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Spin-Exchange Dynamical Structure Factor of the $S = 1/2$ Heisenberg Chain

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We determine the spin-exchange dynamical structure factor of the Heisenberg spin chain, as measured by indirect resonant inelastic x-ray scattering (RIXS). We find that two-spin RIXS excitations nearly entirely fractionalize into two-spinon states. These share the same continuum lower bound as single-spin neutron scattering excitations, even if the relevant final states belong to orthogonal symmetry sectors. The RIXS spectral weight is mainly carried by higher-energy excitations, and is beyond the reach of the low-energy effective theories of Luttinger liquid type.

A remarkable feature of quantum systems in one dimension (1D) is that, in contrast to their higher dimension counterparts, quantum criticality is the norm rather than the exception [1]. 1D and quasi-1D systems as diverse as carbon nanotubes, stripes in cuprate high-temperature superconductors, confined ultracold atomic gases and quantum spin chains all provide realizations of critical quantum liquids. Possibly the most studied prototypical 1D quantum critical system is the Heisenberg $S = 1/2$ antiferromagnetic chain [2]. Its basic excitations are spinons, fractionalized spin excitations that emerge in the critical state [3]. Until recently, the only technique available to investigate these fractional spin excitations was inelastic neutron scattering (INS). In neutron scattering integer spin-flip excitations carrying $S = 1$ are created, so that the INS response involves the pairwise creation of spinons. The INS amplitude is determined by the single-spin dynamical structure factor (DSF) and a precise matching between the observed INS intensity and the dynamical structure factor calculated from theory has recently been achieved [4–6].

A few years ago, x-ray photon scattering emerged as a new tool to measure dispersive magnetic excitations [7]. When the incident photon energy is resonant with an absorption edge of the material, yielding resonant inelastic x-ray scattering (RIXS), the correlations between two neighboring spins factor into the magnetic x-ray scattering processes. The corresponding scattering amplitude is given by the momentum dependent two-spin dynamical structure factor, also referred to as the spin-exchange DSF. When for instance in an antiferromagnetic copper-oxide compound the incident energy is tuned to the copper $K$-edge—so-called indirect RIXS—exclusively this type of magnetic scattering occurs [7–11]. These recent experimental advances in inelastic x-ray scattering allow one to probe one of the most elementary, fractionalized states of critical matter in an entirely new manner, highlighting more elaborate spin-spin correlations.

It is the purpose of this Letter to provide a nonperturbative calculation of the spin-exchange RIXS scattering amplitude for the Heisenberg $S = 1/2$ antiferromagnetic chain with the Hamiltonian $H = J \sum_i (S_i \cdot S_{i+1} - 1/4)$. Our results aim on the one hand to challenge, guide and inspire further experimental magnetic RIXS efforts; on the other hand, the response function that we consider here is of fundamental significance and, as explicitly shown later on, beyond the reach of traditional low-energy based theories because it sits at energies of order of the exchange $J$. Our problem and approach thus represent an example of direct contact between experiment and theory beyond universality.

The model’s integrability [12–15] is well known to permit nonperturbative calculations of equilibrium properties, but it has recently also become possible to accurately compute certain dynamical properties for both finite [16–20] and infinite [21–23] systems. Until now these predictions have been limited to the single-spin DSF measured by INS. For experiments other than INS, governed by different correlators, at present no results are available. Here we fill this gap for RIXS, using the exact Algebraic Bethe Ansatz to calculate the spin-exchange DSF.

Magnetic RIXS cross section.—Before calculating its response function, we briefly describe the magnetic RIXS process. RIXS is a photon-in photon-out scattering technique in which the energy of the incident x-ray photons is tuned to an atomic absorption edge of the material that is studied. X rays can for instance be tuned to the $K$ edge of a transition metal ion ($5–10$ keV), for example, copper. In this case a magnetic scattering process as sketched in Fig. 1 can occur [8,9]. An x ray incoming on site $j$ produces a $1s - 4p$ electronic transition and creates a core-hole on the $j$th copper ion of the chain. In the intermediate state, the presence of the core-hole modifies the $3d$ on-site energy levels through the Coulomb interaction. In the Mott-insulating limit, this perturbation modifies the spin-exchange process with the two neighboring $3d$ electrons,
spin DSF, as measured in neutron scattering.

The cross section for this scattering process is given by the electron.

and thus locally modifies the actual superexchange coupling $J$ between the spins. Following the notation of [8,9] we denote the perturbed coupling $J^\epsilon = (1 + \eta)J$. In the final state, the core hole is filled again by the $4p$ electron, but the spin chain is left behind in an excited state. The cross section for this scattering process is given by the Kramers-Heisenberg relation. As a function of energy loss $\omega = \omega_{\text{in}} - \omega_{\text{out}}$ and momentum transfer along the spin chain $q = q_{\text{in}} - q_{\text{out}}$ the scattering intensity is $I = \sum_j |A_{ji}|^2 \delta(\omega - E_i + E_j)$, where the scattering amplitude

$A_{ji} = \omega_{\text{res}} \sum_{n, \epsilon} \langle f|D_{\text{in}}(n)|D_{\text{out}}\rangle_{\epsilon}$. Here $|i\rangle$, $|n\rangle$, $|f\rangle$ are the initial, intermediate and final states with respective energies $E_i$, $E_n$, $E_f$. The resonance energy is $\omega_{\text{res}}$ and $\hat{D}$ is the dipole operator. Because the 1s core-hole is highly energetic, it quickly decays, leading to an energy broadening $\Gamma$ of the intermediate state—its inverse lifetime. For $\Gamma > E_n$, $A_{ji}$ can be expanded in a power series which resummed to leading-order ultimately provides the x-ray scattering cross section $I \propto \sum_{n, \epsilon} |\langle f|D_{\text{in}}(n)|D_{\text{out}}\rangle_{\epsilon}|^2 S^{\text{exch}}(q, \omega)$. It incorporates the spin-exchange dynamical structure factor

$S^{\text{exch}}(q, \omega) = 2\pi \sum_\alpha \langle 0|X_{\alpha q}|\alpha\rangle^2 \delta(\omega - \omega_{\alpha}), \quad (1)$

where the ground state is $|0\rangle$, the excited states $|\alpha\rangle$, excitation energies $\omega_{\alpha} = E_{\alpha} - E_{\text{GS}}$ and the spin-exchange operator is $X_{\alpha} = \frac{1}{\sqrt{N}} \sum_j e^{i\eta j} (S_{j-1} \cdot S_j + S_j \cdot S_{j+1})$.

Motivated by the successful correspondence between predictions and experiments for the 2D Heisenberg antiferromagnet [9-11,24], we set out to compute this structure factor in the quantum critical 1D case.

**Computing the spin-exchange DSF.**—So far exact calculations for the spin chain have been restricted to the single-spin DSF, as measured in neutron scattering

$S^{\text{single}}(q, \omega) = 2\pi \sum_\alpha \langle 0|S_{\alpha q}^z|\alpha\rangle^2 \delta(\omega - \omega_{\alpha}), \quad (2)$

where $\langle 0|S_{\alpha q}^z|\alpha\rangle$ is the form factor (FF) of the Fourier transformed spin operators $S_{\alpha}^z = (1/\sqrt{N}) \sum_{j=1}^N e^{-i q j} S_j^z$, $\alpha = z, +, -$ between the ground state and eigenstates of excitation energy $\omega_{\alpha}$ (see Fig. 2). The strategy which we adopt for the calculation of the RIXS response function $S^{\text{exch}}$ is that of the ABACUS method [25]. Within this approach eigenstates are explicitly obtained from Bethe Ansatz, matrix elements from the algebraic Bethe ansatz, and the trace over intermediate states is performed numerically using an optimized search through the Hilbert space.

In what follows, we briefly describe each of these three steps needed to compute $S^{\text{exch}}$.

The Hilbert space can be divided into subspaces of fixed magnetization characterized by the number of downturned spins $M$. Eigenstates of the Hamiltonian for an (even) $N$ sites periodic spin chain are completely determined for $M \leq N/2$ by a set of $M$ rapidities $\{\lambda_1, \ldots, \lambda_M\}$ which for the isotropic chain solve the Bethe equations

$\arctan(2\lambda_i) = \frac{\pi}{N} I_i + \frac{1}{N} \sum_{k=1}^M \arctan(\lambda_i - \lambda_k) \quad (3)$

with $I_1, \ldots, I_M$ a set of integers for odd $M$ and half-odd integers for even $M$. Each set of quantum numbers specifies a set of rapidities. The ground state is defined by $|I_k = k - M/2\rangle$, $k = 1, \ldots, M$. The energy and the momentum of an eigenstate are given by

$E = -J \sum_{k=1}^M \frac{1}{2} \sum_{k=1}^M \frac{1}{4 + \lambda_k^2}$;

$P = \pi M - \frac{2\pi}{N} \sum_{k=1}^M I_k (\mod 2\pi)$.

Turning now to the matrix elements of the exchange operators $X_{\alpha}$, we exploit the spin isotropy of the system to express the full spin-exchange DSF matrix element in Eq. (1) as a function of the $S_{j+1}^z S_{j}^z$ FFs only. By globally rotating the $\langle 0|S_{j+1}^z S_j^z|\alpha\rangle$ and $\langle 0|S_{j+1}^z S_{j+1}^y|\alpha\rangle$ FFs appearing in Eq. (1) about the $y$ and $x$ spin axes, respectively, and using the fact that the ground state is a global $su(2)$ singlet, one can show that only singlet excited states contribute to Eq. (1). This conclusion is reached by first noticing that the $S_{j+1}^z S_j^z$ operator creates excited state only in the $S_{\text{tot}}^z = 0, 1, 2$ sectors. Second, the rotation of states belonging to these sectors and with zero magnetization ($S_{\text{tot}}^z = 0$) gives

![Fig. 1](image1.png)

**FIG. 1** (color online). Mechanism by which double-spin flip transitions are created in the indirect magnetic RIXS process.

![Fig. 2](image2.png)

**FIG. 2**. Single-spin dynamical structure factor $S^{\text{single}}(q, \omega)$ of the Heisenberg chain. The DSF is dominated by the 2-spinon spectrum but also 4-spinon and higher excited states contribute (6%). The computation is for $N = 400$ sites.
This is done using the ABACUS algorithm [25] which sums intermediate state contributions in a close to optimal order. By solving the Bethe equations, the eigenstate rapidities give the energy, momentum and the FF value through the determinant expression. Large families of excited states are summed over until satisfactory saturations of sum rules are achieved. More precisely, first we compute the computed integrated intensity with the analytical result in the infinite chain limit: $\int \frac{d\omega}{2\pi} \times \frac{\cos^2(q/2)\pi}{4} \sum_{\alpha \in \text{States}} \left| \sum_{j} e^{i q j} \langle 0| S_{j}^{z} S_{j+1}^{z} | \alpha \rangle \right|^2 \times \delta(\omega - \omega_{\alpha}).$ (4)

In order to calculate the matrix elements $\langle 0| S_{j}^{z} S_{j+1}^{z} | \alpha \rangle$, we make use of algebraic Bethe Ansatz methods [14,26–28] to represent these as explicit functions (taking the form of matrix determinants) of the rapidities involved in the left and right eigenstates. State norms are also given by an explicit determinant [13,29,30]. For brevity, explicit expressions are not presented here [31].

The third step, summation over intermediate states, can now be performed to obtain quantitative results for the DSF in Eq. (1). This is done using the ABACUS algorithm [25] which sums intermediate state contributions in a close to optimal order. By solving the Bethe equations, the eigenstate rapidities give the energy, momentum and the FF value through the determinant expression. Large families of excited states are summed over until satisfactory saturations of sum rules are achieved. More precisely, first we compute the computed integrated intensity with the analytical result in the infinite chain limit: $\int \frac{d\omega}{2\pi} \times \frac{\cos^2(q/2)\pi}{4} \sum_{\alpha \in \text{States}} \left| \sum_{j} e^{i q j} \langle 0| S_{j}^{z} S_{j+1}^{z} | \alpha \rangle \right|^2 \times \delta(\omega - \omega_{\alpha}).$ (4)

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$N = 400$. This ratio is even more drastic than for the INS results where the excitation of four and more spinons is responsible for 6% ($N = 400$) of the signal. Interestingly, the magnetic RIXS response is at first glance rather similar to the neutron scattering one, sharing the same lower bound of the spinon continuum, even if the RIXS and INS final states belong to different symmetry sectors and are orthogonal. This similarity is due to the fact that the dispersion relation of a spinon does not depend on its spin [3]. The existence of the spin-exchange DSF within a continuum is a direct and explicit consequence of the fractionalization of spin excitations in the quantum critical spin chain. Even if the excitation continuum probed by RIXS exactly coincides with the one probed by INS, the spectral weight has a markedly different distribution. This crucial difference is partly caused by the static factor $\cos^2(q/2)$ which originates from the modification of two neighboring exchange couplings in the x-ray scattering process. Excitations are thus associated to a typical length $2a$ (with $a$ the lattice spacing) or equivalently predominantly carry $\pm \frac{\pi}{2}$ (mod $2\pi$) momentum. The figures clearly illustrate this fact: for INS (Fig. 2) the signal is maximal at the antiferromagnetic wave vector $\frac{\pi}{2}$ and for RIXS (Fig. 3) the signal is maximal at the momenta $\frac{\pi}{4}$ and $\frac{3\pi}{4}$.
vector $q = \pi$ at $\omega$ close to zero, whereas the RIXS amplitude vanishes there. Rather it is concentrated above the continuum threshold at $q = \pi/2$, characterized by a vanishing group velocity. The fixed-momentum profiles in Fig. 4 show further differences between RIXS and INS apart from the $\cos^2(q/2)$ factor. While the weight of both responses predominantly sits between the top and bottom of the two-spinon continuum, the weight distribution between these two are clearly distinct, the RIXS response having a much broader shoulder at higher $\omega$ than the INS response. This quantitative difference is sufficiently large to be observable experimentally.

The most promising materials in this regard are Sr$_2$CuO$_4$ and SrCuO$_2$. The magnetically active part of Sr$_2$CuO$_3$ are chains of corner-sharing CuO$_4$ plaquettes. The exchange interaction $J$ between adjacent Cu spins on the chains is extremely large (200–250 meV); the interchain exchange interaction is about 5 orders of magnitude smaller. It is extremely large (200–250 meV); the interchain exchange interaction $J$ is about 5 orders of magnitude smaller. It is therefore a realization of a spin-1/2 antiferromagnetic Heisenberg chain over a wide temperature range, with high-energy magnetic excitations that are in principle accessible to Cu K-edge RIXS, using today’s energy resolutions. In SrCuO$_2$ the Cu-O chains are doubled, but with very weak exchange interactions between the two edge-sharing chains, rendering SrCuO$_2$ the other prime candidate to test our predictions.

The asymptotic behavior of correlations along the spin chain are well described by low-energy effective theories [1], which provide a detailed understanding of the antiferromagnetic singularity at momentum $\pi$ and zero energy in the INS response. The exponent of the singularity at the lower bound of the spectrum is given by the asymptotic decay of the two-spin correlation. Straightforward application of the known results for the four-spin correlator decay of the two-correlation. Straightforward application of the known results for the four-spin correlator [33] to the RIXS intensity fails as the signal vanishes at all low energies. Nonlinear extensions to Luttinger liquid theory [34] or alternatively the quantum group approach [21–23] might bring this within reach in the future.

We have, in conclusion, explicitly calculated the relevant magnetic response function for indirect RIXS on the Heisenberg chain, which involves the $S_i S_{i+1}$ spin-exchange operators. This operator predominantly probes high-energy magnetic excitations and the integrability-based method which we have employed here is then the only one capable of describing RIXS. Alternate approaches based on approximate low-energy theories cannot reach energies of order of the exchange $J$. We have demonstrated in more general terms that via the algebraic Bethe Ansatz formalism a new family of correlators of the type $S_i^a S_{i+1}^b$ with $a, b = \pm, +, -$ is accessible for computation. The challenges of future work are to extend the present exact calculation of the spin-exchange dynamical structure factor to anisotropic spin chains, including also finite magnetic fields.

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