Dynamics of the one-dimensional Heisenberg model and optical absorption of spinons in cuprate antiferromagnetic chains

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We use numerical and analytical results to construct a simple ansatz for the energy dynamical correlation function of the one-dimensional antiferromagnetic Heisenberg model. This is applied to compute the phonon-assisted absorption spectra of magnetic excitations (spinons) in quasi-one-dimensional spin-1/2 insulators and shown to reproduce very well recent infrared measurements in Sr_2CuO_3 . [S0163-1829(97)50206-2]

After the solution by Bethe^{1,2} and many years of intense research, the spin-1/2 Heisenberg chain is one of the most studied many-body problems. In spite of this *little is known* on its dynamical properties. This is because the extreme complexity of Bethe's wave functions makes it too hard to compute dynamical correlations. Much of the progress came from numerical studies which inspired an ansatz for the dynamical structure factor,^{3,4} a recent analytical evaluation of the exact two-spinon contribution⁵ for the same quality, and low-energy bosonization studies.⁶⁻⁹

In this paper we use numerical and known analytical results to obtain a simple ansatz for the energy dynamical correlation function defined below. This correlation function is central to the problem of coupling with phonons⁹ (the spin-Peierls problem) and in optical properties.^{10,11} As an application we compute the line shape for optical absorption of magnetic excitations assisted by phonons,^{10,11} and show how spinon physics appears naturally in the spectra. This is shown to reproduce very well recent measurements of the line shape by Tokura *et al.*^{12,13} in quasi-one-dimensional cuprates (see Fig. 3).

We will consider the one-dimensional (1D) Hamiltonian

$$H = \sum_{i}^{N} JB_{i}, \tag{1}$$

where

$$B_{i} = S_{i}^{x} S_{i+1}^{x} + S_{i}^{y} S_{i+1}^{y} + \Delta S_{i}^{z} S_{i+1}^{z}$$
 (2)

is the bond-energy operator, Δ measures the anisotropy, and we take periodic boundary conditions. We will consider Δ =0 (the XY model) and Δ =1 (the antiferromagnetic Heisenberg model).

The energy correlation function is defined as ${}^9G(p,\omega) = \langle \delta B_p; \delta B_{-p} \rangle(\omega)$ (in Zubarev's notation 14) where δB_p is the Fourier transform of $\delta B_i \equiv B_i - \langle B_i \rangle$.

To warm up we will compute $G(\omega,p)$ for $\Delta=0$. This is easily done by using a Jordan-Wigner tranformation^{6,2} to map the problem to a noninteracting 1D free fermion gas.

Excitations are free fermions and $G(p,\omega)$ becomes a Lindhard-like response describing density-like flucations. We get for the imaginary part

$$-2\operatorname{Im}G_{XY}(p,\omega>0) = D_{XY}(p,\omega)M_{XY}(p,\omega), \qquad (3)$$

where following the approach of Ref. 4 we defined a density of states $D(p,\omega)$ and a matrix element $M(p,\omega)$:

$$D_{XY}(p,\omega) = \frac{N}{2\pi} \frac{\theta(\omega - \omega_1^{XY}(p))\theta(\omega_2^{XY}(p) - \omega)}{\sqrt{[\omega_2^{XY}(p)]^2 - \omega^2}}$$
(4)

with $\omega_1^{XY}(p) = J\sin(p)$, $\omega_2^{XY}(p) = 2J\sin(p/2)$, and

$$M_{XY}(p,\omega) = 2 \frac{[w_2^{XY}(p)]^2 - \omega^2}{N^2 [\omega_2^{XY}(p)]^2}.$$
 (5)

Twofold degenerate states are counted once in Eq. (4) and this is taken into account by the factor of 2 in the matrix element (see Ref. 4).

Notice that Im $G_{XY}(p,\omega)$ is only different from zero between ω_1^{XY} and ω_2^{XY} . For a finite system the spectrum is given by a two-parameter set of discrete states with energy and pole strength given for p>0 by

$$\omega_{XY}(p,q) = \omega_2^{XY}(p)\cos(q-p/2)$$

$$M_{XY}(p,q) = (2/N^2)\sin^2(q-p/2),$$
(6)

where $0 < q \le p/2$. Equation (6) corresponds to particle-hole excitations around the $-\pi/2$ Fermi point.

When we switch to the isotropic limit $(\Delta=1)$ a simple analytic way of computing $G(p,\omega)$ is not available. Next we will follow Ref. 4 and use numerical results in finite chains, known low-energy bosonization results, sum rules, and Bethe ansatz results to construct a phenomenological ansatz for $\text{Im}G(p,\omega)$.

In the isotropic limit there is no upper bound for the support of $\text{Im}G(p,\omega)$; however, the dominant contribution to $\text{Im}G(p,\omega)$ comes from singlet states between $\omega_1(p)$ (the des

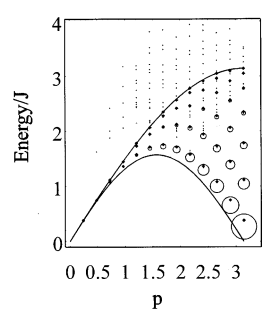


FIG. 1. Exact diagonalization results for $\text{Im}G(p,\omega)$ in the ω -p plane for a 26-site Heisenberg ring. Excitations are given by the position of the circles and matrix elements by the radius. The dots are the XY energies with the renormalization of Eq. (7). The lines are $\omega_1(p)$ (lower) and $\omega_2(p)$ (upper).

Cloizeaux-Pearson¹⁵ dispersion relation) and $\omega_2(p)$ which are obtained from $\omega_1^{XY}(p)$ and $\omega_2^{XY}(p)$ by changing J according to

$$J_{XY} \rightarrow J \pi/2,$$
 (7)

and we attached an *XY* subindex to avoid confusion later. This is analogous to the scenario found for the dynamical structure factor⁴ and corroborated by a recent analytical evaluation which retains only two-spinon excited states.⁵ We illustrate the analogous behaivor for the energy correlation function for a finite system in Fig. 1.

In the thermodynmaic limit these states form a twoparameter continuum degenerate with the triplet states that

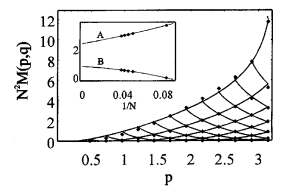


FIG. 2. The dots are the matrix elements (times N^2) vs p from exact diagonalization of a 26-site Heisenberg ring. We had plotted the fit of $N^2M(p,q)$ (the ansatz) as continuous functions of p for each allowed q in the 26-site chain. The intersection fo the continuous curves gives the actual values for the finite chain. The inset shows the finite-size scaling analysis for the constants A and B of the ansatz.

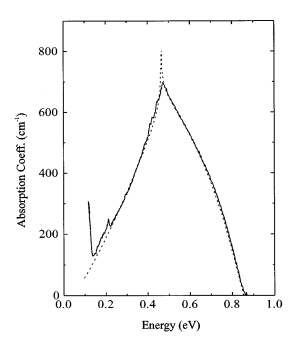


FIG. 3. Experimental abosrption coefficient from Ref. 12 (solid line) and theoretical line shape (dashed line) in $\mathrm{Sr_2CuO_3}$. Parameters are $J{=}0.246$ eV, $\omega_{\parallel}{=}0.08$ eV, and $\alpha_0{=}132$ cm $^{-1}$. ω_{\parallel} was taken similar to the value in the 2D materials. A linear background was subtracted from the experimental data.

dominate the dynamical correlation function and have a density of states, $D(p,\omega) = D_{XY}^{J_{XY} \to J_{\pi/2}}(p,\omega)$.⁴

To construct an ansatz for the energy correlation function of the Heisenberg model we put

$$-2\operatorname{Im}G(p,\omega>0) = D(p,\omega)M(p,\omega). \tag{8}$$

Our ansatz for $M(p, \omega)$ is given by

$$M(p,\omega) = M_{XY}(p,2\omega/\pi) \left(A \frac{\sqrt{J}}{\sqrt{\omega}} + B \frac{J}{\sqrt{\omega^2 - \omega_1(p)^2}} \right). \tag{9}$$

Notice that $M_{XY}^{J_{XY} \to J\pi/2}(p,\omega) = M_{XY}(p,2\omega/\pi)$. For a finite system we use Eq. (6) to construct M(p,q) in the same manner. In Fig. 2 we plot the matrix elements of Fig. 1 as a function of p together with the best fit of M(p,q) from the ansatz. The agreement is very good for all branches with singificant spectral weight.

It is interesting that one can fit all branches except the upper one with $A \approx 3$ and B = 0. The upper branch corresponds to the states closer to $\omega_1(p)$ where the square root becomes important, i.e., the dominant correction to the XY matrix elements in most of the (p,ω) plane is given by the p independent $1/\sqrt{\omega}$ factor.

To determine *A* and *B* for an infinite system we have made a finite-size scaling study (see Fig. 2) and used the sum rule $\sum_{p} \int d\omega (-2) \operatorname{Im} G(p,\omega) = \ln 2 - (\ln 2)^2$ to get A = 2.4, $B = 0.6^{-16}$

Bosonization studies predict⁷⁻⁹ a divergence for $-\operatorname{Im} G(p=\pi,\omega)$ like ω^{-1} , which becomes $(\omega-\omega_1(p))^{-1/2}$ close but not at $p=\pi$. The term proportional to B in

 $M(p,\omega)$ ensures this behaivor and is analogous to the form introduced by Muller *et al.*⁴ for the dynamic structure factor. As mentioned before its importance here is quantitatively minor in most of the (p,ω) plane whereas in the case of the dynamic structure factor it was the dominant contribution.

We next apply our results to the computation of the line shape for phonon-assisted optical absorption of magnetic excitations for a Heisenberg chain. The abosrption coefficient is given by ^{10,11}

$$\alpha(\omega) = \alpha_0 \omega I(\omega - \omega_{\parallel}), \tag{10}$$

where α_0 determines the oscillator strength and is given by

$$\alpha_0 = \frac{4\pi^2 q_A^2}{c\sqrt{\epsilon_1 M V_{\text{CH}} \omega_\parallel}},\tag{11}$$

where M is the O mass, V_{Cu} is the volume associated with a Cu ion, ω_{\parallel} is the frequency of the Cu-O stretching mode phonons, ϵ_1 is the real part of the dielectric constant, q_A is an effective charge associated with this process and is of order $|q_A| \sim e \beta a J/(S \Delta^2)$, where β is the rate of change of the Cu on-site energy due to a nearby O-ion displacement, a is the lattice constant, S=1/2 is the magnitude of the spin, and Δ the on-site energy difference between Cu and O.

The function *I* is given by

$$I(\omega) = -16\sum_{p} \sin^{4}(p/2)\operatorname{Im}G(p,\omega). \tag{12}$$

 $I(\omega)$ has a logarithmic singularity at $\omega = J\pi/2$ which will show up in the optical absorption. The position of the singularity is a very robust feature which comes from the saddle point at $p=\pi/2$ in the exact des Cloizeaux-Pearson¹⁵ dispersion relation (see Fig. 1). The distance between ω_{\parallel} and the singularity in the data provides a quick way to obtain the microscopic parameter J directly from experiments.

The two-dimensional version of this theory 10,11 within an interacting spin-wave approach was recently applied to explain related mid-IR bands measured by Perkins *et al.* in spin-1/2 (Ref. 17) and spin-1 (Ref. 18) insulating materials. Unlike estimations done in higher dimension 10,11,19 the *J* obtained from the position of the singularity in 1D is "exact" since the location of the singularity is known exactly. Here the only approximation is the Heisenberg model itself.

In Fig. 3 we compare the calculated optical absorption line shape to recent optical data by Tokura and collaborators 12,13 in $\mathrm{Sr_2CuO_3}$. The agreement is excellent. We stress that the only fitting parameters are J which fixes the energy scale and α_0 which fixes the intensity scale. The value found for the superexchange J = 0.246 eV agree well with the value obtained in susceptibility measurements $(J = 0.22 \pm 0.03 \text{ eV}).^{20}$

We mention that the agreement, especially close to the singularity, is not so good if *B* is artifically set to zero. This suggests (in accordance with the numerical evidence) that the exponent predicted by bosonization at low energies persists up to energies quite far from its range of applicability.

The value found for α_0 can be used to estimate the effective charge q_A . We find $|q_A|{\simeq}0.4e$ whereas in the 2D cuprates $|q_A|{\simeq}0.1e$ was found. This is not surprising since the processes that give rise to q_A are quite similar to the ones that give rise to J and the latter is found to be a factor of 2 larger in the 1D compounds.

Both the singularity and the cutoff at $\omega_2(\pi) + \omega_{\parallel}$ reflect the dominance of the spectra by two-spinon states analogous to the two-spinon states considered in Ref. 5 for the dynamical structure factor.

In conclusion, we presented a simple ansatz for the energy dynamical correlation function of the Heisenberg model. The ansatz reproduces very well numerical data and the low-energy behavior predicted by bosonization studies and provides new insight on the dynmaical properties of the 1D Heisenberg model. As an application we computed the phonon-assisted optical absorption of a 1D spin-1/2 system and show that it compares very well with recent experimental data.

After this work was completed we became aware of Ref. 13 which contains the experimental data and a qualitative analysis of the line shape. They predicted the same position for the singularity; however, their matrix elements were computed in the XY limit. Although at first glance the theoretical line shape resembles the experiment (but does not fit the data) they have missed the factor ω in Eq. (10) and the p-dependent form factor in Eq. (12). When they are included the resemblance is substantially lost. Additionally they get a cusp singularity instead of a logarithmic signularity.

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