Quantum dynamics of an Ising spin-chain in a random transverse field

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(Dated: March 23, 2022)

We consider an Ising spin-chain in a random transverse magnetic field and compute the zero temperature wave vector and frequency dependent dynamic structure factor numerically by using Jordan-Wigner transformation. Two types of distributions of magnetic fields are introduced. For a rectangular distribution, a dispersing branch is observed, and disorder tends to broaden the dispersion peak and close the excitation gap. For a binary distribution, a non-dispersing branch at almost zero energy is obtained. We discuss the relationship of our work to the neutron scattering measurement in LiHoF$_4$.

The Ising model in a random transverse field has been studied both analytically and numerically$^{9,11,12,13,14,15,16,17,18,19}$ and a great number of results of physical importance have been obtained. However, the dynamical structure factor $S(k, \omega)$ has not been computed for all $k$ and $\omega$ in the random field model, although some analytical and numerical results are available in pure systems for special values of the wave vector. Here we compute the dynamic structure factor in the presence of two types of disorder distributions: a rectangular and a binary distribution.

The one-dimensional lattice Hamiltonian we study is

$$H = -J \sum_i \sigma_i^z \sigma_{i+1}^z - \sum_i h_i \sigma_i^x,$$  (1)

where the $\sigma$’s are Pauli matrices, and $J$ is positive and uniform. We shall choose the energy unit such that $J = 1.0$. The fields $h_i$ are random variables. The first model we study is the rectangular distribution with a mean $h_{\text{ave}}$ and a width $h_w$. The second model is the binary distribution in which $h_i$ is an independent random variable that takes two values: $h_S$ and $h_L$ with probabilities $p$ and $(1-p)$, respectively. In particular, we choose the parameter $p$ to be small such that the chain is almost spatially homogeneous, $h_S < J < h_L$, to ensure that the system is in the paramagnetic phase. Intuitively, this distribution seems to capture a crude adiabatic representation of the electronic spins coupled to a hyperfine spin bath discussed above, where the larger field is the applied transverse field.

We first compute the time-dependent spin-spin correlation functions $C(n, t) = \langle \sigma_i^z(t) \sigma_{i+n}^z \rangle$ at temperature $T = 0$, where the angular brackets denote the average over ground state and the overline an average over disorder configurations. The algorithm for evaluating this quantity is similar to that described in Ref.$^{18}$ except that we are doing a real time calculation, which entails computation over complex variables. We first perform a Jordan-Wigner transformation$^{20}$ and then cast the correlation function $C(n, t)$ in the form of a Pfaffian, which is calculated efficiently, as described below; after averaging over disorder configurations, the dynamical structure

Calculation of real-time dynamics of a correlated quantum system with an infinite number of degrees of freedom are few and far between. Except for isolated examples, construction of real-time behavior from imaginary-time correlation functions (more amenable to numerical methods) by analytic continuation is fraught with various instabilities. The theoretical challenge is particularly acute because neutron scattering experiments often provide a rather detailed map of the frequency, $\omega$, and the wave vector, $k$, dependent dynamical structure factor, $S(k, \omega)$.

The second motivation comes from the desire to study the dynamics of a quantum phase transition involving a zero temperature quantum critical point. In this respect, the Ising spin chain in a transverse field$^{2,3,4,5}$ constitutes a schema from which much can be learned about quantum criticality both with and without disorder.

The third motivation is to examine how the coherence of the quasiparticle excitations is modified in the presence of quenched disorder and is triggered by a recent neutron scattering experiment in LiHoF$_4$, which connects the observed low temperature behavior of $S(k, \omega)$ in terms of the hyperfine coupling of the electronic spins to a nuclear spin bath, where the Hamiltonian of the electronic spins is given by an Ising model in a transverse field. Because the hyperfine splittings are small, we could imagine that, on the time scale of the electronic motion, this bath will appear essentially quenched, modulating the quantum fluctuations characterized by the transverse field. This is still a bit far from the experimental system, as it is three-dimensional, and the Ising couplings are long-ranged and dipolar. Nonetheless, we shall see that there are tantalizing similarities between our calculated structure factor and the experimental one in a given direction of the reciprocal space $\langle 2, 0, 0 \rangle \rightarrow \langle 1, 0, 0 \rangle$ (in reciprocal units). A more appropriate comparison should be with quasi-one dimensional spin systems that are intentionally disordered.

Finally, the role of disorder in a quantum critical system is an important subject in itself and is certainly not fully understood. In the presence of disorder, there are rare regions with couplings stronger than the average, which results in the Griffiths-McCoy singularities.$^{7,8}$ Although the effect is weak in a classical system, it becomes important in quantum systems, especially in low dimension$^{21-24,25}$. Considering the Ising spin chain in a transverse field, the Ising system with an infinite number of degrees of freedom is an important subject in itself and is certainly not disordered.

Quasi-one dimensional spin systems that are intentionally disordered.

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The calculation were performed for a lattice of 160 sites of which 32 sites in the middle were used for evaluating $C(n,t)$. Since we are interested in the gapped paramagnetic phase, the system is still in the paramagnetic regime everywhere, there is only a single dispersive branch, and the excitation remains gapped, as in Fig. 2(a). As we increase $h_w$, the dynamic structure factor can, in principle, have two branches because the DOS can acquire states at zero energy. Clearly, the non-dispersing branch, as shown in Fig. 2(b), has very low spectral weight everywhere. In principle, there could be a strong non-dispersing branch at $\omega = 0$. However, since the excitation gap is closed at this value of disorder strength, and since both branches are significantly broadened due to randomness, the non-dispersing peak may be mixed with the dispersive branch, and perhaps we are not able to observe them separately. In the high disorder limit, $h_w = 2.5$, almost all the intensity concentrates at $(k,\omega) = (0,0)$, but there appears be a ghost of the dispersive branch, see Fig. 2(c). An important feature of Figs. 2(b) and (c) is the horizontal stripe-like patterns, which indicate excitation modes that do not disperse, namely the localized modes. Our code was checked by comparing with the exact result for the pure system for which $S(k,\omega)$ has a single dispersive branch (see below).
the domain $\Delta \Omega$ such that

$$S = \max_{k, \omega}$$

some $(k, \omega)$.

The density of states in Fig. 3 shows zero energy states separated by a gap from the states at higher energy. The calculated $S(k, \omega)$ is shown in Figs. 4(a) through (f), where $h_L = 1.1, 1.2, \ldots, 1.6$. While the dispersing branch is broadened due to disorder, the weight of the central peak around $(k, \omega) = (0, 0)$ is so high that a non-dispersing branch can extend quite far away from the origin.

Let us define the weight of the central peak as

$$I_h = \frac{\int_{\Delta \Omega} S^2(k, \omega) dk d\omega}{\int_{\Omega} S^2(k, \omega) dk d\omega}$$

(7)

where $\Omega$ in the denominator is the entire $(k, \omega)$ domain, while $\Delta \Omega$ is defined as follows: we first find the maximum of the central peak $S_{\max}(k, \omega)$, and then define the domain $\Delta \Omega$ such that $S(k, \omega) > S_{\max}(k, \omega)/\sqrt{2}$ for $(k, \omega) \in \Delta \Omega$. Note that in Fig. 4 we integrate $S^2(k, \omega)$ rather than $S(k, \omega)$. This is because small numerical errors can result in slightly negative values of $S(k, \omega)$ for some $(k, \omega)$, which is obviously unphysical. The dependences of $I_h$ and $\Delta \Omega$ on $h_L$ are plotted in Fig. 6. As we tune $h_L$ from 1.6 to 1.1, $I_h$ increases monotonically, while the region $\Delta \Omega$ shrinks. We conclude that, as the quantum phase transition is approached, the weight is transferred from the dispersing branch to the non-dispersing peak.

In addition to the singularity due to the quantum phase transition, in disordered systems there is also the Griffiths-McCoy singularity: the disorder will drive some rare regions into a phase different from the rest. For the pure Ising chain when $h_i = h$, the dispersion relation is

$$\omega = 2\sqrt{J^2 + h^2 - 2Jh \cos k}$$

(8)

The excitation gap $\Delta = 2|h - J|$ occurs at $k = 0$. As $h \rightarrow J$, the excitation gap closes at the quantum critical point. However, if disorder is strong enough, there is nonzero probability to find some regions in which the
spins are strongly coupled, such that $h_\perp < J$ holds, and thus the cluster is ferromagnetic. Calculations, similar to that given in Ref. 22, show that these clusters give rise to the non-dispersing peak at zero energy, as we describe below.

For the binary distribution, the system is almost homogeneous except for the rare regions of strongly coupled clusters where $h_\perp = h_S$ for all sites inside the cluster. At shorter length scales, the behavior of the pure system dominates, and this leads to the dispersing branch. At longer scales the effects of disorder become important, resulting in the zero energy peak. The autocorrelation function $S(\omega)$, which is the integral of $S(k, \omega)$ over $k$ can be approximated in the following manner. The normalized probability that a given site belongs to a ferromagnetic cluster of length $L$-sites is $P(L) = Lp^{L-1}(1-p)^2$. When $h_\perp = 0$ the two-fold degenerate ground state within a cluster is far away from the excited states of energy of order $2J$; the perturbation $h_\perp = h_S$, will split the ground state by $\tilde{g}e^{-cL}$, where $\tilde{g}$ and $c$ are unknown positive constants determined by the details of the Hamiltonian. The form of the splitting results from large order in perturbation theory, however. If we treat these clusters as independent and average over disorder, or equivalently integrate over the cluster size $L$, we get

$$S(\omega) \sim \int dLP(L)\delta(\omega - \tilde{g}e^{-cL}) \sim \frac{(1-p)^2 \ln(\tilde{g}/\omega)}{\omega} \left( \frac{\tilde{g}}{\omega} \right)^{\ln(p)/c}, \quad (9)$$

which diverges at $\omega = 0$ if $1 > |\ln p|/c$ modulo logarithmic corrections. We have verified that indeed the divergence disappears for sufficiently small values of $p$ (the peak of $S(\omega)$ is then shifted to $\omega$ greater but close to zero, instead), but a detailed verification of this result appears to be difficult.

It is remarkable that a numerically exact solution of a simplified one-dimensional model (binary distribution) can capture some of the experimental features in LiHoF$_4$ and shed interesting light on the role of disorder on the dynamics of a prototypical quantum critical point. It would be of course interesting to experimentally study intentionally disordered systems that are closer to the model discussed here.

We thank Yifei Lou for suggesting to us the powerful calculational scheme described in the text. This work was supported by the NSF under the grant DMR-0411931.

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