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Simplified landscapes for optimization of shaken lattice interferometry

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PAPER

Simplified landscapes for optimization of shaken lattice interferometry

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¹ Author to whom any correspondence should be addressed.E-mail: dana@jila.colorado.edu**Keywords:** atom interferometry, optimization, optical lattices, quantum control

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Abstract

Motivated by recent results using shaken optical lattices to perform atom interferometry, we explore the splitting of an atom cloud trapped in a phase-modulated ('shaken') optical lattice. Using a simple analytic model we are able to show that we can obtain the simplest case of $\pm 2\hbar k_L$ splitting via single-frequency shaking. This is confirmed both via simulation and experiment. Furthermore, we are able to split with a relative phase θ between the two split arms of 0 or π depending on our shaking frequency. Addressing higher-order splitting, we determine that $\pm 6\hbar k_L$ splitting is sufficient to be able to accelerate the atoms in counterpropagating lattices. Finally, we show that we can use a genetic algorithm to optimize $\pm 4\hbar k_L$ and $\pm 6\hbar k_L$ splitting to within $\approx 0.1\%$ by restricting our optimization to the resonance frequencies corresponding to single- and two-photon transitions between Bloch bands. As a proof-of-principle, an experimental demonstration of simplified optimization of $4\hbar k_L$ splitting is presented.

1. Introduction

The control of quantum mechanical systems is of interest in a variety of applications, among them quantum computing and atom interferometry. The pioneering work in [1, 2] showed that by genetic optimization of the lattice phase modulation (or 'shaking') function, one can precisely control the atoms' final state after shaking. We extended this idea to perform so-called 'shaken lattice interferometry' in which the quantized momentum states of the atoms trapped in a shallow optical lattice were transformed and made to undergo a conventional interferometry sequence of splitting, propagation, reflection, reverse propagation, and recombination [3, 4]. Optical lattices have been used for atom interferometry in Raman- or Bragg-based light-pulse schemes [5] and to accelerate interrogated atoms using Bloch oscillations [6]. In a Michelson configuration [7], one-dimensional shaken lattice interferometry was shown to have a sensitivity to applied acceleration that scales as the square of the interrogation time T_1 . Furthermore, its sensitivity can be tuned to the signal of interest (e.g. an AC acceleration signal). Atoms have been held in amplitude-modulated lattices for times on the order of tens of seconds in order to precisely measure gravity [8]. A shaken lattice interferometer thus has the possibility of achieving similar interrogation times.

In this paper we take a different approach than the usual Floquet analysis [9–11] to the dynamics of a shaken lattice system. We wish to explore in detail how one can shake an optical lattice to transform the wavefunction of atoms trapped in the lattice. Specifically, we seek to reduce the dimensionality of the shaking control landscape. The motivation for this is twofold: first, by simplifying the optimization landscape we can improve the efficiency of learning [12]. This is particularly important in experiments limited by drift or finite cycle times. Second, we wish to limit heating and decoherence in the shaken lattice system. Recent work has shown that atoms in a shaken lattice are susceptible to decoherence [13–16] when shaken at certain amplitudes α and frequencies ω , both in the presence and absence of an inertial signal. Furthermore, shaking of a BEC trapped in an optical lattice has been shown to cause heating in the condensate due to atom–atom interactions [10, 11]. Atom scattering into transverse modes has also been shown to be deleterious [17, 18]. Therefore, it is of interest to analyze the lattice shaking protocols that drive these state-to-state transitions and reduce the subspace to eliminate deleterious shaking frequencies.

The desired transformation considered in this paper is the first step of shaken-lattice-based interferometry. That is, we wish to start with atoms in the ground state of the lattice and transform them to a ‘split’ state with an error less than 1%. The split state is defined such that the atoms equally populate two momentum states with the same magnitude but opposite sign. In particular we consider the simplest cases of splitting the atom population equally into the $\pm 2n\hbar k_L$ states for $n = 1, 2, \text{ and } 3$. Here, we define the lattice wavenumber $k_L = 2\pi/\lambda_L$ for a lattice wavelength λ_L . In general the n th order split state $|\psi_n(p, \theta)\rangle$ may have a relative phase θ between the two counterpropagating momentum components. That is, $|\psi_n(2n\hbar k_L, \theta)\rangle = e^{i\theta}|\psi_n(-2n\hbar k_L, \theta)\rangle$. It is important to note that one is not limited to interferometric operations such as beamsplitting when using the shaken lattice technique, but our work focuses primarily on this application, as increased momentum splitting increases interferometer sensitivity, e.g. in the $102\hbar k$ splitting demonstrated by the interferometer developed in [19].

For the simplest case of $2\hbar k_L$ splitting we show that if the lattice is shaken at frequencies near the Bloch band 0 to band 1 transition, we can split the atom wavefunction to within the desired error. This transition gives a relative phase difference of π between the two momentum states in the resulting split state. If we shake the lattice at half of the band 0 to band 2 transition frequency, we can split the atoms to within 1% error with zero relative phase between the two momentum states. In each case, the simulation results are backed up by analytics. This simple shaking scheme is not suitable for higher-order splitting because the transition rate between bands drops precipitously as we transition from band 0 to higher-lying bands. However, we find that if we optimize splitting via a genetic algorithm (GA) using only the band-to-band transition frequencies, we can achieve 4 and $6\hbar k_L$ splitting within 1%. Using this optimization scheme, we have obtained experimental $4\hbar k_L$ splitting with an error of $\approx 10\%$.

We show computationally that after splitting to third order ($\pm 6\hbar k_L$) we can load the atoms into counterpropagating moving lattices and accelerate them, potentially achieving acceleration sensitivities that scale as T_1^3 [6]. In similar schemes atom momentum transfer has also been achieved in optical lattices through the use of Bloch oscillations in a tilted lattice [20, 21]. An interferometer of this type obtaining splitting of $80\hbar k_L$ was demonstrated in [22].

The paper is structured as follows: in section 2 we motivate the description of the lattice dynamics in terms of the Bloch states and describe the split state in terms of these states. Section 3 will describe an analytic treatment of the problem. Section 4 discusses the efficacy of $2\hbar k_L$ and higher-order splitting with single-frequency shaking. Section 5 shows computational and experimental results of optimization wherein we restrict ourselves to shaking at the band-to-band transition frequencies. Section 6 concludes.

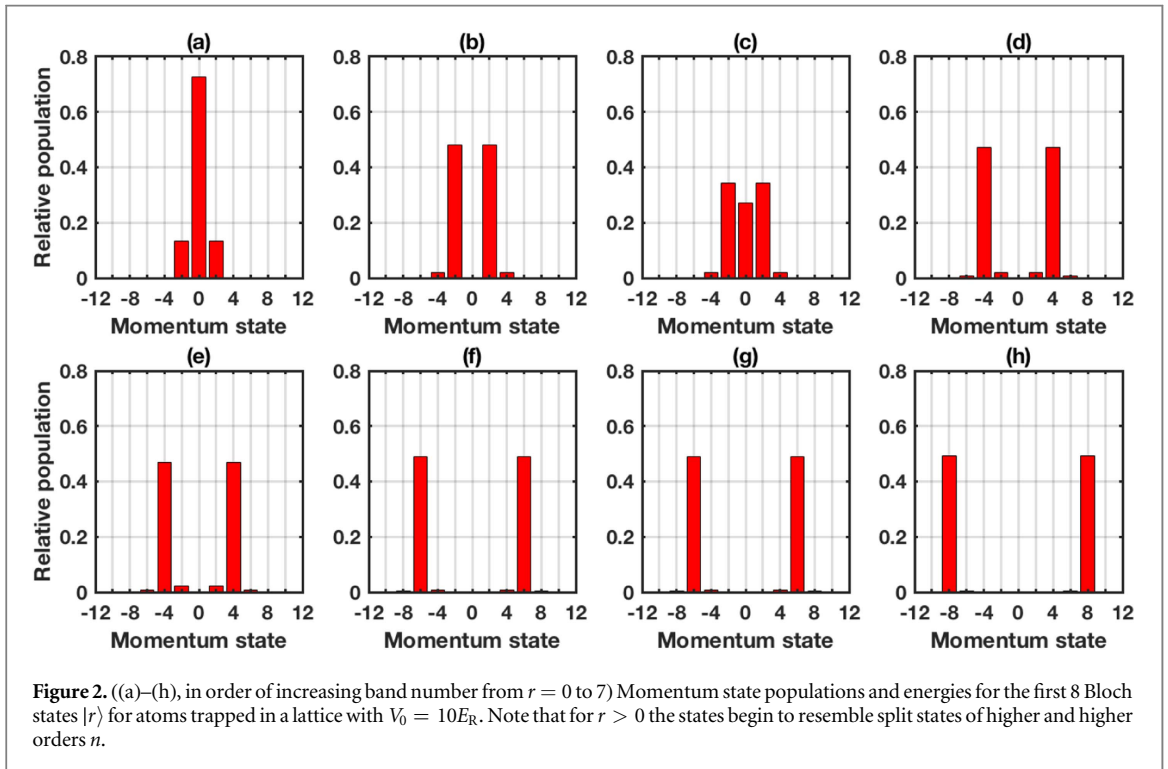
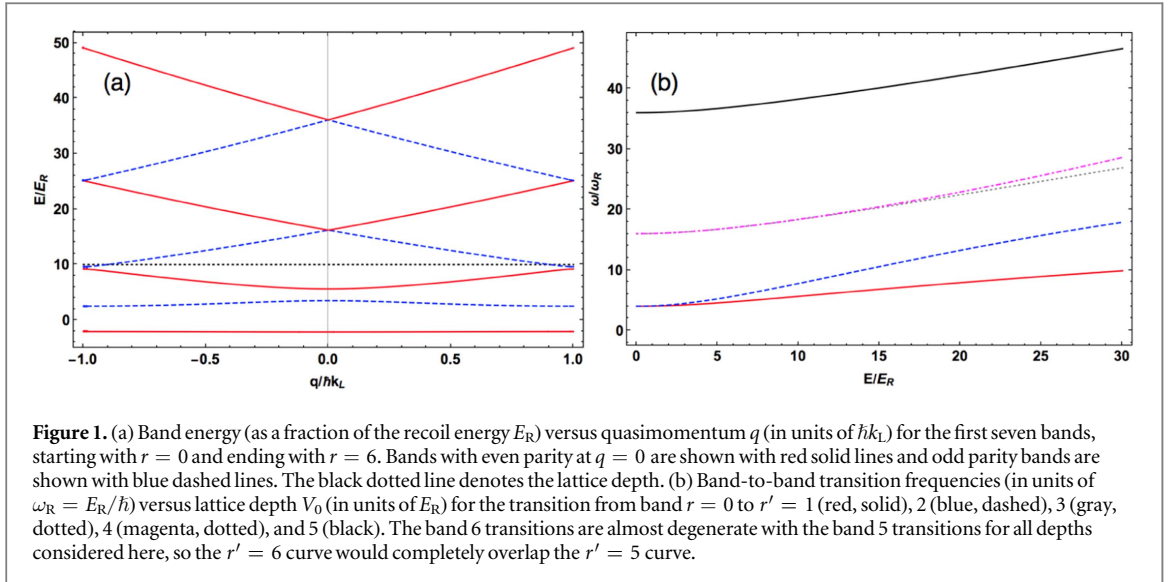
2. Bloch decomposition of the split state

For the simulation results presented in this paper we will make the following assumptions: First, we assume that the atoms are delocalized in the lattice, i.e. in a superfluid state [23]. We will assume that the atoms are initialized with quasimomentum $q = 0$ and this quasimomentum does not change. Finally, we assume the atoms are non-interacting and that the lattice is infinite.

Because we are working in the regime where the atoms are delocalized in an infinite lattice, the Bloch states $|\Psi_r^{(q)}\rangle$ are a convenient basis for calculations, where r denotes the band number and q is the quasimomentum. For the simulations done in this paper the lattice depth was chosen to be $V_0 = 10E_R$, where the recoil energy $E_R = \hbar^2 k_L^2 / 2m$ for atoms with mass m . The band energy E is plotted against the quasimomentum q in figure 1(a). The atoms begin in the state corresponding to the ground Bloch band $r = 0$ with $q = 0$. Since we assume that the quasimomentum is always zero we will suppress the index q in what follows and denote the Bloch states $|\Psi_r^{(0)}\rangle$ as simply $|r\rangle$.

The Bloch states at zero quasimomentum populate only the $2n\hbar k_L$ momentum states (for integer n). We expect then that in our model lattice modulation does not transfer momentum out of these states [1]. This is confirmed by simulation [3]. Therefore a complete description of the atom wavefunction can be given by the amplitudes and (relative) phases of the wavefunction in these quantized momentum states. Of particular interest is the relative phase θ between the two momentum components of the split state, as defined in section 1.

Experimentally one cannot determine these relative phases from time-of-flight absorption images, as we only have access to the probability amplitudes in these experimental measurements. As such, we define a vector \vec{P} with components P_n containing the probability of finding an atom in the $2n\hbar k_L$ momentum state [3]. If we consider an ensemble of atoms, this vector gives the relative population of atoms in each momentum state. In practice because higher-order momentum states are negligibly populated, we can truncate $|n|$ at $N = 5$. We can then define an ‘error’ E_{ab} between two momentum states described by vectors \vec{P}_a and \vec{P}_b as



$$E_{ab} = \left(1 - \frac{\vec{P}_a \cdot \vec{P}_b}{|\vec{P}_a||\vec{P}_b|} \right) \times 100\%. \quad (1)$$

From equation (1), we see that the more similar two states are, the lower the error E_{ab} . Note that if we are comparing any state to the split state, E will be independent of θ and is thus a more useful quantity to look at when considering the results in the context of what is experimentally observable. Thus, we use this expression as an error measure to quantify how well our optimization algorithm is doing.

For bands $r > 0$ there is considerable similarity between the Bloch states and split states of various orders. This is most easily seen when one looks at the momentum state population of the Bloch states corresponding to different bands, as shown in figure 2 [24]. Interestingly, there are two separate Bloch states at different band energies that roughly correspond to each split state. To glean further insight, we calculate the inner product D_{nr} between the n th order split state $\psi_n(p, \theta)$ and the state $|r\rangle$ as

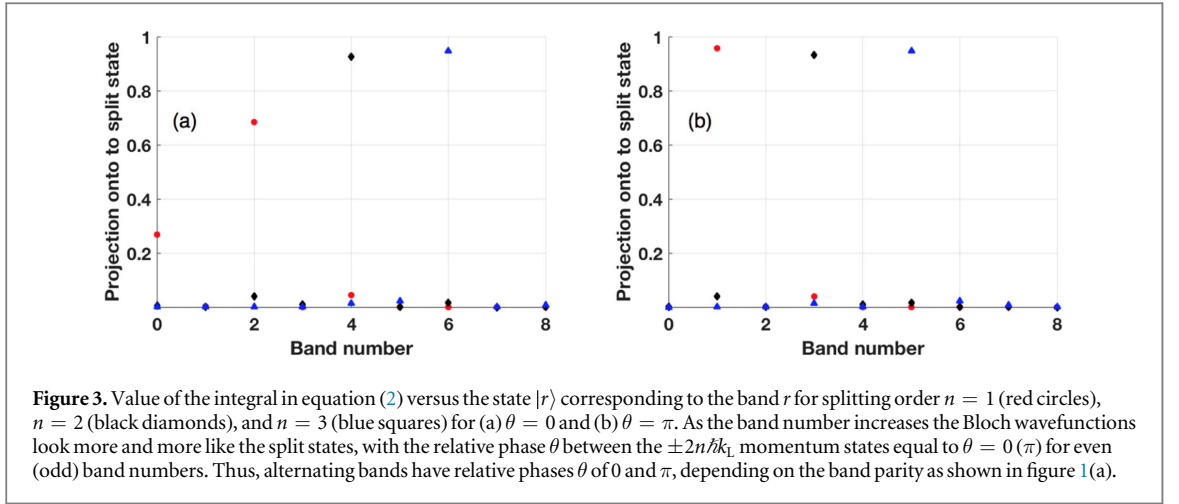


Table 1. Band transition frequencies $f_{r,r'}$, $V_0 = 10E_R$. The single (double) asterisk marks frequencies with matrix element overlaps $M^{(c)}(M^{(s)}) > 0.1$ (see figure 6 and section 3).

Band n	Band m	$f_{r,r'}$ (kHz)
0	1	17.89**
0	2	24.61*
0	3	58.14
0	4	58.25
0	5	121.19
1	2	6.72**
1	3	40.25*
1	4	40.36**
1	5	103.30
2	3	33.53**
2	4	33.64*
2	5	96.58
3	4	0.10
3	5	63.0*
4	5	62.9**

$$D_{nr} = |\langle r | \psi_n(p, \theta) \rangle|^2. \quad (2)$$

From figure 3, we see that the difference between two Bloch states corresponding to bands $l > 0$ and $l' = l + 1$ is that the lower-energy state $|l\rangle$ has a relative phase difference $\theta = \pi$ between the $\pm(l + 1)\hbar k_L$ states, and the higher-energy band $|l'\rangle$ is almost identical, except $\theta = 0$ (and thus the two states are orthogonal). This is commensurate with the fact that states corresponding to adjacent bands have opposite parity.

In the rest of this paper we will be referring to state-to-state transitions between different bands. The transition frequencies for transitions from the ground band to the first 5 excited bands are shown in figure 1(b). For an example of the energy scales at play here, we tabulate the transition frequencies $f_{r,r'}$ between two bands r and r' in table 1. The frequencies given in table 1 assume that we are working with ^{87}Rb atoms (as in section 4.2) at a lattice depth of $V_0 = 10E_R$. We see that the band transitions lie between 0 and 121 kHz, and this is the regime in which our driving is simulated.

The next section will analytically explore the dynamics of atoms trapped in a shaken optical lattice. This case is somewhat difficult to solve analytically, but some basic results can be applied in the simple case of single-frequency shaking at low amplitudes. We verify these results numerically, showing that we can split the atom wavefunction with a phase difference of 0 or π , depending on our driving frequency. Experimental results verify the numerics. From this we gain some insight on how to restrict our optimization parameters and show the results of rapid optimization of higher-order splitting in section 5.

3. Analytics of phase modulation of an optical lattice

The Hamiltonian for a phase-modulated (shaken) lattice with general shaking function $\phi(t)$ is written

$$H = \frac{p^2}{2m} + \frac{V_0}{2} \cos[2k_L x + \phi(t)]. \quad (3)$$

For the specific case where $\phi(t) = \alpha \sin(\omega t)$, we can apply the Jacobi–Anger expansion to equation (3). Using this we can write the potential term $V(x, t)$ in equation (3) as

$$V(x, t) = V_0 \left\{ \cos(2k_L x) \left[J_0(\alpha)/2 + \sum_{k=1}^{\infty} J_{2k}(\alpha) \cos(2k\omega t) \right] - \sin(2k_L x) \sum_{k=1}^{\infty} J_{2k-1}(\alpha) \sin[(2k-1)\omega t] \right\}. \quad (4)$$

Equation (4) shows that we can decompose the phase modulation to a term representing the carrier (first term) and a set of amplitude modulation terms containing both sine and cosine terms. The strength of these amplitude modulation terms are given by the Bessel functions $J_k(\alpha)$ where α is the amplitude of the phase modulation. Because the sine terms are odd, they will drive transitions between opposite parity states while the even cosine terms will drive transitions between states with the same parity [24].

By taking the J_0 term in equation (4) as the bare Hamiltonian $H_0(x)$, we can write the rest of the terms as a perturbation $H'(x, t)$. Using Fermi's golden rule, we can then write down the transition rate $\Gamma_{r,r'}$ from state $|r\rangle$ to $|r'\rangle$ resulting from shaking at a frequency ω as

$$\Gamma_{r,r'} = \frac{2\pi}{\hbar} V_0^2 \sum_{k=1}^{\infty} [J_{2k}^2(\alpha) |\langle r' | \cos(2k_L x) | r \rangle|^2 \times \delta(E_{r,r'} - 2k\hbar\omega) + J_{2k-1}^2(\alpha) |\langle r' | \sin(2k_L x) | r \rangle|^2 \times \delta(E_{r,r'} - (2k-1)\hbar\omega)], \quad (5)$$

where $E_{r,r'} = \hbar\omega_{r,r'} = E_r - E_{r'}$ is the energy difference between states $|r\rangle$ and $|r'\rangle$. The transition rate $\Gamma_{r,r'}$ is governed by the squares of the Bessel functions $J_k^2(\alpha)$ (where α is the amplitude of the phase modulation) and magnitudes of the transition matrix elements $|M_{r,r'}^{(s)}|^2 = |\langle r' | \sin(2k_L x) | r \rangle|^2$ and $|M_{r,r'}^{(c)}|^2 = |\langle r' | \cos(2k_L x) | r \rangle|^2$.

4. Simplified splitting schemes

This section consists of four parts. The first subsection will show first-order single-frequency shaking results via simulation. Next, we will show experimental data that supports the simulation results: namely that we can obtain first-order splitting by shaking with a single-frequency. This will lead into a discussion of the limitations of the single-frequency model, especially with regards to higher-order splitting schemes. Finally, we will discuss how much we need to split to implement an accelerating lattice scheme.

4.1. Theory and simulation of single-frequency splitting

We can use the theory derived in the previous section to make some predictions about the effects of single-frequency shaking. As stated in section 3 the matrix elements $|M_{r,r'}^{(c)}|^2$ connect states with the same parity, and the matrix elements $|M_{r,r'}^{(s)}|^2$ connect states of opposite parity.

For a given value of α the amplitude of the Bessel functions $J_k(\alpha)$ dies off as k increases. For $\alpha \leq 0.3$ we can keep two terms, simplifying the potential in equation (4) to

$$V(x, t) = V_0 [J_0(\alpha) \cos(2k_L x)/2 - J_1(\alpha) \sin(\omega t) \sin(2k_L x) + J_2(\alpha) \cos(2\omega t) \cos(2k_L x)]. \quad (6)$$

As with equation (6) if we keep two terms in equation (5), we obtain

$$\Gamma_{r,r'} = \frac{2\pi}{\hbar} V_0^2 [J_2^2(\alpha) |M_{r,r'}^{(c)}|^2 \delta(E_{r,r'} - 2n\hbar\omega) + J_1^2(\alpha) |M_{r,r'}^{(s)}|^2 \delta(E_{r,r'} - (2n-1)\hbar\omega)]. \quad (7)$$

From this we see that weak shaking of the lattice at $\omega_{r,r'} = 2\pi \times f_{r,r'}$ will drive transitions between Bloch states $|r\rangle$ and $|r'\rangle$ if they have opposite parity, and driving at $\omega_{r,r'}/2$ will drive transitions between $|r\rangle$ and $|r'\rangle$ if they have the same parity. In general, shaking at $\omega_{r,r'}/N$ will drive transitions between $|r\rangle$ and $|r'\rangle$ with parity determined by whether N is odd or even. This is in keeping with the results in [25] for the case of the amplitude-modulated lattice (where only like-parity transitions are allowed) and the phase modulation results in [26]. The

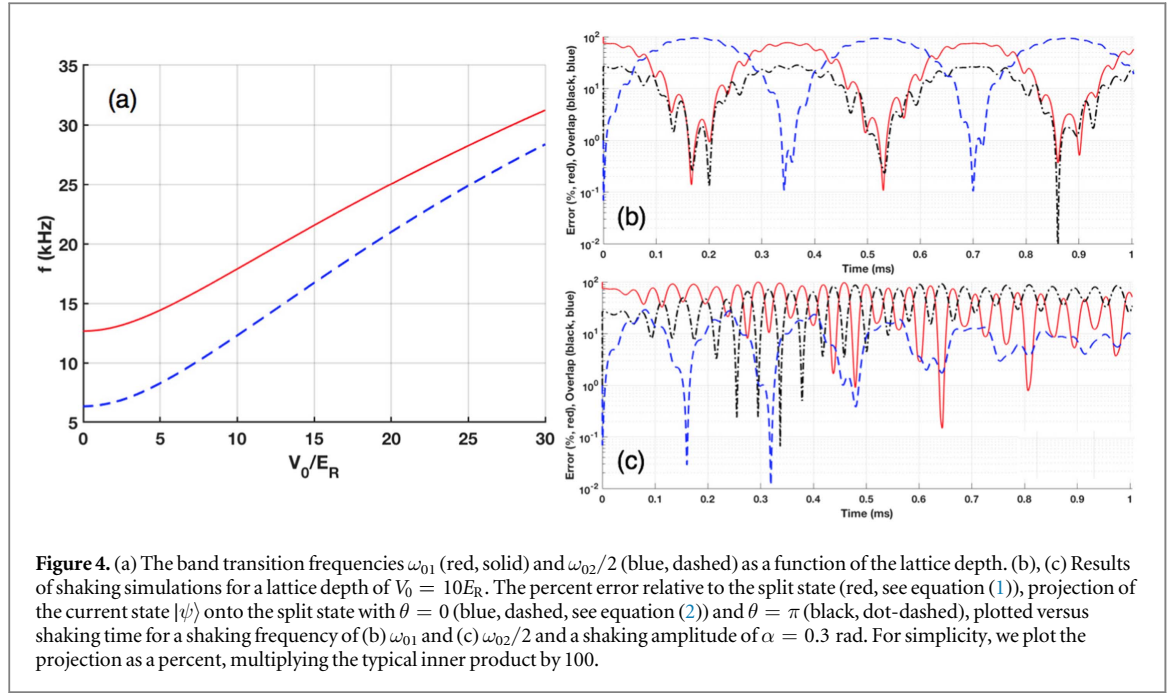


Figure 4. (a) The band transition frequencies ω_{01} (red, solid) and $\omega_{02}/2$ (blue, dashed) as a function of the lattice depth. (b), (c) Results of shaking simulations for a lattice depth of $V_0 = 10E_R$. The percent error relative to the split state (red, see equation (1)), projection of the current state $|\psi\rangle$ onto the split state with $\theta = 0$ (blue, dashed, see equation (2)) and $\theta = \pi$ (black, dot-dashed), plotted versus shaking time for a shaking frequency of (b) ω_{01} and (c) $\omega_{02}/2$ and a shaking amplitude of $\alpha = 0.3$ rad. For simplicity, we plot the projection as a percent, multiplying the typical inner product by 100.

difference in our work is that we approach this problem from a standpoint of inducing band-to-band transitions to perform atom beamsplitting for interferometry.

The above analysis shows that if we begin in the ground state $|r = 0\rangle$ and shake at $\omega = \omega_{01}$ ($\omega_{02}/2$), we will drive odd (even) parity transitions between bands $r = 0$ and $r' = 1$ ($r' = 2$). We simulate both cases using the symmetric split-step method [27] to simulate the time-dependent Schrödinger equation (TDSE) with the Hamiltonian in equation (3) with a single-frequency ω and amplitude $\alpha = 0.3$ for $T \approx 1$ ms.

The band transition frequencies ω_{01} and $\omega_{02}/2$ are plotted in figure 4(a) versus the lattice depth. Given $V_0 = 10E_R$, results for odd parity shaking at $\omega = \omega_{01} = 2\pi \times 17.88$ kHz are shown in figure 4(b), and results for even parity shaking at $\omega_{02}/2 = 2\pi \times 12.3$ kHz are shown in 4(c). For the simulation results presented in figures 4(b) and (c), at each timestep we calculate the percent error relative to the split state as in equation (1) and the inner product between the simulated state $|\Phi(t)\rangle$ at time t and the first-order split state $|\psi_1(p, \theta)\rangle$ as in equation (2).

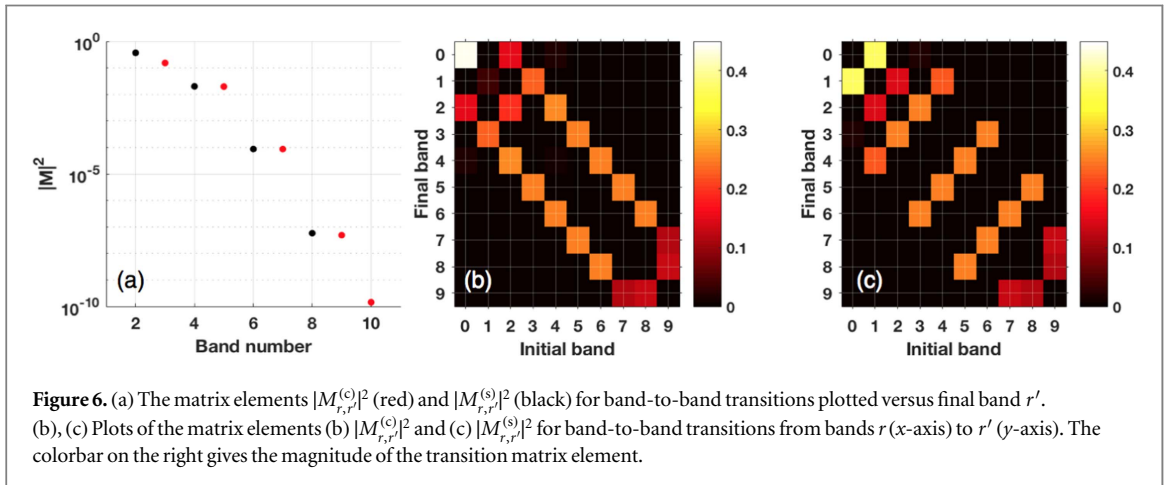
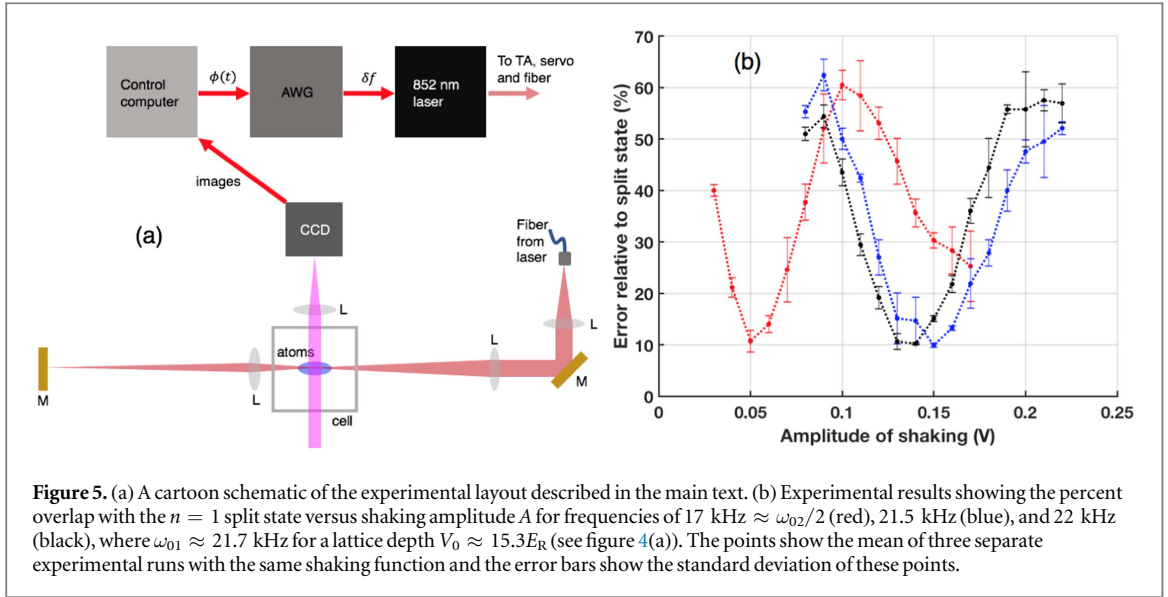
We see that when the percent error is lowest in figure 4(b), the projection of the state $|\psi\rangle$ onto the split states is highest for the split state $|\psi_1(p, \theta = \pi)\rangle$. This shows that we are in fact splitting with relative phase $\theta = \pi$ between the two split arms. Conversely, in figure 4(c) we achieve splitting with $\theta = 0$. Thus, by controlling the shaking frequency we can control the relative phase between the two split arms.

4.2. Experimental results of single-frequency first-order splitting

In this section we demonstrate that the first-order splitting schemes described in the previous subsection and shown in figure 4 are viable experimentally. The experimental scheme is similar to the shaken lattice interferometry experiment described in [4]. In the experiment we start with Bose-condensed ^{87}Rb atoms loaded into the ground state of an optical lattice of (intentionally) unknown depth. The lattice is made by retro-reflecting an 852 nm laser onto itself. By modulating the frequency of the lattice laser [28], we shake the lattice for a time $T = 0.2$ ms with varying amplitude A and frequency f^2 . We use a computer-controlled arbitrary waveform generator (AWG) to generate a pure tone modulated by an envelope of the form $f(t) = \cos^2(\pi t/T)$, which allows for smooth turn-on and turn-off of the shaking, as in [3, 4]. After 20 ms time-of-flight, we take an absorption image of the atoms using a CCD camera and analyze the images to extract the atoms' momentum state.

We find that we can split the atom wavefunction to within $E \approx 10\%$ at frequencies corresponding to ω_{01} and $\omega_{02}/2$. This is shown in figure 5(b). We do not generally obtain splitting to better than 10% due to spurious atoms detected in the $0\hbar k_L$ momentum state (e.g. due to heating and imaging noise), the finite momentum spread of the condensed atoms in the lattice [29, 30], and the finite signal-to-noise ratio in imaging [4]. Our experimental momentum width is about $0.6\hbar k_L$, limited mostly by the tight atom-chip-based trap in which we

² In [4] we used an electro-optic modulator (EOM) to modulate the lattice phase. When we were made aware of the method described in [28], we abandoned the EOM method in favor of the frequency-modulation method. This was done because the EOM was lossy and required relatively high voltages to achieve the phases we required.



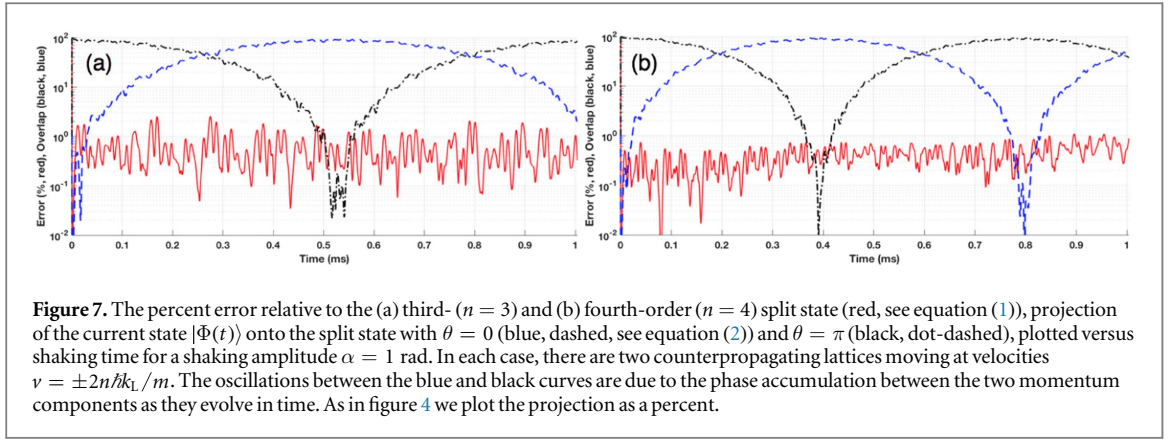
perform our evaporation. Simulations show that for such momentum spreads we are limited to errors of about 4%–8%, depending on our shaking frequency.

From the results shown in figure 5, we estimate the lattice depth to be $V_0 \approx 15.3E_R$. In this case the relevant band-to-band transition frequencies are $\omega_{01} = 21.7$ kHz and $\omega_{02} = 2 \times 17$ kHz. This not only confirms the simulation results from the previous subsection but provides us with a reliable way to approximate the lattice depth. In our current interferometry experiments the absolute lattice depth is less important than the day-to-day and shot-to-shot stability of the lattice depth. Thus, an approximate depth combined with the stability provided by a lattice laser intensity servo is sufficient for our purposes.

4.3. The limitations of the single-frequency model

While the single-frequency model works well for the first-order splitting schemes described in section 4.2, for higher amplitudes first-order perturbation theory becomes less and less applicable. Thus we can no longer use Fermi's Golden rule to accurately describe the physics of the situation. In this case we must keep more terms in the Jacobi–Anger sums of equation (6) and go to higher orders in perturbation theory. Furthermore, we cannot use this simple picture to obtain higher-order splitting. This is due to the fact that the matrix elements $|M_{r,r'}^{(c)}|^2$ and $|M_{r,r'}^{(s)}|^2$ become relatively small as we consider transitions from the state $|r = 0\rangle$ to higher-lying states with $|r > 2\rangle$. This is shown in figure 6(a) where for higher-band transitions the relevant matrix element is at least one order of magnitude below the lower-band transitions. As such, the transition strength is much lower and the transitions become less favorable.

However, we can make transitions from the ground state $|r = 0\rangle$ to an intermediate state in band $r' = 1$ or 2 and then to higher-lying states in bands $r'' > 2$. If we make these intermediate state transitions, the matrix



elements become more favorable. This is shown in figures 6(b), (c). As expected, the cosine transition matrix elements are strongest when making transitions between states in next-to-adjacent bands, but the sine matrix elements are strongest when making transitions between states in adjacent bands. Interestingly, when considering the sine matrix elements we see that it is also favorable to make transitions between states in bands $r = 1$ to $r' = 0, 2$ or 4 . We also observe that transitions between bands $r = 3$ and $r' = 2$ and 6 are favorable, but transitions between bands $r = 3$ and $r' = 4$ are not. This is possibly due to the avoided crossing between bands 3 and 4 at $q = 0$ (see figure 1). We find that the strongest transitions in figure 6 are most influential can be used to simplify the optimization landscape for higher-order splitting in section 5.

4.4. How much must we split?

Due to the fact that the single-frequency shaking does not work as well for higher-order splitting (see figure 6), higher-order splitting is more difficult to obtain. This is the subject of section 5. However, before we dive into the next section it is instructive to demonstrate how much we must split the atoms to be able to accelerate them in a moving lattice.

If we truncate the Bessel function expansion of equation (6) to two terms and do some trigonometry, we obtain

$$V(x, t) = 2V_0 \{J_0(\alpha) \cos(2k_L x)/4 + J_1(\alpha) [\cos(2k_L x - \omega t) + \cos(2k_L x + \omega t)]\}. \quad (8)$$

Equation (8) describes a carrier lattice and two counterpropagating moving lattices with velocity $v = \pm\omega/2k_L$. If we can split the atoms to a high enough order, we can trap the split atoms in one of the two moving lattices. The atoms will then move with the lattice if we accelerate and decelerate the lattice. This will allow us to obtain interferometry with T_1^3 sensitivity to an applied signal [6]. In this case the moving atoms will not be able to ‘see’ the counterpropagating lattices and will thus not be affected by them³. In this case, the atoms in the positive (negative) momentum state will be trapped in the lattice moving with positive (negative) velocity. Then, if the lattices are accelerated by changing the shaking frequency such that the magnitude of the counterpropagating lattice velocity changes, the atoms should follow the lattices in which they are trapped. The atoms will thus accelerate as the lattices are accelerated, given that this is done slowly enough [31, 32].

If we begin with atoms split to third order (that is, $\pm 6\hbar k_L$), we can shake the lattice at $\omega = 12\hbar k_L^2/m = 12\omega_R$ with $\alpha = 1$ such that the lattice is moving with $v = \pm 6\hbar k_L/m$ without disturbing the atom wavefunction appreciably. Here, $\omega_R = E_R/\hbar$ is the recoil frequency of the atoms in the lattice. In this case the atoms maintain their split state to within $\approx 1\%$, as shown in figure 7(a). Furthermore, simulations show that if the atoms begin in the $\pm 8\hbar k_L$ split state and are trapped in a lattice shaken at $\omega = 16\hbar k_L^2/m$, the atoms will continue to maintain their splitting to within 1%, as shown in figure 7(b). This trend continues for even higher splitting orders.

This analysis shows that if we can achieve third-order splitting we can then accelerate the atoms in the lattice with negligible perturbation. In the next section we will show how to optimize such splitting by shaking at frequencies corresponding to band-to-band transitions.

³ The atoms moving with one of the lattices must be in an eigenstate of the lattice, but shaking can always be modified to prepare the split atoms so that they resemble the ground state of the moving lattice with depth $V_0 J_1(\alpha)$.

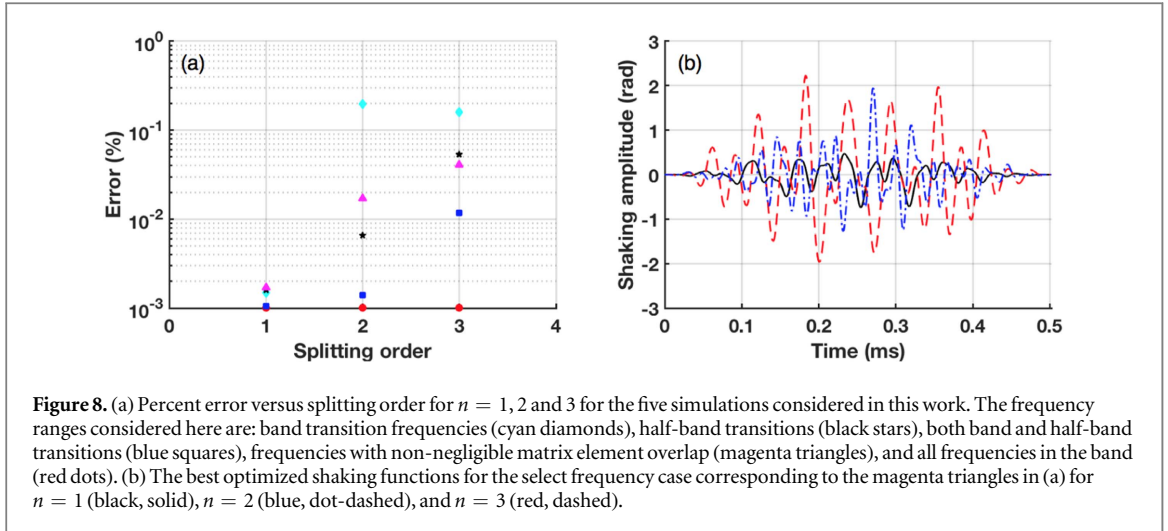


Figure 8. (a) Percent error versus splitting order for $n = 1, 2$ and 3 for the five simulations considered in this work. The frequency ranges considered here are: band transition frequencies (cyan diamonds), half-band transitions (black stars), both band and half-band transitions (blue squares), frequencies with non-negligible matrix element overlap (magenta triangles), and all frequencies in the band (red dots). (b) The best optimized shaking functions for the select frequency case corresponding to the magenta triangles in (a) for $n = 1$ (black, solid), $n = 2$ (blue, dot-dashed), and $n = 3$ (red, dashed).

5. Simplified optimization of higher-order splitting using band-to-band transitions

5.1. Computational results

This section presents results of simulations optimizing splitting protocols up to $n = 3$. Our optimization simulates the TDSE using the Hamiltonian in equation (3) as in section 3 and the optimization tries to minimize the error as given in equation (1). The optimization is done via a GA as detailed in [1, 3]. We chose the GA because the variation that we have developed has been proven to work very well for simulations of atoms in a shaken lattice. However, our GA was not well-suited for our initial experiments in [4] due to the relatively large number of iterations needed for the algorithm to converge.

Our experimental optimizations detailed in section 5.2 make use of the CRAB and dCRAB algorithms [33–35] in a closed-loop system. This choice of algorithms is mostly based on what worked for our initial simulations and experiments, and our choices should not be taken as a hard-and-fast rule, as genetic optimizers have been used in experiments (e.g. [36]), and the CRAB algorithm was originally used to optimize simulation dynamics.

We will compare results of the GA using the full frequency bandwidth up to the $r = 0 \rightarrow r' = 5'$ transition to optimize the lattice shaking to results where only single-photon band-to-band and two-photon half-band transitions are used. By the term ‘half-band transitions’, we mean that we shake the lattice at a frequency $\omega_{r,r'}/2$ to cause even parity transitions. We know from [11] that off-resonant shaking in the presence of atom–atom interactions causes heating. Furthermore, to avoid the transverse scattering described in [17, 18] we want to shake at single- and two-photon band-to-band resonances so that no excess energy is available for transverse scattering. Therefore we wish to restrict our optimization to resonant transitions in order to limit the heating due to these factors. Note that as in [3] our simulations do not take interactions into account.

In the simulations presented here we shook the lattice for $T \approx 0.5$ ms and optimized for $\pm 2n\hbar k_L$ splitting for $n = 1, 2$, and 3 . To ensure smooth turn-on and turn-off of the shaking function, we multiplied each shaking function by an envelope function $f(t) = \cos(2\pi t/T)$ [3, 4]. Due to the inherent randomness in the GA, we ran each class of simulations 10 separate times and took the best result for our analysis.

We ran five different classes of optimization simulations. One class included every frequency in the band from DC up to $r = 0 \rightarrow r' = 5'$, another included only the 10 band transition frequencies in this region, and a third included the 10 half-band transition frequencies. All frequencies used here are tabulated in table 1. A fourth simulation class included all 20 band and half-band transition frequencies, and a fifth chose only the 9 frequencies corresponding to appreciable (>0.1) matrix element overlap in figure 6 (marked with asterisks in table 1). We plot the lowest error achieved after 1000 iterations in figure 8. Note that convergence below $10^{-3}\%$ is limited by phase errors in the split-step method.

From the results presented in figure 9, we see that in all cases we can split the atom wavefunction to better than 0.3%. We see in figure 8(a) that the error is lowest if we include all frequencies, but in this regime the interaction-induced heating (which is not present in our current simulation model) will be highest. For simulations restricted to the select strongest band transitions we can obtain splitting to better than 0.05%. While the use of more complex fitness functions (e.g. that used for first-order splitting in [3]) may further improve this splitting efficiency, we obtain good results by simply minimizing the error in equation (1). In summary, by truncating our search space from ≈ 50 frequencies (limited by the discrete temporal sampling inherent in the numerics) to ≈ 10 frequencies, we can still split with sufficiently low error.

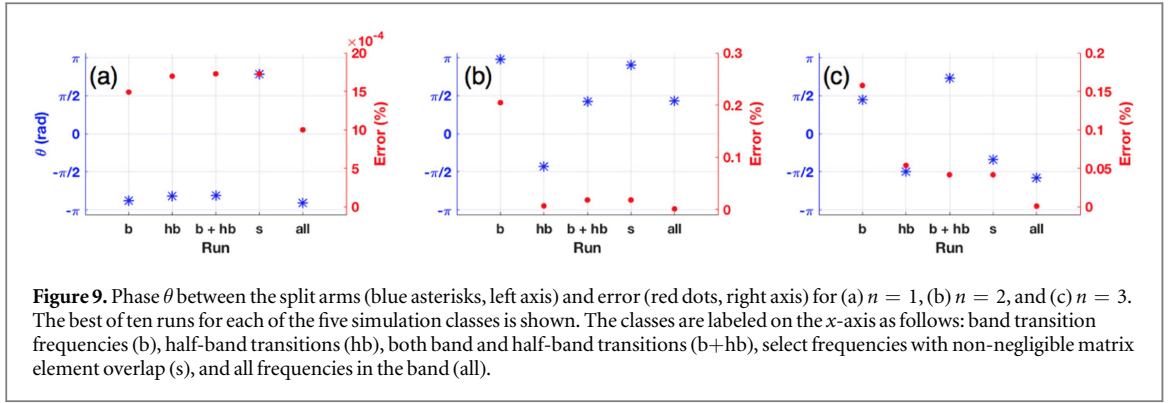


Figure 9. Phase θ between the split arms (blue asterisks, left axis) and error (red dots, right axis) for (a) $n = 1$, (b) $n = 2$, and (c) $n = 3$. The best of ten runs for each of the five simulation classes is shown. The classes are labeled on the x-axis as follows: band transition frequencies (b), half-band transitions (hb), both band and half-band transitions (b+hb), select frequencies with non-negligible matrix element overlap (s), and all frequencies in the band (all).

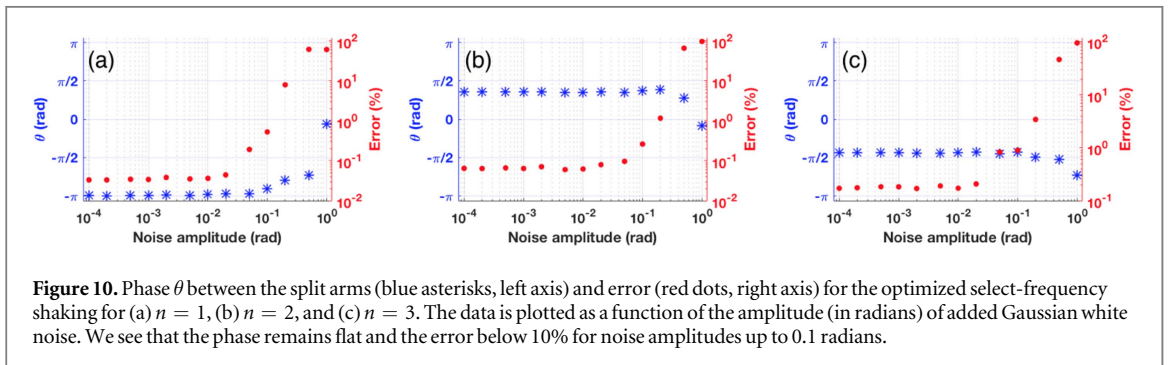


Figure 10. Phase θ between the split arms (blue asterisks, left axis) and error (red dots, right axis) for the optimized select-frequency shaking for (a) $n = 1$, (b) $n = 2$, and (c) $n = 3$. The data is plotted as a function of the amplitude (in radians) of added Gaussian white noise. We see that the phase remains flat and the error below 10% for noise amplitudes up to 0.1 radians.

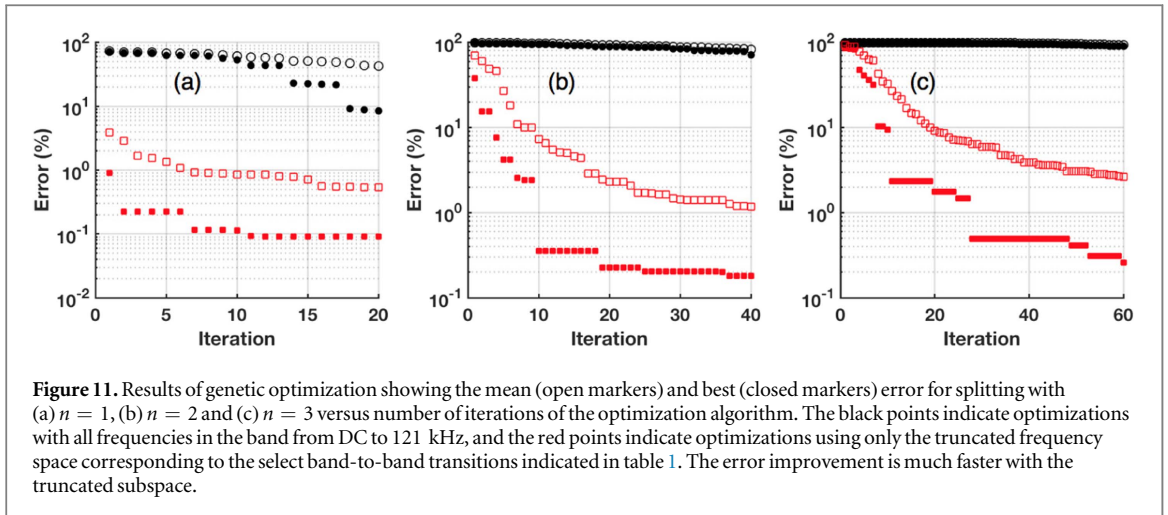
Even though we cannot access the relative phase θ of the two split arms of these optimized split states experimentally [4], it is of interest to examine them in simulation in order to better understand the shaking dynamics. Thus, we plot the final phase θ of the optimized split state for the best results of each of the five simulation classes and three splitting orders in figure 9. We include the error from figure 8 for easy reference and comparison.

The results show that the phase dynamics of multi-frequency splitting are more complex than the simple model presented in section 3 predicts. For example, when we split using the half-band transition frequencies we would expect that the phase θ be near zero. However, we find that this phase is closer to $\theta = -\pi$ for first-order splitting and $-\pi/2$ for higher-order splitting. Therefore, our simple model derived in the case of single-frequency shaking has broken down. Unfortunately it is analytically difficult to consider multiple frequency shaking due to the fact that neither the Jacobi–Anger expansion nor the results of Floquet analysis applies. Furthermore, as shown in figure 8(b) the shaking function amplitude is about an order of magnitude greater than that used to justify the truncation of the sum in equation (4) and apply first-order perturbation theory. Fortunately, for the select case, these relative phases are robust in the face of experimental noise and remain roughly constant as white noise is added to the shaking function up to a noise amplitude of 0.1 radians, as shown in figure 10. This corresponds to roughly 5%–10% of the maximum shaking amplitude; this is consistent with the robustness studies we did for $2\hbar k_L$ splitting in [3] and the phase coherence demonstrated in [4].

We can, however, make some general inferences from our simple model. The fact that $|\theta| \approx \pi$ for the first-order split state likely comes as a result of the fact that the two-photon matrix element $|M_{0,2}^{(c)}|^2$ is about a factor of two lower than the single-photon element $|M_{0,1}^{(s)}|^2$ connecting $r = 0$ and $r' = 1$. Thus, transitions between the odd parity states are more favorable, as can be seen from figure 4 where the dynamics of shaking at ω_{01} are far less complex than those of shaking at $\omega_{02}/2$.

For higher-lying states, the multiple pathways for an atom to get from $|r = 0\rangle$ to the higher-lying states means that the even and odd parity transitions tend to interfere, and the split state will in general be a mixture of the two bands. From figure 1 we see that these higher-lying states $|r' > 2\rangle$ corresponding to splitting with $n > 1$ are nearly degenerate. This interference causes the phase difference between the two split arms to average to $|\theta| = \pi/2$. In these cases the optimized split state is not dominated by population transfer into a single higher-lying band but rather this state is composed of a mixture of states corresponding to two nearly degenerate bands.

From a purely experimental standpoint our results dramatically simplify the optimization landscape that we must explore. This allows for sufficient error minimization within a low number of iterations. That is, even though simulations with more frequencies tend to converge to lower errors, they take longer to do so. For



example, if we run 10 simulations to optimize splitting of the atom wavefunction with the select transition frequencies indicated in table 1, for $n = 1, 2$, and 3 , we can achieve convergence to better than 1% error in (on average) 1, 10, and 28 iterations, respectively. In each case, the error for the simulations with all frequencies in the band is $>70\%$, as shown in figure 11. Our simulations also show that if we start with atoms in the $n = 2$ split state, we can optimize transfer into the $n = 3$ split state within 1% within <10 iterations. In this case the total splitting time will double, but optimization of $6\hbar k_L$ splitting is possible with fewer than 20 total iterations. This learning speedup is extremely important for practical implementations of shaken lattice interferometry in that optimization happens more quickly and effectively, allowing for fast optimization of the interferometer sequence.

In the experiment, if the lattice depth is known (e.g. via the measurement scheme described in section 3 or in [24, 30]) we can restrict our shaking the selected transitions with appreciable transition matrix elements. Then a closed-loop algorithm (e.g. the CRAB or dCRAB algorithm [33–35]) can be used to efficiently optimize the splitting protocol (as was done for first-order splitting in [4]). Thus we have found a reduced subspace that allows for faster optimization of the system and reduces heating due to off-resonant shaking [11]. As an example, we demonstrate optimization of $4\hbar k_L$ splitting in the next section.

5.2. Experimental results

Here we present experimental results of $n = 2$ splitting using the CRAB algorithm using the restricted subspace defined in the previous section. Currently, experimental limitations prohibit us from obtaining $n = 3$ splitting, but with some straightforward improvements to the apparatus (as outlined in [4]), higher-order splitting can be achieved. The results shown here therefore serve as an encouraging proof-of-principle of the efficacy of optimization with the simplified subspace.

Using the apparatus described in section 4.2 and [4], we use the CRAB algorithm to optimize the higher-order split state. For simplicity we do not use the ‘dressed’ CRAB (dCRAB) algorithm, but this could be implemented in the future as a way to improve the optimization. As with the GA, we fix the shaking frequency components corresponding to the 9 select band-to-band transitions described in the previous section. Optimization then occurs on 5 vectors of amplitudes corresponding to each of these frequency components. As in section 4.2, we use an envelope function to ensure smooth turn-on and turn-off of the shaking function.

The system runs in closed-loop, shaking the lattice, taking and analyzing experimental absorption images, then feeding the results (averaged over three separate but identical runs of the experiment) into the optimization algorithm. The algorithm updates the shaking function and the optimization continues. In this way we can converge on the desired state. We set the shaking time to be 0.4 ms, twice the $2\hbar k_L$ splitting time used in [4]. The learning curve is shown in figure 12(a). The error in splitting was measured to be $8.4 \pm 2.4\%$ after averaging over 15 shots. (The value obtained in figure 12(a) is slightly higher due to the fact that it is an average over only 3 shots.)

From this, we have shown that the simplified shaking scheme can quickly optimize $n = 2$ splitting to within 10% in 5 iterations, which is consistent with the results shown in figure 11. Thus, it is expected that one should be able to optimize $n = 3$ splitting within about 10 iterations using the same optimization protocol.

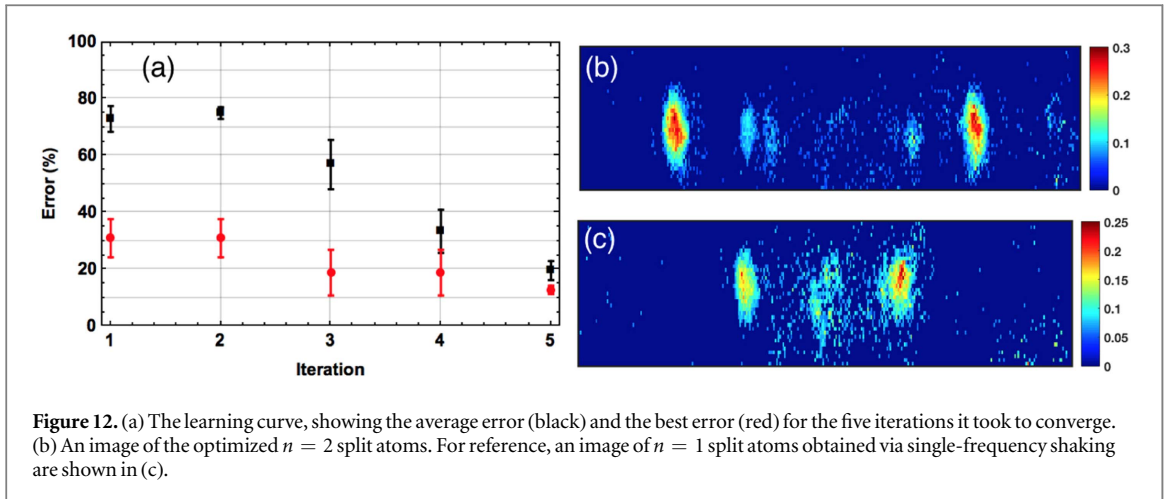


Figure 12. (a) The learning curve, showing the average error (black) and the best error (red) for the five iterations it took to converge. (b) An image of the optimized $n = 2$ split atoms. For reference, an image of $n = 1$ split atoms obtained via single-frequency shaking are shown in (c).

6. Conclusion

In conclusion we demonstrate a simple means of using the band-to-band transitions to implement an atom beamsplitter in an optical lattice. We develop a theoretical model that allows us to use a single shaking frequency to implement low-order splitting. However, at higher frequencies our simple model breaks down and we must incorporate multiple frequencies in order to obtain good splitting. Due to heating caused by atom–atom interactions it is of interest to restrict our shaking frequencies to those resonant with single- and two-photon transitions between bands. We show that we can obtain simulated higher-order splitting up to $n = 3$ with an error $< 0.1\%$ by optimizing shaking with a learning algorithm using a reduced subspace of frequencies corresponding to the strongest band and half-band transition resonances. We show this experimentally for the case of $n = 2$ splitting. This simplification of the optimization landscape allows for faster optimization with less deleterious heating effects due to atom–atom interactions. Finally, we show that higher-order splitting can be implemented by accelerating the atoms in the optical lattice and can potentially allow for interferometry with sensitivity that scales with the cube of the interrogation time. This opens up potential new pathways for improving and expanding interferometry using atoms trapped in a shaken optical lattice.

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