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Local Spectroscopies Reveal Percolative Metal in Disordered Mott Insulators

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We elucidate the mechanism by which a Mott insulator transforms into a non-Fermi liquid metal upon increasing disorder at half filling. By correlating maps of the local density of states, the local magnetization and the local bond conductivity, we find a collapse of the Mott gap toward a V-shape pseudogapped density of states that occurs concomitantly with the decrease of magnetism around the highly disordered sites but an increase of bond conductivity. These metallic regions percolate to form an emergent non-Fermi liquid phase with a conductivity that increases with temperature. Bond conductivity measured via local microwave impedance combined with charge and spin local spectroscopies are ideal tools to corroborate our predictions.

Introduction.—The metal-to-insulator transition (MIT) driven by increasing disorder and the effect of Coulomb interactions on this transition has been a problem of fundamental interest. It is well known that disorder can create a transition from a metallic to an insulating state in both 2D and 3D due to localization effects. In the absence of interactions, all states are localized in one and two dimensions for arbitrarily small potential disorder, while in three-dimensions the MIT occurs at a finite critical disorder strength [1, 2]. In the presence of Coulomb interactions, perturbative calculations show an enhancement of localization in all dimensions [3]. However, the idea that disorder can create an insulator-to-metal transition (IMT) is relatively new. The first hint of an IMT in two dimensions due to long-range Coulomb interactions came from the renormalization group (RG) analysis by Finkel’stein [4, 5, 6] which showed that the critical indices for the correlation length and time scales become frequency-dependent and the RG flows take the system to a strong coupling fixed point. This was followed by an RG analysis of a two-parameter theory for long-range Coulomb interactions and disorder in the limit of large number of valleys that found a quantum critical point for the IMT in two dimensions. This theory was successful in explaining experimental data on thermodynamics and transport in high-mobility silicon metal-oxide-semiconductor field-effect transistors (Si-MOSFETS) [7].

In the opposite limit of strong on-site repulsion for commensurate filling, we have several examples of Mott insulators [8] in narrow band systems in which electrons are localized due to strong repulsion with an energy gap to excitations. The discovery of Mott insulators that can be driven into metallic or superconducting states upon doping has opened the field of competing charge-ordered, spin-ordered, nematic, pseudogap, superconducting and strange metallic phases. To understand the emergent behavior, it is important to separate out the effects of increasing disorder from adding or removing carriers. In this regard, gate tuning is a useful knob that tunes only the chemical potential without necessarily adding disorder, as distinct from chemical doping.

Previous experiments have observed a power-law suppression of the local density of states upon doping the Mott insulator Sr3Ir2O7 with Ru substitution for Ir [9], which indicates that new states are added at the chemical potential. Ru has been experimentally shown to be an isovalent substitution in Sr3Ir2O7, so the IMT in this system may be a good example of a disorder-induced transition. Transport measurements help determine whether the states are localized or extended, and indeed corroborate a metallic state with a finite resistivity extrapolated to $T = 0$ in the Ru substituted compounds. Theoretical methods ranging from inhomogeneous mean field theory [10], DMFT-based approaches [11–17], quantum Monte Carlo [18–20], and exact diagonalization studies [21] of the Anderson-Hubbard model support the presence of insulator-metal transition (IMT). However, the mechanism behind this transition and the nature of the emergent metallic phase remains unclear. Current research aims to develop a local picture that elucidates the nature of this pseudogap transition that is the subject of intense high resolution experiments in correlated materials [22].

In this Letter, we investigate the tension between two localizing tendencies: Mott repulsion and Anderson localization in two dimensions at half filling in the Anderson-Hubbard model, and their roles in driving quantum phase transitions. We investigate the correlations among the local maps of the magnetization, density of states, and the local bond conductivity, for a given realization and strength of disorder. We depict our results in a schematic phase diagram in the interaction-disorder plane in Fig. 1(a) that is based on our inhomogeneous mean field results for the Anderson-Hubbard model. It shows that while disorder and interactions independently enhance localization and promote an insulating state, acting together results in a novel metallic phase sandwiched between the Mott insulator at low disorder and a correlated Anderson insulator at high disorder.
Specifically, our two key new results are:

1. Our earlier study [10] showed that disorder adds spectral weight within the Mott gap resulting in a V-shaped density of states [also shown in Fig. 1(d)]. Here we calculate the dc conductivity and show that, remarkably, the conductivity increases with increasing fraction of disordered sites and also increases with temperature for a fixed disorder fraction [Fig. 1(g)]. Interestingly, this emergent metallic phase shows a non-Drude response in the optical conductivity [Fig. 1(f)].

2. Local dc conductivity profiles show the formation of conducting bonds in regions surrounding disorder sites, and the emergence of percolating metallic networks.

Model.—The Hamiltonian for the Anderson-Hubbard model is given by

\[ H = -t \sum_{(i,j),\sigma} (c_{i\sigma}^\dagger c_{j\sigma} + \text{H.c.}) + U \sum_i n_{i\uparrow} n_{i\downarrow} + \sum_{i,\sigma} (V_i - \mu) n_{i\sigma}, \]

where \( c_{i\sigma}^\dagger \) \( c_{i\sigma} \) is the electron creation (annihilation) operator at site \( i \) with spin \( \sigma \), and \( n_{i\sigma} \equiv c_{i\sigma}^\dagger c_{i\sigma} \). \( t \) represents the hopping amplitude between nearest neighbor sites, and \( U \) is the onsite electron-electron repulsion. \( V_i \) the onsite disorder potential, treated as binary-alloy disorder: \( p \) fraction of randomly chosen sites have \( V_i = V \), and \( 1 - p \) fraction of sites have \( V_i = 0 \). The chemical potential \( \mu \) is adjusted to achieve global half filling. The Hubbard interaction term \( U \) is treated at the Hartree-Fock level in terms of the site-dependent spin and charge density fields. This numerical method has the advantage that it treats the disorder potential exactly, and thus captures the localization physics due to the inhomogeneous potential profile accurately [23].

Global properties.—With increasing disorder, the DOS \( N(\omega) \) shows an evolution from a Mott gapped insulator to a gapless phase leading to a V-shaped pseudogap at the chemical potential [Fig. 1(d,e)] [9, 10]. To get some idea of whether these in-gap states are localized or extended, we plot the inverse participation ratio (IPR) \( \equiv \frac{\langle \sum_r |\psi_r|^2 \rangle}{\langle \sum_r |\psi_r|^4 \rangle} \), where \( \psi_a \) is the real-space wavefunction associated with eigenenergy \( \alpha \) and \( \xi_{\text{loc}} \) is its associated localization length (zero IPR value corresponds to infinitely delocalized state). In-gap states at low disorder fraction are bound states that are localized, as shown by the large value of their IPR shown in Fig. 1(b). As disorder regions grow and extend across the system, energy eigenstates become increasingly delocalized, depicting a transition from a Mott insulator to a metallic state. The IPR and derivative results that follow agree with previously analyzed metallic phases and are robust in the infinite system size limit as seen by finite size scaling pro-
FIG. 2. (a, b, c) Local charge density, conductivity, and antiferromagnetic (AFM) order parameter \(\langle m_i^\dagger \rangle \equiv (-1)^{\mathbf{x}_i\cdot \mathbf{y}_i}(S_i^z)\) as a function of disorder fraction. Data averaged over 40 random disorder realizations at \(V = 3.5t, U = 4.0t, T = 0.01t\) for 40 \times 40 lattice. (b) Local dc conductivity distribution normalized by the total number of sites \(N\), extrapolated from the low frequency conductivity. (d) Site disorder with Gaussian broadening and its correlation with local staggered magnetization; shading shows the standard deviation.

vided in Supplemental Material [Fig. S3(a)].

To understand this emergent phase with a finite density of states at the chemical potential, we evaluate the optical conductivity \(\sigma(\omega)\), shown in Fig. 1(f). Starting with a finite gap at low disorder, we observe the gap closing with increasing disorder, consistent with previous theoretical studies [24, 25]. As the system develops an increasing low-frequency conductivity, the behavior of \(\sigma(\omega)\) is non-Drude, with a peak in the conductivity at a nonzero frequency that moves toward lower frequency with increasing disorder. Rather remarkably, we observe a nonzero dc conductivity that grows with increasing disorder fraction as depicted in low frequency behavior of \(\sigma_\text{dc}(\omega)\) in the inset in Fig. 1(f). Beginning at 30%, the linear behavior of \(\sigma(\omega)\) allows us to extrapolate a nonzero dc conductivity \(\sigma_\text{dc}\) [Fig. 1(g)]. A finite \(\sigma_\text{dc}\) indicates the onset of a metallic phase in which the conductivity grows with increasing disorder. Similar enhancement of the conductivity was found for uniform box potential disorder \([-V,+V]\) [24–26], indicating that the emergence of the metallic phase is ubiquitous.

Local properties.—Insight into how metallicity arises as a result of the competition between disorder and interactions is captured by the distribution of local quantities: antiferromagnetic (AFM) order parameter \(\langle m_i^\dagger \rangle \equiv (-1)^{\mathbf{x}_i\cdot \mathbf{y}_i}(S_i^z)\), LDOS, and transport characteristics \(\sigma_{ij}^{\mu\nu}\), as we discuss below.

(a) Local magnetization.—For positive potential \(V\), it becomes energetically unfavorable to occupy the disorder sites, leading to a reduction in the local moment and charge density on disorder sites. The bimodal charge density distribution shown in Fig. 2(a) depicts disorder sites with relatively fixed mean occupation while non-disordered sites slowly transition away from unit filling, initially only impacting nearest neighbor sites. The spin ordering distribution echoes this nearest neighbor to disorder behavior. In Fig. 2(c), the distribution of AFM order is sharply peaked close to the maximum value at low disorder fractions, and becomes broader and shifts toward zero as the fraction increases, indicating a transition from a uniform AFM phase toward a nonuniform paramagnetic phase. Fig. 2(c) shows a reduction of the local moment on neighboring sites as the occupation increases beyond unit filling: Sites are screened from the effects of disorder at low disorder fraction, seen by the disparate peak and slightly perturbed sharp AFM profile. (b) Local AFM Order

LDOS vs. AFM

FIG. 3. (a-d) Inhomogeneity in LDOS and dependence on \(\langle m_i^\dagger \rangle^2\). Data collected from 40 random disorder realizations on a 40 \times 40 lattice, \((U,V) = (4.0t,3.5t)\), (e) Average local bond conductivity as a function of \(\langle m_i^\dagger \rangle\) taken from 8 disorder realizations for 15%, 25%, 35%, 45% on lattices of sizes 26 \times 26. V is not strong enough to eliminate ordering in the red curve \((U = 4.0t; V = 2.3t)\), so the local AFM remains highly ordered \(\geq 0.6\). See Fig. S4 for full conductivity distribution vs ordering.
for 15%. As the density of disorder in local regions grows, more charge occupies neighboring sites and neutralizes spin ordering. We correlate the magnetic ordering with local disorder density by smoothing the original disorder potential to create an effective disorder profile [Fig. 2(d)]. At low fraction \( (m_i^1)^2 \) decreases linearly with the degree of disorder, so initially \( V_i \) only has localized effect on interacting sites. As the disorder throughout the full lattice grows, the impact becomes increasingly non-local where sites away from disorder become paramagnetic.

(b) Local bond conductivity.—The distribution of local conductivities (current-current correlator between bond \((i,j)\) and all \((k,l)\) bonds) in Fig. 2(b) shows that mostly clean systems have a local dc conductivity distribution that is sharply peaked near zero, as expected for a Mott insulator. As disorder increases, the distribution broadens and the mean increases. Thus bonds become increasingly conducting and sites increasingly paramagnetic as the system becomes more disordered.

(c) Inhomogeneous LDOS.—The magnetization and conductivity hint at an inhomogeneous nature of the emergent disorder-driven metal. We ask how does this local non-uniformity promote charge transport in a Mott insulator. The first insight into how a metal emerges with increasing disorder comes from local spectroscopic analysis. Fig. 3 depicts the local density of states averaged over sites with different ranges of magnetic order. For low disorder fraction [Fig. 3(a)], regions with high AFM order exhibit a Mott-gap around the Fermi energy with almost no states below \( E_F \). For moderate disorder, [Fig. 3(b,c)] show that Mott physics is preserved in magnetically ordered regions, while increasingly disordered regions have enhanced spectral weight within the Hubbard gap with the formation of a V-shaped pseudogap. Such spectroscopic dependence on disorder has been observed experimentally in Mott insulating materials, where the pseudogap behavior is enhanced near impurity atoms [9].

(d) Correlation between local moment and conductivity.—Extending the previous discussion on the correlation of eigenstate delocalization and closing spectral gap with reduced magnetic ordering, to low frequency conductivity in Fig. 3(e), we show that transport and magnetic order are anti-correlated: The less magnetically ordered the region, the more conducting. Introducing few disorder sites decreases the magnetization and drives charge mobility on these sites, see Fig. 3(e). Maximal conductivity occurs at nonzero magnetic ordering in each curve, suggesting that weak correlations are crucial for promoting mobility.

In Fig. 4 we present a real space picture of two representative disorder realizations at 15% and 35% to show how disorder breaks down an initial Mott insulating system. Fig. 4(a-c) provide a spatial map of the random disorder potential and its effect on charge and spin.

To relate disorder and interactions with their effects on the local magnetization and bond conductivity in real-space, we overlay the magnetization profile for a representative disorder realization with the most conducting bonds (shown in red) in Fig. 4(d). The contrast with the non-interacting system [Fig. 4(e)] is remarkable: While disorder reduces conduction as expected in the non-interacting case, the behavior is quite the opposite for the interacting system, where one observes pockets of enhanced conductivity localized around a small fraction of disorder sites \( p = 15\% \), Fig. 4(d, left), expand to a percolating cluster at larger disorder fraction \( p = 35\% \), Fig. 4(d, right).

(e) Formation of percolating metal.—To exhibit the percolative nature of transport, we construct a network with each site as a node and \( \rho_{ij} = 1/\sigma_{ij} \) as the bond weights. The minimum series resistance \( \rho_{min} \) to connect the two ends of the system by the shortest path is obtained by a weighted path analysis of the conductivity graph [Fig. 4(f)]. The addition of 10% disorder leads to a factor of 2 decrease \( \rho_{min} \) while adding 30% disorder leads to a decrease by nearly two orders of magnitude. Above 30–35% disorder the cost remains constant as expected above the percolation threshold. The percolative nature of disorder we have characterized adds to current theoretical descriptions [27–30] and experimental characterization of disorder in materials [31].

In conclusion, our results capture the previously unexplored local properties of the intermediate pseudogapped metallic phase in the half-filled Anderson-Hubbard model. This work illuminates the interplay between disorder and interactions in the simplest fermionic model, and provides an avenue for understanding how non-Fermi liquid behavior arises at the microscopic level.

Our model, which does not include long-range Coulomb interaction, is particularly applicable to disordered systems with isovalent substitutions. The suppression of magnetism in regions near dopants, and subsequent development of in-gap states, which we observe in our results, are still expected in the presence of long-range interaction. Although our work does not address the effects of Coulomb gap on the emergent percolation network, we speculate that these metallic channels, identified in our local conductivity maps, play an important role in the low energy transport properties of the system.

The conductivity maps provide predictions for increasingly powerful spatially resolved spectroscopic techniques such as microwave impedance microscopy, 4-probe STM, and LC-AFM, which only recently been used to study local conductivity profiles on 100nm down to atomic resolution [32–34]. Recent breakthroughs in cold atom experiments now allow for resistivity and optical conductivity experiments, where the highly tunable nature of these experiments provide an ideal testing environment for the results we present here [35].

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