The construction of stationary ensembles for integrable quantum systems

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We reconsider the thermalization of integrable quantum systems. It is shown how the generalized Gibbs Ensemble (GGE) can be constructed as the best approximation to the time dependent density matrix. Our procedure allows for a systematic construction of the GGE by a constrained minimization of the entropy. We apply our method to a quenched hard core Bose gas. In contrast to the standard GGE, our correlated GGE properly describes the higher order correlation functions.

Thermalization is a fundamental feature of closed many-body systems on which much progress has been made in recent years [1–9]. After the early pioneering work by von Neumann [10, 11], the subject was put aside for a long time and thermalization was mainly understood as the result from interactions with an environment. The advent of closed quantum systems with a high degree of controllability, in particular ultracold atomic gases [12], has revived the interest in this subject. The implementation of integrable [13] cold atom systems has also sparked the interest in understanding the role of integrability in thermalization.

The long time behaviour of non-integrable systems is in all cases that have been investigated described by the standard Boltzmann-Gibbs ensemble: \( \rho_{eq} = \exp(-\beta H) \). For integrable systems, that are characterized by the existence of an additional set of conserved quantities \( \hat{I}_\ell \), the canonical Boltzmann-Gibbs ensemble may even fail to describe elementary properties such as the momentum distribution. A particularly striking example is given by one dimensional hard core bosons, that can be mapped onto non-interacting fermions, that have conserved occupation numbers [14, 15].

The formalism to generalize the Boltzmann-Gibbs ensemble was initiated by Jaynes [16], who pointed out that statistical physics can be seen as statistical inference and an ensemble as the least biased estimate possible on the given information. By including the conserved observables in our information, the Boltzmann-Gibbs ensemble is generalized to \( \rho_{eq} = \exp\left(-\sum_\ell \alpha_\ell \hat{I}_\ell\right) \), where the Lagrange multipliers \( \alpha_\ell \) have to be adjusted for the ensemble to predict the correct expectation values. The momentum distribution of quenched hard core bosons has been successfully described by such a generalized Gibbs Ensemble (GGE), that takes into account the conserved fermionic occupation numbers [14]. This clearly shows the relevance of the conserved quantities in constructing ensembles for quantum systems.

Unfortunately, it is not always clear which conserved quantities have to be taken into account. Actually, the linearity of quantum mechanics implies that the set of conserved quantities contains all projectors on eigenstates of the Hamiltonian \( \hat{P}_\alpha \). The number of these projection operators scales with the size of Hilbert space and is exponential in the physical system size. The ensemble that takes into account all expectation values of these projectors is the so-called diagonal ensemble [17]. In order to construct this ensemble, one needs the exponentially many diagonal elements of the density matrix in the eigenbasis of the Hamiltonian.

Clearly, the diagonal ensemble contains much more information than the GGE. The fact that more information than the one included in the GGE may be relevant is easy to see: if \( \hat{I}_I \) is conserved then also \( \hat{I}_I \hat{I}_I \) is conserved. This means that initially non-zero correlations between the different integrals of motion are conserved, where the GGE predicts no correlations between them. This fact has led to criticism on the validity of the GGE [18]. Furthermore, a macroscopic difference in the entropy between the diagonal and generalized Gibbs ensembles was reported by Santos et al. [19].

An elementary question thus arises: which information is essential to be included in the generalized Gibbs ensemble in order to get the correct entropy and correct expectation values for all the observables? The ‘statistical inference approach’ by Jaynes is not very useful in this respect. It would simply tell us that we should use the diagonal elements of the density matrix if we have access to them. This does not answer the question whether it is possible to construct a more economical representation of the density matrix that does not scale exponentially with system size.

In order to address this problem, we take a slightly different view on the task of statistical mechanics. Instead of maximizing the entropy subject to constraints, we wish to minimize the average distance between a stationary trial ensemble \( \sigma \) and the true state \( \hat{\rho}(t) \). This formulation as an optimisation problem will allow us to systematically improve the ensemble by including more observables when needed. We thus set out to minimize the average distance between a stationary trial ensemble \( \sigma \) and the state \( \hat{\rho}(t) \) over time

\[
\bar{D} = \lim_{T \to \infty} \frac{1}{T} \int_0^T dt D(\hat{\rho}(t)\|\sigma) .
\]

As a measure for the distance between density matrices we adopt the Kullback-Leibler distance (also known as
relative entropy)
\[ D = -S_N (\hat{\rho}_0) - \text{Tr} [\hat{\rho}_d \ln (\hat{\sigma})], \tag{2} \]
where \( S_N (\hat{\rho}) = -\text{Tr}(\hat{\rho} \ln \hat{\rho}) \) denotes the von Neumann entropy and \( \hat{\rho}_d \) the diagonal ensemble \[^{[17]}\], i.e.
\[ \hat{\rho}_d = \lim_{T \to \infty} \frac{1}{T} \int_0^T dt \hat{\rho}(t). \]
One readily finds that the distance is minimized for \( \hat{\sigma} = \hat{\rho}_d \) with the minimal distance equal to \( D_{\min} = S_N (\hat{\rho}_d) - S_N (\hat{\rho}_0) \). The diagonal ensemble thus always represents the optimal stationary approximation of the true state and the diagonal entropy quantifies the distance to the instantaneous density matrix.

In the case of a pure state, for which \( S_N (\hat{\rho}_0) = 0 \), the diagonal entropy is equal to the Kulback-Leibler distance between the instantaneous state \( \hat{\rho}(t) \) and the time independent approximation \( \hat{\sigma} \). Entropy is then identified as the information that is lost by describing a time dependent approximation \( \hat{\rho}(t) \) between the instantaneous state \( \hat{\sigma} \). This entropy is then identified as the information that is lost by describing a time dependent approximation \( \hat{\rho}(t) \) between the instantaneous state \( \hat{\sigma} \). This entropy is then identified as the information that is lost by describing a time dependent approximation \( \hat{\rho}(t) \) between the instantaneous state \( \hat{\sigma} \). This entropy is then identified as the information that is lost by describing a time dependent approximation \( \hat{\rho}(t) \) between the instantaneous state \( \hat{\sigma} \).

By unconstrained minimization, we have recovered the diagonal entropy, that requires the specification of a number of parameters that is exponential in system size. Let us now address the problem of constructing a stationary matrix that captures the relevant physics while being more economical in the number of parameters. Consider therefore the ansatz
\[ \hat{\sigma} = Z_\sigma^{-1} \exp \left[ - \sum_j \lambda_j \hat{A}_j \right], \tag{3} \]
where \( \{ \hat{A}_j \} \) is a set of operators, which we leave unspecified for the moment. Minimization of the distance with respect to the set \( \{ \lambda_j \} \) yields the condition that
\[ \text{Tr} [\hat{A}_j \hat{\sigma}] = \text{Tr} [\hat{A}_j \hat{\rho}_d], \tag{4} \]
and the minimal distance is consequently equal to
\[ D_{\min} = S_N (\hat{\sigma}) - S_N (\hat{\rho}_0). \tag{5} \]
Again the difference in entropy between the trial and the initial state quantifies the distance between the two. Furthermore, because the Kulback-Leibler distance is always positive, the entropy of the canonical trial is always bigger than that in the diagonal ensemble and the entropy difference is equal to the distance between the two, i.e.
\[ D (\hat{\rho}_d \| \hat{\sigma}) = S_N (\hat{\sigma}) - S_N (\hat{\rho}_d). \]
It is important to note that, even though our ansatz \[^{[3]}\] and the conditions \[^{[4]}\] are the same as in Jaynes work, our interpretation is very different. In our approach, the entropy has to be minimized under the constraints \[^{[4]}\] by using the best set of observables \( \{ \hat{A}_j \} \). In Jaynes’s approach on the other hand, the entropy is maximized for a set of observables that is fixed beforehand. This difference is important for the following reason. A lower value of the entropy can be reached by including more observables. It therefore makes sense to construct the optimal density matrix \[^{[4]}\] with as few observables as possible. This procedure then allows to identify the physically relevant observables. Jaynes’ maximization of the entropy does not offer this principle, because the maximum entropy is reached when no observables are used (equal occupation of all energy eigenstates).

The physical importance of the Kulback-Leibler distance is brought forward by Pinsker’s inequality \[^{[20]}\], combined with the operation interpretation of the trace distance \[^{[21]}\]. These guarantees that the difference in any projection operator valued measurement (POVM) is bound from above by the Kulback-Leibler distance:
\[ D (\hat{\rho}_d \| \hat{\sigma}) \geq \frac{1}{2} \max_B \left\{ \sum_j \left| \text{Tr} [\hat{B}_j \hat{\sigma}] - \text{Tr} [\hat{B}_j \hat{\rho}_d] \right|^2 \right\}, \tag{6} \]
where the maximum is taken over all sets of operators \( \{ B \} \) that form a resolution of the identity (POVM). Consequently, if a set of operators \( \{ \hat{A}_j \} \) exists that makes the distance between the GGE and the diagonal ensembles small, all POVMs show small differences between the two ensembles.

It is important to note that because of the optimum condition \[^{[4]}\], the operators that are included in the definition of the GGE do not show any deviation between the diagonal and GGE ensembles. This suggests an iterative construction of the GGE, where operators that show the largest deviation between the diagonal and GGE ensembles are included in the set \( \{ \hat{A}_j \} \). Depending on the system, the convergence of this procedure will yield a different number and different kinds of operators to be included in the GGE.

This discussion is valid for both integrable and non-integrable quantum systems. For the latter there is compelling numerical evidence \[^{[19, 22]}\] that they thermalize, such that the standard statistical mechanics ensembles provide a good approximation to the diagonal ensemble for what concerns the description of physical (few-body) observables. It is then sufficient to include the Hamiltonian, number operator and possibly a few other operators related to global symmetries in the trial ensemble.

Integrable quantum systems behave quite differently. As discussed above, additional conserved quantities are essential to be included in the construction of the ensemble. In the following, we will consider non interacting fermions because of their experimental relevance \[^{[12]}\].
and because many other 1D models can be mapped via a Jordan-Wigner transformation on free fermions, e.g. transverse field Ising model and hardcore bosons.

More specifically we numerically study a quench in one dimensional non-interacting lattice fermions by switching on an additional periodic potential. The system under consideration is identical to that in Ref. [19], where it was used to describe hard-core bosons. The Hamiltonian reads

\[
H = -\sum_{j=1}^{N} \left( \hat{c}_{j+1}^\dagger \hat{c}_{j} + \text{h.c.} \right) + J \sum_{j=1}^{N} \cos \left( \frac{2\pi}{\lambda} \right) \hat{c}_{j}^\dagger \hat{c}_{j}.
\]

The period \( \lambda = 5, t = 1 \) throughout the article and we consider the strength of the potential \( J \) as the quench parameter. In all quenches we start in the ground state of \( \hat{H} \) with \( J = 0 \) and quench to different values of \( J \).

Surprisingly, it was found in Ref. [19] that in this simple system, the entropies calculated in the GGE and diagonal ensembles show an extensive difference. According to [19], this means that there may be macroscopic observables that show large differences between the diagonal ensemble and the GGE. Indeed, it was pointed out by Gangardt and Pulstilnik [18] that correlations in momentum space are wrongly described by the GGE, casting doubt on its validity.

The reason for the failure of the GGE can be seen by noting that all information about the time-dependent state \( \hat{\rho}(t) \) is contained in the one-body density matrix

\[
\rho_{ij}^{(1)}(t) = \hat{a}_{i}^\dagger \hat{a}_{j},
\]

where \( \hat{a}_{j} \) annihilated a particle in a post-quench eigenstate, i.e. Hamiltonian [23] can be written as \( \hat{H} = \sum_{j} \epsilon_{j} \hat{a}_{j}^\dagger \hat{a}_{j} \). Only the diagonal is conserved over time such that it is the only part of the operator with a non-trivial time averaged expectation value. It should therefore be included in the trial ensemble. If one only includes these operators one immediately arrives at the standard generalised Gibbs ensemble

\[
\hat{\rho}_{\text{GGE}} = Z^{-1} \exp \left[ - \sum_{j} \lambda_{j} \hat{a}_{j}^\dagger \hat{a}_{j} \right].
\]

All off-diagonal elements vanish because their phases rotate at a frequency \( \epsilon_{i} - \epsilon_{j} \). However, the amplitude of the elements is conserved over time and that information is completely lost in the GGE while retained in the diagonal ensemble. As a consequence the GGE predicts a wrong value for the 2-body correlation:

\[
\text{Tr} \left[ \hat{a}_{j}^\dagger \hat{a}_{i}^\dagger \hat{a}_{i} \hat{a}_{j} \left( \hat{\rho}_{\text{GGE}} - \hat{\rho}_{d} \right) \right] = \left| \rho_{ij}^{(1)} \right|^{2} - n_{i}^{2} \delta_{i,j}.
\]

As the difference is significant we must include a 2-body interaction term in the trial ensemble to remove the discrepancy between the GGE and the DE. We therefore propose the improved correlated generalised Gibbs ensemble (CGGE)

\[
\hat{\rho}_{\text{CGGE}} = Z^{-1} \exp \left[ - \sum_{j} \lambda_{j} \hat{a}_{j}^\dagger \hat{a}_{j} - \sum_{i,j} V_{i,j} \hat{a}_{j}^\dagger \hat{a}_{i}^\dagger \hat{a}_{i} \hat{a}_{j} \right],
\]

which includes a density-density interaction \( V_{i,j} \). In order to compare all ensembles we calculate their entropy which is depicted in Fig. 1. It is immediately clear that in contrast to the GGE, the CGGE has the same slope as the diagonal ensemble. This implies that, while there is an extensive difference in entropy between the GGE and the diagonal ensemble, the difference between CGGE and diagonal ensemble is sub-extensive. Moreover, both entropies are so close that they put a strong bound on the trace distance between the two ensembles. This implies that almost all observables are perfectly described in the CGGE. The CGGE should therefore recover the diagonal probabilities with great accuracy. At first sight, it could seem surprising that the proper ensemble for quenched fermions is non-Gaussian, so that correlation functions cannot be computed with Wick’s theorem. This is however a simple consequence of the mathematical fact that the time average of a time-dependent Gaussian density matrix is not Gaussian.

In Fig. 1 we compare the different ensembles directly by plotting the diagonal elements for the density matrix. In order to show the whole interval of occupied states, we have made a coarse graining of the energy with \( \Delta E = t \). The CGGE coincides well with the diagonal ensemble and both are indistinguishable by eye. In contrast, the prediction of the GGE is rather poor. The total variation distance between the coarse grained GGE and the DE is 0.28, where the distance between the CGGE and DE is only 0.01. The inset in Fig. 1 shows the probabilities of individual eigenstates over a small energy interval. The CGGE correctly captures all the probabilities of the significantly occupied states, while the GGE does not.
Figure 2. (Color online) The coarse grained ($\Delta E/t = 1$) energy distribution for the $J = 12$, $N = 30$ quench. The green bars show the exact distribution. Red circles are the GCE prediction and the blue crosses is the present result. The inset shows some non-course grained matrix elements on a semi-logarithmic scale; blue crosses are the CGGE, red circles the GCE and green boxes the DE.

Correctly predict any occupation of the individual eigenstates. In fact for $J = 12$ and $N = 30$ the trace distance $D_1(\hat{\rho}_d, \hat{\rho}_{GCE}) = 0.867$, which is still much larger than the distance of 0.28 that we obtained after coarse grainning in energy. The trace distance between the DE and the CGGE is only $D_1(\hat{\rho}_d, \hat{\rho}_{V}) = 0.028$. This small difference between the CGGE and DE means that they are hardly distinguishable by any POVM. All observables, including higher order correlation functions are consequently correctly described by the CGGE.

The associated interaction potential $V_{i,j}$ is depicted in Fig. 3. As expected, its diagonal is zero, as the diagonal correlations were already properly predicted by the GGE. In fact the overall intra-band interactions are weak. Furthermore, the interaction between bands with odd quantum number, i.e. $V_{11}, V_{33}, V_{13}, V_{31}$, are extremely small and tend to zero with increasing $J/t$. This can immediately be understood from the nature of the bands. While the even bands are particle-like, the odd bands are hole-like and can thus not be occupied in the high $J/t$ limit. The interaction is dominated by inter-band repulsion in the electronic bands.

In conclusion we have formulated the construction of the generalized Gibbs ensemble as an optimization problem. Entropy is interpreted in our approach as the information that is lost by approximating the time dependent state by a stationary density matrix. The diagonal ensemble was shown to be the optimal ensemble when the evolution time tends to infinity. Other ensembles that depend on less parameters can be constructed by minimizing the Kullback-Leibler distance to the diagonal ensemble. We have applied this ideas to a quench in a non interacting Fermi system, which shows significant discrepancies between the diagonal and standard GGE. We have constructed a correlated GGE that shows excellent agreement with the diagonal ensemble, providing accurate predictions for all observables and not only for the single particle distribution.

Since our approach does not rely on any assumption of thermal equilibrium, it could also be applied to construct GGEs for systems out of equilibrium, such as periodically driven systems [24] or driven-dissipative many body systems [25].

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