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SPECTRAL DENSITIES AND DIAGRAMS OF STATES OF ONE-DIMENSIONAL IONIC PAULI CONDUCTOR

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Спектральні густини та діаграми станів одновимірного іонного провідника Паулі

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Спектральні густини та діаграми станів одновимірного іонного провідника Паулі

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Анотація. Робота присвячена вивченню енергетичного спектру та діаграм станів, отриманих методом точної діагоналізації для скінченного іонного ланцюгового провідника в періодичних граничних умовах. Одновимірний іонний провідник описується ґратковою моделлю, де іони розглядаються як частинки Паулі, при цьому враховується іонний перенос і двочастинкова взаємодія між найближчими сусідами. Було розраховано та проаналізовано спектральні густини та діаграми стану такої системи для різних температур та величин взаємодії. Проаналізовано умови переходу системи з однорідного (стану т.зв. моттівського діелектрика) у модульований стан через стан типу фази з бозе-конденсатом (подібної до надплинної фази в моделях жорстких бозонів).

Spectral densities and diagrams of states of one-dimensional ionic Pauli conductor.

I.V.Stasyuk, O.Vorobyov, R.Ya.Stetsiv

Abstract. We focus on the features of spectra and diagrams of states obtained via exact diagonalization technique for finite ionic conductor chain in periodic boundary conditions. One dimensional ionic conductor is described with the lattice model where ions are treated in frames of 'mixed' Pauli statistics. The ion transfer and nearest-neighbour interaction between ions are taken into account. The spectral densities and diagrams of states for various temperatures and values of interaction are obtained. The conditions of transition from uniform (Mott insulator) to the modulated (charge density wave state) through the superfluid-like state (similar to state with the Bose-Einstein condensation observed in hard-core boson models) are analyzed.

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1. Introduction

Ionic conductors are a wide class of physical and biological objects ranging from ice to DNA membranes. One of the most interesting subclass of all are superionic conductors that exhibit high temperature phase with high conductivity that arises due to the motion of ions [1] or protons [2].

Theoretical description of systems with ionic conductivity is most frequently based on the lattice models. Some of them treat ions as Fermiparticles focusing on different aspects of the ionic subsystem like longrange interactions [3–5] or interaction with phonons [6,7]. Some recent attempts have also paid some attention to short-range interaction between particles [8–12].

However more correct consideration of ions should be based on the mixed statistics of Pauli [13] since these particles are bosons by nature but they also obey the Fermi rule. Because of the special commutation rules the utilization of Pauli operators generates additional mathematical complexities. But from the other hand this approach might be very effective. For instance it has been shown that the lattice model of Pauli particles is capable to describe the appearance of superfluid-like state (that correspond to superionic phase) in the system even in the absence of interaction between particles [14–16]. From the other hand the lattice model of Pauli particles is similar to hardcore Bose-Hubbard model widely used for the description of ionic conductivity phenomena as well as for the modeling of energy spectrum of absorbtied ions on crystal surface and intercalation in crystals [17]. Bose-Hubbard model also exhibits the transition from Mott insulator state to superfluid-like state [18–24]. Some of the authors also observe the possibility of formation of intermediate "supersolid" phase that may appear on the phase diagrams alongside the transition from dielectric (CDW) to superfluid phase.

In this work we focus on the diagrams of state for one-dimensional ionic conductor described by the system of Pauli particles. Our lattice model include ion transfer as well as the interaction between nearestneighbouring ions. We calculate the single-particle spectral densities of the finite system in periodic boundary conditions and obtain the diagrams of state analyzing the features of this spectra. The conditions of transition from Mott insulator (MI) like state to the modulated charge density wave (CDW) one through the superfluid(SF)-like state (similar to state with the Bose-Einstein (BE)condensation observed in hard-core boson models) are discussed.

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Figure 1. The model for one-dimensional ionic conductor. Large circles denote heavy ionic groups while the small one denote light movable ions.

2. The model for ionic conductor

Let us consider the chain of heavy immobile ionic groups (large circles on Fig. 1) and light ions that move along this chain occupying positions denoted with small circles on Fig. 1. The subsystem of light ions is described with the following Hamiltonian

$$\hat{H} = t \sum_{i} (c_i^+ c_{i+1} + c_{i+1}^+ c_i) + V \sum_{i} n_i n_{i+1} - \mu \sum_{i} n_i.$$
(1)

This model takes into account the nearest-neighbour ion transfer (with hopping parameter t) and interaction between ions that occupy nearest-neighbouring positions (with corresponding parameter V).

If this Hamiltonian is considered in frames of Fermi statistics the corresponding model is known as spinless-fermion model. This model is widely used in the theory of strongly correlated electron systems [25] as well as for the description of ionic conductors [26]. More complex two-sublattice case of this model can be applied to proton conductor [27].

More correct consideration of ions should be based on "mixed" Pauli statistics and this approach is used onwards. In this case the model (1) is equivalent to the extended hard-core boson model, i.e. boson Hubbard model with repulsive interaction between nearest neighbours and infinite on-site repulsion [28]. The latter is often applied to the investigation of the problems of BE-condensation and superfluidity.

3. Exact diagonalization technique

We calculate the spectral densities of one-dimensional ionic Pauli conductor using exact diagonalization technique. For the chain of N sites we introduce the many-particle states $|n_{1,a}n_{1,b}...n_{N,a}n_{N,b}\rangle$. The Hamiltonian matrix on the basis of these states is the matrix of the order $2^N \times 2^N$ and is constructed according to the following way

$$H_{mn} = \sum_{i=1}^{N} \left[t \left(H_{mn}^{(1)} + H_{mn}^{(2)} \right) + \widetilde{V} H_{mn}^{(3)} - \mu H_{mn}^{(4)} \right], \qquad (2)$$

where

$$\begin{split} H_{mn}^{(1)} &= \langle n_1 \dots | c_i^+ c_{i+1} | n'_1 \dots \rangle = \delta(n_i - n'_i - 1) \delta(n_{i+1} - n'_{i+1} + 1) \\ &\times \prod_{l \neq i; i+1} \delta(n_l - n'_l), \\ H_{mn}^{(2)} &= \langle n_1 \dots | c_{i+1}^+ c_i | n'_1 \dots \rangle = \delta(n_i - n'_i + 1) \delta(n_{i+1} - n'_{i+1} - 1) \\ &\times \prod_{l \neq i; i+1} \delta(n_l - n'_l), \\ H_{mn}^{(3)} &= \langle n_1 \dots | n_i n_{i+1} | n'_1 \dots \rangle = \delta(n_i - 1) \delta(n'_i - 1) \delta(n_{i+1} - 1) \\ &\times \delta(n'_{i+1} - 1) \prod_{l \neq i; i+1} \delta(n_l - n'_l), \\ H_{mn}^{(4)} &= \langle n_1 \dots | n_i | n'_1 \dots \rangle = \delta(n_i - 1) \delta(n'_i - 1) \prod_{l \neq i} \delta(n_l - n'_l). \end{split}$$

This matrix is diagonalized numerically

$$U^{-1}HU = \widetilde{H} = \sum_{p} \lambda_{p} \widetilde{X}^{pp}, \qquad (3)$$

where λ_p are eigenvalues of the Hamiltonian, \widetilde{X}^{pp} are Hubbard-operators. The same transformation is applied to the creation and annihilation operators

$$U^{-1}c_i U = \sum_{pq} A_{pq} \widetilde{X}^{pq} \quad , \quad U^{-1}c_i^+ U = \sum_{pq} A_{rs}^* \widetilde{X}^{rs} \tag{4}$$

which are required to construct one-particle Green's function $\ll c_{i,a}|c_{i,a}^+ \gg$ that contains information about one-particle energy spectrum of the system. For Pauli creation and annihilation operators this Green's function can be constructed in two ways, i.e. commutator Green's function

$$\ll c_i(t)|c_i^+(t') \gg = -i\Theta(t-t')\langle [c_i(t), c_i^+(t')]\rangle$$
(5)

and anticommutator Green's function

$$\ll c_i(t) | c_i^+(t') \gg = -i\Theta(t - t') \langle \{ c_i(t), c_i^+(t') \} \rangle.$$
(6)

Imaginary part of these Greens' functions are one-particle spectral densities (also referred to as densities of states or DOS)

$$\rho(\omega) = -\frac{1}{\pi N} \sum_{i=1}^{N} \operatorname{Im} \ll c_{i,a} | c_{i,a}^{+} \gg$$
$$= -\frac{1}{\pi N} \sum_{i=1}^{N} \operatorname{Im} \left[\frac{1}{Z} \sum_{pq} A_{pq} A_{pq}^{*} \frac{e^{-\beta\lambda_{p}} - \eta e^{-\beta\lambda_{q}}}{\omega - (\lambda_{q} - \lambda_{p})} \right], \quad (7)$$



Figure 2. Commutator spectral densities of non-interacting ($V_{ef} = w_{ef} \equiv V = 0$) ionic Pauli conductor for different values of chemical potential (right figure) compared to exact results obtained in [14] via fermionization procedure (left figure). $t = 1, T = 0.2, \Delta = 0.4$. Spectral density on the left figure is scaled to 2 π , while on the right figure it is scaled to unity.

where $Z = \sum_{p} e^{-\beta \lambda_{p}}$. Spectral densities in (7), obtained from commutator $\eta = 1$ (5) and anticommutator $\eta = -1$ (6) Greens' functions respectively, exhibit discrete structure, i.e. consist of several δ -peaks due to the finite size of a cluster. Therefore we apply the periodic boundary conditions to the cluster and introduce small parameter Δ to broaden the δ -peaks according to Lorentz distribution

$$\delta(\omega) \to \frac{1}{\pi} \frac{\Delta}{\omega^2 + \Delta^2}.$$
 (8)

4. Results and discussion

We perform calculations of one-particle spectral densities of one-dimensional ionic Pauli conductor (1) for the chain of ten sites (N = 10) in periodic boundary conditions. To test the results we compare them with the exact solution obtained by means of fermionization procedure [14] in the absence of nearest neighbour interaction (V = 0) for different values of chemical potential (Fig. 2). The redistribution of statistical weight with the change of chemical potential level can be observed and the good level of agreement is achieved.

The analysis of the spectral densities is the way to distinguish the different states of ionic subsystem and the corresponding conditions. In the case of half-filling, as we turn on the interaction starting from spectral



Figure 3. Commutator spectral density of ionic Pauli conductor for different values of nearest-neighbour interaction ($V_{ef} = w_{ef} \equiv V$) at half filling ($\mu = 0$). $t = 1, T = 0.2, \Delta = 0.4$.

density which shape corresponds to SF state, we observe the development of the gap in the spectra (Fig. 3). The similar effect was found for ionic and proton conductors described by the analogous models in frames of Fermi statistics. It was connected with the splitting of spectra due to charge ordering with doubling of lattice period [11,12]. The detailed analysis of spectral densities of Fermi and Pauli models of ionic conductor can be found in [29]. So, we have here a transition from SF to CDW state (at $T \neq 0$ this transition manifests itself as a crossover), described analytically in [30,31].

The existence of the gap in the spectra which separates the bottom of energy band from the chemical potential level is also the sign of the presence of homogenous MI state. The other specific feature of the spectra is the appearance of negative branch that points to the transition to SF-like state (see, for example, [32]). According to these criteria we analyze the spectral densities at different temperatures and values of interaction and build the corresponding diagrams of state (Fig. 4). The system is in MI homogenous state at high temperatures and far away from half-filling (at large δ). As the temperature decreases or one comes closer to half-filled case the system undergoes transition to SF-like state that corresponds to the appearance of the negative branch on the spectral density. At farther temperature decrease and closer to half filling we observe transition to the state with the gap on the spectral density that corresponds to the CDW-ordering. As the interaction strength Vincreases such a region becomes broader while the region of SF-like state becomes smaller. From the other hand, with the decrease of V the CDW state diminishes and disappears at $V \approx t$. It should be mentioned that



25-

2.0

1.5

1.0

0.5

0.0 -

0.00 0.05

⊢



Figure 4. Diagrams of state for different interactions and the spectral densities that correspond to each state. $t = 1, \Delta = 0.25$. $\delta = \langle n \rangle - 1/2$ denotes the deviation from half-filling.



Figure 5. The curves that separate SF- and MI- states at different temperatures and values of interaction. SF-state is on the left side of the curves and below them. $t = 1, \Delta = 0.25, \delta = < n > -1/2$ denotes the deviation from half-filling.

0,35

sequence of states, the system is going through at the increase of mean occupancy δ , corresponds to phase diagram obtained in [33]

We have also performed the detailed analysis of the transition to SFlike state at different temperatures and values of interaction (Fig. 5). It is interesting that at weak interactions (V < 1) the increase of interaction strength facilitates the formation of SF-like state while farther increase of V suppresses this transition.

5. Conclusions

We have performed analysis of diagrams of states of one-dimensional Pauli ionic conductor with exact diagonalization technique. We have shown that the system undergoes transition from conducting to superfluidlike state and then to insulating sate. At weak interaction the latter transition may vanish.

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