

Velocity selective trapping of atoms in a frequency-modulated laser field

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The wave function of a moderately cold atom in a stationary near-resonant standing light wave delocalizes very fast due to wave packet splitting. However, we show that frequency modulation of the field may suppress packet splitting for some atoms having specific velocities in a narrow range. These atoms remain localized in a small space for a long time. We propose that in a real experiment with cold atomic gas this effect may decrease the velocity distribution of atoms (the field traps the atoms with such specific velocities while all other atoms leave the field).

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I. Introduction. Laser cooling and trapping of atoms and ions is a rapidly developing field of modern physics. Cold particles in a laser field are a common physical substrate used in numerous fundamental and applied issues such as Bose–Einstein condensates, quantum chaos, single-atom laser, quantum computer, etc. A significant number of methods of atomic cooling in a laser field were developed in the recent decades (the Doppler cooling [1, 2], the Sisyphus cooling [3, 4], the velocity selective coherent population trapping (VSCPT) [5], dynamical localization and trapping [6], etc. [7]). Modern sophisticated methods provide temperatures of the order of 100 picokelvin [8].

In this paper we suggest a method of coherent laser cooling in the absence of spontaneous emission. When an atom moves in a near-resonant standing light wave, two periodic optical potentials form in the space [9]. When the atom crosses a standing wave node, it may undergo the Landau–Zener (LZ) transition between these two potentials. Such transitions cause splitting of the wave packets [10] and rapid delocalization of the wave function [11]. In this paper we show that frequency modulation of the field may suppress the splitting of wave packets for atoms that have velocities in the specific narrow range (determined by the field modulation parameters). We suppose that in a real experiment, this may significantly decrease the energy distribution of moderately cold atoms. This method does not pretend to establish any temperature record, however, it might be useful in some experiments due to its conceptual simplicity.

The ideology of this method is similar to VSCPT and dynamical trapping in some aspects. The analogy

with VSCPT is rather gentle. Both VSCPT and our method do not cool initially “hot” atoms, they only trap the atoms that already have specific velocities. However, in our method, this velocity is non-zero, and the particular trapping mechanism differs from VSCPT radically. Our method is not based on “the dark states”. It is based on the synchronization between the LZ transitions and the field modulation. The analogy with dynamical localization and trapping is more deep. Dynamics of cold atoms in a periodically modulated (and kicked) standing wave has been studied both theoretically and experimentally for 20 years by the groups Raizen and Zoller [6, 12, 13]. A lot of effects related to dynamical chaos and quantum-classical correspondence were reported. In particular, it was shown that in a modulated field, some atoms with special initial positions and momentums can be dynamically trapped (without obvious energy conditions for such trapping). In terms of dynamical system theory, these atoms are trapped in a resonance islands embedded in a chaotic sea (in a phase space) [6]. In our study, resonance between field modulation and atomic mechanical oscillations plays similar role. However, cited works describe semiclassical atomic motion far from atom-field resonance. Therefore, there is only one effective optical potential (with modulated amplitude). In our study, there are two optical potentials and LZ tunnelings between them. This physical situation differs significantly.

In our study the reported effect was initially proposed theoretically (semiclassical model) and then confirmed numerically (purely quantum model). However, we have organized this paper in an alternative order for better understanding. First, we demonstrate the numerical manifestations of the velocity selective trapping, and then explain the effect theoretically.

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II. Equations of motion. Let us consider a two-level atom (with the transition frequency ω_a and mass m_a) moving in a strong standing laser wave with the modulated frequency $\omega_f[t]$. Let us assume that the depth of modulation is negligible in comparison with the average value of frequency $\langle\omega_f[t]\rangle$ (but not with the detuning $\omega_f[t] - \omega_a$), so we can consider the corresponding wave vector k_f a constant. In absence of spontaneous emission (the atomic excited state must have long lifetime, or some experimental methods must be used to suppress the decoherence) the atomic motion may be described by the Hamiltonian

$$\hat{H} = \frac{\hat{P}^2}{2m_a} + \frac{1}{2}\hbar(\omega_a - \omega_f[t])\hat{\sigma}_z - \hbar\Omega(\hat{\sigma}_- + \hat{\sigma}_+)\cos k_f\hat{X}, \quad (1)$$

where $\hat{\sigma}_{\pm,z}$ are the operators of transitions between the atomic excited and ground states (the Pauli matrices), \hat{X} and \hat{P} are the operators of the atomic coordinate and momentum, and Ω is the Rabi frequency. This Hamiltonian was used in [10, 11, 14], though for a constant field without modulation.

Let us use the following dimensionless normalized quantities: momentum $p \equiv P/\hbar k_f$, time $\tau \equiv \Omega t$, position $x \equiv k_f X$, mass $m \equiv m_a \Omega / \hbar k_f^2$ and detuning $\Delta[\tau] \equiv (\omega_f[\tau] - \omega_a)/\Omega$. Let us suppose that the field modulation is harmonic,

$$\Delta[\tau] = \Delta_0 + \Delta_1 \cos[\zeta\tau + \phi], \quad (2)$$

and apply the following conditions: $\zeta \ll 1$, $\Delta_0 \lesssim \Delta_1 \ll 1$. Using these approximations we obtain the equations for the probability amplitudes to find an atom with the normalized momentum p in the excited or ground state, $a[p, \tau]$ and $b[p, \tau]$, correspondently:

$$\begin{aligned} i\dot{a}[p, \tau] &= \left(\frac{p^2}{2m} - \frac{\Delta[\tau]}{2}\right)a[p] - \frac{1}{2}(b[p-1] + b[p+1]), \\ i\dot{b}[p, \tau] &= \left(\frac{p^2}{2m} + \frac{\Delta[\tau]}{2}\right)b[p] - \frac{1}{2}(a[p-1] + a[p+1]). \end{aligned} \quad (3)$$

Here the dot designates the differentiation with respect to τ . For every value of p , there is its own pair (3).

Let us choose the values of the parameters and initial conditions in order to perform the numerical simulation. The average initial atomic momentum $\langle p[0] \rangle$ will be a variable condition for the purpose of this paper. All other conditions will be fixed: normalized mass $m = 10^5$ (by order of magnitude this corresponds to the experiments with Cs [15] and Rb [16] atoms, but for a stronger field $\Omega \sim 10^{9-10}$ Hz), field parameters $\Delta_0 = -0.02$, $\Delta_1 = 0.047$, $\zeta = 0.00508$, $\phi = 0$, and the initial form of wave packet

$$\begin{aligned} a[p, 0] &= b[p, 0] = \\ &= \frac{1}{\sqrt{2\sigma_p[0]}\sqrt{2\pi}} \exp\left[\frac{-(p - \langle p[0] \rangle)^2}{4\sigma_p^2[0]}\right]. \end{aligned} \quad (4)$$

Therefore, the initial wave packet has a Gaussian form with $\langle x[0] \rangle = 0$ and the initial probability to find the atom in the excited state 0.5. Here σ_p is the standard deviation of the atomic momentum (equal to the half-width of the packet by order of magnitude). At $\tau = 0$ we fix it by the value of $\sigma_p[0] = 5\sqrt{2}$. Therefore, in accordance with the Heisenberg relation, the standard deviation of the initial coordinate is $\sigma_x[0] = 1/(2\sigma_p[0]) = 0.1/\sqrt{2}$ (it is much less than the normalized optical wavelength 2π).

In numerical experiments, we use these initial conditions to simulate the system of 8000 equations (3) with $-1000 \leq p \leq 1000$. For larger values of $|p|$, we put $a[p, \tau] = b[p, \tau] = 0$ due to the energy restrictions. Obtaining the solution in the momentum space we perform the Fourier transform and get the wave function in the coordinate space in the range of $-4\pi < x \leq 4\pi$ (see figures in the next section).

III. Numerical results. In [10, 11], the atomic motion was studied in absence of field modulation. The following basic modes of motion were reported.

At $\Delta = 0$ and $|\Delta| \gtrsim 1$ the atomic motion is simple. Atoms move in constant spatially periodic potentials. Slow atoms are trapped in potential wells and fast atoms move ballistically through the wave.

At $0 < |\Delta| \ll 1$ the atomic motion is more complex. The slowest atoms ($|\langle p[0] \rangle| < \sqrt{2m}$) are trapped in potential wells. Faster atoms ($\sqrt{2m} \leq |\langle p[0] \rangle| < 2\sqrt{m}$) perform a kind of random walk. Their wave packets split each time they cross standing-wave nodes, and this causes fast delocalization of the wave functions. The fastest atoms ($|\langle p[0] \rangle| > 2\sqrt{m}$) move ballistically through the wave. Their wave packets split, but all products move in the same direction, so the overall delocalization is slow. In Fig. 1 we calculate the variance of the atomic position σ_x^2 after a relatively long time span of coherent evolution $\tau = 5000$ as a function of the initial atomic momentum $\langle p[0] \rangle$. For the constant field (solid curve) this function shows fast delocalization of all atoms in the range of $\sqrt{2m} \simeq 440 \lesssim \langle p[0] \rangle \lesssim 2\sqrt{m} \simeq 640$ (cold atoms with velocities of the order of 1 m/s). Local peak at $\langle p[0] \rangle \simeq 630$ is produced by moderately fast atoms having an uncertain scenario of either random walking or flying ballistically.

Now let us “switch on” the field modulation and see the changes. In Fig. 1 the analogous function of σ_x^2 is shown with triangles. This function has a more complex

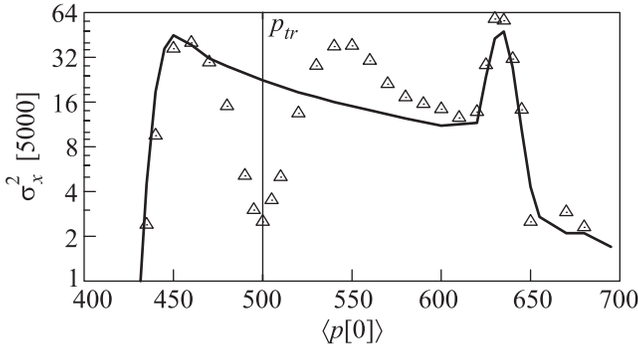


Fig. 1. The variance of atomic position σ_x^2 at $\tau = 5000$ as a function of initial atomic momentum $\langle p[0] \rangle$: curve – constant field $\Delta(\tau) = -0.02$, triangles – modulated field $\Delta(\tau) = -0.02 + 0.047 \cos[0.00508\tau]$

structure. In particular, it has a prominent additional minimum at $\langle p[0] \rangle = p_{tr} \simeq 500$. These atoms are not trapped in potential wells in a strict sense (their energy is too high, see the theory in the next sections), but some mechanism significantly suppresses the delocalization of their wave functions (note that both functions are shown in a logarithmic scale).

Let us consider the evolution of the corresponding wave packets in a coordinate space. In Fig. 2 we show the evolution of wave functions with $\langle p[0] \rangle = 600$ and 500 (other parameters are the same as in Fig. 1). In both cases wave packets split. The first splitting occurs near the first node, $x \simeq 1.57$ (products overlap at $\tau = 400$, but become completely independent at $\tau = 800$). However, the proportion of splitting radically differs for $\langle p[0] \rangle = 600$ and 500. In Fig. 2a fission products have similar “weights”, while in Fig. 2b they are radically different: a single large packet regularly oscillates in the range of $-2 \lesssim x \lesssim 2$ “emitting” very small packets in both directions.

We conclude that the slow delocalization of the wave function with $\langle p[0] \rangle \simeq 500$ is caused by the prominent asymmetry of wave packet splitting. Some mechanism suppresses the splitting of packets, and the atom is almost completely trapped in the range of $-2 \lesssim x \lesssim 2$ (the variance of its position x is even smaller, see Fig. 1). This suppression is significant only for atoms with $490 \lesssim \langle p[0] \rangle \lesssim 510$ (a comparatively narrow momentum and velocity range).

IV. Explanation of the effect. In the previous section we used quantum equations to simulate atomic dynamics. In this and the further section, in order to explain the effect of velocity selective trapping, let us mention some semiclassical analytical results from [10, 11] (obtained for the stationary field).

In a stationary field with $|\Delta| \ll 1$ the atomic motion can be described in terms of two potentials

$$U^- = -\sqrt{\cos^2[x] + \frac{\Delta^2}{4}}, \quad U^+ = \sqrt{\cos^2[x] + \frac{\Delta^2}{4}} \quad (5)$$

(Fig. 3a, dashed lines). An atom moves in one of these potentials when it is far from the standing wave nodes. When an atom crosses the node, the potential may change the sign (atom undergoes the Landau–Zener tunneling between potentials U^\pm). The probability of tunneling depends on Δ as $\exp[-A\Delta^2]$ (A is a combination of other parameters). At $0 < |\Delta| \ll 1$ the tunneling causes splitting of the wave packet (observed in numerical experiments). At $\Delta = 0$ potentials coincide at nodes, so the probability of tunneling is equal to 1 and wave packets do not split. The correspondent potential takes the simplest form $U = \pm \cos[x]$ (Fig. 3a, solid line).

What happens, if we “switch on” the field modulation? When an atom moves far from the nodes nothing radically changes. It moves in a constant potential that does not depend much on the value of Δ . Far from nodes we may neglect the term $\Delta^2/4$ in (5) and put $U \simeq \pm \cos[x]$ with good accuracy.

There are two possible scenarios when an atom crosses the node (at time τ): 1) $\Delta[\tau] \neq 0$, therefore, the packet splits significantly; 2) $\Delta[\tau] \simeq 0$, therefore, the splitting is suppressed.

The first scenario is more typical if the modulation is not synchronized with the atomic mechanical motion (because most of the time $\Delta[\tau] \neq 0$). Second scenario may occur sometimes, but does not change the overall statistics of the atomic motion. The evolution of the wave function shown in Fig. 2a is typical for moderately small detunings $|\Delta| \sim 0.01$ (both for the stationary and the modulated field).

However, the evolution radically changes if the field modulation is synchronized with the atomic mechanical motion. In particular, it is possible to choose such modulation parameters and atomic momentum (the particular values are estimated in the next section) that $\Delta[\tau]$ takes zero values each time an atom crosses the node. With our parameters such synchronization occurs at $\langle p[0] \rangle = p_{tr} \simeq 500$ (Fig. 2b). Note that packet splittings are suppressed, but not completely. Slight splittings are caused by the Landau–Zener transitions that occur not exactly at a standing wave node, but in its small vicinity (when $\Delta[\tau]$ is small but does not equal to zero).

V. Estimation of trapping conditions. Let us obtain the analytic relationship between trapping momentum p_{tr} and field parameters. Trapping occurs, if $\Delta[\tau] = 0$ each time atom crosses the nodes of the standing wave. In other moments of time $|\Delta[\tau]| \ll 1$.

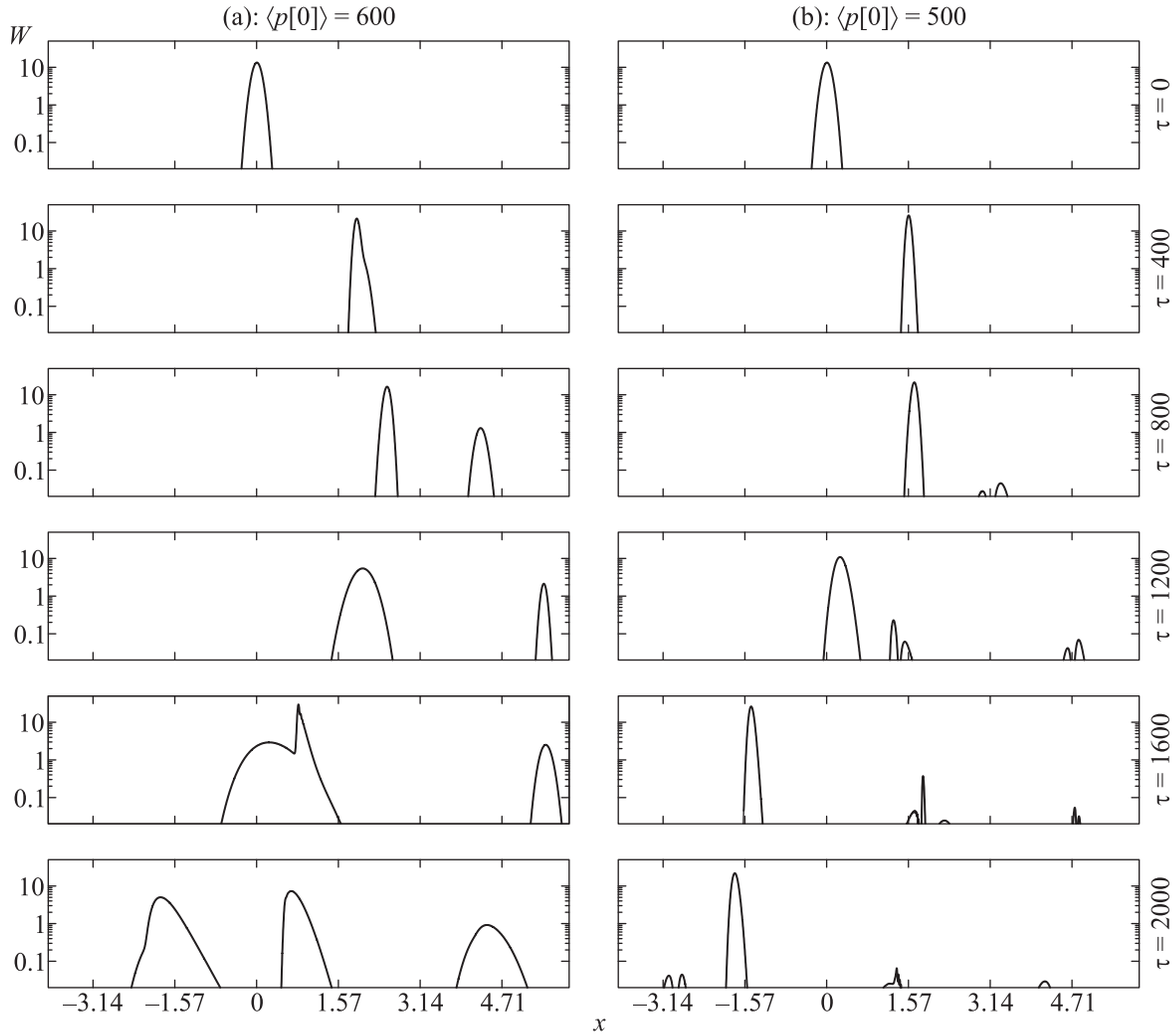


Fig. 2. Atomic wave packet splitting during quantum evolution (in the coordinate space). (a) – Fast delocalization of typical wave function ($\langle p[0] \rangle = 600$). (b) – Slow delocalization of wave function in the velocity selective trapping mode ($\langle p[0] \rangle = 500$). Here $W[x]$ is the probability density to find an atom at coordinate x . Note: due to logarithmic scale, it may seem that the norm of wavefunction is not conserved. However, our computations have shown that it is conserved with a good accuracy

Therefore, the term $\Delta^2/4$ in (5) is always negligible, and the trapped atom moves in the effective potential $U \simeq -\cos[x]$ (we choose the negative sign of U , because in this paper atoms with initial position $x[0] = 0$ start their motion from the potential well). Therefore, the atomic center-of-mass motion may be described by the semiclassical equations of motion [14]

$$\dot{x} = \frac{p}{m}, \quad \dot{p} = -\text{grad}[U] = -\sin[x], \quad (6)$$

with the trapping energy

$$E_{tr} \equiv \frac{p^2}{2m} - \cos[x] = \frac{p_{tr}^2}{2m} - \cos[\langle x[0] \rangle] \quad (7)$$

being the integral of motion (determined by the initial conditions). The trapping energy must be in the range

of $0 < E_{tr} < 1$ (for $x[0] = 0$, this corresponds to $\sqrt{2m} < |p[0]| < 2\sqrt{m}$). Slower atoms cannot reach the standing wave node, and faster atoms move ballistically.

Let us calculate the atomic traveling time between the two successive crossings of nodes in the negative and the positive segments of potential $-\cos[x]$ by integrating (6) (for $0 < E_{tr} < 1$)

$$\begin{aligned} \tau^- &= 2k\sqrt{m}, \quad k \equiv \sqrt{\frac{2}{1+E_{tr}}}, \\ \tau^+ &= 2k\sqrt{m} \left(F \left[\frac{\pi - |\arccos[E_{tr}]|}{2}, k \right] - 1 \right). \end{aligned} \quad (8)$$

Here F is the elliptic integral of the first kind.

In order to synchronize the modulation with the atomic mechanical motion the time intervals τ^\pm must

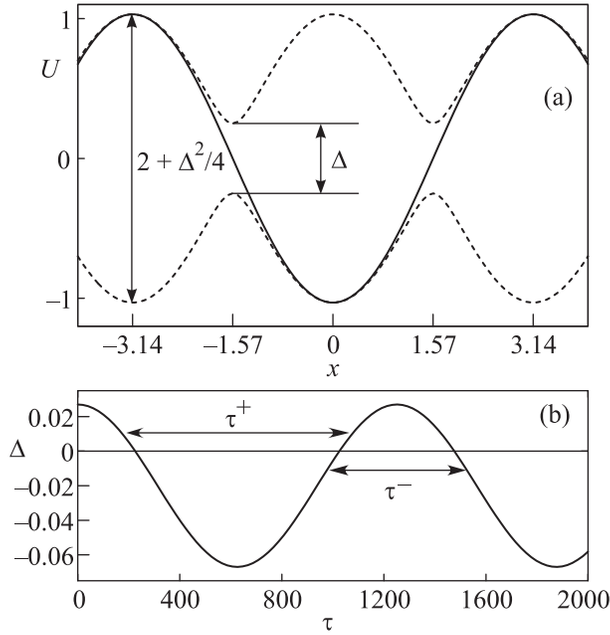


Fig. 3. (a) – Periodic potentials in the space: dashed line – non-resonant potentials U^\pm , solid line – resonant potential $-\cos[x]$. (b) – Illustration of the trapping condition: the modulation of detuning $\Delta[\tau]$ must be synchronized with atomic mechanical motion ($\Delta = 0$ each time a trapped wave packet crosses the standing wave node)

be equal to time intervals between successive zeros of $\Delta[\tau]$ (Fig. 3b). Therefore, using (2) and (8), we get

$$\zeta = \frac{2\pi}{\tau^- + \tau^+}, \quad \frac{\Delta_0}{\Delta_1} = -\cos\left[\frac{\pi\tau^-}{\tau^- + \tau^+}\right]. \quad (9)$$

These formulae are true for atoms with any initial positions (not only $x[0] = 0$ used in (4)). At any value of atomic energy in the range of $0 < E_{tr} < 1$ (and appropriate initial momentum) the velocity selective trapping of atoms can be achieved with appropriate values of $\Delta_{0,1}$, ζ calculated by these formulae. E.g., in order to observe trapping at $\langle p[0] \rangle = 500$, $x[0] = 0$, the field must have parameters $\zeta = 0.00508$, $\Delta_0/\Delta_1 = -0.4248$. We use them in numerical experiments, additionally fixing $\Delta_0 = -0.02$.

VI. Conclusions. In this paper we report the effect of velocity selective trapping of atoms in a frequency-modulated standing laser wave.

Intensive coherent light produces significant mechanical action on cold atoms having velocities of the order of 1 m/s. There is a wide range of field parameters at which atom performs a kind of random walk accompanied with wave packets splitting and fast delocalization of wave function. In this paper we report a specific field modulation mode that suppresses wave packet splitting

for atoms with precisely selected velocities. These atoms oscillate in potential wells, and their wave functions are almost completely localized.

This effect cannot cool atoms in the sense of achieving zero velocity, but it can decrease their mechanical energy distribution. E.g., if we have a cloud of moderately cold atoms having wide position and momentum distribution we can switch on the modulated standing wave and wait for some time. Most of atoms will leave the wave, and only small fraction will be trapped. These trapped atoms will have similar mechanical energy determined by field parameters (see formulae (8), (9)), and only the phase of their mechanical oscillations in wells will differ (because initial position distribution is random). In future study, we plan to simulate numerically large atomic ensemble cooled by the modulated laser. This will demonstrate explicitly that our effect not only traps but also cools the atoms.

The effect of velocity selective trapping of atoms, being theoretically predicted with the semiclassical apparatus, has been confirmed by purely quantum numerical modeling. Therefore, it is not just an artifact of semiclassical analytics but a real possibility. The drawback of this result is that it is obtained in absence of dissipation. However, we believe that this is just a quantitative technical limitation that may be overcome by an appropriate choice of atoms and hi-Q cavities.

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