Ground State Magnetization of Polymerized Spin Chains

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We investigate the ground state magnetization plateaus appearing in spin 1/2 polymerized Heisenberg chains under external magnetic fields. The associated fractional quantization scenario and the exponents which characterize the opening of gapful excitations are analyzed by means of abelian bosonization methods. Our conclusions are fully supported by the extrapolated results obtained from Lanczos diagonalizations of finite systems.

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Models of low-dimensional magnets, such as strongly correlated quantum spin chains \[\text{1}\] and ladders \[\text{2}\], are currently receiving renewed and systematic attention for a variety of reasons. Amongst the most remarkable are the spin-Peierls dimerization instability \[\text{3}\], the Hal-dane conjecture \[\text{4}\], and novel concepts such as fractional quantization and topological energy gaps \[\text{5}\]. These rather complex phenomena are largely owing to quantum fluctuations of individual spins which tend to restore the rotational symmetry of the ground state. Depending on the exchange interactions, fluctuations can manifest themselves collectively into many possible ground states, particularly in lower dimensions where their effects are most severe \[\text{6}\].

A wealth of issues have been addressed experimentally to confirm these expectations in a series of quasi-one-dimensional compounds \[\text{7}\]. After a vast body of research, it is by now well established that half-integer-spin chains are massless whereas integer ones are gapful (see also \[\text{8}\]). In spite of the availability of a number of excellent realizations of one dimensional Heisenberg antiferromagnets \[\text{9}\], detailed measurements of spin excitations however, have remained confined within the limits of applied magnetic fields which are low with respect to the interplay between explicit breaking of full translational symmetry and applied magnetic fields, say along the \(z\)-axis, in spin-1/2 Heisenberg chains at \(T = 0\). This can be conveniently described by a set of Heisenberg antiferromagnets in which the exchange coupling interactions \(J_n\) are all equal but one every \(p\) sites, i.e.

\[
J_n = \begin{cases} J(1 - \delta), & n/p \in \mathbb{Z}, \\ J, & \text{otherwise}. \end{cases}
\]  

(1)

It should be stressed that periodic arrays of couplings are relevant to the study of ferrimagnetic materials \[\text{10}\] and that also one dimensional dimerized and trimerized materials are known to exist \[\text{11,12}\].

The Hamiltonians of our polymerized chains in the presence of a (dimensionless) magnetic field \(h\) applied along the \(z\)-axis are thus given by:

\[
H_p = \sum_{n=1}^{L} J_n \vec{S}_n \cdot \vec{S}_{n+1} - \frac{h}{2} \sum_{n=1}^{L} S_n^z.
\]

(2)

Here, the \(\vec{S}_n\) are spin-1/2 operators, whereas periodic boundary conditions are assumed along the \(L\) sites of the chain \((\frac{L}{p} \in \mathbb{Z})\). Despite their simplicity, we will show however that these Hamiltonians entail a highly non-trivial magnetic behavior controlled solely by the chain periodicity \(p\) and the external field \(h\).

As is known \[\text{13}\], the full translationally invariant (FTI) \(S = \frac{L}{2}\)-chain remains gapless for all magnetizations \(\langle M \rangle \equiv \frac{1}{L} \sum_n S_n^z\), up to a saturation field where each of the \(L\) individual spins becomes fully polarized. On general grounds however, the Lieb-Schultz-Mattis theorem \[\text{14}\] indicates that FTI Hamiltonians of arbitrary spin-\(S\), can be gapful provided the magnetization per spin \(\langle M \rangle\) satisfies \((S - \frac{1}{2} \langle M \rangle) \in \mathbb{Z}\). Such gapful excitations should be reflected through the presence of magnetization plateaus, in principle at these special values of \(\langle M \rangle\). However, notice that the above theorem does not prove the existence of this quantization scenario as it refers to non-magnetic excitations, i.e. modes which preserve the total magnetization. Nevertheless, magnetization plateaus have been extensively conjectured and observed in both Peierls dimerized \[\text{15}\] and trimerized \[\text{16}\] spin chains, as well as in frustrated \[\text{17,18}\], anisotropic spin-1 systems \[\text{19}\] and ladders models \[\text{20}\]. They all are examples of a rather subtle phenomenon namely, fractional quantization of a macroscopic physical quantity under external varying fields. Here, we examine this situation for the whole class of non-homogeneous chains \[\text{3}\].
most relevant perturbation term is given by the interaction. After some algebra, in the limit of weak $\equiv$ one. Since the Fermi level is given by $k = \cdots$ (see e.g. [23]). The functional dependence of $R$ can be obtained from the exact Bethe Ansatz solution by solving a set of differential equations obtained in [24,25] (for a fuller derivation consult for instance Ref. [16]). Exploiting this knowledge, the bosonized expression of the low-energy effective Hamiltonian $\|\|\|$ in the homogeneous case $\delta = 0$, can be readily shown to adopt the form

$$H = \int dx \frac{\pi}{2} \left\{ \Pi^2(x) + R^2(\langle M \rangle)(\partial_x \phi(x))^2 \right\}, \quad (3)$$

with $\Pi = \cdots$ and $\phi = \phi_L + \phi_R$, $\dot{\phi} = \dot{\phi}_L - \dot{\phi}_R$. Here, the effect of the magnetic field $h$ enters through the radius of compactification $R(\langle M \rangle)$. This radius governs the conformal dimensions, in particular the conformal dimension of a vertex operator $e^{i\beta \phi}$ is given by $\frac{2}{\pi \beta}^2$. Within the framework of the theory of Luttinger liquids, it is worth pointing out also that the compactification radius is related to the parameter $K$ by $R^2 = \frac{4K}{\pi\beta}$.

The bosonized expressions for the spin operators read:

$$S^z_i(x) \approx \frac{1}{\sqrt{2\pi}} \frac{\partial\phi_i}{\partial x} + \text{const.} : \cos(2k_F^i x + \sqrt{4\pi \phi_i}) : + \frac{\langle M \rangle}{2}, \quad (4)$$

and

$$S^-_i(x) \approx e^{-i\sqrt{\pi} \phi_i} (1 + \text{const.} : \cos(2k_F^i x + \sqrt{4\pi \phi_i}) :), \quad (5)$$

where the colons denote normal ordering with respect to the ground state with magnetization $\langle M \rangle$. Now we apply this methodology to compute the effective form of the interaction. After some algebra, in the limit of weak polymerization $\delta \ll 1$, it can be readily shown that the most relevant perturbation term is given by

$$H_{\text{int}} \approx \delta \sum_{x'}^{L/P} \cos(2k_F(px' + \frac{1}{2}) + \sqrt{4\pi \phi}). \quad (6)$$

This operator will survive in passing from the lattice to the continuum model, assuming that the fields vary slowly, only when the oscillating factor $\exp(i2px'k_F)$ equals one. Since the Fermi level is given by $k_F = \frac{\sqrt{2}}{\pi}(1 - \langle M \rangle)$, this in turn will happen when the condition

$$\frac{P}{2}(1 - \langle M \rangle) \in \mathbb{Z} \quad (7)$$

is satisfied.

We can now study when a plateau will appear in the magnetization curve for the polymerized chain $\|\|$. To do that, we first have to see which are the values for the magnetization where there could be a plateau, for a given value of the period $p$, (i.e. solve for [3]) and then we need to evaluate the scaling dimension of the operator $\|\|$, which at zero loop is given by

$$d = \frac{1}{4\pi R^2}, \quad (8)$$

which is in turn governed by the radius of compactification as we already stressed. By virtue of the lower bound of the compactification radius $\|\|$, namely $R(\langle M \rangle) \geq r(\pm 1) = 1/(2\sqrt{\pi})$, it follows from Eq. (8) that $d < 2$ for all magnetizations $|\langle M \rangle| < 1$. This ensures the relevant character of the operator $\|\|$ which in turn survives in the continuum limit whenever Eq. (8) is satisfied. Therefore, we can conclude that constraint $\|\|$ constitutes a sufficient condition ultimately responsible for the appearance of magnetization plateaus and massive spin excitations. This is our main result.

We now turn to a numerical finite-size analysis. In Figs.1(a)–(e) we display a variety of magnetization regimes as a function of both polymerization parameters $J'/J \equiv 1 - \delta$ and applied magnetic fields $h$. This is a rather compact form of representing conventional magnetization curves for different polymerization strengths. Here, each line is associated to successive values of $\langle M \rangle$ which decrease monotonically from top to bottom, as they should for a non-frustrated system. The results were obtained from exact diagonalization of finite systems via a recursion type Lanczos algorithm [26] applied on each magnetization subspace $S^z = \{0, 1, ..., L/2\}$. To avoid the formation of spurious interfaces, even multiple lengths of the lattice periodicity were taken throughout. Using fully isotropic chains up to $L = 24$ sites with periodic boundary conditions, our numerical analysis supports entirely the quantization constraint $\|\|$. As expected, the ground state 'phase diagrams' exhibit bands of empty regions corresponding to the magnetization plateaus of $S^z$ referred to above, while regions filled with magnetization lines reflect smooth magnon excitations arising in the thermodynamic limit $L \to \infty$. It can be readily observed that for chains of periodicity $p > 1$ (dimers, trimers, etc), a plateau-like structure emerges precisely at the rational magnon densities $\langle M \rangle = 1 - 2q/p$. (Here $q = 0, 1, ..., p$) which is implicit in the general scenario of Eq. (8). It is worth mentioning the robustness of this topological constraint as similar results continue to hold for anisotropic (XXZ) chains, the plateaus always appearing at the same values of $\langle M \rangle$.

We can also predict the behavior of the mass gap (width of the plateau), with the polymerization strength $\delta$ by means of a simple zero loop computation [27]. Aside logarithmic corrections to the case $\langle M \rangle = 0$, this yields

$$g \propto \delta^{1/(2-d)}, \quad (9)$$
with \( d \) given as in \( [3] \).

To enable an independent check of this result, we now turn to the issue of extrapolating the numerical finite size estimates of the mass gaps \( g_L \) towards their corresponding thermodynamic limits. Note, on one hand, that any extrapolation procedure by necessity assumes that the asymptotic behavior applies to the values of \( L \) within reach. However, it is known \([24]\) that finite size corrections to the gap in the excitation spectrum of the homogeneous Heisenberg chain vary slowly as \( \ln (\ln L)/\ln^2(L) \), thus affecting the results over a wide range of sizes. In fact, as can be seen in Figs. \( \), this turns out to be the case also for weak polymerization regimes, \( \delta \to 0 \), where finite size effects are more pronounced. Therefore, in studying numerically the mass gap behavior obtained in Eq. \( [4] \) we are confronted to restrict considerations to the non-critical region \( 0 < |\delta| \leq 1 \), yet suitable to test independently the correctness of our bosonization approach.

To estimate the actual masses in the limit \( L \to \infty \), we fitted the whole set of finite-size results (even integer multiples of \( p \) within the range \( 4 \leq L \leq 24 \)), using both linear, and logarithmic type methodologies of convergence \([25]\), i.e.

\[
 g_L \simeq g + A e^{-B L}, \quad g_L \simeq g + A/L^B.
\]  

Either extrapolation procedure yields basically the same result with at least 3 significant digits. This latter variation ultimately gives an estimative idea of the lower bound of the extrapolation error. The reliability of our results was checked also by comparing the trend arising from smaller systems \( (L \leq 20) \). When the critical region is approached however, the accuracy differs widely, particularly for \( |\delta| < 0.2 \).

Although there are alternative extrapolation algorithms which do not involve fits to specific forms \([25]\) we should hasten to add however, that their efficiency depends strongly on the abundance of data. In our case, this is translated in the availability of matching sizes, already constrained by both the periodicity \( p \) and the antiferromagnetism. Nevertheless, we were able to find a remarkable agreement with the compactification radius comprehended in Eq. \( [3] \) and the exponents of Eq. \( [4] \). The results are shown in Fig. 2 where we display respectively the gap openings around \( \langle M \rangle = 0, 1/3, 1/2, \) for \( p = 2, 3, \) and 4. The dimerized case reproduces the well known \( 2/3 \) exponent predicted in \([3]\) and corroborated subsequently by diverse numerical studies \([20]\). To our knowledge however, opening exponents for \( p \geq 3 \) (see Fig. 2), have not been elucidated yet by other investigations.

Finally, it is instructive to comment further on the role of quantum fluctuations namely, the tendency of spins to spontaneously tilt occasionally due to the Heisenberg uncertainty relations, and their relevance to our results \([3] \). For classical spins, e.g. Ising and \( n \)-vector models, the interplay between dimensionality and statistical fluctuations, though crucial in determining phase transitions, is not sufficient to entail the fractional behavior studied so far. In fact, an elementary transfer matrix calculation shows that the Ising equivalent of \([2]\) wipes out all but two magnetization plateaus, namely \( \langle M \rangle = 0, 2/p \), (even \( p > 2 \)), their widths behaving linearly with \( \delta \). Thus, it is worth pointing out that Eqs. \( [3] \) and \( [4] \), in contrast, constitute a genuine macroscopic quantum effect.

In summary, we have presented a bosonization picture that accounts for the fractional quantization observed in a class of non-homogeneous Heisenberg antiferromagnets. All low-energy exponents which characterize the opening of gapful excitations have been obtained and treated on an equal footing while checked with Lanczos diagonalizations of finite systems. Aside these theoretical pursuits, we trust our study will help to convey a clearer understanding of the many characteristics present in real low dimensional magnets. A similar analysis in polymerized ladder systems is in progress.

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5126 (1997); Phys. Rev. B58, 6241 (1998).
[17] T. Tonegawa et al, to be published in Physica B (Proc. 5th Int. Symp. on Research in High Magnetic Fields, Sydney, 1997), preprint [cond-mat/9712297]. B.S. Shastry and B. Sutherland, Phys. Rev. Lett. 47, 964 (1981).
[18] K. Totsuka, Phys. Rev. B57, 3454 (1998).
[19] S. Yamamoto, S. Brehmer, H.-J. Mikeska, preprint [cond-mat/9710332] and references therein.
[20] R.B. Griffiths, Phys. Rev. 133, A768 (1964); C.N. Yang and C.P. Yang, *ibid.* 150, 321 (1966).
[21] E. Lieb, T. Schultz, D. Mattis, Ann. Phys. 16, 407 (1961); I. Affleck and E.H. Lieb, Lett. Math. Phys. 12, 57 (1986).
[22] H.J. Schulz, Phys. Rev. B34, 6372 (1986).
[23] F.D.M. Haldane, Phys. Rev. Lett. 45, 1358 (1980).
[24] V.E. Korepin, N.M. Bogoliubov, A.G. Izergin, *Quantum Inverse Scattering Method and Correlation Functions*, Cambridge University Press, Cambridge (1993).
[25] N.M. Bogoliubov, A.G. Izergin, V.E. Korepin, Nucl. Phys. B275, 687 (1986).
[26] See, for example G.H. Golub and C.F. Van Loan, *Matrix Computations*, 3rd edition (Johns Hopkins University Press, Baltimore 1996).
[27] I. Affleck, et al., J. Phys. A 22, 511 (1989).
[28] For a review, consult A.J. Guttmann in *Phase Transitions and Critical Phenomena*, edited by C. Domb and J. Lebowitz (Academic Press, New York 1990), Vol. 13. See also C.J. Hamer and M.N. Barber, J. Phys. A14, 2009 (1981).
[29] G. Spronken, B. Fourcade and Y. Lépine, Phys. Rev. B 33, 1886 (1986), and references therein.
FIG. 1. Magnetization contours of finite polymerized chains for (a) $p = 2$, $L = 24, 20, 16$; (b) $p = 3$, $L = 24, 18, 12$; (c) $p = 4$, $L = 24, 20, 16$; (d) $p = 5$, $L = 20, 10$ (full and dotted lines respectively) and; (e) $p = 6$, $L = 24, 18, 12$. Except for (d), full, dashed, and dotted lines stand respectively for large, medium and small sizes. They denote all accessible magnetizations, whereas their values decrease from top to bottom. Though numerical accuracy in $h/J$ is bounded by $10^{-7}$, size effects become evident for $J' = J$, as no plateaus (empty wide bands), should occur in the thermodynamic limit.

FIG. 2. Extrapolated values of the gap for $p = 2$ around $\langle M \rangle = 0$, $p = 3$ for $\langle M \rangle = 1/3$ and, $p = 4$ with $\langle M \rangle = 0, 1/2$. Solid lines are guide to the eye whereas slopes of dashed lines denote the estimated opening exponents, namely (in descending order), 0.8(1), 0.77(10) and, 0.66(10), ($\langle M \rangle = 0$). To improve the clarity of the figure, the uppermost curve was shifted multiplying the gap by a scale factor 2.5.