We study the Holstein-Double Exchange (H-DE) model in three dimensions in the presence of substitutional disorder. Using a new Monte Carlo technique we establish the phase diagram of the clean model and then focus on the effect of varying electron-phonon coupling and disorder at fixed electron density. We demonstrate how extrinsic disorder controls the interplay of lattice polaron effects and spin fluctuations and leads to widely varying regimes in transport. Our results on the disorder dependence of the ferromagnetic $T_C$ and metal-insulator transitions bear direct comparison to data on the 'optimally doped', $x = 0.3-0.4$, manganites. We highlight disorder induced polaron formation as a key effect in these materials, organise a wide variety of data into a simple 'global phase diagram', and make several experimental predictions.

The Holstein-Double Exchange (H-DE) model provides a minimal description of strongly coupled charge, spin, and lattice degrees of freedom, typical of the perovskite manganites $A_{1-x}A'_xMnO_3$ [1], and captures the interplay of polaronic tendency and spin disorder that occurs in these materials. Although a detailed solution has not been available in three dimensions (3D), the H-DE model is believed to describe the trend towards stronger localisation and weakened ferromagnetism observed with reducing $A$ site ionic radius $(r_A)$ in the manganites.

The role of quenched disorder in these systems, arising, for example, from the cation size mismatch $(\sigma_A)$ is only partially understood. Most of the disorder related theory [2,3] has focused on effects near a first order phase boundary, i.e., the 'bicritical' region [4], motivated by the observation of cluster coexistence in some materials [5]. Phase competition and bicriticality, however, occurs only in a limited part of the phase diagram. Studies with controlled variation in $\sigma_A$ indicate that quenched disorder has a dramatic effect on the ferromagnetic $T_C$ and resistivity [6–9], and optical response [10] even far from bicriticality. The origin of these effects remain unexplained.

Our primary contribution in this paper is to demonstrate how this unusual sensitivity arises due to a positive feedback of quenched disorder on the polaron formation tendency, and its interplay with spin fluctuations. Through a detailed real space solution of the disordered H-DE model on large 3D lattices we (i) demonstrate disorder controlled metal-insulator transitions (MIT), in the ground state and at finite temperature, that are absent in the 'clean' system, (ii) explain the rapid suppression of $T_C$ with increasing disorder, (iii) compare our results in detail with data on the manganites at $x = 0.3 - 0.4$, constructing a 'global phase diagram', and (iv) make testable predictions about the $\sigma_A$ dependence of optical spectral weight, lattice distortion, and tunneling density of states in the optimally doped manganites.

We study the adiabatic disordered $(d)$ H-DE model:

$$H = -t \sum_{\langle ij \rangle,\sigma} (c_{i\sigma}^\dagger c_{j\sigma} + h.c) + \sum_i (\epsilon_i - \mu) n_i$$

$$- \lambda \sum_i n_i x_i - J_H \sum_i S_i \sigma_i + K 2 \sum_i x_i^2 \tag{1}$$

The $t$ are nearest neighbour hopping on a cubic lattice, $\epsilon_i$ is the quenched binary disorder, with $\bar{\epsilon}_i = 0$ and value $\pm \Delta$, $\lambda$ is the electron-phonon (EP) interaction, coupling electron density to the local distortion $x_i$, $K$ is the stiffness, $\mu$ is the chemical potential, and $J_H$ is the Hund’s coupling. The parameters in the problem are $\lambda/t$, $\Delta/t$, electron density $n$, and temperature $T$. We treat the phonons and spins as classical (the adiabatic limit), assume $J_H/t \rightarrow \infty$, and set $|S| = 1$. We also set $K = 1$ and measure energy, frequency, $T$, etc, in units of $t$.

The clean H-DE model was proposed early on [11] as a minimal model for the manganites, and a mean field study [11] suggested strong decrease of $T_C$ and increasing localisation with increase in EP coupling. More recently the H-DE model with ‘cooperative phonons’ has been analysed [12] in 3D, yielding a thermally driven MIT at low $n$ but not near optimality. The effect of disorder has been considered on EP-DE models but with focus mainly on phase coexistence [13] and bicriticality [3]. These studies yield insight on how disorder leads to cluster formation near a first order phase boundary [2,13], or its impact on a commensurate charge ordered phase [3], but do not have much bearing on the effects observed, for example, at $x = 0.3$, where there is no obvious phase competition. In our understanding, the effects of disorder in the manganites, away from bicriticality, await an explanation. In this paper we overcome a key technical limitation of the earlier studies and provide new results on transport combining the effects of strong EP coupling, thermal spin fluctuations, and quenched disorder, on lattices upto $12^3$ in size.
Since we work with $J_H/t \to \infty$, the electron spin is 'slaved' to the core spin orientation leading to an effectively spinless, hopping disordered model: $H = -\sum_{\langle ij \rangle}(t_{ij}\gamma_i^\dagger \gamma_j + h.c.) + \sum_i(\epsilon_i - \mu - \lambda_2 \mu_i + (K/2) \sum_i x_i^2).$ The hopping amplitude depends on the spin orientations via $t_{ij}/t = \cos \frac{\theta_i \theta_j}{2} + \sin \frac{\theta_i \theta_j}{2} e^{i(\phi_i - \phi_j)}$, where $\theta_i$ and $\phi_i$ are respectively the polar and azimuthal angles of the spin $S_i$. We use our recently developed “travelling cluster approximation” (TCA) [14] for solving the annealing problem on large lattices, by computing the energy cost of a MC update by constructing a ‘cluster Hamiltonian’, of $N_c = L^2$ sites, around the update site. We will use $N_c = 4^3$ and, as documented earlier [14], this is sufficient for equilibrating the spin and lattice variables. The electronic properties, $e.g.$ resistivity, density of states, and optical response, are obtained by diagonalising the full electron Hamiltonian on the equilibrium $\{x_i, S_i\}$ configurations, averaging thermally and finally over $8 - 10$ realisations of disorder $\{c_i\}$. The transport results are obtained using a formula described in detail previously [15]. We will use system size $N = 8^3$, but have checked our results on $N$ up to $12^3$. Our resistivity results are in units of $\hbar a_0/(\pi e^2)$, where $a_0$ is the lattice spacing. As a rough estimate, $\rho = 1$ in our units would correspond to $\sim 30\mu\Omega cm$.

Let us summarise the phases of the clean H-DE model before addressing the effects of disorder. (a). Structurally, a phase can be either charge ordered (CO), or charge disordered, i.e. a polaron liquid (PL), or homogeneous, i.e. a Fermi liquid (FL). (b). Magnetically it can be either ferromagnetic (F) or paramagnetic (P), and (c). in terms of electrical response it can be a ‘metal’, with $d\rho/dT > 0$, or an insulator, with $d\rho/dT < 0$.

Combining these features, the ground state can be either a ferromagnetic Fermi liquid (F-FL), or F-PL, or F-CO, depending on $\lambda$ and $n$, while finite $T$ allows P-FL, P-PL, and P-CO phases as well. Notice that while the threshold for single polaron trapping, i.e. $n = 0$, in 3D, is $E_p = \lambda^2/2K \approx 5.4t$, i.e. $\lambda_c(n = 0) \sim 3.3$ [16], at finite density the threshold for a collective PL is lower, e.g. $\lambda_c(n = 0.3) \sim 2.6$. These thresholds are further affected by structural and spin disorder.

For $\lambda \lesssim 2.0$, Fig.1.(a), the phases are (i) F-FL at low $T$ away from $n = 0.5$, (ii) P-FL at high $T$, and (iii) a F-CO phase at $n = 0.5$, separated from the F-FL, at low $T$, by a window of phase separation. For $\lambda = 2.0$ the $T_c$ at $n = 0.5$ is suppressed with respect to the pure DE problem, but is still larger than the CO scale $T_{CO}$. At stronger coupling, of which $\lambda = 2.5$ is typical, Fig.1.(b), much of the phase diagram is taken up by insulating phases. As $\lambda$ is increased beyond $2.5$ the metallic window at low $n$ shrinks quickly and all the phases turn insulating. The ground state, however, continues to be ferromagnetic since we have not included any competing antiferromagnetism (AF). From the resistivity we confirmed that the clean model does not show a thermally driven MIT except for $n \lesssim 0.1$, as noted earlier [12].

Even weak structural disorder dramatically affects the electronic state at strong coupling, leading to metal-insulator transitions which are not present in the clean limit. In the rest of the paper we focus on $n = 0.3$, typical of the high density metallic regime, and study the effect of varying disorder and EP coupling. Since the $r_A$ controlled bandwidth variation in real materials is relatively small we use only a ±10% variation in $\lambda/t$ about $\lambda = 2.0$ and study the weak disorder regime, $\Delta = 0 - 1.0$.

Disorder creates density inhomogeneities in the FL which are amplified by the EP coupling, leading to a rapid rise in residual resistivity, and also opens up a window for the crucial F-FL to P-PL crossover. Fig.2.(a) – (c) show $\rho(T)$ with varying $\lambda$ and $\Delta$ illustrating the three broad transport regimes. At $\lambda = 1.8$, which is towards the lower end of our coupling range, moderate $\Delta$ increases $\rho(0)$ but the overall $\rho(T)$ has the same nature as in the simple DE model [15] and neither EP coupling nor disorder seems to have any qualitative effect. This is true of $\Delta \lesssim 0.6$ and, we suggest, is akin to what one observes in La$_{0.7}$Sr$_{0.3}$MnO$_3$. At $\Delta = 1.0$, where $\rho(0) \gtrsim 100 \sim \rho_{MIT}$ we see hints of a weak downturn in $\rho(T)$ at high $T$. Overall, $\rho(T)$ at $\lambda = 1.8$ is mainly characterised by a F-FL to P-FL crossover. The density of states (DOS), Fig.2.(d), remains featureless at all $T$ and the lattice distortions, Fig.2.(g), are weak with no significant variation across $T_c$. This “DE like” character in $\rho(T)$ survives upto $\lambda = 2.2$ in the clean limit, see Fig.2.(b) – (c). However, for $\lambda \gtrsim 2.0$ addition of weak disorder (i) sharply increases $\rho(0)$ and creates a pseudogap in the DOS, (ii) maintains a regime with $d\rho/dT > 0$ for $T \lesssim T_c$, but (iii) crosses over to an ‘insulating’ PL regime with $d\rho/dT < 0$ for $T \gtrsim T_c$. As Fig.2.(h) – (i) indicate, these changes are associated with large lattice distortions and their variation across $T_c$. For $\Delta \sim 1.0$ the ground state itself turns insulating for $\lambda \gtrsim 2.0$. We provide a compact picture of the various MIT in Fig.3, discussed later.

To gain insight on the F-FL to P-PL crossover we also calculated the ‘effective carrier number’, $n_{eff}(\omega, T) = \int ...$
The combination of $n_{eff}$, Fig.4.(a), the spatial patterns (not shown here) and the DOS, Fig.2.(e), suggest the following scenario at strong coupling and moderate disorder: (i) there is polaronic localisation of a fraction of carriers (states at $\omega < \mu$) at $T = 0$, lowering $n_{eff}$, but the states at the chemical potential remain extended, (ii) the polaronic states are further localised with increasing $T$ due to DE spin disorder, while the delocalised electrons near $\mu$ experience scattering from the spin fluctuations, leading to $dp/dT > 0$, (iii) by the time $T \sim T_c$ the net disorder arising from $\epsilon_i$, the lattice distortions, and spin disorder is so large that the system is effectively at a localisation transition on the complex $\{\epsilon_i, x_i, S_i\}$ background: the states at $\mu$ also get localised, (iv) the $T > T_c$ phase has a modest ‘mobility gap’ (at moderate $\lambda, \Delta$) so there is no simple activated behaviour. The pseudogap in the DOS, Fig.2.(e), deepens initially and gradually fills up for $T > T_c$.

Let us consider the relevance of these results to manganites at $x \approx 0.3 - 0.4$, away from commensurate filling and charge ordering instabilities. The mean hopping amplitude between Mn ions, via the oxygen orbitals, is controlled by $r_A$ through the Mn-O-Mn bond angle. The mismatch between different A site cations is quantified through the variance, $\sigma_A$, of the ionic size, and acts as disorder on the electronic system. For a material of composition $A_{1-x}A'_x\text{MnO}_3$, say, $r_A$ and $\sigma_A$ depends on the radius of the A, A’ ions as well as $x$. To disentangle the effect of varying carrier density ($x$) from that of varying ‘coupling constants’, there have been systematic studies of the manganites at fixed $x$, with controlled variation in $r_A$, related to our $\lambda/t$ ratio, and $\sigma_A$, related to our $\Delta/t$.

Focusing initially on $x = 0.3$, the data from a wide variety of sources can be organised in terms of the transport response, $\rho(T)$, and magnetic $T_c$. We choose a simple characterisation where we construct the MIT phase diagram in terms of the low $T$ to high $T$ crossover observed in $\rho(T)$. For example, La$_{0.7}$Sr$_{0.3}$MnO$_3$ (LaSr), is a metal at both low $T$ and $T > T_c$, so it shows M-M crossover. In that spirit LaCa shows M-I crossover, while PrCa is I-I. These simple ‘binary’ cationic systems are well studied, have relatively small $\sigma_A$, and the transport response can be argued to arise mainly from variations in $r_A$. However, more complex cationic combinations have been used, notably by Attfield and coworkers [6,7], to vary $\sigma_A$ at fixed $r_A$ and $x$. Our organisation of the experimental data, Fig.3.(a) uses three such families, at $r_A = 1.26, 1.23$ and $1.20\AA$, as well as data from other sources [8,9] to highlight the combined impact of $r_A$ and $\sigma_A$ variation on transport character. We have used the 9 fold coordinated values of ionic radius from Shannon [17] in computing $r_A$ and $\sigma_A$ for about 30 compositions.

Four features are notable in the experimental MIT and $T_c$ data, Fig.3.(a) – (b): at $x = 0.3$: (i) the ‘clean’ limit allows only a narrow window, $r_A \sim 1.18 - 1.20\AA$, for M-I
discovered that it leads to a sharper drop in 

$$d$$

present in the manganites. We repeated the interactions and possibly a sharper MIT across 

$$\sigma$$

or $$\Delta$$. The qualitative results of this paper, however, remain unaffected.

Finally, if our argument about disorder induced polaron formation is correct then the following features should be experimentally observed with increasing $$\sigma_A$$: (a) systematic suppression of low frequency optical spectral weight, $$n_{eff}$$ at $$\omega \lesssim 0.1$$eV, at low $$T$$, (b) increase in the incoherent lattice distortion (oxygen Debye-Waller factor), progressively larger at smaller $$r_A$$, and (c) appearance of an increasingly deeper pseudogap in the tunneling DOS at low $$T$$. Fig.2 and Fig.4 quantify these effects. Experiments in the recent past do claim disorder induced polaron formation in La$_{0.54}$Ba$_{0.46}$MnO$_3$ [19] but more direct confirmation for $$x \sim 0.3$$ is desirable.

To conclude, we have studied a model with Holstein phonons, double exchange and quenched disorder, and illustrated the disorder enhancement of polaronic tendency and its interplay with spin fluctuations. Our results compare well with the disorder dependence of transport and $$T_c$$ observed in the ‘optimally doped’ manganites. We make three testable predictions regarding the behaviour of optical spectral weight, oxygen Debye-Waller factor and tunneling density of states for controlled variations of $$\sigma_A$$ in these materials.

We thank HRI for use of the Bawulf cluster and Saurabh Madaan for analysis of the experimental data.

* Present address: Institute for Physics, Theoretical Physics III, Electronic Correlations and Magnetism, University of Augsburg, 86135 Augsburg, Germany.

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FIG. 4. (a) Computed optical spectral weight, $$n_{eff}$$, at cut-off $$\omega = 0.5t$$ and $$T = 0.01$$ for varying $$\Delta$$ and $$\lambda$$. Inset: data from Ref[10] on LaSr, etc, at $$x = 0.4$$, $$\omega = 0.1$$eV, plotted versus $$\sigma_A$$. The $$r_A$$ values, LaSr downwards, are 1.256 Å, 1.232 Å, 1.22 Å and 1.202 Å. (b) Our results on disorder induced lattice distortion at $$T = 0.01$$. Inset shows the $$\sigma_A$$ increase in the mean square oxygen displacement at low $$T$$ in a sequence of $$x = 0.3$$ and $$r_A = 1.23$$Å manganites, from Ref[6].

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(c) The oxygen atoms are corner shared among MnO$_6$ octahedra so the phonon variables are cooperative. We have not explored this aspect but believe the main effects would be lower critical EP coupling for localisation [12] and possibly a sharper MIT across $$T_c$$. (iii) There are AF interactions present in the manganites. We repeated the d-H-DE calculation in the presence of AF coupling and discovered that it leads to a sharper drop in $$T_c$$, shifts the MIT phase boundaries somewhat, and leads to an AF-I phase at large $$\lambda$$ or $$\Delta$$. The qualitative results of this