External electric field mediated quantum phase transitions in one-dimensional charge-ordered insulators: a density matrix renormalization group study

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Abstract

We perform density matrix renormalization group calculations extensively on one-dimensional chains with on-site \((U)\) as well as nearest-neighbour \((V)\) Coulomb repulsions. The calculations are carried out in full parameter space with explicit inclusion of the static bias and we compare the nature of spin-density-wave (SDW) and charge-density-wave (CDW) insulators under the influence of external electric fields. We find that, although the SDW \((U > 2V)\) and CDW \((U < 2V)\) insulators enter into a conducting state after a certain threshold bias, CDW insulators require much higher bias than the SDW insulators for insulator–metal transitions at zero temperature. We also find the CDW–SDW phase transition on application of an external electric field. The bias required for the transitions in both cases decreases with increase in system size.

Strongly correlated electronic systems in low dimensions are always very interesting because of their unique characteristics like quantum fluctuations, lack of long range ordering etc [1–7]. The correlation effects in such low-dimensional systems lead to Mott or charge-ordered insulating states. The one-dimensional extended Hubbard model \([8, 9]\) with on-site Hubbard repulsion term, \(U\), along with the nearest-neighbour Coulomb repulsion term, \(V\), is a standard model which exhibits these different phases. This is perhaps the simplest possible model which can capture many interesting properties of strongly correlated systems. In the strong coupling limit, this model gives rise to two insulating phases, the spin-density-wave (SDW) and charge-density-wave (CDW) phases, which are separated by a first-order phase transition line at \(U \simeq 2V\). On the other hand, in the weak coupling limit, the perturbative analysis shows that the transition is continuous at \(U = 2V\) [10, 11].

The effect of electric field on such insulating phases has attracted much interest in recent times due to the practical applications in tuning their dielectric and piezoelectric properties [12]. Many electronic conduction processes seem to suggest electric field induced phenomena, such as negative differential resistance in molecular electronics [13, 14]. Experiments on low-dimensional Mott insulators with spin-density-wave and charge-ordered phases as ground states, suggest a collapse of the insulating phase in the presence of an external electric field at finite temperatures [15–25]. However, the breakdown in charge-ordered phases is not due to Joule heating. Rather it has been argued that it is the applied bias which generates a conduction pathway resulting in metallic characteristics [18]. However, a tractable computational method which can take into account the static electric field and its response to the correlated extended electronic systems is still lacking.

In this paper, we use the density matrix renormalization group (DMRG) [26–28] method which is known to be highly accurate for low-dimensional interacting systems with high precision. We have included an electric field in the DMRG algorithm and have obtained ground and excited eigenstate behaviours of one-dimensional CDW and SDW insulating systems with various system sizes. The static electric field is included as a ramp potential and we find that the electric field can induce an insulator–metal–insulator transition in...
both cases, with a strong dependence on the Hamiltonian parameters. Instead of the field per unit length, we consider the total field, applied between two ends of a one-dimensional chain, irrespective of the chain length. Similar consideration was applied for SDW insulators, where we extrapolated our finite size results to very large (effectively infinite) systems to obtain thermodynamic behaviour [29]. In the present study on the insulating CDW phase, we find the CDW–SDW quantum phase transition along with the breakdown of insulating phase on application of bias.

We consider the one-dimensional strongly correlated chain described by the extended Hubbard Hamiltonian

\[ H = t \sum_i (a_i^\dagger a_{i+1} + \text{h.c}) + U \sum_i n_i^+ n_i^- + V \sum_i (n_i - n_{\text{av}})(n_i^+ - n_{\text{av}}) \]  

where \( t \) is the hopping term, \( U \) is the on-site Hubbard repulsion term and \( V \) is the nearest-neighbour Coulomb repulsion term. We set \( t = 1 \) in all our calculations and express every energy in units of \( t \). \( a_i^\dagger \) (\( a_i \)) is the creation (annihilation) operator and \( n \) (\( n_{\text{av}} \)) is the number operator (average electron density on every site).

The external electric field applied to the system has the form of a ramp potential, distributed over all the sites in such a way that the potential \( F_i \) at site \( i \) becomes \( -F_i/F_{\text{max}} \), where \( F \) is the total applied bias and \( N \) is the total number of sites in the 1D chain. This form of the potential ensures that the bias varies between \(-F/2\) and \( F/2\) across the molecule. The potential adds an extra term \( \sum_i F_i a_i^\dagger a_i \) to the above Hamiltonian. In DMRG, we use a density matrix cut-off, \( m = 140 \). For \( U = 0 \) and \( V = 0 \), the problem can be exactly solved and we obtain the ground state and excitation spectrum in the presence of bias using the tight-binding one-electron formalism. We perform our calculations keeping a fixed value of \( U (=5) \) and varying the \( V \) from 0 to 4 with bias from 0 to 10 volts in steps of 0.5 volts.

With increase in the value of \( V \), one encounters three different regions, namely, the spin-density-wave one (\( 2V < U \)), the spin-density-wave to charge-density-wave crossover one (\( 2V = U \)) and the charge-density-wave one (\( 2V > U \)). To understand the effect of electric field on the excited states of the system, for a given value of \( U \) and \( V \), we have computed the energetics of the systems with one extra \( E(N + 1) \) and one less electron \( E(N - 1) \) than for half-filling \( E(N) \) and have calculated the many-body charge excitation gap as the difference between the energy required to add (\( \mu_+ \)) and remove (\( \mu_- \)) electrons from the ground state [30]:

\[ \Delta_{\text{charge}} = \mu_+ - \mu_- \]  

where \( \mu_+ = E(N + 1) - E(N) \) and \( \mu_- = E(N) - E(N - 1) \).

To investigate the effect of polarization on the applied electric field, i.e., the screening of the external electric field by the shifted electron density, we solve the self-consistent Poisson equation for SDW as well as for CDW phases. We have considered the boundary conditions so that the final electrostatic potential field extending from one electrode to another becomes different for different atomic sites and the atomic electron density will adjust to the field in such a way that the charges are stabilized locally. We start our self-consistent calculation by assuming that the electrostatic field is a linear ramp function across the interface of the electrode and the system. By solving the Schrödinger equation within the many-body limit we obtain the charge density at every site and use that as an input in the one-dimensional Poisson equation. The diagonal elements of the Hamiltonian are then modified with the modified bias, obtained from Poisson’s equation. Solving the modified Hamiltonian we obtain the charge densities and again use them as input in Poisson’s equation. This is continued until all the charge densities and all the site potential fields converge. In figure 1 we plot both the on-site atomic charge densities and the spatial distribution of potentials for both SDW and CDW phases with explicit consideration of the screening of the applied electric field and show the same, without considering the screening, for comparison. From figure 1 it is clear that, in the presence of strong electronic correlations, the ramp nature of the external electric field is retained without any significant change of the charge density or potential profile with the consideration of the polarization effect. Here we consider the system with \( N = 10 \) and use the exact diagonalization method, and we present the results for two different bias values. The ground state of the CDW phase is a linear combination of two configurations (having alternating doubly occupied sites) with the same weighting. So we take the average of the charge densities and the potentials of the two degenerate ground states in the case of the CDW phase. Moreover, for the CDW phase we consider a fairly large value of the bias for investigating the breakdown region, which we discuss later. We infer that large correlations reduce Poisson’s equation to the Laplace equation with no significant variation of the charge-density distribution which we observed in our earlier studies also [31, 32]. The inclusion of the effect of polarization can change the quantitative estimation of the results without any significant change of the physics. So, to avoid the computational expense of the self-consistent Poisson equation.
calculations, we ignore the polarization effect in our further DMRG calculations with larger system sizes.

In figures 2(a)–(c) we have plotted the charge gap for 1D chains with different $N$ as a function of bias for CDW, SDW–CDW crossover and SDW phases, respectively. For the SDW phase and the SDW–CDW crossover region, the charge gap decreases with increasing bias up to a minimum, whereas in the case of the CDW phase, the charge gap increases initially with increasing bias. In the latter case, the nearest-neighbour Coulomb repulsion term, $V$, is higher than in the previous two cases, and as a consequence, the electrons favour remaining paired in alternate sites over increasing kinetic energy by hopping to the neighbouring sites. The SDW phase has a charge gap of $U$ in the $U \rightarrow \infty$ limit. But the CDW phase with double occupancy at alternate sites for a half-filled 1D chain has smaller charge gap than $U$ and at the $U, V \rightarrow \infty$ limit, the charge gap is $U - V$. Therefore, in the CDW phase, $V$ is the stabilizing term and this can be proved unambiguously from the model Hamiltonian. Bias, however, ‘tries’ to make the electrons hop in its direction and as a consequence the site occupancies start changing, leading to a higher charge gap. It is thus the hopping which creates a larger gap in the charge-ordered phase inducing a positive contribution from $V$. However, this initial increase is followed by a decrease in charge gap at quite a higher bias, although the charge gap minimum cannot be seen for all the system sizes in the bias window that we have considered. In the case of the SDW phase, the bias overcomes the effect of $U$ by forcing the electrons to hop in its direction and leads the system to a charge gap minimum corresponding to a conducting state. So, it is clear from figure 2 that the CDW phase needs much higher bias to pass through a charge gap minimum than the SDW phase. The SDW–CDW crossover region shows a combination of the properties of the SDW and CDW phases. It can also be seen from figure 2 that the increase in system size reduces the initial charge gap for all three cases and reduces the bias corresponding to the first charge gap minimum in the case of the SDW phase and the SDW–CDW crossover region.

To understand the response of the ground state as well as the excited states under the influence of external electric field, we plot $E(N+1)$, $E(N)$ and $E(N-1)$ for a system with 40 sites as a function of the bias in figure 3 for the CDW phase ($V = 3$), the SDW–CDW crossover region ($V = 2.5$) and the SDW phase ($V = 1$). For all three cases, the ground state energies do not change much up to a threshold bias. However, beyond that, the systems start to stabilize. The slope of the ground state energy is different from those for the excited states for all three phases and this difference in slope gives rise to a collapse of the insulating phase at some threshold bias, which is different for different phases and strongly depends on the system size. In contrast to the first two cases, although the CDW phase with one less electron starts getting stabilized with increasing bias, the system with one more electron destabilizes slightly in low bias and this is followed by a reduction in energy at higher bias values. This is due to the fact that one more electron in the CDW phase leads to more prominent electronic repulsion for bias driven hopping because of the higher $V$ value. As a consequence, the charge gap increases initially; however, beyond a certain bias value, required to nullify the effect of $V$, the gap reduces.

To provide a clear understanding of the charge gap variation in the CDW insulating phase with increasing bias, in
The dipole moment as a function of bias for (a) $V = 3$ and (b) $V = 1$ for $N + 1$ (dotted lines), $N$ (solid lines) and $N - 1$ (dashed lines) electron states for a system size of $N = 40$.

Figure 5. The dipole moment as a function of bias for (a) $V = 3$ and (b) $V = 1$ for $N + 1$ (dotted lines), $N$ (solid lines) and $N - 1$ (dashed lines) electron states for a system size of $N = 40$. 

In our previous study [29] on the SDW phase, the initial increase in charge gap is only wider with increase in the size of the system as observed in figure 4 it is clear that the period of oscillation becomes narrower with increase in the size of the system as observed in our previous study [29] on the SDW phase.

To provide an understanding of the underlying reasons for such behaviour of the charge gap, in the inset of figure 4 we plot average charge-density shift per site for the $N = 40$ one-dimensional chain with $N + 1$, $N$ and $N - 1$ electrons as a function of bias. As can be seen, although all the three states show staircase-like behaviour, for the system with one extra electron the nature of the shift of the charge densities is completely different from that for the other two cases. The charge-density shifts slowly at lower bias in the case of a half-filled system and the system with one less electron, whereas the charge-density shift for the system with one extra electron is almost zero up to a certain bias, termed $V_{\text{trans}}$, for this system, beyond which the charge density shows a drastic shift. This is due to the fact that the extra electron faces a more prominent effect of Coulomb repulsion terms, and hopping of electrons is possible only at some bias value which can overcome the electronic repulsion. After $V_{\text{trans}}$, the CDW phase again goes to an insulating phase, characterized by the plateau in the plot, which we anticipate to be attributable to a SDW phase. If we further increase the bias, after certain values the shift in charge densities for all the three systems shows a sudden jump, which indicates kinetic stabilization, followed by plateaus, indicating the reappearance of the insulating phase. We argue that this repetitive period for the charge gap with increase in bias is due to the charge stiffness. The external bias sweeps the charge densities towards one direction with nullification of repulsion terms up to the first minima. Beyond this, the bias 'tries' to shift the charge densities further and the electronic repulsion terms again reappear and start opposing the kinetic stabilization, and as a consequence, the charge gap again increases and the system enters into an insulating phase. Thus, this repetitive metallic and insulating behaviour in different bias regions is a consequence of the interplay between the external bias and the electron–electron interaction terms. However, this oscillation of the charge gap with increase in bias is due to finite size effects. With increase in system size, the periodicity narrows down and we can anticipate that, at infinite system size, all the minima will converge to the same bias value as was found from our previous study [29]. We termed this bias value at the thermodynamic limit the critical bias, $V_c$. Note that, as we discussed earlier, we do not consider the effect of polarization on the applied electric field in our calculations. It can only change the quantitative estimation of $V_{\text{trans}}$ and the critical bias, $V_c$, without any qualitative change in the basic physics.

To obtain a better insight, we plot the dipole moment of the ground state and the states with one extra and one less electron for both SDW and CDW phases as a function of bias in figure 5. It can be seen clearly that the ground state dipole moment shows periodic jumps at the bias values, corresponding to the charge gap minima, and the plateaus indicate the successive insulating phases. So, whenever there is a shifting of charge densities, i.e., sudden increase in the dipole moment, the system allows the electrons to hop, leading to breakdown of insulating phase. These periodic jumps of the dipole moment are purely due to finite size effects. However, the CDW–SDW phase transition cannot be understood from the ground state dipole moments.

To understand the CDW–SDW phase transition, we calculate the order parameter [10] in terms of on-site charge densities $\langle n_i \rangle$ for a finite system, consisting of 40 sites with half-filling and with one less and one more electron than half-filling:

$$ m = \frac{1}{N} \sum_i (-1)^i \langle n_i \rangle $$

and plot its absolute value in figure 6. As can be seen, after a certain bias, the $|m|$ value for the system with one extra electron drops, indicating clearly a phase transition to the SDW phase. The $|m|$ values for the other two cases also show drops at some bias values, which indicates an insulator to metal phase transition. From the above observations, we can conclude that the system with one extra electron can directly indicate the CDW–SDW phase transition on application of an external bias. Although the $|m|$ values show oscillations in figure 6, we can...
CDW phase decreases with the increase in bias and after a certain bias it behaves exactly like the SDW phase, depicted in figure 7(b). This resemblance in behaviour of the correlation function strongly indicates a CDW–SDW phase transition.

In conclusion, we have studied the application of external electric field to both the SDW and CDW insulating phases with on-site and nearest-neighbour Coulombic repulsion. We find that the external electric field can induce an insulator–metal transition in both cases. Moreover, the electric field induces a CDW–SDW quantum phase transition as well. Increase in system size increases the sharpness of both the transitions. Our DMRG calculations allow us to study large systems with explicit application of an electric field, with microscopic understanding of the insulator–metal transition in spin-density-wave and charge-density-wave insulators.

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Figure 6. Absolute value of the order parameter, \( m \), for the CDW phase \((V = 3)\) with \( N + 1 \) (triangle), \( N \) (square) and \( N - 1 \) (circle) electrons as a function of bias.

Figure 7. The charge–charge correlation as a function of the correlation length for (a) \( V = 3 \) at bias values 0.0 (circle), 0.5 (square), 1.0 (triangle) and 5.5 (solid line without symbol) volts and for (b) \( V = 1 \) at bias values 0.0 (circle) and 6.0 (square) volts for a system with 40 sites.
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