Exploiting B-site disorder for phase control in the manganites

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Abstract – Disorder on the active $d$ element site is usually very disruptive for conduction and long-range order in perovskite transition metal oxides. However, in the background of phase competition such “B-site” dopants also act to promote one ordered phase at the expense of another. This occurs either through valence change of the transition metal or via creation of “defects” in the parent magnetic state. We provide a framework for understanding the complex variety of phenomena observed in B-site-doped manganites and identify the key parameters that control the physics. Using a spatially resolved analysis of B ions in various manganite phases we explain the existing data and predict new situations where highly polarisable phase separated states can be created.

Correlated electron systems like the cuprates and manganites involve competition between various long-range ordered phases [1,2]. The interplay of this phase competition with weak disorder underlies phenomena like cluster coexistence, percolative transport, and colossal response. The nature of disorder seems to be crucial for these effects, as observed in the manganites [2], and a low concentration of impurities on the active $d$ element site is an effective trigger for phase separation [3–12] and the associated percolative effects.

The results of such substitution depend on the reference state and the chemical nature of the impurity. For manganites, with rare-earth (RE) and alkaline-earth (AE) combination $\text{RE}_1-\text{AE}_x\text{MnO}_3$, several intriguing results exist for Mn-site doping. i) Magnetic dopants like Cr [3–7], Co or Ni [8] (but not Fe) on the Mn site in a $x = 0.5$ charge ordered insulating (CO-I) manganite promote a percolative ferromagnetic metal (FM-M), while non magnetic dopants of the same valence do not. ii) The orbital ordered A-type antiferromagnet (AF) at $x = 0$ is destabilised in favour of a ferromagnetic state [9] by both magnetic and non-magnetic dopants. iii) In contrast to the cases above, where charge-orbital order is suppressed, doping Fe on a ferromagnetic metal [10,11] at $x \sim 0.4$ promotes a charge ordered insulating state! On spatial imaging most of these systems reveal phase separation (PS) and many of them also exhibit enormous magnetoresistance. It is vital to uncover the organising principle behind this diversity of effects, if we are to exploit B-site disorder as a tool for phase control.

There is unfortunately no microscopic model, let alone a theory, for randomly located B dopants in the manganites. In this paper we write down the first detailed model for B impurities in a manganite host, and study the effect of these dopants in a variety of manganite phases using a real space Monte Carlo technique.

Our principal results are the following: i) We discover that the following hierarchy of effects arise in all B doping cases: a) change of the effective valence on the Mn sites, b) percolation of the metallic phase through impurity free regions, and c) “reconstruction” of the background magnetism and charge order by magnetic dopants. ii) By exploring the prominent manganite states, and different B dopants, we are able to explain most of the outstanding experimental results. iii) We suggest a new experiment to test out an unexplored insulator-metal transition driven by B-site disorder. iv) We demonstrate how B impurity locations determine the percolation pattern and may allow atomic level control of current paths in a material.

The simplest classification of B-site dopants is in terms of their valence in the manganite host. Among the usual dopants Zn, Mg, and Co are divalent, i.e. in a $2^+$ state, Ni, Cr, Fe, Sc, and Al are trivalent, while Ru, Sn, and Ti are tetravalent. Some elements can exist in multiple valence states, e.g., Ni can also be +2, and Ru can be +5, but that will not affect our qualitative arguments. The valence, $\alpha$, of the dopant affects the effective carrier density on the Mn sites through the charge neutrality requirement on the
compound RE$_{3+}$$_{1-x}$AE$_x$$_{3+}$Mn$_{1+\eta}$+$\nu$B$_{\eta}$O$_{3-}$, where $\eta$ is the % of B-site doping, and we write the Mn valence as $3 + \nu$. This yields $\nu(y, \alpha, x) = (x + \eta(3 - \alpha))/(1 - \eta)$. The effective $e_g$ electron count on Mn is $n = 1 - \nu$, modified from $n_0 = 1 - x$ at $\eta = 0$. This change of effective carrier density can itself drive phase change as we will see later. Secondly, dopants with same valence can have different effects depending on their magnetic character. Non magnetic dopants only affect the Mn valence, while those with partially filled $d$ shells can have magnetic coupling to the neighbouring Mn moments. Experiments suggest that Cr has strong AF coupling [13] to the Mn ions, Ni couples ferromagnetically [13], while Fe, despite its magnetic $d^5$ configuration, couples rather weakly.

Based on the inputs above we construct the following model for manganites with B-site dopants: $H_{tot} = H_{ref} + H_{imp} + H_{coup}$, with

$$H_{ref} = \sum_{(ij),\sigma} t_{ij}^{\alpha\beta} c_{i\alpha\sigma}^\dagger c_{j\beta\sigma} - J_H \sum_i S_i \cdot \sigma_i + J \sum_{(ij)} S_i \cdot S_j$$

$$-\lambda \sum_i Q_i \cdot \tau_i + \frac{K}{2} \sum_i Q_i^2,$$

$$H_{imp} = V \sum_{n\alpha\sigma} d_{n\alpha\sigma}^\dagger d_{n\alpha\sigma},$$

$$H_{coup} = \sum_{(ij),\sigma} t_{ij}^{\alpha\beta} c_{i\alpha\sigma}^\dagger c_{j\beta\sigma} + J' \sum_{(ij)} s_n \cdot S_j + V_c \sum_{(ij)} q_n q_j.$$

The reference “manganite model” $H_{ref}$ is constructed to reproduce the correct sequence of phases in the “clean” limit. It involves the nearest neighbour hopping of $e_g$ electrons with amplitude $t_{ij}^{\alpha\beta}$ [14]. Hund’s coupling $J_H$, AF superexchange $J$ between Mn spins, and Jahn-Teller (JT) interaction, $\lambda$, between the electrons and the phonon modes $Q_i$. The stiffness of the JT modes is $K$. We will not consider RE-AE cation disorder in $H_{ref}$ since such “A-site” disorder masks the B-site effects. The sites $R_i$, $R_j$ and operator $c, c^\dagger$ refer to Mn locations, and $S_i$, etc., are Mn spins. The local physics of the B ions is contained in $H_{imp}$ where $R_i$ refers to the B locations and the operators $d, d^\dagger$ refer to the B ion $e_g$ states at an energy $V$ above the center of the Mn band. The sites $R_n$ are random, with only the constraint that two B dopants are not nearest neighbours, to minimise electrostatic repulsion. $H_{coup}$ involves i) $e_g$ hopping matrix elements $t_{i\beta\sigma}$, which we keep the same as between the Mn, ii) for magnetic B ions, a superexchange coupling $J'$ between the B moment $s_n$ and the neighbouring Mn moments, and iii) a nearest neighbour Coulomb repulsion $V_c$ between the B dopant and the neighbouring Mn. The total charge $q_n$ on the Mn ion is $4 - n_j$, where $n_j$ is the $e_g$ occupancy, and $q_0$ is the (fixed) B ion valence. Figure 1 is a schematic, showing the relevant levels on Mn and B, and the coupling between these atoms. Earlier attempts at modelling B dopants (or Mn vacancies) only employed a random onsite potential [15].

The overall carrier density is controlled through the chemical potential term: $-\mu(\sum_{i\alpha\sigma} c_{i\alpha\sigma}^\dagger c_{i\alpha\sigma} + \sum_{n\alpha\sigma} d_{n\alpha\sigma}^\dagger d_{n\alpha\sigma}$). We use the standard limit $J_H/t > 1$, and set $K = 1$. In studying magnetic field effects we will use a coupling $H_{mag} = -h \cdot (\sum_i \sigma_i + \sum_n s_n)$, where $h = \lambda$ is the applied field. We treat all spin and phonon degrees of freedom as classical [16], and measure all energies in units of the Mn-Mn hopping $t$ [14]. The spins, $S_i$, etc., are treated as unit vectors, and the magnitude of the spin is absorbed in the couplings $J$ and $J'$. The parameter space of the problem is very large. We have to contend with manganite states at different doping ($x$), different (inverse) bandwidth ($\lambda/t$), and AF strength ($J/t$). The impurities are characterised by their concentration ($\eta$), the $e_g$ level ($V/t$), and the magnetic coupling ($J'/t$). We use $\lambda/t = 1.6$ as a typical JT coupling, $J/t = 0.1$ (to capture the correct phase competition around $x \sim 0.5$), and $V_c = 0.1 t$ [17]. We will explore several doping levels $x = 0.25, x = 0.4$ and $x = 0.5$ and study the effect of a low density of B dopants.

We use a Monte Carlo (MC) technique based on the “travelling cluster approximation” (TCA) [18]. It allows ready access to system size $\sim 40 \times 40$ using a moving cluster of size $8 \times 8$, and handles disorder accurately. The method, and the associated transport calculation, has been extensively benchmarked by us and used successfully in several earlier studies [14,19].

The first principle that we wish to illustrate is phase separation driven by change in effective carrier density. Figure 2 shows the phases in $H_{ref}$ for varying “hole density” $x = 1 - n$. The $T = 0$ phases, in increasing order of $x$, are an orbital ordered insulator at $x = 0$ [19], a ferromagnetic charge ordered insulator (FM-CO-I) at $x = 0.25$, a FM-M window between $x = 0.30 - 0.42$, the CE-CO insulator at $x = 0.50$, and a magnetic phase (“A-2D”), with structure factor peaks at $q = \{0, \pi\}, \{\pi, 0\}$, between $x \sim 0.55 - 0.60$. Between these phases are the shaded windows of PS. If the carrier density is in one of these PS windows the system would
The percolation of the FM-M patches creates a “global” phase at short distance but loses coherence at larger scales. These can be exploited by choosing dopants of suitable valence. The tendency towards large-scale PS separations at the edge of PS columns indicates phases of interest to us are the ferromagnetic CO insulator at $x = 0.25$, the ferromagnetic metal for $x \sim 0.30–0.42$, and the CE-CO phase at $x = 0.50$. Shaded regions indicate phase separation.

Let us start with $x = 0.25$. In the clean limit the orbital ordered (OO) JT insulator at $x = 0$ [19] is separated from the FM-CO-I at $x = 0.25$ by a wide PS window. The FM-CO-I can be looked upon as the charge ordering of the FM-M at $x = 0$. To “metallise” the $x = 0.25$ phase we need impurities that decrease the effective electron density, pushing it towards the FM-M. From our expression for $n = 1 - \nu$ at low $\eta$ we obtain $n \approx (1 - x) - \eta(3 + x - \alpha) + O(\eta^2)$, so any impurity with valence $\leq 3$ will serve to lower $n$ and push the system into the $x \sim 0.25–0.30$ PS window. The system could then phase separate, with FM-CO-I and FM-M clusters whose pattern is controlled by the B ion locations.

Figure 3 top row shows a snapshot of the charge density field $n_x$ in the reference CO state at $\eta = 0$ (left), followed by the impurity locations (center), and $n_x$ (right) in the presence of the B ions. The impurities are non-magnetic ($J' = 0$), divalent, with $V = 5$ (which is roughly half the Mn bandwidth) and $\eta = 2\%$. Trivalent impurities yield similar results. The emergence of homogeneous regions of $n \sim 0.7$ ($x \sim 0.3$) coexisting with short range (SR) CO patches is clear. The FM-M live in the “impurity free” regions, while the SR-CO can coexist with the impurities at short distance but loses coherence at larger scales. The percolation of the FM-M patches creates a “global” metallic state. Figure 4(a) show the evolution of the resistivity $\rho(T)$ with $\eta$. Since both the reference CO phase and the emergent metal are FM we cannot get any magnetoresistance in this regime. It would be interesting to choose $y$ in the $(La_{1-y}Pr_y)_{0.75}Ca_{0.25}MnO_3$ family so that the material is at the $T = 0$ insulator-metal phase boundary, and explore the impact of 2+ or 3+ B dopants.

Let us shift to $x = 0.4$ which is next to the PS window separating the FM-M from the $x = 0.5$ CE-CO-I phase. The reference state is a homogeneous metal. We explore the impact of non-magnetic divalent dopants with $V = 5$ on this state, and push $n$ into the PS window. Such a situation had been explored experimentally early on by doping Mg [12] onto $Pr_{0.7}Ca_ySr_{0.3-y}MnO_3$. A metal-insulator transition in the ground state was seen with increasing Mg doping. Similar results were observed on doping Fe [10] (which is a 3+ dopant with weak magnetic coupling) into $La_{1-x}Ca_xMnO_3$ at $x = 0.37$.

The left panel in the middle row in fig. 3 shows the “flat” profile of $n_x$ in the undoped FM-M state. The central panel shows the dopant locations at $\eta = 6\%$ and the right panel shows the $n_x$ in the presence of dopants. The final $n_x$ shows local charge order arising out of “disordering” the FM-M! This has indeed been reported [11] recently. The FM order in the ground state is significantly suppressed with increasing $\eta$, and fig. 4(b) shows the low $T$ metal-insulator transition driven by B-site doping.
has two parallel and two antiparallel neighbours, using to the reference CE phase with strong AF coupling to the neighbouring Mn. In contrast to the electron density on the CE state at separated state, we can explore the effect of magnetic metal at "metallised" by a modest field. The disorder driven insulating state can be readily suppressed with 2+ dopants at \( \eta = 0.06 \), \( V = 5 \), \( J' = 0 \). Bottom row, \( x = 0.5 \) with 3+ dopants at \( \eta = 0.06 \), \( V = 5 \), \( J' = 0.2 \). The fields are (left to right) \( h = 0 \), 0.002, 0.005. The uniform (red) regions are FM while the striplike (green) regions are AF.

Figure 4(c) shows the dramatic suppression in resistivity of this PS system, at \( \eta = 6\% \), in response to a magnetic field. The disorder driven insulating state can be readily "metallised" by a modest field.

Just as depleting the electron density converts the ferromagnetic metal at \( x = 0.4 \) to a FM-M+AF-CO-I phase-separated state, we can explore the effect of increasing the electron density on the CE state at \( x = 0.5 \). Since in the undoped CE-CO-I at \( x = 0.5 \) (dotted line), "metallisation" at \( \eta = 6\% \) with 3+ dopants (\( V = 5 \), \( J' = 0.2 \)) and field response.

Figure 5: (Colour on-line) Nearest neighbour \( \mathbf{S}_i \cdot \mathbf{S}_j \) from MC snapshots in the presence of an applied field. Top row, \( x = 0.4 \) with 2+ dopants at \( \eta = 0.06 \), \( V = 5 \), \( J' = 0 \). Bottom row, \( x = 0.5 \) with 3+ dopants at \( \eta = 0.06 \), \( V = 5 \), \( J' = 0.2 \). The fields are (left to right) \( h = 0 \), 0.002, 0.005. The uniform (red) regions are FM while the striplike (green) regions are AF.

Fig. 4: (Colour on-line) Temperature and field dependence of resistivity \( \rho(T) \). (a) Metallisation of the \( x = 0.25 \) FM-CO-I with non magnetic dopants of valence 2+ and \( V = 5 \), the curves are for different \( \eta \). (b) Ferromagnetic metal to (CO) insulator transition at \( x = 0.4 \) with increasing concentration of non magnetic 2+ dopants with \( V = 5 \). (c) Field response at \( x = 0.4 \) and \( \eta = 6\% \): \( h_0 = 0 \), \( h_1 = 0.002 \), \( h_2 = 0.005 \). (d) Resistivity in the undoped CE-CO-I at \( x = 0.5 \) (dotted line), "metallisation" at \( \eta = 6\% \) with 3+ dopants (\( V = 5 \), \( J' = 0.2 \)) and field response.

We highlight the case where the dopant, e.g., Cr, has a strong AF coupling to the neighbouring Mn. In contrast to the reference CE phase with \( J = 0.1 \), where a spin has two parallel and two antiparallel neighbours, using B ions with \( J' = 0.2 \) forces all four nearest neighbours (NN) to be antiparallel to the B spin. Due to its low \( e_g \) occupancy the B site acts locally like the “corner site” of a CE chain, and the four \( NN \) spins are also antiparallel to the B spin. This “1+8” cluster \([20]\) has a large moment but at low \( \eta \) distant clusters are uncorrelated so the magnetic “reconstruction” by itself does not create a global FM state. The presence of small NN Coulomb repulsion \( V_c \) suppresses the \( e_g \) charge density on Mn sites which neighbour the B ions. To conserve the electron count, the charge that is pushed out creates small FM-M regions of locally high density, \( n \sim 0.6 \), dominated by the double exchange interaction. These FM-M droplets connect the otherwise disconnected “1+8” clusters. The resulting pattern is a complex mix of FM-M, FM-CO and AF regions. The ferromagnetism in the magnetic B ion doped case emerges from a combination of i) breakup of the CE pattern by the magnetic dopants, creating tiny FM clusters, and ii) their coupling via FM-M droplets created by charge pushed out due to the Coulomb interaction.

In addition to converting insulators to metals, and vice-versa, B-site doping effects are interesting because the final state is necessarily inhomogeneous, with possibly huge magnetoresistance (MR). For example if the PS is between FM-M and AF-CO-I the randomly located B ions fragment the PS state and the FM-M domains in such a situation are weakly linked to each other at zero field. The large "moments" of the FM-M domains, see fig. 5, can be aligned by a small field leading to enormous increase in conductance. This large polarisability and MR are key signatures of a PS state. The large changes in resistivity driven by an applied field are shown in fig. 4(c), (d). Figure 5 shows the spatial evolution of the magnetic state for two systems doped into PS windows. The dramatic transport response correlates with a growth in volume of the FM-M phase. The effect is stronger for the \( x = 0.4 \)
case, but is also visible for the more complex $x = 0.5$ case with 3+ magnetic dopants.

Since we have attempted to model a complex phenomenon, and had to make several approximations to make progress, let us list out some of the checks [20] we have performed to establish the robustness of our results. i) Any real material will have some degree of A-site disorder, while we have assumed the reference manganite to be “non-disordered”. We have checked that our qualitative results survive even if we include binary “A-site” disorder of magnitude up to 0.1 in our model. In the experiments also most B-site results are on the La-Pr-Ca family, where the cation disorder $\sigma_A$ is small. ii) Impurity valence fluctuation: in reality it is not the valence of the impurity, but its electronic level, that is fixed. We have assumed B dopants to be of integer valence, but that is true only if the B-site potential $V$ is large, and the conduction electron density $n_\sigma$ on the B site is $\ll 1$. We tried out “softer” B potentials, down to $V = 1$, and that leads to $n_\sigma \sim 0.25$ on the B site. For an assumed valence of 3, say, this implies a modest change to 2.75, not significantly affecting our argument. iii) Weak localisation (WL) effects: we have described “metals” in 2D in the presence of B-site disorder. On large sizes such a system will show WL effects, and a genuine metallic phase will occur only in three dimensions.

Let us conclude. We have illustrated how one might engineer phase separation in low cation disorder manganites by doping B ions of suitable valence and magnetic character. The most promising regimes are close to commensurate filling, $x = 0, 0.25, 0.50$, etc., which are typically ordered insulating states close to a metal. Most of the effects can be understood in terms of the valence change of Mn, but the B-Mn magnetic coupling is also crucial, particularly for dopants on the CE phase. The percolative state that emerges is typically highly polarisable and has a large low field magnetoresistance. The current paths in the phase separated regime are dictated by avoidance of B dopant locations. Since presently available techniques allow atomic scale manipulation [21] of dopant locations, B-site doping opens up the prospect of controlling the nanoscale current paths in a manganite.

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