Control of atomic currents using a quantum stirring device

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Abstract – We propose a BEC stirring device which can be regarded as the incorporation of a quantum pump into a closed circuit: it produces a DC circulating current in response to a cyclic adiabatic change of two control parameters of an optical trap. We demonstrate the feasibility of this concept and point out that such device can be utilized in order to probe the interatomic interactions.

The realization of Bose-Einstein Condensation (BEC) of ultra-cold atoms in optical lattices and atom chips [1] and the availability of conveyor belts [2] are considered to be a major breakthrough with potential applications in the arena of quantum information processing [3], atom interferometry [4–6], lasers [1,7] and atom diodes and transistors [8–10]. A major advantage of BEC-based devices, as compared to conventional solid-state structures, lies in the extraordinary degree of precision and control that is available, regarding not only the confining potential, but also the strength of the interatomic interactions, their preparation and the measurement of the atomic cloud. Accordingly it is envisioned that the emerging field of “atomtronics” will provide a new generation of nanoscale devices.

The possibility to induce DC currents by periodic (AC) modulation of the potential is familiar from the context of electronic devices. If an open geometry is concerned, it is known as “quantum pumping” [11], while for closed geometries we use the term “quantum stirring” [12]. We consider below the stirring of condensed ultra-cold atoms due to the periodic variation of the on-site potentials and of the tunneling rates between adjacent confining traps. We show that the nature of the transport process depends crucially on the sign and on the strength of the interatomic interactions. We distinguish between four regimes of dynamical behavior: For strong repulsive interaction the particles are transported one by one, which we call sequential crossing; for weaker repulsive interaction we observe either gradual crossing or coherent mega crossing; finally, for strong attractive interaction the particles move together as one composite unit from trap to trap.

Model. – The simplest model that captures the physics of quantum stirring is the three-site Bose-Hubbard Hamiltonian (BHH) [10,13–15] (see fig. 1)\textsuperscript{1}. We call the $i=0$ site “gated-well” since we late assume that we have control over its potential energy $v_0 = \varepsilon$. The $i=1,2$ sites form a two level “double-well” with potential energy $v_1 = v_2 = 0$.

\textsuperscript{1}The three-site BHH is a prototype system for many recent studies [13]. A classical analysis of this system was performed in [15] where it was shown that for appropriate system parameters and initial conditions, chaotic dynamics would emerge. In this work we consider adiabatic driving of ground-state preparation and therefore chaotic motion is not an issue.
The $N$ boson BHH is

$$\hat{\mathcal{H}} = \sum_{i=0}^{2} \left[ v_i \hat{n}_i + \frac{U}{2} \hat{n}_i (\hat{n}_i - 1) - k_i \hat{b}_i^\dagger \hat{b}_i - \sum_{i'=1,2} k_{i'} \hat{b}_{i'}^\dagger \hat{b}_0 + \text{h.c.} \right]$$

(1)

We set $\hbar = 1$ which corresponds to measuring energies in units of frequency. Furthermore, without loss of generality we choose time units such that $k_c = 1$. Accordingly the two single-particle levels of the double-well are $\varepsilon_{\pm} = \pm 1$. The annihilation and creation operators $\hat{b}_i$ and $\hat{b}_i^\dagger$ obey the canonical commutation relations $[\hat{b}_i, \hat{b}_{i'}^\dagger] = \delta_{i,j}$, while the operators $\hat{n}_i = \hat{b}_i^\dagger \hat{b}_i$ count the number of bosons at site $i$. The interaction strength between two atoms in a single site is given by $U = 4\pi a_s V_{\text{eff}}/m$, where $V_{\text{eff}}$ is the effective volume, $m$ is the atomic mass, and $a_s$ is the $s$-wave scattering length which can be changed by applying an additional magnetic field [16]. The on-site potentials $v_i$ as well as the coupling strengths $k_i$ are controlled by changing the confining potential. The main assumption underlying the BHH is that the single-particle ground-state wave functions are sufficiently localized at the sites, and that for the temperatures involved they are well separated in energy from the excited single-particle levels. Experimentally, such deep trapping potentials can accommodate several hundred particles [17].

The couplings between the gated-well and the two ends of the double-well are $k_1$ and $k_2$. We assume that both are much smaller than $k_c$ (for the two-mode BEC dynamics see, for example, [18]). It is convenient to define the two control parameters of the pumping as $X_1 = (1/k_2) - (1/k_1)$ and $X_2 = \varepsilon$. By periodic cycling of the parameters $(X_1, X_2)$ we can obtain a non-zero amount ($Q$) of transported atoms per cycle. The pumping cycle is illustrated in figs. 1 and 2. Initially all particles are located in the gated-well which has a sufficiently negative on-site potential energy ($X_2 < 0$). In the first half of the cycle the coupling is biased in favor of the $k_1$ route ($X_1 > 0$), while $X_2$ is raised until (say) the gated-well is empty$^2$. In the second half of the cycle the coupling is biased in favor of the $k_2$ route ($X_1 < 0$), while $X_2$ is lowered until the gated-well is full. Assuming $U = 0$, the gated-well is depopulated via the $k_1$ route into the lower energy level $\varepsilon_-$ during the first half of the cycle, and re-populated via the $k_2$ route during the second half of the cycle. Accordingly the net effect is to have a non-zero $Q$. If we had a single particle in the system, the net effect would be to pump roughly one particle per cycle. If we have $N$ non-interacting particles, the result of the same cycle is to pump roughly $N$ particles per cycle$^3$. We would like to know: what is the actual result using a proper quantum-mechanical calculation?

And furthermore we would like to investigate what the effect of the interatomic interaction $U$ on the result is.

**Methods.** – Within the framework of linear response theory the induced current is $I = -G_1X_1$ if we change $X_1$ and $I = -G_2X_2$ if we change $X_2$. The coefficients $G_1$ and $G_2$ in these linear relations are defined as the elements of the geometric-conductance matrix, and can be calculated using the Kubo formula approach (see below). Integrating the current over a full cycle we get

$$Q = \oint \text{cycle} \, I \, dt = -\oint (G_1 \, dX_1 + G_2 \, dX_2).$$

(2)

In order to calculate the geometric conductance we use the Kubo formula approach to quantum pumping [19] which is based on the theory of adiabatic processes [20]. It turns out that in the strict adiabatic limit, $G$ is related to the vector field $B$ in the theory of the Berry phase which is known as the “Berry Curvature”. The adiabatic slowness condition on $\dot{X}$ in the present context, taking into account the two-orbital approximation, is discussed in sect. 4 of [21].

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$^2$Later we estimate the required $X_2$ variation in order to have all the particles transferred from the gated-well to the double-well. We also analyze smaller pumping cycles (see fig. 3b) for which only a fraction of particles gets through.

$^3$Through the driving cycle the total number of bosons remains constant. The energy is not a constant of motion, but in the adiabatic limit the system comes back to the same state at the end of each cycle.
the notations $B_1 = -G_2$ and $B_2 = G_1$ it is illuminating to rewrite eq. (2) as

$$Q = \oint B \cdot d\vec{s},$$

(3)

where we define the normal vector $d\vec{s} = (dX_2, -dX_1)$ as illustrated in fig. 2. The calculation of the so-called Kubo-Berry Curvature is done using the following formula [19]:

$$B_j = \sum_{n \neq n_0} \frac{2 \text{Im}[\mathcal{V}_{n_0 n}] \mathcal{F}_j^{n_0 n}}{(E_n - E_{n_0})^2}.$$  

(4)

Above $I = \int \mathcal{S} \mathcal{H} \mathcal{S}^{-1}$ is the averaged current via the 0\rightarrow 1 and 2\rightarrow 0, while $\mathcal{F}_j = -\partial \mathcal{H} / \partial X_j$ is the generalized force associated with the control parameter $X_j$. The index $n$ labels the eigenstates of the many-body Hamiltonian. We assume from now on that $n_0$ is the BEC ground state.

The advantage of the above, so-called “geometric” point of view is in the intuition that it gives for the result: Formally the field $B$ is like a projection of a fictitious magnetic field in an embedding three-dimensional $X$ space (the third coordinate $X_3$ if formally defined such that $I = -\partial \mathcal{H} / \partial X_3$). The flux of this fictitious magnetic field through any out-of-plane surface which is enclosed by the pumping cycle gives the so-called Berry phase, while the line integral over this fictitious magnetic field, i.e. eq. (3)$^4$, gives $Q$. As implied by inspection of eq. (4) the sources of $B$ are located at points where the ground level $n_0$ has a degeneracy with the next level. A simple argument implies that this “magnetic charge” is quantized like Dirac monopoles, else the Berry phase would be ill defined. For details see [19]. In our model system for $U = 0$ all the “magnetic charge” is concentrated in one point. As the interaction $U$ becomes larger the $(N + 1)$-fold degeneracy of the levels is lifted, and this “magnetic charge” disintegrates into $N$ elementary “monopoles” (see fig. 2). We further discuss the energy spectrum in the next section.

Regimes. – We define the average coupling as $\kappa = (k_1 + k_2)/\sqrt{2}$. In the zeroth-order approximation $k_1$ and $k_2$ are neglected, and later we take them into account as a perturbation. For $\kappa = 0$ the number $(n)$ of particles in the gated-well becomes a good quantum number hence we can associate the level index $n$ with the number of particles in the gated-well. Furthermore, we adopt a “two-orbital approximation”$^4$: we assume that there is non-zero occupation only in the gated-well and in the lower double-well level, which is valid if $NU \ll k_c$. Note also that we assume an adiabatic process$^3$, and accordingly

$^4$Due to the continuity equation the same result for $Q$ is obtained irrespective of whether we measure the current via the 0\rightarrow 1 bond or via the 2\rightarrow 0 bond. The advantage of using the “symmetrized” version for $Z$ is that the same amount of particles is being transported during both halves of the cycle, allowing us to focus on (say) the first half and then double the result.

$^3$In the $X_3 = 0$ plane the third component of the fictitious magnetic field is zero due to the time-reversal symmetry [19].
\[ \kappa_n = (n-1) |\mathcal{H}|n\] 

The calculation involves the matrix elements of \( b_i^\dagger b_0 \), leading to \( \kappa_n = [(N + 1 - n)n]^{1/2} \). An analogous expression applies to the current operator where \( \kappa \) is replaced by \( (k_1 - k_2)/\sqrt{2} \). For large \( U \), as \( \varepsilon \) is varied, we encounter (say for \( N = 3 \)) a sequence of distinct Landau-Zener transitions \((|3\rangle \rightarrow |2\rangle \rightarrow |1\rangle \rightarrow |0\rangle)\). The distance between avoided crossings is of order \( U \), while their width is \( \delta \varepsilon_n = \kappa_n \). The widest crossings are at the center with \( \delta \varepsilon_n \sim N \kappa \). This should be contrasted with the energy scales \( U \) and \( NU \) that describe the span of the crossings. Accordingly we deduce that for repulsive interaction there are three distinct regimes: for \( U \ll \kappa/N \) we have a “mega crossing”; for \( U \gg \kappa \) we have the “sequential-crossing regime”, while in the intermediate regime we have a “gradual crossing”. We observe that the regime of behavior depends on the ratio \( \kappa/U \). If \( N \) is not too large one can resolve a sequence of two-level crossings. Below we summarize the results in the various regimes. In particular, eq. (7) is obtained from eq. (4) with a two-level approximation for each crossing. We also related briefly to the \( U < 0 \) regime.

**Results.** – Most of the contribution to the line integral in eq. (2) comes as we change \( X_2 \) during the avoided crossings that have been discussed in the previous section. If we close the pumping cycle outside of this limited range, then the \( X_1 \) variation can be safely neglected. Accordingly we refer from now on to \( G = G \) only. An overview of the numerical results for the conductance is shown in fig. 3, where we plot \( G \) as a function of \( X_2 \) for various interaction strengths \( U \). In the same figure we report the normalized amount of particles \( Q \) for various driving cycles. As the shape of \( G \) changes, the dependence of \( Q \) on the \( X_2 \) span of the pumping cycle becomes of importance. Thus, by measuring \( Q \) we obtain information on the strength of the interatomic interactions.

In fig. 4 more details are presented: besides \( G \) we also plot the \( X_2 \)-dependence of the energy levels, and of the site population. Four representative values of \( U \) are considered including also the \( U < 0 \) case. Let us discuss the observed results. For \( U = 0 \) all the particles cross “together” from the gated-well orbital to the \( \varepsilon \)– double-well orbital. We call this type of dynamics “mega crossing”. The outcome is just \( N \) times the single-particle result:

\[ G = \frac{1}{N} \frac{(k_1^2 - k_2^2)/2}{[(\varepsilon - \varepsilon_n)^2 + 2(k_1 + k_2)^2]^{3/2}}, \] (5)

which can be expressed in terms of the control parameters \((X_1, X_2)\). This result approximately holds as long as \( U \ll \kappa/N \). Integrating over a full cycle one obtains

\[ Q = N \frac{1 + (\kappa R)^2}{[1 + (\kappa R)^2]^{1/2} - 1} \frac{\kappa R}{\kappa}, \] (6)

where \( R \) is the radius of the pumping cycle (see fig. 2). For small cycles we get \( Q \approx N \kappa R/2 \), while for large cycles we get the limiting value \( Q \approx N \). In the other extreme, for very repulsive interaction \((U \gg \kappa/N)\) we get

\[ G = \left( \frac{k_1 - k_2}{k_1 + k_2} \right) \sum_{n=1}^{N} \frac{(\delta \varepsilon_n)^2}{[(\varepsilon - \varepsilon_n)^2 + (2 \delta \varepsilon_n)^2]^{3/2}} \] (7)

For intermediate values \( U \in [\kappa/N, N \kappa] \), we find neither the sequential crossing of eq. (7), nor the mega crossing.

![Fig. 4: (Color online) Evolution of the energy levels, the site occupation and the conductance. Further details relating to the data of fig. 3. We refer to four representative values of \( U \), which are indicated on top of each set of panels. Upper panels: the lowest \( N + 1 \) energy levels \( E_n \) which dominate the conductance \( G \) are plotted as a function of \( X_2 = \varepsilon \). The insets represent magnifications of the indicated areas. Middle panels: the site occupations \( n_0 \) (blue △), \( n_1 \) (black ○), \( n_2 \) (red □). Lower panels: the corresponding conductance \( G \) as a function of \( \varepsilon \). Numerical results are represented by solid black lines, while the dotted red line corresponds to the analytical result (5) in (b) and to (7) in (c), (d).](image)
of eq. (5), but rather a gradual crossing. Namely, in this regime, over a range $\Delta X_2 = (3/2)(N - 1)U$ we get a constant geometric conductance:

$$G \approx -\frac{k_1 - k_2}{k_1 + k_2} \frac{1}{3U}$$

(8)

which reflects in a simple way the interaction strength. This formula was deduced by extrapolating eq. (7) and then was validated numerically (lower panel of fig. 4c).

As discussed above for large positive $U$ the $(N+1)$-fold “degeneracy” of the $U = 0$ Landau-Zener crossing is lifted, and we get a sequence of $N$ Landau-Zener crossings (for schematic illustration see the lower panel of fig. 2, and compare with the numerical results in the upper panels of fig. 4). Also for $U < 0$ this $(N+1)$-fold “degeneracy” is lifted, but in a different way: the levels separate in the “vertical” (energy) direction rather than “horizontally” (see upper panels of fig. 4). Accordingly all the particles execute a single two-level transition from the gated-well to the double-well (see fig. 4a). This direct Landau-Zener transition from the $n = 0$ level to the $n = 0$ level is very sharp because it is mediated by an $N$-th order virtual transition via the intermediate $n$ states. Accordingly, for sufficiently strong attractive interaction all the particles move together from the gated-well to one of the double-well sites. When the sign of $X_1$ is reversed they are transported from one end of the double-well to the other end (not shown). This should be clearly distinguished from the $(N+1)$-fold degenerated transition to the lower double-well level which is observed in the $U = 0$ case.

Summary. – The theoretical [10,18,22] and experimental [23] study of driven dynamics in single- and double-site systems is the state of the art. Study of three-site systems adds the exciting topological aspect: controlled atomic current can be induced using optical-lattice technology [24]. The actual measurement of induced neutral currents poses a challenge to experimentalists. In fact there is a variety of techniques that have been proposed for this purpose. For example one can exploit the Doppler effect in the perpendicular direction, which is known as the rotational frequency shift [25]. The analysis of the prototype trimer system reveals the crucial importance of interactions. The interactions are not merely a perturbation: rather they determine the nature of the transport process. We expect the induced circulating atomic current to be extremely accurate, which would open the way to various applications, either as a new metrological standard, or as a component of a new type of quantum information or processing device.

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