Delocalization of ultracold atoms in a disordered potential due to light scattering

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We numerically study the expansion dynamics of ultracold atoms in a one-dimensional disordered potential in the presence of a weak position measurement of the atoms. We specifically consider this position measurement to be realized by a combination of an external laser and a periodic array of optical microparticles arranged in a waveguide. The position information is acquired through the scattering of a near-resonant laser photon into a specific eigenmode of one of the cavities. The time evolution of the atomic density in the presence of this light scattering mechanism is described within a Lindblad master equation approach, which is numerically implemented using the Monte Carlo wave function technique. We find that an arbitrarily weak rate of photon emission leads to a breakdown of Anderson localization of the atoms.

I. INTRODUCTION

The realization of potentials with controlled disorder for ultracold atoms by means of optical speckle fields 1,2 or bichromatic optical lattices 3 has recently led to the observation of Anderson localization with Bose-Einstein condensates 1,3. In those experiments, atomic Bose-Einstein condensates, prepared in a harmonic trap, were released into one-dimensional optical waveguides which were superimposed with disordered potentials realized with speckle fields 1 as well as with bichromatic optical lattices 3. Absorption images of the atomic cloud after the expansion process within the waveguide clearly revealed an exponential decrease of the average atomic density with the distance from the center of the former trap, which is the characteristic signature of Anderson localization 1,3. While interaction effects did not play a role in those pioneering experiments, more recent studies specifically focus on the interplay of atom-atom interaction and localization in disordered potentials (e.g. Ref. 2, 8). Current research directions include the exploration of Anderson localization with ultracold atoms in three spatial dimensions 3, with the particular aim to study the Anderson metal-insulator transition 10.

Clearly, a key condition for the observability of Anderson localization with ultracold atomic gases is the overall coherence of the atomic cloud. Any mechanism of decoherence would compromise the phenomenon of destructive wave interference that lies at the heart of Anderson localization 1,3 and thereby give rise to delocalization. This also concerns any in-situ monitoring of the evolution of the atomic cloud during its expansion, by intermediate measurements of the positions of atoms. Evidently, the strong refocusing of the atomic wavefunction that results from a precise position measurement would destroy the coherence of the atom, enhance its kinetic energy, and eventually let the atom behave as a classical particle when being performed several times.

FIG. 1: (Color online) Sketch of the configuration under consideration. An atom, which is initially localized within a harmonic trap at the center, is expanding within a one-dimensional waveguide (indicated by the green horizontal lines) to which a disordered potential is superimposed. A periodic array of optical microparticles (symbolized by the red arcs) is used to measure the position of the atom, on a length scale that is comparable to its localization length (indicated by the shadow plot in the waveguide, which shows the density of a localized state). For this purpose, the waveguide is considered to be irradiated by a near-resonant laser beam, which may induce spontaneous emissions of photons into one of the cavities, possibly to be measured by photodetectors. The sketch is to scale with the parameters considered in this study, as far as the horizontal length scales are concerned.

The situation is less obvious for “weak” measurement processes, in which the position of the atom is determined with a large spatial uncertainty that is of the order of the expected localization length within the disordered potential. Such weak measurements might still preserve coherence to a certain extent, while, at the same time, providing some rough in-situ information on the position of the atom. One could, for this purpose, conceive e.g. a periodic array of optical microparticles placed around the waveguide in which the atoms propagate, as depicted in Fig. 11. A near-resonant laser beam which irradiates this configuration can be used to transfer the atoms to an electronically excited state, from which they can relax to the ground state by a spontaneous emission of a photon into one of the cavities, which in turn could be measured by photodetectors placed behind the cavities.
The whole configuration could possibly be fabricated on “atom chips” [11], in which case the disordered potential could arise from imperfections in the current-carrying wires that generate the magnetic waveguide potential of the atoms [12,13]. Our results below are, however, more general and we expect to see the same effects on the localization in the presence of any mechanism of similar position measurement.

The aim of this study is to investigate to which extent this approximate realization of a “Heisenberg microscope” gives rise to delocalization of an atom in a one-dimensional disordered potential. For the sake of simplicity, we shall restrict our consideration to the propagation of one single atom, and thereby discard collective processes arising within Bose-Einstein condensates due to atom-atom interactions or superradiance. We shall, moreover, assume that the atom will emit photons into one single mode of the cavities only. Such an emission will then give rise to a recoil that is mainly perpendicular to the direction of propagation of the atom and does therefore not dramatically enhance its longitudinal kinetic energy. We neglect effects of transverse excitations within the waveguide due to this recoil and assume that neither the effective waveguide confinement nor the disordered potential are affected by temporary populations of the excited electronic state of the atom.

The dynamics of the atom is modeled via a one-dimensional master equation for its density matrix $\hat{\rho}(t)$, which can be unraveled using the Monte-Carlo wavefunction technique [14,15]. This master equation accounts both for the coherent motion within the disordered potential and the incoherent scattering of photons [16]. We shall, in Section II, first account on the expansion and localization dynamics of a single atom in a one-dimensional disordered potential in the absence of any decoherence mechanism. In Section III, we outline the Monte-Carlo wavefunction approach that is used to integrate the master equation for the special case of an atom that propagates in a homogeneous, disorder-free waveguide. Decoherence and disorder are finally put together in Section IV in which we discuss the expansion of an atomic wave packet in the presence of disorder and spontaneous emission. We show that even very rare position measurements of the atom give, on average, rise to a gradual delocalization of the wave packet, and we provide numerical evidence for superballistic expansion in the presence of strong emission rates.

II. WAVE PACKET EXPANSION IN DISORDER

In this section, the expansion of an initially trapped wave packet in a weak one-dimensional disordered potential is discussed. For the sake of simplicity, we model the disorder by a Gaussian correlated random potential $V(x)$ defined along the $x$-axis, with the properties $V(x) = 0$ and $\langle V(x)V(x') \rangle = U^2 \exp[-(x-x')^2/(2\sigma^2)]$ for the mean spatial correlation function. Here, $U$ characterizes the typical size of the fluctuations of the potential, and the correlation length $\sigma$ controls the average width of fluctuations.

In Fig. 2 we show the time evolution of the disorder-averaged spatial density of wave packets propagating in such disorder configurations. These wave packets are initially prepared in the ground state of a harmonic trap with the oscillator length $a_0 = \sqrt{\hbar/m\omega}$. After the trapping potential is switched off, the wave packet expands within the disordered potential until it approaches, on average, a stationary profile. The convergence to the average density distribution happens faster at the center than in the wings. This is a consequence of the quadratic growth of the localization length as a function of the wave vector, as described in Eq. (3) below.

The final density profile shown in Fig. 2 is fairly well reproduced by a theory as described, for example, in Ref. [17], which is based on the assumption that the asymptotic probability distribution is an incoherent sum of individually localized plane waves with momentum $p$. This consideration yields the spatial density

$$\rho_{\text{loc}}(x) = \int dp \frac{\rho_0(p)}{2\xi(p)} \exp[-|x|/\xi(p)]$$

where $\rho_0$ denotes the momentum density of the wave packet at the initial time $t = 0$. The key ingredient for the evaluation of Eq. (2) is the localization length $\xi(p)$.
that can be calculated using diagrammatic theory\textsuperscript{18} as
\[ \xi(p) = 2l_B(p), \]
with \( l_B \) the Boltzmann mean free path (see Refs.\textsuperscript{13,19} for other approaches). For the Gaussian correlated random potential under consideration, we obtain
\[ \xi(p) = \frac{1}{\sqrt{2\pi}} \frac{\hbar^2 p^2}{m^2 U^2 \sigma} \exp[2(p\sigma/h)^2]. \tag{3} \]

In the regime of short correlation lengths \( \sigma \ll h/p \), we can approximate \( \exp[2(p\sigma/h)^2] \approx 1 \) and the localization length depends only on the effective strength \( U^2 \sigma \) of the disorder. Using
\[ \rho_0(p) = \frac{a_0}{\sqrt{\pi h}} \exp[-(a_0 p/h)^2] \tag{4} \]
and introducing the characteristic localization length scale of the wave packet as \( \xi_0 \equiv \xi(h/a_0) \), we then obtain the prediction
\[ \rho_{\text{loc}}(x) = \frac{1}{2 \sqrt{\xi_0 |x|}} \exp\left(-2 \sqrt{|x|/\xi_0}\right) \tag{5} \]
for the localized density. As shown in Fig.\textsuperscript{2} this approximate expression is, apart from a global prefactor, in good agreement with the numerically computed mean density at the final time \( t = 800/\omega \).

In the above numerical simulations, we effectively assumed that the atomic cloud is prepared in a clean harmonic trap in absence of any disorder. At \( t = 0 \) the trapping potential is suddenly switched off and the disorder is ramped on at the same time. The initial state is then a perfect Gaussian wavefunction [see Eq.\textsuperscript{4}] which expands within the disordered potential. This procedure is, in general, not precisely in accordance with expansion experiments on Anderson localization such as Ref.\textsuperscript{1} in which the disordered potential is already present during the formation of the Bose-Einstein condensate in the harmonic trap. The initial state of the atomic wavefunction is, in that case, given by the ground state of an effective trapping potential that consists of a harmonic confinement modulated by the disorder. A numerical comparison of these two expansion scenarios, however, displays no significant difference in the asymptotic density profile for the case of weak disordered potentials with \( U \simeq 0.1 \hbar \omega \) and \( \sigma = 0.2 a_0 \).

A convenient numerical observable for measuring localization is the participation ratio \( \text{Pr}(t) \) which for a wave packet with the density \( \rho(x,t) \) is defined by
\[ \text{Pr}(t) = \left( \int_{-\infty}^{\infty} dx |\rho(x,t)|^2 \right)^{-1}. \tag{6} \]

In practice, \( \text{Pr}(t) \) represents a measure for the spatial extent of the wave packet, yielding large values for rather extended distributions \( \rho(x,t) \) and going to zero for strongly peaked wavefunctions. It therefore exhibits a similar behavior to the spatial root mean square (rms) width \( \Delta x = \sqrt{\langle x^2 \rangle - \langle x \rangle^2} \) of the wave packet. This latter quantity is, however, rather sensitive to the evolution of the (experimentally inaccessible) wings of the wave packet. This is shown in Fig.\textsuperscript{2} where we display the time dependence of the disorder-averaged rms width and participation ratio. While the rms width continuously increases with time, due to the long-time dynamics in the wings of the averaged density distribution (see Fig.\textsuperscript{2}), the participation ratio, which is much less sensitive to the behavior of the wings, saturates at a finite length scale. This length scale can be used in order to define an effective localization length \( L_{\text{wp}} \) of the wave packet.

Fig.\textsuperscript{3} shows the time evolution of the participation ratio for different initial kinetic energies \( E \), resulting from different confinement frequencies of the initial trapping potential. A linear increase of the participation ratio with \( E \), corresponding to a linear increase of the effective localization length \( L_{\text{wp}} \) of the wave packet, is found for \( E_0 < E < 2E_0 \).

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig3.png}
\caption{(Color online) Root mean square (rms) width \( \Delta x = \sqrt{\langle x^2 \rangle - \langle x \rangle^2} \) (red line) and participation ratio \( \text{Pr}(t) \) (black line) as a function of the evolution time \( t \), showing the expansion and localization of a wave packet for the disorder strength \( U = 0.15 \hbar \omega \) and the correlation length \( \sigma = 0.2 a_0 \). The dashed lines show, for comparison, the rms width and the participation ratio of a free wave packet that expands in the absence of disorder.}
\end{figure}

III. Master Equation Dynamics

To account for spontaneous emissions of photons into the cavities, we model the dynamics of the atom via a one-dimensional master equation for its density matrix \( \hat{\rho}(t) \), including coherent interactions with a disordered potential and the incoherent scattering of light\textsuperscript{16}. This master equation is given by
\[ \frac{d}{dt} \rho(t) = -\frac{i}{\hbar} \left[ \hat{H}, \hat{\rho}(t) \right] + \gamma_{\text{eff}} \int_{-k}^{k} dq \left( \hat{C}_{q} \hat{\rho}(t) \hat{C}_{q}^\dagger - \hat{\rho}(t) \right). \tag{7} \]

Here, \( \hat{H} = \frac{\hat{p}^2}{2m} + V(\hat{x}) \) describes the Hamiltonian for a particle that propagates in the disordered potential. \( \hat{C}_{q} = e^{-iq\hat{x}} \) is the decay or jump operator representing one spontaneous emission event, which exerts a recoil on the atom with longitudinal momentum \( h\hat{q} \) which is
assumed to be equidistributed between $-\hbar k$ and $+\hbar k$. This model considers off-resonant inelastic scattering, in which a laser couples the electronic ground state to an excited state from which spontaneous emission back to the ground state can occur. It assumes a low spontaneous decay rate $\gamma$ as compared to the detuning $\delta$ of the laser with respect to the intra-atomic transition frequency, and a low Rabi frequency $\nu$ for laser-induced transitions between the ground state and the excited state as compared to the spontaneous decay rate $\gamma$, i.e. we assume $\nu \ll \gamma \ll \delta$. We then obtain $\gamma_{\text{eff}} = \gamma \nu^2 / (\gamma^2 + 4\delta^2)$ as the effective decay rate that enters the master equation.

To solve the time evolution generated by the master equation, we employ the Monte Carlo wave function method. Here, the evolution of the density matrix is decomposed into the non-unitary evolution of a large number $N = 100$ of wave functions. A single trajectory $|\psi_i(t)\rangle$ evolves according to $i\hbar \partial_t |\psi_i(t)\rangle = \hat{H}_{\text{eff}} |\psi_i(t)\rangle$, with $\hat{H}_{\text{eff}} = \hat{H} - i\gamma_{\text{eff}} / 2$, until the exponentially decaying norm $|||\psi_i(t)\rangle||^2 = e^{-\gamma_{\text{eff}} t}$ equals a random number chosen between 0 and 1. At this point, a jump operator $\hat{C}_q$ acts on the Monte Carlo wave function: $|\psi(t + \delta t)\rangle_i = \hat{C}_q |\psi(t)\rangle_i$. This jump operator is determined by randomly choosing $q$ from the interval between $-k$ and $k$.

To relate this light scattering process to the position measurement under consideration, we note that the Lindblad master equation is invariant under unitary transformations on the set of decay operators. Indeed, it was shown in Ref. [24] that the Fourier transformation $\int_{-1}^{1} du \exp(iu k x / 2) \hat{C}_u$ with integers $u \in \mathbb{Z}$ allows one to switch to decay operators

$$\hat{C}_\nu = \sqrt{2} \frac{\sin(k x - \nu)}{k x - \nu}$$

In this picture, the application of the decay operator induces a localization of the wavefunction within a spatial region whose extent is on the order of $k^{-1}$. For the sake of simplicity, we assume that these decay operators exactly correspond to the longitudinal structure of the cavity modes into which the atom may emit the photon. The spatial period of the array of cavities is then given by $\lambda = 2\pi / k$.

In Fig. 5 we show the time evolutions of the expectation value of the position $\langle x \rangle$ and its rms width $\Delta x = \sqrt{\langle x^2 \rangle - \langle x \rangle^2}$ for a single quantum trajectory in free space, $V(x) = 0$. In this particular trajectory, the first spontaneous emission took place in one of the two wings of the wave packet, which is mainly constituted by plane-wave components with high momenta. The subsequent localization process projects the wavefunction on those high-momentum components, which gives rise to a permanent drift. The rms width, however, remains small during this evolutions, which is due to the fact that the atom emits photons at a rate that is faster than the inverse dispersion time of the wave packet. The rms width would freeze for sufficiently high emission rates, which is reminiscent of the quantum Zeno effect.

It is of great advantage to work in a regime where $k$ is small compared to fluctuations of the density matrix in momentum space $\rho(p, p', t)$. To study the momentum density distribution, we can then approximate the integrand of Eq. (7) by its Taylor expansion to first order, as done in Ref. [24]. Taking the integral over $k$ leads to the diffusion equation

$$\partial_t \rho(p, t) = \frac{1}{6} \gamma_{\text{eff}} k^2 \partial_p^2 \rho(p, t)$$

for the diagonal elements of the density matrix, with the effective diffusion coefficient $D = \gamma_{\text{eff}} k^2 / 6$. Hence, the
wave packet will undergo diffusive spreading in momentum space. Noting that the variance of the momentum distribution is nothing but the kinetic energy, we obtain

$$\langle \hat{T} \rangle = \text{Tr}\left\{ \frac{\hat{p}^2}{2m} \hat{\rho}(t) \right\} = E_0 + \frac{\hbar^2 k^2}{6m} \gamma_{\text{eff}} t$$

for the growth of the mean kinetic energy of the wave packet.

IV. DISSIPATIVE EXPANSION IN DISORDER

Having introduced the necessary tools, we now study wave packet expansion in the presence of disorder and dissipation. In Fig. 6 we plot the participation ratio as a function of time for different effective emission rates $\gamma_{\text{eff}}$. In accordance with the sketch shown in Fig. 1, we have chosen the photon wavelength to be very long compared to the initial extension $a_0$ of the wave packet. The amount of kinetic energy given to the wave packet at each emission event is thereby rather reduced.

The most important observation is a delocalization of the wave packet at any emission rate. Instead of saturating to a stationary value, the participation ratio linearly increases with time after the typical time scale that is needed for developing an Anderson-localized density profile in the absence of spontaneous emission. Quantitatively, this linear growth is very different from a ballistic expansion in free space, which takes place with much faster expansion velocities (see the dashed lines in Fig. 3). It is different from simple diffusion which one would naively expect to prevail for a quantum particle that propagates within a disordered potential in the presence of a decoherence mechanism. We attribute this difference to the fact that the spontaneous emission of a photon gives rise to a recoil of the atom and thereby increases its energy. Hence, the effective diffusion constant should also gradually increase with time.

It is, in this context, interesting to note that the expansion velocity $dPr/dt$ depends only on the product of the effective rate of emission $\gamma_{\text{eff}}$ and the recoil energy $\hbar^2 k^2/(2m)$. This can be seen by comparing the two blue lines in the two panels of Fig. 6 showing expanding participation rates for $k = 0.07/a_0$ and $\gamma_{\text{eff}} = 0.05\omega$ (left panel) as well as for $k = 0.035/a_0$ and $\gamma_{\text{eff}} = 0.2\omega$. There appears, furthermore, no change in the behavior when we tune the rate of emissions across the scale $1/T_{\text{loc}}$, with $T_{\text{loc}}$ the time at which the unperturbed evolution shows localization.

It is tempting to relate the linear increase of the participation rate with time to the combination of a linear growth of the kinetic energy due to spontaneous emission with the approximately linear scaling of the wave packet’s localization length with its mean kinetic energy in the absence of spontaneous emission, as shown in Fig. 1. This reasoning essentially assumes that in between two subsequent spontaneous emission events the wave packet has enough time to approach its asymptotic stationary profile within the disordered potential. Extracting from Fig. 3 the approximate scaling $Pr/a_0 \sim 100 E_0/\hbar \omega$ and using $dE/dt = \hbar^2 k^2 \gamma_{\text{eff}}/(6m)$ for the growth rate of the energy according to Eq. (10), we obtain the prediction

$$\frac{dPr}{dt} \approx 100 \frac{a_0}{E_0} \frac{dE}{dt} \approx 400 \frac{\gamma_{\text{eff}} \hbar^2 k^2}{6m} \frac{a_0}{\hbar \omega}$$

for the expansion velocity $dPr/dt$ of the participation rate, using $E_0 = 0.25 \hbar \omega$.

Figure 6 shows, however, that this expansion velocity increases more strongly with the rate of increase of the kinetic energy than predicted by Eq. (11). As a matter of fact, $dPr/dt$ is found to scale as a square root of $dE/dt$ in the parameter regime in which we carried out our numerical investigations. One may attribute this behavior to the fact that the above reasoning rather applies to an individual quantum trajectory in the spirit of Fig. 1.

The energy of the wavepacket corresponding to each individual trajectory increases linearly and its participation ratio increases on average as described by Eq. (11). However, while different trajectories describe similar narrow wavepackets, each wavepacket will be centered around a different point in space. Thus the full (incoherent) density will be spreading faster over a larger region than a single wavepacket (as is obvious from Fig. 5 for the case of disorder-free propagation). This effect is obviously not accounted for in the considerations leading to Eq. (11).

Let us finally investigate the regime of strong dissipation for which it is expected that the expansion becomes independent of the disordered potential. In Fig. 5 the momentum recoil is set to $k = 0.35/a_0$, with...
represents the prediction of Eq. (11). The dashed line. The dash-dotted straight line in the main panel

dt \propto \frac{\alpha}{dE/dt}$: we have $dE/dt$ the participation ratio as a function of the energy growth rate $\omega$. For different effective emission rates $\gamma_{\text{eff}}$ for $k = 0.35/a_0$. In summary, we have shown that even a very weak rate of photon scattering gives rise to a breakdown of Anderson localization of an atom that propagates in a one-dimensional disordered potential. This breakdown is most conveniently quantified in terms of the disorder-averaged participation ratio of the atomic density, which represents a measure for the spatial width of the atomic wave packet. While this participation ratio saturates, within a characteristic time scale, to a finite value in the case of a perfectly coherent expansion process within the disordered potential, it is found to linearly grow with time beyond that time scale in the presence of spontaneous photon scattering.

Finally, we expect similar findings in the presence of other mechanisms that can behave as a position measurement of the propagating atom. Such mechanisms include noise on the lattice beams as well as collisions with background gas atoms, to mention two examples. Undesired effects of this type are therefore also expected to induce a delocalization of the atom in the disorder potential.

**V. CONCLUSION**

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