Statistical properties of eigenstates in three-dimensional mesoscopic systems with off-diagonal or diagonal disorder

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(Received 20 November 2000; published 13 June 2001)

The statistics of eigenfunction amplitudes are studied in mesoscopic disordered electron systems of finite size. The exact eigenspectrum and eigenstates are obtained by solving numerically Anderson Hamiltonian on a three-dimensional lattice for different strengths of disorder introduced either in the potential on-site energy (“diagonal”) or in the hopping integral (“off-diagonal”). The samples are characterized by the exact zero-temperature conductance computed using real-space Green function technique and related Landauer-type formula. The comparison of eigenstate statistics in two models of disorder shows sample-specific details which are not fully taken into account by the conductance, shape of the sample, and dimensionality. The wave function amplitude distributions for the states belonging to different transport regimes within the same model are contrasted with each other as well as with universal predictions of random matrix theory valid in the infinite conductance limit.

DOI: 10.1103/PhysRevB.64.014203  PACS number(s): 72.15.Rn, 73.23.--b, 05.40.--a

The disorder induced localization-delocalization (LD) transition in solids has been one of the most vigorously pursued problems in condensed matter physics since the seminal work of Anderson. In thermodynamic limit, strong enough disorder generates a zero-temperature critical point in $d>2$ dimensions as a result of quantum interference effects (in $d\leq 2$ even weakly disordered metal turns into an Anderson insulator for sufficiently large sample size). Thus, research in the “premesoscopic” era was mostly directed toward the viewpoint provided by the theory of critical phenomena. The advent of mesoscopic quantum physics has unearthed large fluctuations, induced by quantum coherence and randomness of disorder, of various physical quantities (e.g., conductance, local density of states, current relaxation times, etc.), even well into the delocalized phase. Thus, complete understanding of the LD transition requires to examine full distribution functions of relevant quantities. Especially interesting are the deviations of their asymptotic tails, a putative signature of incipient localization, from the (usually) Gaussian distributions expected in the limit of infinite dimensionless conductance $g = G/G_\text{Q}$ ($G_\text{Q} = 2e^2/h$ is the conductance quantum).

This paper presents the study of such type—numerical computation of the statistics of eigenfunction amplitudes in finite-size three-dimensional (3D) nanoscale (composed of ~1000 atoms) mesoscopic disordered conductors. The 3D conductors are often “neglected” in favor of the more popular and tractable playgrounds—two-dimensional systems (2D), where one can study states resembling 3D critical wave functions in a wide range of systems sizes and disorder strengths, or quasi-one-dimensional systems where analytical techniques can handle even nonperturbative phenomena (like the ones at small $g$). For example, in $d = 2 + \epsilon$ dimensions LD transition occurs at weak disorder (weak-coupling regime of the corresponding field-theoretical description), while in $d = 3$ small parameter needed for analytical treatment is lacking. In 3D systems critical eigenfunctions, exhibiting multifractal scaling, are expected only at the mobility edge $E_c$ which separates extended and localized states inside the energy band.

The essential physics of disordered conductors is captured by studying the quantum dynamics of a noninteracting (quasi)particle in a random potential. This problem is classically nonintegrable, thereby exhibiting quantum chaos. The concepts unifying disordered electron physics with the standard concepts of quantum chaos come from statistical approaches to the properties of energy spectrum and corresponding eigenstates, which cannot be computed analytically. While level statistics of disordered systems have been explored to a great extent, investigation of the statistics of eigenfunctions has been initiated only recently. These studies are not only revealing peculiar spectral properties of random Hamiltonians, but are relevant for the thorough understanding of various unusual features of quantum transport in diffusive metallic conductors (including the ones which are proximity coupled to a superconductor). The standard examples are long-time tails in the relaxation of current or log-normal tails (in $d = 2 + \epsilon$) of the distribution function of mesoscopic conductances. Distribution of eigenfunction amplitudes is found to be relevant for tunneling measurements on quantum dots probing the coupling to external leads, which depends sensitively on the local properties of wave functions. Experiments which are the closest to directly delving into the microscopic structure of quantum chaotic or disordered wave functions exploit the correspondence between the Schrödinger and Maxwell equations in microwave cavities.

The study of fluctuations and correlations of eigenfunction amplitudes in mesoscopic systems has led to a concept of the so-called “prelocalized” states. In 3D delocalized phase, this notion refers to the states which have sharp amplitude peaks on the top of an extended background. These kind of states appear even in the diffusive, $l \ll L < \xi$, metallic ($g \gg 1$) regime, but are anomalously rare in such samples (here $l$ is the elastic mean free path and $\xi$ is the localization length). In order to get “experimental”
feeling for the structure of states with unusually high amplitude spikes, an example is given on Fig. 1; this state is found in a special realization of quenched disorder (out of many randomly generated impurity configurations) inside the sample characterized by large average conductance. Thus, prelocalized states are putative precursors of LD transition and determine asymptotics of some of the distribution functions, studied in open or closed mesoscopic systems. In 2D, where all states are considered to be localized, prelocalized states have anomalously short localization radius when compared to "ordinary" localized states in low-dimensional systems. They underlie the multifractal scaling in weakly localized (g ≫ 1) 2D conductors of size L smaller than exponentially large ξ (which plays the role of a phase transition correlation length ξc = ξ in d = 2). In 3D, the correlation length ξc [defined as the size of a hypercube for which g(ξc) = O(1)] is always microscopic (ξc ≈ λF) in good metallic samples, and no multifractal scaling is expected. The appearance of small regions inside disordered sample gives the same classical path in the opposite directions. This simple quantum interference effect leads to a coherent back-scattering (i.e., suppression of conductivity) in a time-reversal invariant systems without spin-orbit interaction. However, in 3D systems WL correction is not "strong" enough to provide a full microscopic picture of complicated quantum interference processes which are responsible for LD transition, and facilitate the expansion of quantum intuition.

The paper presents the statistics of eigenfunction intensities |ψα(r)|² in isolated 3D mesoscopic conductors characterized by two different types of microscopic disorder. Numerical methods employed here make it possible to treat phenomena in both semiclassical (described by Bloch-Boltzmann formalism and perturbative quantum corrections) and fully quantum transport regime (dominated by nonperturbative effects, where semiclassical concepts, like l, lose their meaning), as well as in the crossover region. Since mesoscopic physics has provided efficient techniques for "measuring" exactly the transport properties of finite-size samples on the computer, this study connects the eigenstates statistics of a closed sample to its zero-temperature conductance. The statistical properties of eigenstates are described by the disorder-averaged distribution function

\[ f(t) = \frac{1}{\rho(E)N} \sum_{r,\alpha} \delta(t - |\psi_\alpha(r)|^2 V) \delta(E - E_\alpha) \]  

on N discrete points r inside a sample of volume V. Here \( \rho(E) = \langle \Sigma_\alpha \delta(E - E_\alpha) \rangle \) is the mean level density at energy E. Averaging over disorder is denoted by \( \langle \cdots \rangle \). Normalization of eigenstates gives \( \tilde{t} = \int dt t f(t) = 1 \). A finite-size disordered sample is modeled by a tight-binding Hamiltonian (TBH) with nearest neighbor hopping integral \( t_{nm} \)

\[ \hat{H} = \sum_m e_m |m\rangle\langle m| + \sum_{(m,n)} t_{mn} |m\rangle\langle n|, \]  

on a simple cubic lattice \( 16 \times 16 \times 16 \) of lattice constant a. Each site \( m \) contains a single s-orbital \( \langle r|m\rangle = \phi(r - m) \). Periodic boundary conditions are chosen in all directions. In a random hopping (RH) model the disorder is introduced by taking the off-diagonal matrix elements to be a uniformly distributed random variable \( 1 - 2W_{RH} < t_{mn} < 1 \), while diagonal elements are zero \( e_m = 0 \). The strength of the disorder is measured by \( W_{RH} \). The other system studied is described by a diagonally disordered (DD) Anderson model with potential energy \( e_m \) on site \( m \) drawn from the uniform distribution, \( -W_{DD}/2 < e_m < W_{DD}/2, \) and \( t_{mn} = 1 \) is the unit of energy. The Hamiltonian (2) is a real symmetric matrix because time-reversal symmetry is assumed. The results for \( f(t) \) in the samples described by the RH and DD Anderson models are shown on Figs. 2, 3 and Fig. 4, respectively. Although some of the samples are characterized by similar values of conductance, the eigenstates in the two models show different statistical behavior.

In what follows the meaning of these findings is explained in the context of statistical approach to quantum systems.

FIG. 1. An example of eigenstates in the band center of a delocalized phase. The average conductance at half filling is \( g(E_F = 0) \sim 17 \), entailing anomalous rarity of the "prelocalized" states. The disordered conductor is modeled by an Anderson model with diagonal disorder \( W_{pq} = 4 \) on a simple cubic lattice \( 12^3 \). For plotting of the eigenfunction values in 3D, the sites \( m \) are mapped onto the lattice site numbers \( \{1, \ldots , 1728\} \) in a lexicographic order, i.e., \( m = (m_x, m_y, m_z) \rightarrow 144(m_x - 1) + 12(m_y - 1) + m_z \).
with nonintegrable classical dynamics. In particular, the results are compared to the universal predictions of random matrix theory (RMT).

In the statistical approach of RMT, the Hamiltonian of a quantum chaotic system is replaced by a random matrix drawn from an ensemble defined by the symmetries under time reversal and spin rotation. This leads to the Wigner-Dyson (WD) statistics for eigenvalues and Porter-Thomas (PT) distribution for eigenfunction intensities. For the Gaussian orthogonal ensemble (GOE), relevant for studies of time-reversal-invariant Hamiltonians like Eq. (2), the PT distribution is given by

$$f_{PT}(t) = \frac{1}{\sqrt{2\pi t}} \exp(-t/2).$$

The function $f_{PT}(t)$ is plotted as a reference in Figs. 2, 3 and Fig. 4. The predictions of RMT are universal, depending only on the symmetry properties of the relevant ensemble. They apply to the statistics of real disordered systems in the limit $g \to \infty$ where $\Delta = 1/p(E)$ is the mean energy level spacing and $E_{Th} = hD/L^2$ is the Thouless energy, set by the classical diffusion across a sample of size $L$ with diffusion constant $D$.

The spectral correlations in RMT are determined by logarithmic level repulsion which is independent of true dynamics. All sample-specific details are absorbed into the mean level spacing. Also, the level correlations are independent of the eigenstate correlations. The RMT answer (3) for the distribution function (1) was derived by Porter and Thomas by assuming that the coordinate-representation eigenstate $\langle \Psi_P | r \rangle$ in a disordered (or classically chaotic system) is a Gaussian random variable. The behavior of $\Psi_P(r)$, even within the framework of RMT, is simple only in the systems with unbroken or completely broken time-reversal symmetry [the only difference between the two limiting ensembles is the functional form of $f(t)$]. Thus, RMT implies statistical equivalence of eigenstates which equally test the random potential all over the sample—typical wave function has more or less uniform amplitude up to inevitable Gaussian fluctuations.

Microscopic theory brings corrections to the RMT results in the case of samples with finite $g$. In the finite-size systems level statistics follow RMT predictions in the ergodic regime, i.e., on the energy separation scale smaller than $E_{Th}$. Nonuniversal corrections to the spectral statistics or eigenfunction statistics (which describe the long-range correlations of wave functions) depend on dimensionality, shape of the sample, and conductance $g$. These deviations from RMT predictions grow with increasing disorder (i.e., lowering of $g$). At the LD transition wave functions acquire multifractal properties, while the critical level statistics become scale-independent. For strong disorder or, at fixed disorder, for energies $|E|$ above the mobility edge $|E_c|$, wave functions are exponentially localized. The simple (and usually invoked) picture is that of a wave function which decays as

![Image](image1.png)

**FIG. 2.** Statistics of wave function intensities in the RH Anderson model, with $W_{RH}=1$, on a simple cubic lattice $16^3$. The distribution function $f(t)$ in Eq. (1) is computed for the states around the following energies: (a) $E=0$, (b) $E=1.5$, (c) $E=2.55$, and (d) $E=2.75$. Disorder averaging is performed over $N_{Ens}=40$ different samples. The Porter-Thomas distribution (3) is denoted by RMT. The part of the distributions inside the dashed box is enlarged on Fig. 3.

![Image](image2.png)

**FIG. 3.** Statistics of wave function intensities in the RH Anderson model, with $W_{RH}=1$, on a simple cubic lattice with $N=16^3$ sites. This figure plots the same distributions $f(t)$ as the ones plotted in Fig. 2, in the range defined by the dashed square in Fig. 2. The same labels apply to both figures.

![Image](image3.png)

**FIG. 4.** Statistics of wave function intensities in the DD Anderson model on a simple cubic lattice $16^3$. The distribution function $f(t)$, Eq. (1), is computed for the states around following energies. Upper panel, $W_{DD}=10$: (a) $E=0$, (b) $E=6.0$, (c) $E=7.45$, and (d) $E=7.85$. Lower panel, $W_{DD}=6$: (a) $E=0$, (b) $E=4.1$, (c) $E=6.56$, and (d) $E=6.7$. Disorder averaging is performed over $N_{Ens}=40$ different samples. The Porter-Thomas distribution (3) is denoted by RMT.
\( \Psi (r) = p(r) \exp(-r/\xi) \) from its maximum centered at some point inside the sample of size \( L > \xi \). Here \( p(r) \) is a random function and approximately spherical symmetry of decay is assumed. Since two states close in energy are localized at different points in space, there is almost no overlap between them. Therefore, the levels become uncorrelated and obey Poisson statistics. If \( p(r) = \epsilon \) is simplified to a normalization constant, the distribution function of intensities is given by

\[
f_{\xi}(t) = \frac{4 \pi}{V} \int_{0}^{L/2} \, dr \, r^{2} \delta(t - |\Psi(r)|^{2}) = \frac{\pi \xi^{3} \ln(2c^{2}V/t)}{4V},
\]

where a spherically symmetric sample of radius \( L/2 \) is assumed. In the localized phase \( \xi \ll L \), \( f_{\xi}(t) \) is expected to be insensitive to the assumed shape of the sample, and is determined by the ratio of these two relevant length scales.

The distribution function \( f(t) \) is equivalently given in terms of its moments \( b^{\text{PT}}_{q} = \int d\tau \int d\tau' f(t) \) (this statement is not rigorous since examples of different distribution functions which possess exactly the same sets of moments are encountered in various statistical problems\(^{39,40} \)). For GOE, the PT distribution \( (3) \) has moments \( b^{\text{PT}}_{q} = 2^{q} V^{-q+1} \Gamma(q + 1/2)/\Gamma(1/2) \). They are related to the moments \( I_{q}(q) = \int d\tau |\Psi_{\sigma}(\tau)|^{2q} \) of the wave function intensity. In the universal regime \( q \rightarrow \infty \) wave functions cover the whole volume with only short-range correlations (on the scale \( r \approx L \) persisting between \( \Psi_{\sigma}(\tau_{1}) \) and \( \Psi_{\sigma}(\tau_{2}) \). This means that integration in the definition of \( I_{q}(q) \) provides self-averaging, and \( I_{q}(q) \) does not fluctuate in the universal limit,\(^{37} \) i.e., \( I_{q}(q) = b^{\text{PT}}_{q} \). On the other hand, at finite \( g \) spatial correlations of wave function amplitudes at distances comparable to the system size are non-negligible. Therefore, \( I_{q}(q) \) fluctuates from state to state and from sample to sample.\(^{22,41} \) Although these long-range spatial correlations necessitate to study the full distribution function\(^{37,41} \) of \( I_{q}(q) \), for the subsequent analysis in this study it is enough to use an ensemble average of \( I_{q}(q) \), i.e., following Wegner\(^{42} \)

\[
I(q) = \Delta \sum_{\tau} \| \Psi_{\sigma}(\tau) \|^{2q} \delta(E - E_{\sigma}).
\]

The moment \( I_{q}(2) \) is usually called inverse participation ratio (IPR). It is a one-number measure of the degree of localization (i.e., it measures the portion of space where the amplitude of the wave function differs markedly from zero). This becomes obvious from the scaling properties of the average moments \( I(q) \) with respect to the system size

\[
I(q) \approx \begin{cases} 
L^{-d(q-1)} & \text{metal,} \\
L^{0} & \text{insulator,} \\
L^{-d^{*}(q-1)} & \text{critical.}
\end{cases}
\]

Here \( d^{*}(q) \leq d \) is the fractal dimension. Its dependence on \( q \) is the hallmark of multifractality of wave functions. The multifractal wave functions are delocalized, but extremely inhomogeneous occupying only an infinitesimal fraction of the sample volume in thermodynamic limit. The IPR is affected by mesoscopic fluctuations which scale in metallic samples as\(^{22} \) \( \delta I_{q}(2) \sim 1/g^{2} \times L^{4-2d} \). In the critical region \( g \sim 1 \) fluctuations\(^{22} \) are of the same order as the average value, which is then not enough to characterize the critical eigenstates (even though multifractal wave function extend throughout the whole sample, their IPR is not self-averaging\(^{41} \)).

I use \( I(2) \) as a rough guide in selecting eigenstates with different properties in the delocalized phase (Fig. 5). The second parameter used in the selection procedure is the conductance \( g(E_{F}) \) computed for a band filled up to the Fermi energy \( E_{F} \) equal to the state eigenenergy (see Fig. 6). The conductance as a function of band filling allows one to delineate delocalized from localized phase as well as to narrow down the critical region around LD transition point which is defined by \( g(E_{F}) \sim 1 \). Upon inspection of these two parameters, a small window is placed around chosen energy, and \( f(t) \) is computed for all eigenenergies whose eigenvalues fall inside the window. This provides more detailed information on the structure of eigenstates than is embedded in IPR.

The average IPR for both RH and DD Anderson model is shown in Fig. 5. The models with random hopping have been around for a long time,\(^{33,44} \) but have attracted considerable attention only recently, inasmuch as they show a disorder induced quantum critical point in less than three dimensions\(^{30,45-47} \) where delocalization occurs in the band center. Furthermore, several models used for strongly correlated electron systems share similar mathematical structure with the random hopping 2D models formulated in the framework of (noninteracting) disorder electron physics.\(^{48} \) Real solids which could be described by TBH (2) with off-diagonal disorder include doped semiconductors,\(^{44} \) such as P-doped Si, where hopping integrals \( t_{\text{nn}} \) vary exponentially with the distances between the orbitals they connect, while

FIG. 5. Inverse participation ratio \( I(2) \), averaged over both 40 different conductors and small energy bins, of eigenstates in the RH and DD Anderson models on a simple cubic lattice with \( N = 16^{3} \) sites. Top: diagonal disorder with (a) \( W_{\text{DD}} = 6 \) and (b) \( W_{\text{DD}} = 10 \). Bottom: off-diagonal disorder with (a) \( W_{\text{RH}} = 0.25 \), (b) \( W_{\text{RH}} = 0.375 \), and (c) \( W_{\text{RH}} = 1.0 \).
diagonal on-site energies \( \epsilon_{nm} \) are nearly constant. The behavior of low-dimensional RH Anderson model goes against the standard mantra of the scaling theory of localization for non-interacting systems\(^5\) that all electron states are localized in \( d \leq 2 \) for arbitrarily weak disorder. The possibility of delocalization transition in one dimension\(^49\) goes back to the work of Dyson\(^50\) on glasses. Also, the scaling theory for quantum wires with off-diagonal disorder requires two parameters\(^51\) which depend on the microscopic model, thus breaking the celebrated universality in disordered electron problems. In 3D case explored here, the states in the band center are less extended than other localized states inside the band (Fig. 5). However, the off-diagonal disorder is not strong enough\(^52\) to localize all states in the band, in contrast to the usual case of diagonal disorder where whole band becomes localized\(^53\) for \( W_{\text{DD}} \geq 16.5 \).

The mobility edge for the strongest RH disorder \( W_{\text{RH}} = 1 \), as well as for DD models, is found by looking at an exact zero-temperature static conductance. This quantity (which is a Fermi surface property) is computed from the Landauer-type formula\(^54\)

\[
g(E_F) = \text{Tr}[t(E_F)t^\dagger(E_F)],
\]

where transmission matrix \( t(E_F) \) is expressed in terms of the real-space (lattice) Green functions\(^29,55\) for the sample attached to two clean semi-infinite leads. To study the conductance in the whole band of the DD model, \( t_{nn} = 1.5 \) is used\(^55\) for the hopping integral in the leads. This mesoscopic computational technique ‘‘opens’’ the sample, thereby smearing the discrete levels of an isolated system. Thus, the spectrum of sample + leads = infinite system becomes continuous and conductance can be calculated at any \( E_F \) inside the band. However, the computed conductance, for not too small disorder\(^55,56\) or coupling to the leads\(^57\) (which are of the same transverse width as the sample\(^56\)) is practically equal to the ‘‘intrinsic’’ conductance \( g = \pi F_{\text{Tr}}/\Delta \) determined by the spectral properties of a closed sample.

The conductance and density of states (DOS),

\[
N(E) = \frac{2 \rho(E)}{V},
\]

are plotted in Fig. 6 for the samples whose eigenstates are investigated (the factor of two is for spin degeneracy). The DOS is obtained from the histogram of the number of eigenvalues which fall into equally spaced energy bins along the band. The conductance and DOS of the RH model have a peak at \( E = 0 \), which becomes a logarithmic singularity in the limit of infinite system size.\(^50\) To get an insight into the ‘‘general weakness’’ of the off-diagonal disorder, Fig. 7 plots \( g(E_F) \) and \( N(E) \) for the low \( W_{\text{RH}} = 0.25 \). In this case \( N(E) \) still resembles the DOS of a clean system, even after ensemble averaging. On the other hand, the conductance is a smooth function of energy since discrete levels of an isolated sample are broadened by the coupling to leads. The same is true for the DOS computed from the imaginary part of the Green function for an open system. The mobility edge is absent at low RH disorder (\( W_{\text{RH}} = 0.25 \) and \( W_{\text{RH}} = 0.375 \)) for system sizes \( L \leq 16a \). This means that localization length \( \xi \) is greater than \( 16a \) for all energies inside the band of these systems. For other samples on Fig. 6 the mobility edge appears inside the band. This is clearly shown for \( W_{\text{RH}} = 1 \) case where band edge \( E_b \) \( \left[ N(E_b) = 0 \right] \) differs from \( E_c \). The mobility edge is located at the minimum energy \( |E_c| \) for which \( g(E_c) \) is still different from zero. The conductance of finite samples is always finite, although exponentially small at \( |E_F| > |E_c| \). The approximate values of \( |E_c| \) listed in Fig. 6 are such that conductance satisfies \( g(E_F) < 0.1 \), for \( |E_F| > |E_c| \); typically \( g(E_c) \approx 0.2 \). The mobility edge extracted in this way will not change\(^59\) when going to larger system sizes if \( \xi < L \) for all energies \( |E| > |E_c| \).

The distribution \( f(t) \) of eigenfunction intensities has been studied analytically for diffusive conductors close to the universal RMT limit (where conductance is large and localization effects are small) in Refs. 9 and 23 using the supermatrix \( \sigma \) model (NLSM),\(^12\) or by means of a ‘‘direct optimal fluctuations method’’ of Ref. 60. Numerical studies of statistics of eigenstates in DD Anderson model were conducted in

\[\text{FIG. 6. Conductance and density of states in the RH and DD Anderson models on a simple cubic lattice}}\]

\[\text{FIG. 7. Conductance and density of states in the RH Anderson models on a simple cubic lattice}}\]
2D for all disorder strengths,\textsuperscript{10} while in 3D the focus has been on the states appearing in the semiclassical transport regime where comparison with analytical predictions (parameterized by semiclassical quantities, like $k_F l$) may be made in the regions of small\textsuperscript{61} and large\textsuperscript{62} deviations of $f(t)$ from PT distribution. Here I show how $f(t)$ evolves with the strength of (different types of) disorder in 3D samples, where genuine LD transition occurs in the strong coupling regime of the corresponding field-theoretical formulation.

In the weakly disordered ($k_F l > 1$) metallic ($g > 1$) conductors prelocalized states are extremely rare. For example, in an ensemble of 20,000 samples ($W_{\text{DD}} = 4$), whose typical transport properties are well-described by semiclassical theories, only four states would show up in the band center which exhibit similar amplitude splashes like the one in Fig. 1. Thus, to get a far tail (where deviations from PT distribution are large) of $f(t)$ in such conductors one has to search through enormous ensemble and locate special configurations of a random potential.\textsuperscript{62} The maximum wave function amplitudes which can be observed in this pursuit are, plausibly, determined by the strength of disorder, i.e., conductance $g$. It is, however, interesting that tails at small but finite $g$ in the strong disorder (like $W_{\text{DD}} = 10$) can be longer than the tails of states in the localized phase, where $g$ is vanishingly small, for $|E| > |E_c|$ at some smaller fixed disorder (like in the case of $W_{\text{DD}} = 6$; compare the two panels in Fig. 4 using respective conductances from Fig. 6). For strong enough disorder long tails of $f(t)$ are found, even by investigating small ensembles of disordered conductors (as shown below), since the frequency of appearance of prelocalized states is greatly enhanced (while system is still on the delocalized side of LD transition).

The complete eigenproblem of a single particle random Hamiltonian is solved exactly by numerical diagonalization. Then, $f(t)$ is computed as a histogram of intensities for the chosen eigenstates in delocalized (|$E| > |E_c|$), localized phase (|$E| > |E_c|$), and critical region around the mobility edge ($E_c$). The two delta functions in Eq. (1) are approximated by a box function $\delta(x)$. The width of $\delta(E - E_c)$ is small enough at a specific energy that $\rho(E)$ is constant inside that interval. For each sample, 5–10 states are picked by the energy bin, which effectively provides additional averaging over the disorder (according to ergodicity\textsuperscript{17} in RMT). The amplitudes of wave functions are sorted in the bins defined by $\delta(-|\Psi_{n}(x)|^2V)$ whose width is constant on a logarithmic scale. The function $f(t)$ is computed at all points inside the sample, i.e., $N = 16^3$ in Eq. (1).

The evolution of $f(t)$, when sweeping the band through the interesting states, is plotted in Figs. 2 and 3 for the RH disordered sample. Since prelocalized states generate slow decay of $f(t)$ at high wave function intensities (where PT distribution is negligible),\textsuperscript{12} this region is enlarged on Fig. 3. This is obvious from the prelocalized example in Fig. 1 where state with large amplitude spikes, highly unlikely in the framework of RMT, was found in a very good metal. The same is trivially true for the localized states which determine extremely long tails of $f(t)$. Thus, the long asymptotic tails of $f(t)$, appreciably deviating from PT distribution, are signaling the onset of localization. It is interesting that states in the band center of RH model, which define the largest zero-temperature conductance $\varrho(E_F = 0) \approx 10.2$, $\text{Var} \varrho(E_F = 0) \approx 0.63$, are in fact mostly prelocalized. Moreover, both the frequency of their appearance and high amplitude splashes resemble the situation at criticality (NLSTM-type calculation\textsuperscript{47} shows that 3D wave functions, sufficiently close to the band center, are always extended for any disorder strength). It might be conjectured that these prelocalized states would generate multifractal scaling of IPR in the band center. This result, together with the DOS and conductance from Figs. 6 and 7, shows that phenomena in the band center of 3D conductors with off-diagonal disorder are as intriguing as their much studied counterparts in low-dimensional systems.\textsuperscript{30,45,46} The origin of these phenomena can be traced back to a special sublattice, or ‘‘chiral,’’ symmetry obeyed by TBH (2) with random hopping and constant on-site energy (leading to an eigenspectrum which for $E_{\alpha}$ contains $-E_{\alpha}$, with a special role played by $E = 0$). In the $W_{\text{RH}} = 0.25$ and $W_{\text{RH}} = 0.375$ cases all states are extended. Their $f(t)$ looks similar to the distribution function for the delocalized states at $E = 1.5$ in the sample characterized by $W_{\text{RH}} = 1$. The distribution function $f(t)$ defined in Eq. (4) fits reasonably well the numerical $f(t)$ generated by the states around $E = 2.75$, where $\xi = 1.2a$ is extracted for the localization length. Thus, one can measure approximately $\xi$ in this way\textsuperscript{10} even though the structure of localized eigenstates can be more complicated\textsuperscript{2} than the simple radially symmetric exponential decay used to derive $f(t)$ [e.g., in the case of DD disorder] $\xi = 1.3a$ is obtained for examined localized states in Fig. 4 for $W_{\text{DD}} = 10$ ensemble, while corresponding states in $W_{\text{DD}} = 6$ ensemble seem to be too close to the mobility edge to follow $f(t)$. The same statistical analysis is performed for the eigenstates of DD Anderson model—a ‘‘standard model’’ in the localization theory. Figure 4 plots $f(t)$ at specific energies $E_i$ in samples characterized by different conductances $g(E_F = E_{\alpha})$. The conductance $g(E = 0)$ of TBH with $W_{\text{DD}} = 6$ is numerically close to the conductance of RH disordered samples with $W_{\text{RH}} = 1$. Nevertheless, comparison of the corresponding distribution functions reveals disorder-dependent features\textsuperscript{37} which are beyond universal (i.e., independent of the details of random potential) corrections accounted by the properties of a classical diffusion operator\textsuperscript{14,37} (the spectrum of $-D\nabla^2$, with appropriate boundary conditions, depends on $g$, shape of the sample and dimensionality; note that conductors at $W \approx 6$ with half-filled band lie on the boundary of applicability of such semiclassical concepts\textsuperscript{38}). To get the far tail of the eigenstate statistics in the band center a much larger statistics is needed than used here.\textsuperscript{62} This then makes the observed $f(t)$ in a special case of the band center of $W_{\text{RH}} = 1$ disordered conductor even more spectacular because of very large amplitudes found in the small ensemble of conductors. Thus, the strong dependence on the microscopic details of random potential demonstrated in this study is somewhat different from the disorder-specific short length scale (‘‘ballistic’’ contributions\textsuperscript{9,62} to the standard picture of diffusive NLSM. Namely, here it seems that special features of off-diagonal disorder in the band center generate...
completely different functional form of the far tail, and not just some disorder-specific values of the parameters\textsuperscript{9,62} in the \( \exp(\log\text{-}\text{cube}) \) asymptotics.\textsuperscript{9,23,60}

In both models, all computed \( f(t) \) intersect PT distribution (from below) around \( 6 \leq t \leq 10 \), and then develop tails far above PT values. The length of the tails is defined by the largest amplitude exhibited in the prelocalized state (like that in Fig. 1). For strong DD, \( W_{\text{DD}} = 10 \), the conductance \( g(E_F) \) is smaller than 3.5. In this regime transport becomes “intrinsically diffusive,”\textsuperscript{28} but one can still extract resistivity from the approximate Ohmic scaling of disorder-averaged resistance\textsuperscript{28} [for those fillings where \( g(E_F) > 2 \)]. However, the close proximity to the critical region \( g \sim 1 \) induces long tails of \( f(t) \) at all energies throughout the band—a sign of increased frequency of appearance of highly inhomogeneous states. This provides an insight into the microscopic structure of eigenstates which carry the current in a non-semi-classical transport regime\textsuperscript{9} (characterized by the lack of some intuitive concepts, like mean free path \( l \), since unwarranted use of the Boltzmann theory would give \( f \sim a \) in this transport regime although the sample is still far away from the LD transition).

In conclusion, the statistics of eigenstates in 3D mesoscopic conductors, modeled by the tight-binding Hamiltonian on the finite-size simple cubic lattice, have been studied. The disorder is introduced either in the potential energy (diagonal) or in the hopping integrals (off-diagonal). Also calculated are the average inverse participation ratio of eigenfunctions and the exact zero-temperature conductance as a function of Fermi energy. This comprehensive set of parameters makes it possible to compare the eigenstates in nanoscale samples with different types of disorder, but characterized by similar values of conductance. Disorder-specific details, which are not parameterized by the conductance alone, are found. This is in spite of the fact that dimensionality, shape of the sample, and conductance are expected to determine the finite-size corrections to the universal predictions of random matrix theory, at least in the samples which are in the semiclassical transport regime. The appearance of states with large amplitude spikes on the top of RMT like background is clearly demonstrated even in good metals. At criticality, such “prelocalized” states are directly related to the extensively studied multifractal scaling of IPR. However, even in the delocalized phase with good metallic properties \( (g > 1) \), where the correlation length \( \xi_c \sim \lambda_F \) defined by the sample conductance \( g(\xi_c) \sim 1 \) is microscopic \( (L < \xi_c \) would naturally account for the multifractal scaling.\textsuperscript{6} like in 2D), prelocalized state with unexpectedly high amplitudes are found in the band center of random hopping disordered systems. They are inhomogeneous enough to generate extremely long tails of the distribution of eigenfunction amplitudes, akin to the ones observed at criticality.

Inspiring discussions with V. Z. Cerovski are acknowledged. Valuable guidance has been provided by P. B. Allen. The important improvements of the initial cond-mat report have resulted from criticism provided by B. Shapiro and I. E. Smolyarenko. This work was supported in part by NSF Grant No. DMR 9725037.
Note that random Hamiltonians of real disordered solids are tied to the real-space representation [i.e., matrix elements in this representation are spatially dependent and, e.g., TBH (2) is a band diagonal matrix]. Therefore, they do not satisfy the statistical assumptions of standard RMT ensembles since all elements of random matrices in RMT are nonzero and spatially independent (Ref. 32). Nevertheless, the connection to RMT statistics is provided by Efetov’s supersymmetric approach (Ref. 12).

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