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Citation for published version (APA):
https://doi.org/10.1103/PhysRevA.101.042703

DOI:
10.1103/PhysRevA.101.042703

Document status and date:
Published: 15/04/2020

Document Version:
Publisher’s PDF, also known as Version of Record (includes final page, issue and volume numbers)

Please check the document version of this publication:
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High-precision analysis of Feshbach resonances in a Mott insulator

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(Received 5 December 2019; revised manuscript received 21 February 2020; accepted 13 March 2020; published 15 April 2020)

We show that recent high-precision measurements of relative on-site interaction energies $\Delta U$ in a Mott insulator require a theoretical description beyond the standard Hubbard-model interpretation, when combined with an accurate coupled-channels calculation. In contrast to more sophisticated lattice models, which can be elaborate especially for parameter optimization searches, we introduce an easy to use effective description of $U$ valid over a wide range of interaction strengths modeling atomic pairs confined to single lattice sites. This concise model allows for a straightforward combination with a coupled-channels analysis. With this model we perform such a coupled-channels analysis of high-precision $^7$Li spectroscopic data on the on-site interaction energy $U$, which spans over four Feshbach resonances and provide an accurate and consistent determination of the associated resonance positions. Earlier experiments on three of the Feshbach resonances are consistent with this analysis. Moreover, we verify our model with a more rigorous numerical treatment of the two atom system in an optical lattice.

DOI: 10.1103/PhysRevA.101.042703

I. INTRODUCTION

Precise knowledge and control of the atomic interactions is essential to a wide range of modern ultracold atom experiments. Coupled-channels models can accurately describe ultracold collision properties [1] of a two-atom system by detailed interaction potentials that are fine tuned by just a few parameters, to match atom loss spectra [2], photoassociation rates and collisional cross sections [3], interference patterns between $s$-wave and $d$-wave collisions [4], and rf molecule association data [5]. Accurate information on the interaction strength close to Feshbach resonances [6] is needed to determine the three-body parameter of a strongly interacting Bose gas [7,8]. Also the collisional energy dependence can be substantial, when operating a cesium microgravity clock [9], or for the determination of the nature of a spinor condensate [10,11]. Approximate or effective descriptions like a contact interaction or a harmonic trapping potential are often required to embed the microscopic collisional properties of two atoms into the macroscopic environment of the ultracold gas, where atoms may be held by magnetic or optical traps [12–14], or by optical lattices [15]. Given the collisional properties of two atoms one can then correctly account for collisional energy shifts; however, the collisional properties themselves depend on those energy shifts via the relative kinetic energy of the colliding atoms that is linked to the total energy of the two particles, which is shifted by the collisional energy shift. Therefore a self-consistent approach is needed and the accuracy of the coupled-channels interaction parameters depends crucially on the accuracy of the model connecting the microscopic processes to the macroscopic environment. Using a less accurate model may result in inconsistencies with respect to the two-body parameters when comparing between different experiments, or even when comparing data within a single experiment.

In this paper, we want to utilize a coupled-channels model in combination with an accurate but easy to use theoretical lattice model description of the on-site interaction energy $U$, to find a consistent description of high-precision spectroscopic data of a two component Mott insulator with two particles per site, where the on-site energy was varied by using Feshbach resonances [16]. In this experiment, atoms in different spin states were confined in a cubic optical lattice geometry. Such systems can implement a spin-Heisenberg model [17], which is of interest to study quantum magnetism. In the Mott insulating phase with doubly occupied sites individual atom pairs localize around each lattice site [18,19]. At the same time the interactions between different spin components of the atoms can be controlled utilizing Feshbach resonances. This condition results in an ideal situation to study the isolated two-particle system. Then, the strength of the atomic interaction characterized by the scattering length $a_S$ can be linked to the energy per localized atom pair $E(a_S)$ and thus to the on-site interaction energy $U$ defined as $U = E(a_S) - E(0)$. In Ref. [16] high-precision data on the on-site interaction energy have been acquired in such a two component Mott insulator using interaction spectroscopic techniques.

We show that the standard Hubbard model approximation [18] leading to a linear relation between the on-site interaction energy $U$ and the scattering length $a_S$ is not accurate enough to describe the experimental data to the required precision even for moderate scattering lengths. Instead we offer a simple effective description to approximate the effects of a cubic lattice potential for two identical particles of mass $m$, in...
which the on-site interaction energy $U$ is parametrized by a harmonic oscillator model with contact interactions [12]. The harmonic oscillator frequency $\omega_{\text{eff}}$ should be adjusted to best represent the on-site potential. Note that this is not necessarily the frequency related to the second-order approximation around a lattice potential minimum. We choose $\omega_{\text{eff}}$, such that the harmonic model matches the linear description from the Hubbard model close to the noninteracting case, where the Hubbard model should give the correct description up to first order. The effective trapping frequency $\omega_{\text{eff}}$ is then completely determined by the noninteracting lattice parameters

$$h\omega_{\text{eff}} = 2\pi E_r \left( \frac{1}{k_c} \int |\psi_0(r)|^2 dr \right)^2,$$

where $\psi_0$ is the lowest band Wannier function of the one-dimensional single-particle system, $E_r = h^2 k_c^2 / (2m)$ is the recoil energy with $k_c = 2\pi / \lambda$, and $\lambda$ is the wavelength of the lattice light. All in all our effective model is meant to describe the two-particle system with contact interactions including the full lattice potential. This system has been theoretically studied in detail [15,20]. To make the connection to a more complete description of the interparticle interactions, the strength of the contact interaction is gauged to match the free scattering properties of the full coupled-channels system for a fixed collision energy $\epsilon$. In ultracold experiments it is often sufficient to take $\epsilon \approx 0$ and thus to match the scattering length $a_S$. To achieve a more accurate description we want to match the full coupled-channels and contact system at the correct relative collision energies $\epsilon$ in a self-consistent manner. We find an approximate expression $\epsilon(a_S) \approx E(a_S) - E_0$, which is given by the total energy per localized atom pair depending on the scattering length minus the energy in the center-of-mass (c.m.) mode. For example, in a harmonic oscillator potential of frequency $\omega$ the energy in the c.m. mode would be $E_0 = 3\hbar \omega / 2$.

By gauging the contact interaction strength and matching the collision energy we can thus find the on-site interaction energy $U$ for a full coupled-channels description in a lattice environment. To check the validity and limitations of the effective description we compare to the full contact system in the lattice using a direct diagonalization approach as presented in [20,21]. In addition we investigate the collision energies from the full contact bound-state wave functions to match the contact to the coupled-channels solution at the correct relative energies. This analysis also enables us to identify regimes in which the simple contact approximation is likely to fail, for example, close to narrow resonances, where the full systems scattering properties show strong dependence on the relative collision energy. Finally we apply our approach to perform a full coupled-channels analysis of recent experimental data on $^7\text{Li}$ [16].

II. THEORY

We start with some remarks on the system we aim to model. All in all we have a many-body system in mind, but since we assume the system to be in a Mott insulating phase with two atoms per lattice site in a regime where the interaction with atoms at neighboring sites may be neglected we restrict to a simple two-particle model system with both particles well localized at a single lattice site. The two-particle system in a lattice is described by the following Hamiltonian:

$$H = \frac{p_1^2}{2m} + \frac{p_2^2}{2m} + V_{\text{opt}}(\mathbf{r}_1, \mathbf{r}_2) + V_{\text{int}}(\mathbf{r}_1 - \mathbf{r}_2).$$

with $p_i$ and $\mathbf{r}_i$ the momentum and position operators of particle $i \in \{1, 2\}$, $m$ the mass of the particles, $V_{\text{int}}$ the interaction between the particles, and $V_{\text{opt}}$ the lattice potential.

To simplify the system further one can mimic the interactions between the particles with a contact interaction of variable interaction strength. By adjusting this interaction strength we can correctly represent the physics at length scales bigger than the range $r_0$ of the real interaction potential. Note, however, that this approximation will only be good around some fixed relative energy $\epsilon$ between the particles.

In the following we give a detailed analysis of the contact interaction case and focus in the second part on the connection of contact to coupled-channels model. We include a discussion of effects related to the nonconserved relative collision energy in the lattice scenario.

A. Contact interaction

To mimic the interactions between the particles we introduce a contact interaction of variable interaction strength, implemented with a Bethe-Peierls boundary condition [22]. We parametrize the interaction strength by the scattering length $a_S$ of the interaction such that we have

$$V_{\text{int}}(\mathbf{r}_1 - \mathbf{r}_2) = \frac{4\pi \hbar^2 a_S}{m} \delta(\mathbf{r}_1 - \mathbf{r}_2).$$

Let us first consider the case of a weakly interacting system where $a_S \rightarrow 0$ and a perturbative analysis is possible. We closely follow the derivation of the Hubbard model and thus also make the connection to the many-body theory. We start from the noninteracting case ($a_S = 0$), such that we can restrict to the single-particle scenario. We consider a cubic optical lattice

$$V_{\text{opt}}(\mathbf{r}_1, \mathbf{r}_2) = \sum_{i \in \{1, 2\}} V_0 E_i \left[ \sin(k_x x_i)^2 + \sin(k_y y_i)^2 + \sin(k_z z_i)^2 \right],$$

with $V_0$ the lattice depth in recoil energies $E_r$. In this case the band structure as well as the single-particle wave functions are known and can be expressed in terms of solution to Mathieu’s equation [19]. However, the so-called Bloch solutions to the problem are not localized, but one can combine the Bloch waves of each band $n$ to find the band’s Wannier function $\psi_n(\mathbf{r}_j)$ [23]. The Wannier function is real valued and well localized around $r_j = 0$ for deep lattices $E_r \rightarrow \infty$. Together with versions $\psi_n(\mathbf{r}_j - \mathbf{R}_i)$ shifted by a lattice vector $\mathbf{R}_i$ the Wannier function provides a complete basis set for the $n$th band subspace. In addition the Wannier functions $\psi_n$ approach eigenfunction solutions of a harmonic approximation around the lattice sites in the deep lattice limit and they are approximate solutions to the Schrödinger equation up to the width of the respective bands. To approximate the system’s low-energy part around a single lattice site it is therefore reasonable to project onto the lowest band Wannier function $\psi_0(\mathbf{r}_j)$ such
that the first-order correction in energy will be
\[ U = (W|V_{\text{int}}|W), \]  
where \( W = w_0(r_1)w_0(r_2). \) For the contact interaction we therefore obtain
\[ U = \frac{4\pi \hbar^2 a_S}{m} \int |w_0(r)|^4 dr, \]
which leads to a linear correspondence between the energy correction \( U \) and the scattering length \( a_S. \) Note that this corresponds exactly to the on-site interaction energy in the Hubbard model. Further we note that this treatment assumes the two-body wave function and therefore also the atomic density around a given lattice site to stay unaltered. To include the deformation of the wave function or atomic density caused by the interaction potential a higher-order treatment is necessary.

To go beyond the linear regime Eq. (6) we can approximate around a single lattice site with an interacting harmonic model [12]:
\[ H_0(\omega_{\text{eff}}) = \frac{p_i^2}{2m} + \frac{p_j^2}{2m} + \frac{4\pi \hbar^2 a_S}{m} \delta(r_1 - r_2) + \frac{1}{2m} \omega_{\text{eff}}^2 r_1^2 + \frac{1}{2m} \omega_{\text{eff}}^2 r_2^2. \]

The harmonic oscillator frequency \( \omega_{\text{eff}} \) should be adjusted to best represent the lattice potential. Note that this is not necessarily the frequency related to the second-order approximation around a lattice potential minimum. We rather adjust the harmonic oscillator frequency such that it matches up with the first-order result from the Hubbard model close to vanishing interaction strength [see Eq. (1)]. The spectrum of this effective harmonic model can be determined analytically and the shift in ground-state energy \( U \) is determined by [12]
\[ \frac{\sqrt{2} \Gamma \left( \frac{9}{2} \right)}{\ell_{10} \Gamma \left( \frac{9}{2} \right)} = \frac{1}{a_S}, \]
where \( \Gamma \) denotes the gamma function, \( E_{\text{HO}} = h\omega_{\text{eff}}, \) and \( \ell_{10} = \sqrt{\hbar/(m_0a_0^2)}. \) Since the solutions to the harmonic model Hamiltonian are exact it naturally includes also the deformation of the wave function caused by the interaction, which manifests itself in the radial s-wave component of the relative wave function \( \psi_{n,\ell=0}(r) \) given explicitly below in Eq. (16).

To analyze the accuracy and limitations of such an effective harmonic approach we calculate the spectrum of the contact system described by \( H \) using a direct diagonalization approach similar to [20,21,24] making use of the eigenstates of the harmonic model. Such a numerical approach is accurate only around a single lattice site, therefore the population of neighboring lattice sites might be underestimated. This can lead to slight deviations between the numerical and Hubbard model description of the system, which we will discuss later.

We start by singling out the harmonic Hamiltonian in the full system by adding and subtracting the harmonic potential:
\[ H = H_0(\omega_{\text{eff}}) + V_{\text{int}}(r_1, r_2) - \frac{1}{2} m \omega_{\text{eff}}^2 (r_1^2 + r_2^2) \]
\[ = H_0(\omega_{\text{eff}}) + V_\Delta. \]

We will project on the eigenvectors of \( H_0(\omega_{\text{eff}}) \), therefore the major task is to determine the coupling matrix elements resulting from the deviation from the lattice potential \( V_\Delta \). But before analyzing those in more detail let us first change to relative \( r \) and c.m. coordinates \( R \) with
\[ r_1 = \frac{1}{\sqrt{2}} (R - r) \quad \text{and} \quad r_2 = \frac{1}{\sqrt{2}} (R + r). \]

Note that this is not the standard definition of relative and c.m. coordinates. We adopted the conventions of [12], where relative and c.m. directions are rescaled to have equal effective mass. In addition we introduce units natural to the harmonic model system \( H_0(\omega_{\text{eff}}) \), so all energies will be given in multiples of \( h\omega_{\text{eff}} \) and all lengths in multiples of \( \sqrt{\hbar/(m_0a_0^2)}. \) The difference in potentials \( V_\Delta \) separates into \( x, y, \) and \( z \) directions:
\[ V_\Delta = v_\Delta(X, x) + v_\Delta(Y, y) + v_\Delta(Z, z). \]

where the components \( v_\Delta \) are given by
\[ v_\Delta(X, x) = \frac{V_0}{2\sqrt{V_{\text{eff}}}} \left[ \sin^2 \left( \frac{X - x}{\sqrt{2}V_{\text{eff}}} \right) + \sin^2 \left( \frac{X + x}{\sqrt{2}V_{\text{eff}}} \right) \right] - \frac{1}{2} x^2 \]
\[ = \frac{V_0}{\sqrt{V_{\text{eff}}}} \left[ \sin^2 \left( \frac{X}{\sqrt{2}V_{\text{eff}}} \right) \cos^2 \left( \frac{x}{\sqrt{2}V_{\text{eff}}} \right) \right] \]
\[ + \cos^2 \left( \frac{X}{\sqrt{2}V_{\text{eff}}} \right) \sin^2 \left( \frac{x}{\sqrt{2}V_{\text{eff}}} \right) - \frac{1}{2} x^2 \]
\[ = \sum_{i,j} \alpha_{ij}(V_0, V_{\text{eff}}) x_i^2 X_{-j}. \]

In the last step we performed a Taylor series expansion and have introduced the effective lattice depth parameter \( V_{\text{eff}} = h^2\omega_{\text{eff}}^2/(2E_r)^2. \) Note that in the series expansion only even powers of \( x \) as well as \( X \) occur. This leads to the symmetry properties of the system that have been discussed in a more general setting in [21].

We now change to a basis of eigenstates \( |N_x, N_y, N_z, n, \ell, m\rangle \) of the effective harmonic system. The \( N_\ell \) are integers labeling the Harmonic oscillator eigenstates in the respective c.m. direction, whereas \( n, \ell, \) and \( m \) are quantum numbers in the relative direction, with \( \ell \) the angular momentum quantum number, \( m \) the magnetic quantum number, and \( n \) the solutions in the relative separation \( r \). For \( \ell \neq 0 \) the quantum numbers \( n, \ell, \) and \( m \) just describe the usual noninteracting harmonic oscillator states, while, for \( \ell = 0, \) \( n \) labels the solutions of the interacting harmonic model in the relative separation \( r \). The general solution with correct behavior for \( r \to \infty \) and relative energy \( \epsilon_{rel}(n) \) is given up to a normalizing constant by
\[ \psi_{n,\ell=0}(r) \propto e^{-\frac{r}{\sqrt{2}}} r U \left[ \frac{3}{2} - \frac{\epsilon_{rel}(n)}{2} \right] \frac{3}{2}, r^2], \]
where \( U \) is Tricomi’s confluent hypergeometric function. The quantization condition determining \( \epsilon_{rel}(n) \) is then given by the boundary condition at \( r = 0 \) and can be expressed in terms of
the scattering length $a_S$: 
\[
\frac{\sqrt{2}\Gamma\left(\frac{1}{2}\left[3 - 2\epsilon_{\text{el}}(n)\right]\right)}{\Gamma\left(\frac{1}{2} - \epsilon_{\text{el}}(n)\right)} = \frac{1}{a_S}.
\] 
(17)

We can now give the Hamiltonian $H_0(\omega_{\text{eff}})$ in its diagonal form:
\[
H_0(\omega_{\text{eff}}) = \sum_{N_x, N_y, N_z, n, \ell, m} \left[ E(N_x, N_y, N_z, n, \ell, m) \times |N_x, N_y, N_z, n, \ell, m\rangle \langle N_x, N_y, N_z, n, \ell, m| \right],
\] 
(18)

where eigenenergies corresponding to the states $|N_x, N_y, N_z, n, \ell, m\rangle$ are given by
\[
E(N_x, N_y, N_z, n, \ell, m) = \hbar\omega_{\text{eff}}(3 + N_x + N_y + N_z + \ell + \delta_{\ell,0}[\epsilon_{\text{el}}(n) - 3/2] + [1 - \delta_{\ell,0}]2n).
\] 
(19)

To reduce the states we project on, we can use the symmetry properties of the Hamiltonian as we mentioned earlier. We have reflection symmetries in $X \rightarrow -X$, $Y \rightarrow -Y$, and $Z \rightarrow -Z$, therefore there will be just coupling between even or odd values of $N_x$, $N_y$, and $N_z$ respectively. We also have the symmetry of inversion of $r \rightarrow -r$ equivalent to particle exchange, which leads to separation between even and odd values of $\ell$. For bosons we obviously need the even $\ell$ values. The reflection symmetry in $(x, y) \rightarrow (-x, -y)$ leads to the restriction to even or odd values in $m$. Finally we could also change to a base
\[
|N_x, N_y, N_z, n, \ell, |m\rangle\rangle_{S/A}
\] 
(20)
of symmetric and antisymmetric combinations in the sign of $m$. The reflection symmetry in $z \rightarrow -z$ guarantees the separation in $S/A$. We are interested in the component with $N_x$, $N_y$, $N_z$, $\ell$, and $|m\rangle$ even and symmetric combinations ($S$) in $|m\rangle$, which corresponds to the ground state of the interacting harmonic model $H_0(\omega_{\text{eff}})$ and thus to our solution up to zeroth order in the corrections $V_\Delta$ introduced by the full lattice potential.

We want to determine the spectrum of the full Hamiltonian close to the ground-state energy of the noninteracting full system. To do so we restrict to a finite expansion of $V_\Delta(X, x) \approx \sum_{i,j} \alpha_{ij}(V_0, V_{\text{eff}})\times x^i X^j$. We choose to expand up to tenth order in $2i + 2j$. This leads to a deviation of less than $0.05V_0E_r$ to the full potential when restricting to a region up to $0.7d$ around the lattice site at the origin. Here $d$ denotes the lattice constant. We then project on a finite set of states $|N_x, N_y, N_z, n, \ell, |m\rangle\rangle_{S/A}$ and obtain the energy spectrum by diagonalizing the resulting Hamiltonian matrix. The base set is chosen big enough to obtain a converged spectrum around the energy of the lowest band.

In Fig. 1 we compare the energy shifts $U$ from the first-order perturbative and the effective description to our full numerical solution for an optical lattice of depth $V_0 = 35E_r$. We assume here a lattice constant $d$ of $532nm = 100.5\lambda_0$, but the result will depend only on the ratio $a_S/d$. We get good agreement with the full result for the effective harmonic model with a maximal relative deviation of about $<0.8\%$ for $|a_S| < 0.3d \approx 50\lambda_0$. Note that the numerical results presented here rely on the two-particle wave function to be localized around a single lattice site $\pm 0.7d$, since we needed to cut the expansion in $v_\Delta$. We want to estimate the error caused by this approximation. Therefore we compare the effective harmonic models determined from the numerical approach and the Wannier function of the system [see Fig. 1(b)]. The models differ since the Wannier function of the system possesses side peaks at neighboring lattice sites. Those are absent in our numerical approach since the neighboring lattice sites lie in a classically forbidden region due to the finite expansion in $v_\Delta$. This leads us to an estimate of the theoretical accuracy of our model.
indicated by the gray shaded area in Fig. 1(b). With that we still have a maximal relative systematic error in $U$ of 1.3% for $|a_s| < 0.05d$.

When interpreting experimental data on $U$ the refined harmonic description leads to corrections in the position and width of Feshbach resonances as compared to the linear model. With the refined model one finds resonance positions shifted towards the direction of negative scattering lengths as well as increased resonance widths. The shifts in resonance positions result from non-anti-symmetric behavior of $U$, while the increase in width is related to the flattening off of $U$ for diverging $a_s$.

B. Connection to a coupled-channels model

So far we considered particles interacting via a contact potential. In ultracold gas experiments the collisions are usually considered to happen at zero energy and thus the scattering lengths of the coupled-channels and contact systems need to be matched. For the considered lattice geometry, however, we can estimate the relative energy at which the particles collide to be around the energy $\epsilon$ of the lowest single-particle band, which is for our parameters $\epsilon/k_B \approx 20 \mu K$. This is estimated from two noninteracting particles in the lowest band with energy $2\epsilon$. Approximately half of this is relative energy, which is also the collision energy at zero separation. Adding a repulsive interaction leads to an increase in collision energy since the wave function needs to shift to higher energies, which also leads to an increase in size of the wave function in relative direction. Despite the wider spatial and therefore narrower momentum distribution at low momenta of the two-particle wave function, there is an increase in average relative kinetic energy due to an increase in the high momentum tail.

In general a contact model can be adjusted to correctly represent the s-wave component of any short-range potential for far separation of the particles at a fixed collision energy $\epsilon = \hbar^2 k^2/(2\mu)$. To do so one has to choose the following scattering length for the contact system [25]:

$$\frac{1}{a_s(k)} = -k \cot[\delta_s(k)],$$  \hspace{1cm} (21)

with $\delta_s(k)$ the scattering phase shift of the full coupled-channels interaction. The right-hand side can be expanded in powers of $k$ with

$$k \cot[\delta_s(k)] = -\frac{1}{a_s^2} + \frac{R_c}{2} k^2 + O(k^4),$$  \hspace{1cm} (22)

with $a_s^2$ the scattering length of the coupled-channels model and $R_c$ the effective range. This leads us to the implications of including collisions at a finite energy. Effects due to finite collision energy $\epsilon$ are relevant especially close to narrow resonances, where the effective range $R_c$ can take large values. For narrow resonances the resonance position at finite collision energy $B_0(\epsilon)$ defined by

$$k \cot[\delta_s(B_0, k)] = 0$$  \hspace{1cm} (23)

can thus be subject to shifts of order $\Delta B_0(\epsilon) \sim \epsilon/\mu_B \approx 0.3$ G.

Similar to the free case Eq. (21) we want to arrive at a matching condition for the lattice environment. Since the lattice system cannot be reduced to relative and c.m. motion there will be no well-defined collision energy. Instead the collision is happening in different c.m. channels simultaneously, where each of those c.m. channels has an assigned collision energy. The c.m. channels are not to be confused with the internal spin channels of the coupled-channels model, which describes the collision in the relative direction. To see that the collision is happening in different c.m. channels we first split the Hamiltonian into three components:

$$H = \frac{p^2}{2m} + \frac{p^2}{2m} + V_{\text{int}}(r) + V_{\text{opt}}(r, R)$$  \hspace{1cm} (24)

$$= \frac{p^2}{2m} + V_{\text{opt}}(0, R) + \frac{p^2}{2m} + V_{\text{int}}(r) + \hat{V}_R(r, R)$$  \hspace{1cm} (25)

$$= H_R + H_c + \hat{V}_R(r, R),$$  \hspace{1cm} (26)

one acting solely on the c.m. component $H_R$, one acting solely on the relative coordinate $H_c$, and a part acting on both $V_{\text{opt}}(r, R)$. Here $H_R$ describes a lattice system in the c.m. coordinate [see Eq. (28)]. We note that the c.m. lattice is effectively four times deeper then the single-particle lattice when transformed to lattice units with all lengths in multiples of $\hbar$ and all energies in multiples of $E_k = E_r/2$.

We can get an approximate expression for the full Hamiltonian valid around a single lattice site by projecting on the c.m. Wannier functions $|i\rangle$ at that given site, thereby neglecting the coupling to other lattice sites in the c.m. direction:

$$H \approx \sum_{i,j} |i\rangle \langle j'| \delta_{ij}(H_c + \epsilon_i) + \langle i| \hat{V}_R |j'| \rangle.$$  \hspace{1cm} (27)

Here $i$ labels the c.m. bands with associated c.m. energies $\epsilon_i$ that we set to be in the center of the corresponding c.m. bands. In addition we find that the coupling term $\hat{V}_R$ vanishes up to second order in $r$ as $r \to 0$. For clarity we give the terms $V_{\text{opt}}(0, R)$ and $\hat{V}_R(r, R)$ explicitly:

$$V_{\text{opt}}(0, R) = 2E_r V_0 \sin(k_X/\sqrt{2}) + \sin(k_{Z}/\sqrt{2})^2$$  \hspace{1cm} (28)

$$\hat{V}_R(r, R) = E_r V_0 [1 - \cos(\sqrt{2}k_X)\cos(\sqrt{2}k_Z) + [1 - \cos(\sqrt{2}k_Y)\cos(\sqrt{2}k_Z)$$  \hspace{1cm} (29)

$$\approx (k_r)^2 \hat{V}_R^{(2)}(\hat{r}, R) + O((k_r)^4)).$$  \hspace{1cm} (30)

For a contact interaction the inner boundary condition in all c.m. channels is determined by the scattering length. Upon investigating the full numerical contact solutions $\Psi$ we can get a lower bound for the population of the lowest c.m. channel $|0\rangle$ by

$$|\langle \Psi |(|0\rangle \otimes \mathbb{1}_r) |\Psi \rangle| \geq \alpha^2 \beta^2 - 2\alpha \beta \sqrt{1 - \alpha^2} \sqrt{1 - \beta^2},$$  \hspace{1cm} (31)

with $\alpha = |\langle 0 | \langle 0 \rangle \otimes \mathbb{1}_r) |\Psi \rangle|$, and $|\langle N_X = 0, N_Y = 0, N_Z = 0 \rangle |\Psi \rangle|$, where $|0\rangle$ denotes the lowest harmonic oscillator state in the c.m. component. For the bound state considered we find that the lowest c.m. channel is always populated to more than 99% in the considered scattering length regime. Therefore we expect to get a good approximation by matching
the contact solution in this c.m. channel to the full coupled-channels result.

The solution $\Psi$ thus approximately obeys the following Schrödinger equation when the separation between the particles is small:

$$E\Psi \approx H\Psi$$

$$\approx |0\rangle \langle 0| [(H_e + \varepsilon_0) + \langle 0| \hat{V}_\Delta |0\rangle] \Psi$$

$$\rightarrow |0\rangle \langle 0| [(H_e + \varepsilon_0)] \Psi,$$

where $H_e$ describes the free scattering process in the relative direction. The energy of this free collision is determined by the energy $E$ of the full state $\Psi$ minus the energy $\varepsilon_0$ stored in the center-of-mass component of the system. Therefore the collision energy of the particles is determined by

$$\frac{\hbar^2 k^2}{2\mu} \approx E - \varepsilon_0 = U(k) + E(0) - \varepsilon_0.$$ (35)

The strength of the contact interaction and thus the scattering length $a_S(k)$ can then be determined from a coupled-channels model by justifying the condition [see Eq. (8)]

$$\frac{\sqrt{2} \Gamma(-\frac{U(k)}{\Delta_{1\phi}})}{\ell_{HO} \Gamma(-\frac{U(k)}{2\Delta_{1\phi}} - \frac{1}{2})} = -k\cot[\delta_S(k)].$$ (36)

This ensures that the full contact and the coupled-channels wave function are properly matched in the lowest c.m. band. The boundary condition in the other populated collision channels is, however, not exactly satisfied. Therefore this limits the contact model approach to regimes where either the effective range correction terms are small or the lowest c.m. channel is dominating.

To sum up in Eq. (36) we combine an effective harmonic parametrization of $U$ representing a contact interaction model with the phase-shift behavior $\delta_S(k)$ of a full coupled-channels calculation. This enables us to perform a coupled-channels analysis of on-site interaction data [16] in the following section.

### III. COMPARISON TO EXPERIMENT

Our model is captured by Eq. (36). We apply it to recent experimental data taken for $^7$Li [16]. For $^6$Li and $^7$Li the ultracold interatomic interactions already have been accurately characterized by different experiments; however, also discrepancies are known to exist directly related to the two-body interactions, for instance, in the determination of three-body parameters near an $|f, m_f\rangle = |1, 1\rangle$. Feshbach resonance of $^7$Li [5,26]. In [16] a sample of ultracold $^7$Li atoms is prepared in a Mott insulating state with doubly occupied sites in a cubic optical lattice with a depth of $35E_r$. The experiment involves the two lowest hyperfine states labeled as $|a\rangle$ and $|b\rangle$. More precisely, at each doubly occupied lattice site one of the three symmetric combinations in spin $|aa\rangle$, $|ab\rangle_S = (|ab\rangle + |ba\rangle)/\sqrt{2}$ or $|bb\rangle$ can be realized and correspond to one of three different interaction channels. With the help of radio-frequency pulses transitions between the different spin states can be driven. From the resonance frequency positions the difference in on-site interaction energy between the scattering channels ($U_{ab} - U_{aa}$) and ($U_{bb} - U_{ab}$) can be inferred. This can be done for a wide range of external magnetic fields $B$.

We note that the analysis presented here relies on the effective harmonic model using the oscillator frequency determined from the numerical model [orange line in Fig. 1(a)]. We include the gray shaded area in Fig. 1(b) as our theoretical error estimate.

As a first test we want to verify that the effective model indeed leads to an improvement compared to the linear Hubbard description. For that purpose, we compare the experimental interaction spectroscopic data to a coupled-channels model gauged to earlier experiments [5]. We use the scattering length data determined from this model and map it with either the linear Eq. (6) or the effective harmonic description Eq. (8) onto the experimental data. In Fig. 2 we show that the effective harmonic model leads to improved agreement with the experimental data. There we compare the differences in on-site interaction energy $\Delta U$ by showing the deviations $\Delta U_{exp} - \Delta U_{cc}$ between experiment and theory for better visibility. We find as expected that the resonance positions in $\Delta U$ for the linear conversion appear at too high magnetic-field values.

We use the effective harmonic approximation including finite collision energy effects Eq. (36) to map coupled-channels phase-shift data $\delta_{cc}$ to on-site interaction energy $U$. We take the coupled-channels model presented in detail in [5], where it was used to interpret rf-spectroscopy data taken for $^7$Li. The most crucial parameters in the coupled-channels model are the van der Waals coefficient $C_6$ and the adjustments in the singlet $S$ and triplet $T$ boundary conditions parametrized in the form of phase parameters $\Delta \phi_S$ and $\Delta \phi_T$ [5]. We take those as free parameters that we fit to the experimental data by performing a $\chi^2$ minimization. Our fit results are presented in Fig. 3 and Table I. The theoretical error estimate has been included into our error analysis by adding the theoretical error determined from the model of Ref. [5] to the experimental error bars prior to the fit. We find good agreement with the experimental data (see Fig. 3); just close to the narrow resonance our model seems to underestimate the width of the resonance. Note that we identified the regime close to the narrow Feshbach resonances to be less well approximated by the contact model. In Table I we compare to the results obtained in [16] with a linear Hubbard model in combination with dispersive shapes to parametrize $a_S$ in the different interaction channels. For the positions of the broad resonances (at 738, 795, and 894 G) we find values corrected by 0.2 to 0.7 G to higher magnetic fields. We attribute the major contribution to these corrections to the effective harmonic model. However, for the narrow resonance (at 845 G) we find a correction to lower magnetic fields of 0.1 G as a result of the finite collision energy effects included. Comparing to previous results we find improved agreement for the broad resonances, while the deviation in the position of the narrow resonance increased.

Similar good agreement can be achieved by fitting with the effective harmonic model without finite collision energy corrections Eq. (8). For comparison we give those results also in Table I. The resulting resonance positions for the broad resonances are generally in agreement within the two different models while the narrow resonance is significantly shifted to lower values for the energy corrected model.
FIG. 2. Comparison of the experimental to the coupled-channels model determined in [5]. We show the residuals between experimental data and theoretical data for $\Delta U$ in the $|aa\rangle$ to $|ab\rangle$ transition (top) and the $|bb\rangle$ to $|ab\rangle$ transition (bottom). The plots on the left-hand side have been obtained with the linear and the plots on the right-hand side with the effective harmonic model. The color scale indicates the shift $\delta B_{\text{res}}$ required in the theoretical resonance positions to reach agreement. Thus the color scale can serve as a simple measure for the quality of agreement between the unfitted coupled-channels model and the experimental data. The color map has been obtained starting from a dispersive model fitted to the coupled-channels result.

IV. CONCLUSION AND OUTLOOK

We presented a full coupled-channels description of the on-site interaction energy $U$ of a Mott insulator state with two atoms per lattice site. Our description is based on a parametrization of the on-site interaction energy with an effective harmonic model adjusted to match the linear behavior at small interaction strengths that can be determined from the Hubbard model. A matching condition has been obtained in Eq. (36) that combines the effective harmonic parametrization of $U$ with the relative collision energy in the lowest c.m. band from the noninteracting scenario. We verified that for moderate scattering lengths up to $0.05a_d$ and a lattice depth $V_0 = 35 E_r$ our effective approach gives good agreement with the full contact scenario. We applied our effective description to perform a successful coupled-channels analysis of recent

FIG. 3. Experimental data $U_{ab} - U_{aa}$ [orange (light gray) points] and $U_{bb} - U_{ab}$ [blue (dark gray) points] and best fit with the coupled-channels description to the combined data set [orange (light gray) and blue (dark gray) lines]; the lower half shows the difference of the data points and the fitted model along with the experimental error bars taken to be one $\sigma$ of the fit to the resonance spectra, while the systematic error on the experimental side has been estimated to 0.1 kHz. Our estimate of the theoretical uncertainties is indicated by the orange (lighter gray) and blue (darker gray) shaded areas.
TABLE I. Comparison of the coupled-channels (CC) fit results with collision energy corrections Eq. (36) and without collision energy corrections Eq. (8) to the results obtained in [16] where the linear Hubbard model is used in combination with dispersive shapes to parametrize $a_0$ and to previous works. Only the resonance positions have been determined directly from our coupled-channels fit. The error in the resonance positions is taken to be one standard deviation and has been determined from the fit to the experimental data. We give the background scattering lengths $a_{bg}$ and resonance widths $\Delta$ for completeness. $a_{bg}$ and $\Delta$ have been determined by fitting a dispersive model as in [16] to the scattering length data obtained from our best fit with the coupled-channels code. For that we chose magnetic-field values between 650 and 950 G and restricted to points with $|a_0^2/a_0^1| < 1000$. The errors we give on $a_{bg}$ and $\Delta$ are the errors of the fit to the best theoretical model. Note, however, that the values of $a_{bg}$ and $\Delta$ depend strongly on the magnetic-field range and the maximum scattering length used for the fit, since the dispersive shape is just an effective description.

<table>
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<tr>
<th>Channel</th>
<th>$a_{bg}/a_0$</th>
<th>$\Delta$ (G)</th>
<th>$B_{res}$ (G)</th>
<th>$a_{bg}/a_0$</th>
<th>$\Delta$ (G)</th>
<th>$B_{res}$ (G)</th>
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</thead>
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<td>CC fit energy corrected</td>
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<td>Previous works</td>
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<tr>
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<td>$-168.0(1)$</td>
<td>$737.81(2)$</td>
<td>$-20.98$ [5]</td>
<td>$-171.0$ [5]</td>
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<tr>
<td>$ab$</td>
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<td>$-133.4(1)$</td>
<td>$795.20(2)$</td>
<td>$-20.0$ [26]</td>
<td>$-174$ [26]</td>
<td>$737.69(12)$ [26]</td>
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<td>Taken from [16]</td>
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<td>$-20.62(2)$</td>
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<td>$bb$</td>
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<td>$893.98(4)$</td>
<td>$-23.0(1.4)$</td>
<td>$-172.7(10.0)$</td>
<td>$893.34(12)$</td>
</tr>
</tbody>
</table>

| $aa$     | $-17.60(3)$  | $-19.03(3)$ | $845.50(5)$   | $-23.0(1.4)$ | $-172.7(10.0)$ | $893.34(12)$ |
| $ab$     | $-20.95(2)$  | $-133.6(1)$ | $795.31(2)$   | $-29.8(1.3)$ | $-90.5(4.0)$ | $794.64(07)$ |
| $bb$     | $-17.60(3)$  | $-227.8(4)$ | $893.98(4)$   | $-23.0(1.4)$ | $-172.7(10.0)$ | $893.34(12)$ |

There is no error bar given in [5].

Measurements of the on-site interaction energy shift for different lattice depths could reveal the dependence of the resonance positions as a function of collision energy. Also note that the points in magnetic field, where the scattering lengths of two channels are equal, should be independent of the conversion model, in a regime where the scattering lengths determines the interaction. These are the points where $(U_{ab} - U_{aa})$ or $(U_{bb} - U_{ab})$ cross zero or each other. A precise determination of those points might be valuable information in addition to the resonance positions.

ACKNOWLEDGMENTS

We thank Paul Mestrom, Victor Colussi, Wouter Verhoeven, Silvia Musolino, Denise Ahmed-Braun, and Jinglun Li for fruitful discussions. This research is financially supported by the Netherlands Organisation for Scientific Research under Grant No. 680-47-623, and is part of the research program of the Foundation for Fundamental Research on Matter. We also acknowledge support from the NSF through the Center for Ultracold Atoms and Award No. 1506369, from ARO-MURI Non-Equilibrium Many-Body Dynamics (Grant No. W911NF14-1-0003), from AFOSR-MURI Quantum Phases of Matter (Grant No. FA9550-14-10035), from ONR (Grant No. N00014-17-1-2253), and from a Vannevar-Bush Faculty Fellowship. J.A-G. acknowledges support from the NSF Graduate Research Fellowship under Grant No. DGE 1144152.
