How to characterize the dynamics of cold atoms in non dissipative optical lattices?
Daniel Hennequin, Philippe Verkerk

To cite this version:
Daniel Hennequin, Philippe Verkerk. How to characterize the dynamics of cold atoms in non dissipative optical lattices?. 2008. hal-00394331v1

HAL Id: hal-00394331
https://hal.archives-ouvertes.fr/hal-00394331v1
Submitted on 11 Jun 2009 (v1), last revised 27 Jan 2010 (v3)

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
How to characterize the dynamics of cold atoms in non dissipative optical lattices?

BY D. HENNEQUIN AND PH. VERKERK

Laboratoire PhLAM, UMR CNRS, CERLA, Université de Lille 1, 59655 Villeneuve d’Asq, France

Abstract. We examine here the classical dynamics of cold atoms in square optical lattices, i.e. lattices obtained with two orthogonal stationary plane waves. Contrary to much of the past studies in this domain, the potential is here time independent and non dissipative. We show that, as a function of the experimental parameters, a very different behavior is obtained, both for the dynamics of atoms trapped inside individual sites, and for atoms travelling between sites: inside the sites, chaos may be a main regime or, on the contrary, may be negligible; outside the sites, chaos sometimes coexists with other regimes. We discuss what are the consequences of these differences on the macroscopic behavior of the atoms in the lattice, and we propose experimental measurements able to characterize these dynamics and to distinguish between the different cases.

1. Introduction

The cooling of atoms to extremely low temperatures, through the use of magneto-optical traps (MOT), opened since the mid eighties fantastic possibilities to increase our experimental knowledge of the quantum world. The most spectacular realization was the achievement of the Bose-Einstein condensation, and thus of macroscopic quantum objects. However, even in the classical world, the possibility to study the dynamics of atoms not “blurred” by the Doppler effect is very exciting. This requires to develop tools to manipulate the atoms, for e.g. guiding them or “designing” their phase space.

Optical lattices provide such tools: their versatility allows to manipulate atoms with an extreme precision and a relative ease (Guidoni & Verkerk 1999). Because of these qualities, they represent an outstanding toy model, and have recently attracted increasing interest in various domains. In statistical physics, cold atoms in optical lattices, through their tunability, made possible the observation of the transition between Gaussian and power-law tail distributions, in particular the Tsallis distributions (Douglas et al. 2006). Condensed matter systems and strongly correlated cold atoms in optical lattices offer deep similarities, as in the superfluid-Mott insulator quantum phase transition (Greiner et al. 2002), in the Tonks-Girardeau regime (Paredes et al. 2004) or for the observation of Anderson localization (Billy et al. 2008, Chabe et al. 2009). In quantum computing, optical lattices appear to be an efficient implementation of a Feynman’s universal quantum simulator (Jaksch & Zoller 2005), and are among the most promising candidates for the realization of a quantum computer (Mandel et al. 2003, Vollbrecht et al. 2004).

Cold atoms appear also to be an ideal model system to study the dynamics of a system in its classical and quantum limits. Indeed, in non dissipative optical lattices, it is possible experimentally to reach the two situations, and even to change quasi continuously from a regime to the other (Steck et al. 2000). Moreover, the extreme
flexibility of the optical lattices makes it possible to imagine a practically infinite number of configurations by varying the complexity of the lattice and the degree of coupling between the atoms and the lattice. Many results have been obtained during these last years in the field of quantum chaos (Steck et al. 2000, Lignier et al. 2005). However, all these works have used very simple potentials, mainly 1D. Chaos is obtained only with a periodic (or quasi-periodic) temporal forcing of the amplitude or frequency of the lattice (Steck et al. 2000, Lignier et al. 2005), and only the temporal dynamics of the individual atoms is studied. The introduction of this external clock and the restriction to 1D potentials reduce considerably the generality of these results and the type of possible dynamics. In particular, the behaviors related to the appearance of new frequencies or to a frequency shift (quasi-periodic and homoclinic bifurcations, for example) are impossible.

If we want to break these limitations, several problems have to be examined: what type of time-independent lattice will lead to a reasonably complex dynamics? What are the relevant quantities to characterize this dynamics? And what are those which can be implemented experimentally? In this paper, we try to give some answers to these questions. In section 2, we give some facts about cold atoms and optical lattices for those who are not familiar with this domain. Section 3 is devoted to the dynamics of atoms inside the wells, while section 4 deals with the dynamics of atoms traveling between several wells. Finally, we discuss in section 5 of the possible implementation of experimental measurements.

2. Context: cold atoms and optical lattices.

Cold atoms refer here to atoms cooled through a magneto-optical trap (MOT). The cooling is mainly obtained through an exchange of the momentum between an atom and a counter-propagating optical beam: while the absorption of a photon by the atom leads to a deceleration of the atom in the direction of the beam, the spontaneous re-emission of the photon arises in a random direction, and so does not change, in average, the atom velocity. To slow down atoms in 3D, three pairs of counter-propagating laser beams are necessary. Obviously, a moving atom is decelerated by the photons traveling in a direction opposite to its own, but is accelerated by the photons traveling in the same direction as its own. But the frequency of these trap beams is detuned to the red of the atomic transition, so that, because of Doppler effect, the front photons are closer to resonance, and thus the deceleration process is more efficient than the acceleration one. This Doppler cooling process is coupled to an inhomogeneous magnetic field, which enhances the cooling process through the Zeeman levels splitting, and adds a restoring force to increase the atomic density of the cloud of cold atoms. MOTs lead typically, for Cesium atoms, to a 2 mm diameter cloud of $10^9$ atoms at $5 \mu K$. Such a cloud of cold atoms can exhibit spatio-temporal instabilities and chaos (Wilkowski et al. 2000, di Stefano et al. 2003, Hennequin 2004, di Stefano et al. 2004), but an adequate choice of the experimental parameters leads to a stable cloud, with atoms whose residual motion is the thermal agitation.

When atoms are dropped in a stationary wave, they undergo a force $F$, the potential $U$ of which is proportional to the wave intensity $I$, and inversely proportional to the detuning $\Delta$ between the wave frequency and the atomic transition

Article submitted to Royal Society
How to characterize the dynamics of cold atoms in non dissipative optical lattices?

Figure 1. a) Layout of the laser beams. b) Spatial distribution of the intensity in the \((X,Y)\) space. Black corresponds to the minimum value (zero intensity), while white corresponds to the maximum. The dotted square delimits the elementary mesh of the lattice, and the white crosses are the saddle points.

frequency:

\[
F = -\nabla U \tag{2.1}
\]

\[
U \propto \frac{I}{\Delta} \tag{2.2}
\]

Thus, atoms accumulate in bright (resp. dark) sites for \(\Delta < 0\) (resp. \(\Delta > 0\)). When the atoms are cooled with the MOT, the atomic density in these optical lattices is small enough to neglect the collisions between atoms, and so the only source of dissipation is the spontaneous emission. As spontaneous emission is proportional to \(I/\Delta^2\), it is relatively easy to build conservative optical lattices.

The atom dynamics in the lattice depends on the dimensionality of the lattice. For example, in a 1D lattice, atoms have only two dynamical degrees of freedom, and thus even if the potential is not harmonic, the dynamics cannot be complex. It is needed to add at least a periodic forcing in such a lattice to observe chaos. On the contrary, a 2D lattice can exhibit chaos, without external forcing.

But the atom dynamics also depends on the lattice geometry, and numerous lattice geometries can be obtained, as e.g. a vertical stack of ring traps (Courtaude \textit{et al.} 2006), five-fold symmetric lattice (Guidoni & Verkerk 1999) or even quasiperiodic lattices (Guidoni \textit{et al.} 1997). In this paper, we will focus on the case of two orthogonal stationary plane waves with the same polarization. The configuration of the laser beams is shown on Fig. 1a. The total field is \(E = \cos kx + e^{i\phi} \sin ky\), where \(x\) and \(y\) are the two space coordinates, \(\phi\) a phase, \(k = 2\pi/\lambda\) the wave vector and \(\lambda\) the wavelength of the laser beam. The intensity can be written as:

\[
I = \cos^2 kx + \cos^2 ky + 2\alpha \cos kx \cos ky \tag{2.3}
\]

where \(\alpha = \cos \phi\). With the adequate normalization, the potential is

\[
U_\pm = \pm I \tag{2.4}
\]

where the explicit sign is that of \(\Delta\). When \(\alpha = 0\), the coupling between \(x\) and \(y\) disappears, and the problem becomes separable. In all the other cases, the coupling between \(x\) and \(y\) could induce complex dynamics. It is easy to see that in these
cases the elementary mesh of the potential is turned of $\pi/4$ as compared to the $(x, y)$ axes, and thus it is natural to introduce the following new coordinates:

$$X = kx + ky$$
$$Y = ky - kx$$

The intensity and the potential can now be written:

$$I = U_+ = -U_- = 1 + \alpha (\cos X + \cos Y) + \cos X \cos Y$$

Before studying the potential, let us concentrate on the intensity. As an example, Fig. 1b shows the spatial distribution of the intensity for $\alpha = 0.5$. The elementary mesh is indicated through the dotted line. Assuming $\alpha > 0$, the intensity $I$ has an absolute maximum $2 (1 + \alpha)$ at coordinates $(n2\pi, m2\pi)$, where $m$ and $n$ are integers. It has also a relative maximum $2 (1 - \alpha)$ in $(\pi + n2\pi, \pi + m2\pi)$. Once again, we see that $\alpha = 0$ is a special case because the absolute and relative maxima have then the same height. Note that $\alpha = 1$ is another special case, where the intensity at the relative maximum vanishes and, thus, is equal to the minimum value. In this special case, we have black lines along $X = \pi + n2\pi$ and $Y = \pi + n2\pi$. We will not consider these cases in the following. On the other hand, the intensity goes to zero in $(\pi + n2\pi, m2\pi)$ and $(n2\pi, \pi + m2\pi)$. Two neighboring zeros are separated by a saddle point where the intensity has the value $I = 1 - \alpha^2$. It is important to note that these saddle points are on the bisectors, connecting on a straight line an absolute maximum to a relative one and again to the next absolute maximum. On the contrary, the saddle points do not stand on the straight line that connect two neighboring zeros. This will induce a huge difference in the dynamics of atoms in the lattice obtained for red detunings ($\Delta < 0$), where the atoms are attracted in high intensity regions and the one for blue detunings ($\Delta > 0$), where the atoms are repelled from these same regions. The bisectors are clearly escape lines for the atoms when $\Delta < 0$, while it is not the case for $\Delta > 0$.

Optical lattices appear to be an exciting tool to study the dynamics of a conservative complex system, but how to characterize this dynamics in the experiments? What are the experimentally accessible quantities? The typical size of a lattice mesh is $\lambda/2$, i.e. 426 nm for Cesium. As the diameter of a cold atom cloud is typically 2 mm, the $10^8$ atoms are dropped in $22 \times 10^6$ sites for a 2D lattice, which lead to 5 atoms/site. At these scales, it is clear that there is no way to isolate an atom, and thus no way to follow its trajectory. Moreover, to see an atom, we need light, and thus the measure introduces a dissipation and destroy the atomic state. A typical measure consists in illuminating the atoms with a laser flash, and recording the fluorescence of the atoms through a camera. This destructive measure gives snapshots of the atom distribution in the space. We examine in the following if it is possible to extract informations about the atom dynamics from this type of measurement.

3. Dynamics of atoms inside the wells

Before we search for signatures of the dynamics in the experimental measurements, let us investigate in more details what are the relevant parameters and characteristics of the atom dynamics in a lattice. To illustrate this approach, let us consider again the two lattices introduced in Section 2. Although these two lattices differ
only by the sign of their potential, they are deeply different. $U_-$ has its wells where the intensity is maximum, while $U_+ $ has its wells where the intensity vanishes. Let us denote $E_T$ the value of the potential energy at the saddle point of the intensity. Atoms, the energy $E$ of which is smaller than the threshold $E_T$, are trapped into one site because they cannot climb up to the saddle point. On the contrary, atoms with $E > E_T$ can travel between sites, if they move in the good direction.

Inside a trap site, the energy of the atom plays the role of a stochastic parameter. Indeed, for low energies, the atoms remain located close to the bottom of the well, and their dynamics can be approximated by a harmonic motion. As the energy increases, the potential becomes more and more anharmonic, the nonlinearities increase, and the dynamics can become more and more complex. To be able to compare the behavior of atoms in different potentials, we take in the following the origin of the energy at the bottom of the wells, and normalize the energy so that $E_T = 1$. The potential energy then takes a different form for red and blue detunings.

\[
U_+ = \frac{I}{1 - \alpha^2} \quad (3.1)
\]

\[
U_- = \frac{2(1 + \alpha) - I}{(1 + \alpha)^2} \quad (3.2)
\]

Let us now examine in details the dynamics of the atoms in our two potentials. The most relevant way is to look at the evolution of the Poincaré sections as a function of the energy. Our phase space is 4-dimensional, with directions $\left(X, Y, \dot{X}, \dot{Y}\right)$, but because of the energy conservation, the accessible space reduces to a 3D surface. We choose to consider Poincaré section at $Y = 0$ with increasing values, and thus, Poincaré sections are in the 3D space $\left(X, \dot{X}, \dot{Y}\right)$, and they lie on a 2D surface $S_p$, which has the shape looking like a semi-ellipsoid. To represent the Poincaré sections we can project them on the $(X, Y)$ plane or on the more usual $(X, \dot{X})$ plane. The latter shows the Poincaré sections viewed from the vertex of the semi-ellipsoid. However here, because of the stiff sides of $S_p$, the projection in this plane leads to a confuse map, as many curves are projected at the same location, and thus are superimposed. On the contrary, the projection on the $(X, Y)$ plane gives more details, and thus in the following, we often choose it. However, let’s keep in mind that we look at a lateral projection of a “bell”, and thus that we superimpose its front and rear faces.

As pointed out before, because of the normalization we choose for the energy, the form of the potential energy differs in the cases of blue or red detuned lasers. We investigate each cases separately.

In the case of red detuned lasers, the potential energy takes the form:

\[
U_- = \omega_0^2 (1 - \cos X) + \omega_0^2 (1 - \cos Y) - \frac{(1 - \cos X)(1 - \cos Y)}{(1 + \alpha)^2} \quad (3.3)
\]

with $\omega_0^2 = (1 + \alpha)^{-1} \quad (3.4)$

This potential appears to be the sum of two simple pendula coupled through the third term. The frequency for oscillations with small amplitude is the same.
for the two directions. This degeneracy together with the coupling term leads to a strong synchronization of the motion in the two directions (Beaute et al. 2002). However, in contrast with the Huygens clocks, we do not have any dissipation here, so a frequency locking cannot occurs.

However, it is interesting to identify the resonances of the system. A very simple approach is to restrict the problem to the first anharmonic terms, similar to the undamped Duffing oscillator. We then look for a periodic harmonic solution in the form \( X = X_0 \cos(\omega t) \) and \( Y = Y_0 \cos(\omega t + \varphi) \), with \( \omega \) close to \( \omega_0 \). We drop terms at other frequencies (i.e. \( 3\omega \)) and we end up with six families of solutions: the first two are the trivial ones: motion along the \( X \) or the \( Y \) directions (\( Y_0 = 0 \) or \( X_0 = 0 \)). The other four are obtained for \( X_0 = Y_0 \) and for \( \varphi = 0, \pi, \pm \pi/2 \). For a given energy \( E \), the relations giving \( X_0 \) and \( \omega \) are not simple, and it is beyond the aim of this article to write them explicitly. For the large amplitudes considered in the following, the motion is not any more harmonic and we cannot keep only the lower order terms, but the main result remains: we have six periodic trajectories, leading to points in the Poincaré section (except for the trajectory \( Y_0 = 0 \), that we cannot catch in a Poincaré section at \( Y = 0 \)). In the 3D space, these points have the coordinates \((0, -Y_0, 0), (\pm X_0, -X_0, 0)\) and \((0, -X'_0, \pm \omega X'_0)\).

In Fig. 2, we show the dynamics in the \( U_- \) potential for different normalized energies in the case of \( \alpha = 0.5 \). These results have been obtained through numerical resolution of the equations of motion which are derived from the potential (3.3), without the addition of any random quantity. All the described behaviors are thus deterministic. For each value of the energy, we project the Poincaré section on the \((X, Y)\) plane (top figures) and on the \((X, \dot{X})\) plane (bottom figures). For low enough energies (e.g. \( E = 0.8 \) in Fig. 2a and 2b), we see four distinct domains separated by an X-shaped separatrix. In each of these domains, the Poincaré section is cycling around one of the non-trivial resonances found above. As the motion along \( X \) and \( Y \) is governed by the same frequency, and because of the coupling between these two pendula, a synchronization between the two directions occurs, through a phase locking between the two motions. The corresponding behavior can be described as mainly a \( \omega \) periodic cycle perturbed by small sidebands.

The dynamics in \( U_- \) evolves only slightly when \( E \) is increased. The Poincaré surfaces are always organized around the separatrix delimiting 4 areas. In each area, the nature of the motion is the same, namely phase locking between the motions in the \( X \) and \( Y \) directions. Chaos appears close to the separatrix for \( E \simeq 0.88 \) (Fig. 2c and 2d), but it remains marginal, even when \( E = 1 \) (Fig. 2e and 2f). Its very small development is probably due to the original degeneracy of the frequencies of the coupled pendula and to the strong coupling between them.

For blue detunings (\( \Delta > 0 \)), the bottom of the well corresponds to \( I = 0 \), i.e. \((X = 0, Y = \pi)\) sites. For the sake of simplicity, we shift the origin in \( Y \) by \( \pi \), in order to have a trapped motion centered at the origin. Thus, we can write:

\[
U_+ = \omega^2_{0X}(1 - \cos X) + \omega^2_{0Y}(1 - \cos Y) - \frac{(1 - \cos X)(1 - \cos Y)}{(1 - \alpha^2)} \tag{3.5}
\]

with \( \omega^2_{0X} = (1 + \alpha)^{-1} \quad \text{(3.6)} \)

\( \omega^2_{0Y} = (1 - \alpha)^{-1} \quad \text{(3.7)} \)

\[ Article \ submitted \ to \ Royal \ Society \]
Once again, this potential appears to be the sum of two coupled pendula. But now, the two frequencies for oscillations with small amplitudes are different: for the value $\alpha = 0.5$ chosen here, the ratio $\sqrt{3}$ of these two frequencies is irrational.

For very small energies (Fig. 3a), the dynamics consists essentially in a regular motion around the bottom of the well, along a quasiperiodic trajectory with frequencies $\omega_X$ and $\omega_Y$ close to $\omega_{0X}$ and $\omega_{0Y}$. At the top of Fig. 3a, Poincaré sections are those of atoms, the motion of which is essentially along the $X$ axis. In $Y = 0$, the trajectory is a periodic cycle along the $X$ direction (edge of the semi-ellipsoid). At the opposite, the periodic cycle at the bottom of the figure corresponds to the situation where the atomic motion is exclusively along the $Y$ axis (vertex of the semi-ellipsoid). Note that the nature of the motion along these quasiperiodic cycles is deeply different from those described with $\Delta < 0$. Indeed, as $\omega_{0X}$ and $\omega_{0Y}$ are very different, no locking occurs. In particular, in the spectrum of the motion, the two main frequencies are close to $\omega_{0X}$ and $\omega_{0Y}$.

As the energy of the atom is increased, the atom can climb more and more in the well, the frequencies $\omega_X$ and $\omega_Y$ change because of the anharmonicity of the

Figure 2. $(X, Y)$ (top) and $(X, \dot{X})$ (bottom) Poincaré sections of the atomic dynamics in the $U_-$ potential. (a) and (b): $E = 0.80$; (c) and (d): $E = 0.88$; (e) and (f): $E = 1.00$. 

Article submitted to Royal Society
potential, but the dynamics does not change fundamentally until $E \simeq 0.6$. At that point a new feature appears: a stable periodic trajectory shows up as a cycle close to the bottom of Fig. 3b, obtained for $E = 0.63$. In fact, for amplitudes large enough, the frequencies $\omega_X$ and $\omega_Y$ depart so much from their initial values $\omega_{0X}$ and $\omega_{0Y}$ that a new resonance appears at $\omega_Y = 2\omega_X$.

For higher energies, the $\omega_Y = 2\omega_X$ resonance grows and comes closer to the centre of the figure and influence a non-negligible fraction of the trajectories. In Fig. 3c, for $E = 0.8$, the resonance is clearly visible in $Y \simeq -0.74$. In the $(X,Y,\dot{X})$ space, its Poincaré section consists in 2 points (superimposed in the projection of fig. 3c), explored alternatively by the trajectory. Around this point, the Poincaré sections are a double closed loop. The corresponding quasiperiodic motion consists in a perturbed $\omega_Y = 2\omega_X$ phase locked periodic cycle, where the perturbation consists in small sidebands of $\omega_X$ and $\omega_Y$ in the spectrum. Thus the separatrix appears here to be the limit between this phase locked and the unlocked behaviors. The central domain and the two linked lateral domains (bottom left and right) correspond to the phase locking. The difference between these two domains is the
relative phase on the motion along $X$ and along $Y$. In the two other domains (top and bottom), there is no locking between the $\omega_X$ and $\omega_Y$ frequencies.

In $E = 0.8$ (Fig. 3c), all the trajectories are still periodic cycles or quasiperiodic tori. When the energy is increased further, chaos appears at $E \approx 0.88$, starting in the vicinity of the separatrix (Fig. 3d). Then, it expands with some quasiperiodic islands remaining (Fig. 3e), but finally, for $E = 1$ (Fig. 3f), the only significant quasiperiodic domains are those around the $X$ and $Y$ periodic cycles. Around the locked periodic cycles, a narrow area with tori remains, but chaos appears really to be dominant.

We have shown in this section that it is relatively easy to find two slightly different lattices with fundamentally different dynamics. These two configurations are easy to reach experimentally, as they differ only by the sign of the detuning. It would be interesting now to examine how to measure experimentally these differences, and if these differences have an impact on the dynamics of atoms when they jump between sites of the lattice. The next section deals with the latter.

### 4. Dynamics of atoms visiting several wells

To travel from site to site, an atom needs to have an energy $E \geq 1$, but this is not a sufficient condition. Only atoms with an adequate trajectory will effectively escape from a well. This implies that for a given energy $E \geq 1$, at least two classes of atoms can exist: trapped atoms remaining in a single well, and traveling atoms, which escape the wells. In fact, the situation is more complex, as we will see now.

Let us first examine the dynamics of traveling atoms in the blue case ($\Delta > 0$). We are here interested by atoms with an energy $1 < E < 4$. Indeed, atoms with $E > 4$ have an energy larger than the potential maximum, and thus they “fly” above the potential, and their trajectory is purely ballistic. On the contrary, the dynamics of the atoms with an intermediate energy consists in complex trajectories visiting a large number of sites, as in a random walk. As our model is fully deterministic, it involves in fact chaotic trajectories. Fig. 4a illustrates such a chaotic diffusion: it reports the trajectories followed by 100 atoms. Such a trajectory is in fact an...
alternation of oscillations inside wells and of jumps between wells. Here, we know that chaos dominates inside the wells, and thus the chaotic nature of the diffusion is not surprising. However, as we will see below, the existence of chaos inside the wells is not a necessary condition to observe a chaotic diffusion.

To think of an experimental characterization of this chaotic diffusion, a simple way would be to characterize the diffusion function, and to evaluate a diffusion coefficient. Fig. 5a reports the distance covered by atoms as a function of time. There are clearly two groups of atoms: trapped atoms remain within a short distance (smaller than the mesh, i.e. $2\pi$) of their initial location, while diffusing atoms move away to distances of the order of $10^5$. Although there is small dependence of these curves as a function of the energy of the atom, the orders of magnitude remain the same for all energies $1 < E < 4$.

In the red detuned situation ($\Delta < 0$), the maximum of the potential is at $E = 1.33$. As in the blue case, atoms with an energy $E > 1.33$ have ballistic trajectories, and atoms with $1 < E < 1.33$ exhibit a diffusive chaotic behavior (Fig. 4b). The origin of chaos now is clearly in the jumps between wells, as the dynamics in the wells is regular. And in fact, there is a main difference as compared to the blue case: the diffusion scale is larger by one order of magnitude, on the whole interval $1 < E < 1.33$. We did not check if the slower diffusion originates effectively in the chaotic trajectories followed by the atoms inside the wells, but it would be interesting to check in a future work how these chaotic behavior could slow down the atoms. However, the difference of one order of magnitude in the diffusion speed reveals that the macroscopic behavior of atoms could effectively be used to characterize the nature of the dynamics in optical lattices.

But there is another important difference between the two lattices: in the red case a third regime exists, neither trapping neither diffusing. It is illustrated on Fig. 4b, where trajectories appear following the two bisectors. These trajectories correspond to atoms traveling along the escape lines of the lattice, as they were described in section 2. These atoms follow in fact a ballistic trajectory, where they travel very rapidly along the bisectors. For example, in Fig. 4b, the ballistic trajectories reach $10^6$ in all directions, while the diffusive atoms reach only $2 \times 10^4$ of the same units in the same time. Note that the ballistic trajectories we discuss here occur as soon as the threshold $E = 1$ is reach, and only along the escape lines of the potential.

Fig. 5b shows the distance covered by the atoms as a function of time. We have now clearly three groups of trajectories: trapped trajectories at bottom, diffusive trajectories for distances of about $10^4$, and ballistic trajectories at the top, for distances larger than $10^5$. The main difference as compared to the $\Delta > 0$ case is the cohabitation of ballistic and diffusive trajectories, even just above threshold. This put in evidence three specific time scales of the dynamics of atoms with a given energy, associated respectively with the trapped, the chaotic diffusive and the ballistic trajectories.

In this section, we examined the dynamics of atoms, whose energy is large enough to escape the potential wells, but remains smaller than the potential maxima. We focused on atoms traveling between wells, and found a different behavior for our two lattices. For the red lattice, atoms can be classified following two types of dynamics: the diffusive atoms exhibit a chaotic dynamics carrying them off their initial location; the ballistic atoms move away rapidly from their initial location.
These behaviors are associated with two different time scale. But is it sufficient to identify these different regimes in a real experiment? We have also shown that the dynamics of atoms in the blue lattice is quite different, both for the diffusive regime and the ballistic one: the time scale of the former is one order of magnitude smaller, while the latter simply does not exist. Can we use these properties to characterize and distinguish experimentally the two lattices? These questions are discussed in the next section.

5. Macroscopic signatures of chaos

Our aim is to characterize the dynamics of the cold atoms in the optical lattice. As we are concerned by conservative lattices, we cannot hope to “film” in real time the atoms in the lattice, as it would introduce dissipation. Thus we have to find other techniques. As the specificity of each lattice concerns the traveling atoms, an experimental measurement aiming at characterizing these lattices should characterize these traveling atoms.

Experimentally, the lattice is finite. So the traveling atoms will reach the edge of the lattice, and finally leave the lattice. Therefore a simple measure of the lifetime of the atoms in the lattice give informations about the trapped and traveling atoms. However, as there are several types of traveling atoms, the simple measure of a lifetime is not sufficient, and the lifetime curve itself, in particular its shape, must be analyzed. Thus we will plot now the number of atoms in the lattice as a function of the time. The shape of the curve and the lifetime itself should give informations about the traveling atoms, while the baseline gives the percentage of trapped atoms.

In the experiment, all the atoms have not the same energy, but on the contrary, they exhibit a distribution of energies linked to their temperature. Thus the results shown below have been obtained by using a sample of atoms with an appropriate distribution of energy.

Fig. 6a shows the number of atoms in the blue lattice versus time. To simulate the finite size of the lattice, atoms are removed as soon as they reach a distance $D_L = 1000$. The curve exhibits a plateau at short times, followed by an exponential-like decreasing to an asymptote. The plateau corresponds to the time needed by the first atoms to reach the edge of the lattice (all the atoms are supposed to be initially at the center of the lattice). The decreasing corresponds to the diffusing atoms escaping the lattice, and the asymptote to the number of atoms trapped in wells. This behavior does not depend on the lattice size $D_L$, except that the life time of atoms increases. In fact, the distance $D_L = 1000$, i.e. about 150 lattice
meshes or 70 μm for a Cs trap, is smaller by one order of magnitude than a typical experimental realization. However, a value of $D_L = 10^4$ leads, for the data presented in Fig. 4a and 5a, to an almost flat curve, because the time series are not long enough. To reach such a distance, one should increase the evolution time by two orders of magnitude.

Fig. 6b shows the number of atoms in the red lattice versus time, for $D_L = 10^4$. The shape of the curve is qualitatively different from that obtained for the blue lattice. At short times, a fast decreasing appears, corresponding to the loss of the ballistic atoms. At large times, not visible on the figure, an asymptote is reached, corresponding to the trapped atoms. The intermediate decreasing correspond to the loss of the diffusing atoms. Note that the decreasing appears to be more or less linear. In fact, the shape of this part of the curves is the sum of the diffusing losses of different classes of atoms differing by their energy. As a function of $D_L$, this sum can exhibit very different shapes, from an exponential-like shape, as in Fig. 6a, to an almost linear shape, as in Fig. 6b.

Fig. 6 shows that the measure of the lifetime of atoms in a conservative optical lattice provides qualitative and quantitative informations about the nature of the lattice and the nature of the dynamics of the atoms in the lattice, in particular about the chaotic diffusion. Therefore the measure of the atom lifetime, in particular the existence of several characteristic times in the decreasing of the atom number, appears to be a signature of the chaotic dynamics of atoms in the lattice.

6. Conclusion

We have shown in this paper that optical lattices are a good toy model to study experimentally the dynamics of conservative systems, provided that relevant experimental measures are found to characterize this dynamics. In particular, we show that changing a simple experimental parameter can lead to two deeply different lattices, where atoms exhibit very different dynamical behaviors. We have shown that these differences exist both in the local dynamics of atoms inside a well, and in the non local dynamics of atoms traveling between wells. We searched numerically for signatures of these different dynamics in the experimentally accessible quanti-
ties, and found that the measure of atom lifetimes in the lattice gives numerous informations about the existence and the type of chaotic diffusion of the atoms.

It would be interesting now to characterize more precisely the diffusion function, as a function of the experimental parameters, in particular the atom temperature and the lattice size, and obviously to test these results on a real experiment.

We thank Frédéric Carlier for fruitful discussions. The Laboratoire de Physique des Lasers, Atomes et Molécules is Unité Mixte de Recherche de l’Université de Lille I et du CNRS (UMR 8523). The Centre d’Études et de Recherches Lasers et Applications (CERLA) is supported by the Ministère chargé de la Recherche, the Région Nord-Pas de Calais and the Fonds Européen de Développement Économique des Régions.

References


Article submitted to Royal Society