deviations from the BCS result. One can understand this discrepancy by simply looking at the overlaps plotted in Fig. 2. It was shown \(^{17,21}\) that the equilibrium static correlation functions are unequally spaced from the BCS-like wave function, but for any finite system size, the small coupling limit leads to a ground state formed by uncorrelated electron pairs and the mean-field BCS treatment loses validity. Nevertheless, in the regime of relatively small quenches \((\Delta_{g_0}/\Delta_g \rightarrow 1)\), one still sees that the impact of quantum fluctuations (which are not present in the BCS mean-field treatment) on the long time-average gap is strongly suppressed even for the relatively small system sizes treated here.

The two additional panels in Fig. 7 show the time evolution (and its Fourier transform) of the off-diagonal order parameter. Those results are given for \(M=16, N=32\) which therefore necessitates the use of the previously described truncated eigenbasis. We focus on quenches from the uncorrelated ground state to various finite values of the coupling.

For general quenches, the mean-field treatment is known to lead to integrable classical dynamics, for which, in the steady state reached for quenches from small to large coupling, the time evolution of the BCS gap is given by the Jacobi elliptic function,\(^{19,20}\)

\[
\Delta(t) = \Delta_0 \text{sn}[\Delta_0(t - \tau_0), k], \quad k = 1 - \frac{\Delta_0^2}{\Delta_g^2},
\]

with parameters \(\Delta_0 \sim \Delta_g\) and \(\Delta_0 \rightarrow 0\) for a weakly coupled \(g \rightarrow 0^+\) initial state. This gives rise to nonharmonic persistent oscillations which are periodic in time with a period given by the complete elliptic integral of the first kind,

\[
T = \frac{2}{\Delta_g} \int_{0}^{\pi/2} \frac{d\phi}{\sqrt{1 - k^2 \sin^2 \phi}}.
\]

The mean-field solution should therefore show a Fourier transform made of equally spaced peaks which, as one can see, is quite different from the results obtained here. For small values of the final \(g\), this difference is no surprise since the mean-field treatment assumes a BCS-like wave function, which, for a finite-size system in sufficiently weak coupling, is not realized. However, when \(g \approx g^* = (2 \ln N)^{-1}\), it was shown\(^{17,21}\) that the equilibrium static correlation functions are undistinguishable from the BCS correlations. For the largest final \(g\) values shown here, this inequality is respected but the quench dynamics are still qualitatively different from the expected BCS result.

One can understand this discrepancy by simply looking at the overlaps plotted in Fig. 2. It was clear from the start that an instantaneous quench can, in principle, create excitations at arbitrarily large frequencies. What our calculations show is that the excitation is done in a highly nonthermal way, by creating preferentially excitations in the midrange energy spectrum. On the other hand, static correlation functions depend mostly on low-energy properties. Since the equispaced low energy bands which characterize the mean-field BCS regime are already well formed at \(g \approx g^* = (2 \ln N)^{-1}\) (see the energy distribution on Fig. 2), it is therefore expected that low energy properties are quite similar to the BCS ones. The quench dynamics, however, by probing higher energy properties which still differ strongly from the BCS behavior, will retain a visible impact of the
quantum fluctuations that the mean-field treatment cannot capture. As it was proposed in other models, the experimental accessibility to quantum quenches could provide a powerful spectroscopic tool to study quantum fluctuations in the pairing properties of fermions.

VI. DISCUSSION

In this work we have reported a new approach to the study the dynamics of a quantum system driven out of equilibrium by a global interaction quench. It uses integrability which provides numerically tractable ways to compute eigenstates, eigenenergies, and form factors of various operators. We were able to exploit the peculiar property of the Richardson model in order to find simple expressions giving the overlaps of the eigenstates of the pre- and postquench Hamiltonians. The obtained results show that the particular structure of these overlaps also allows one to design a simple but very efficient truncated Hilbert space giving access to nearly exact results in systems too large to allow the full Hilbert space to be treated.

The generalization to other integrable models such as quantum spin chains or atomic Bose gases would be a highly desirable feat. However, in other general integrable models, the current lack of known tractable expressions for the elements of the quench matrix hinders such progress for now. Such a representation would allow exact calculation of quench behavior for a large variety of potential experiments.

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