Multi-orbital and density-induced tunneling of bosons in optical lattices

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Abstract. We show that multi-orbital and density-induced tunneling have a significant impact on the phase diagram of bosonic atoms in optical lattices. Off-site interactions lead to density-induced hopping, the so-called bond-charge interactions, which can be identified with an effective tunneling potential and can reach the same order of magnitude as conventional tunneling. In addition, interaction-induced higher-band processes also give rise to strongly modified tunneling, on-site and bond-charge interactions. We derive an extended occupation-dependent Hubbard model with multi-orbitally renormalized processes and compute the corresponding phase diagram. It substantially deviates from the single-band Bose–Hubbard model and predicts strong changes of the superfluid-to-Mott-insulator transition. In general, the presented beyond-Hubbard physics plays an essential role in bosonic lattice systems and has an observable influence on experiments with tunable interactions.

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1. Introduction

Hubbard models are extremely successful in describing a variety of systems ranging from electrons in solids to bosonic quantum gases in optical lattices. While in solids all kinds of complex lattice systems are realized, optical lattices offer simple geometries in combination with precisely controllable experimental conditions. The striking idea that ultracold bosonic atoms in optical lattices constitute an excellent tool for studying the superfluid (SF)-to-Mott-insulator (MI) transition \([\textsuperscript{1}]\) has initiated an intensive investigation of bosonic lattice models. It has caused a large number of theoretical studies on the Bose–Hubbard model ranging from mean-field treatments \([\textsuperscript{2–4}]\) and quantum Monte Carlo \([\textsuperscript{5}]\) to density-matrix renormalization group (DMRG) \([\textsuperscript{6}]\) approaches. In addition, the influence of disorder \([\textsuperscript{2, 7–11}]\) and the effect of confining potentials have been discussed \([\textsuperscript{12–15}]\). However, the first-order corrections to the Bose–Hubbard model itself, namely higher orbital tunneling (figure 1(a)) and bond-charge interactions (figure 1(b)), have so far been mostly neglected in these approaches. Bond-charge interactions known from solids \([\textsuperscript{16–19}]\) constitute a density-induced tunneling process caused by the scattering of particles on neighboring sites. This extension to the Hubbard model was discussed in the context of superconductivity \([\textsuperscript{16}]\) and ferromagnetism \([\textsuperscript{19}]\). While for fermionic systems these effects are partly suppressed due to the Pauli principle, they have a large impact on bosonic systems. The unique experimental access in optical lattices allows for a detailed investigation of this interaction-assisted tunneling. This density-dependent hopping was recently discussed also for boson–fermion mixtures \([\textsuperscript{20, 21}]\), where the Bose–Fermi–Hubbard model does not fully cover effects arising from the interspecies interaction \([\textsuperscript{20–24}]\). In this
context, the role of higher orbitals in the tunneling processes [20–22] was investigated. For bosonic systems, modifications of tunneling and on-site interaction due to higher bands were studied by variational mean-field methods [25–28] and by numerical exact methods restricted to either double- or triple-well systems [29–32] or to the on-site interaction [33–36]. However, in the correlated regime present in the vicinity of the MI transition, the applicability of mean-field methods to describe higher-band processes is doubtful. Thus, the challenging problem is to find a comprehensive description including both bond-charge hopping [16–20, 37, 38] and higher band processes [25–28, 30, 31, 33–36, 39, 40] that is valid also for strongly correlated systems. The recent experimental progress in accessing the occupation-dependent on-site interactions [35, 41–43] and the tunneling matrix element [24] necessitates an accurate theoretical method for calculating these parameters. Furthermore, the SF–MI transition in experiments with tunable interactions [23, 42] is directly affected by bond-charge tunneling as well as multi-orbital extensions to conventional and bond-charge tunneling.

Here, we show that interaction-induced processes cause two substantial modifications of the Bose–Hubbard model for bosonic atoms in three-dimensional (3D) optical lattices. Firstly, the so-called bond-charge or density-induced tunneling combining interaction and hopping (figure 1(b)) is a notable contribution to tunneling. Secondly, higher bands are occupied due to the interaction of particles on a lattice site. This increases the tunneling significantly as the tunneling in higher bands (figure 1(a)) is enhanced strongly. We present a general scheme to calculate renormalized tunneling processes by effectively incorporating higher-band contributions. On this basis, we derive an effective occupation-dependent model where
Figure 2. Phase diagrams showing the SF–MI transition for interaction strengths $a_s$ normalized to the lattice spacing $a$ (a) $a_s/a = 0.042$ and (b) $a_s/a = 0.014$. The latter corresponds to the background scattering length $a_s = 100 \, a_0$ of $^{87}$Rb at a lattice spacing of $a = 377$ nm [23]. The phase boundaries are plotted for the Bose–Hubbard model (black) and the full Hamiltonian equation (1) using mean-field (blue) and Gutzwiller (GW) (dashed red). The green line corresponds to the occupation-independent Hamiltonian equation (2).

the renormalization solely depends on the solution of the on-site problem. The well-known phase diagram for bosons in lattices describing the SF–MI transition is drastically altered. The presented results have direct relevance for optical lattice experiments, which can therefore serve as an ideal testing ground for beyond Hubbard physics.

After presenting the resulting phase diagram in section 2, we discuss the extensions of the Hubbard model individually starting with the effect of bond-charge interactions in section 3. The multi-orbital renormalization of the operators for conventional tunneling, bond-charge tunneling and on-site interaction is elaborated in sections 4–7. The basic idea is to separate the strongly correlated on-site problem from the inter-site dynamics. Firstly, we solve the on-site problem numerically exactly using the Wannier states of the lattice. Subsequently, we derive renormalized matrix elements for the multi-orbitally dressed operators. Finally, we calculate the phase diagram by treating the many-site problem with mean-field theory in section 8. Note that the multi-orbital renormalization can be used for various methods to treat the many-site problem and that the results are expected to be similarly affected.

2. Phase diagram and model Hamiltonian

We now first explain as one central result the phase diagram for bosonic atoms (figure 2), which is derived and further discussed afterwards. The black lines depict the Mott lobes predicted by the single-band Bose–Hubbard model applying mean-field theory [4]. Our calculations (colored lines) take into account the corrections to the tunneling as mentioned above as well as multi-orbital modifications of the on-site interaction. The phase diagrams are obtained for bosonic atoms of arbitrary species in cubic sinusoidal optical lattices with $V(\mathbf{r}) = V_0 \sum_i \cos^2(\pi r_i/a)$, where $i = \{x, y, z\}$, $a$ is the lattice spacing and $V_0$ is the lattice depth. We predict that the phase area of the MI is considerably reduced even for moderate interaction strengths and substantially deformed for stronger interactions. This corresponds to a significant shift of the critical lattice

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Figure 3. Critical lattice depths of the SF–MI transition for the Hubbard model (dashed), the full multi-orbital Hamiltonian equation (1) and the occupation-independent lowest-band Hamiltonian equation (2). The results are plotted as a function of the scattering length $a_s/a$ in units of the recoil energy $E_R = \hbar^2/8ma^2$.

depth of the SF–MI transition (figure 3) due to an effectively increased tunneling and reduced on-site interaction. The discrepancy can reach more than $5E_R$ for $n = 3$ particles per lattice site. We expect the shift to be observable in experiments with a filling $n \geq 2$ and strong or tunable interactions.

The mentioned extension to the Hubbard model leading to the phase diagram in figure 2 (red and blue lines) can be included in an effective Hamiltonian

$$\hat{H}_{\text{full}} = - \sum_{\langle i,j \rangle} \tilde{b}_i^\dagger \tilde{b}_j J_{\text{tot}}^{\tilde{n}_i,\tilde{n}_j} + \frac{1}{2} \sum_i U_{\text{MO}}^{\tilde{n}_i} \tilde{n}_i (\tilde{n}_i - 1)$$

with multi-orbitally renormalized creation and annihilation operators $\tilde{b}_i^\dagger$ and $\tilde{b}_i$, respectively, and the operator $\tilde{n}_i = \tilde{b}_i^\dagger \tilde{b}_i$ corresponding to the number of particles on site $i$. This Hamiltonian has the same structure as the Bose–Hubbard Hamiltonian [1, 2] but with multi-orbitally renormalized tunneling matrix elements $J_{\text{tot}}^{\tilde{n}_i,\tilde{n}_j}$ and on-site interactions $U_{\text{MO}}^{\tilde{n}_i}$ that explicitly depend on the site occupation $n_i$. In the following, we calculate these occupation-dependent parameters with exact diagonalization and derive the phase boundaries using a mean-field (MF) and a GW approach. Furthermore, we present a simplified Hamiltonian with occupation-independent parameters (green lines in figure 2), which is in good agreement with the full model above. This Hamiltonian is restricted to conventional and bond-charge tunneling in the lowest band

$$\hat{H}_{\text{occ.--indep.}} = - \sum_{\langle i,j \rangle} \left[ J \hat{b}_i^\dagger \hat{b}_j + J_{\text{BC}} \hat{b}_i^\dagger (\hat{n}_i + \hat{n}_j) \hat{b}_j \right] + \frac{\tilde{U}_2}{2} \sum_i \hat{n}_i (\hat{n}_i - 1)$$

with single-band operators $\hat{b}_i^\dagger$ and $\hat{b}_i$, the Hubbard tunneling $J$, lowest-band bond-charge interaction $J_{\text{BC}}$ and renormalized on-site interaction $\tilde{U}_2$ for two particles, as is elaborated in

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detail below. In general, the results for Hamiltonian (2) show that the lowest-band bond-
charge tunneling is an important contribution and its influence increases with the number of 
particles per site. Note that the mean-field and the GW approach fully coincide for occupation-
dependent models (green and black lobes), whereas there is a minimal discrepancy for 
the Hamiltonian (1) with occupation-dependent tunneling. In the mean-field calculation, the 
decoupling of the lattice sites requires the use of $J_{n_i,n_j}^\text{tot}$ as an approximation for $J_{n_i,n_j}^\text{tot}$ and 
$J_{n_i,n_j,\pm 1}^\text{tot}$ (see appendices F and G).

3. Bond-charge interactions

The two-particle interaction of bosonic particles in the lowest band of the lattice is given by 
$\hat{H}_{\text{int}} = \frac{1}{2} \sum_{ijkl} U_{ijkl} \hat{b}_i \hat{b}_j \hat{b}_k \hat{b}_l$ with interaction integrals

$$U_{ijkl} = g \int \text{d}^3r \, w_i^*(r)w_j^*(r)w_k(r)w_l(r)$$

and the Wannier functions $w_i(r) = w_i^0(r - r_i)$ of the lowest band, which are maximally 
localized at site $i$. Here, a $\delta$-interaction potential with repulsive interactions $g = \frac{4\pi \hbar^2}{m} a_s > 0$ is 
assumed. One of the key ingredients of the Hubbard model [1, 2, 44] is to restrict the two-
particle interaction to the dominating term, namely the on-site interaction $U = U_{iii}$. However, 
some of the neglected off-site processes correspond to scattering accompanied by hopping to 
a neighboring site (e.g. $\hat{b}_i^\dagger \hat{b}_j^\dagger \hat{b}_k \hat{b}_l$) known as bond-charge tunneling (figure 1(c)). It is essential 
that this process constitutes a hopping process which physically modifies the overall tunneling in 
the system. Thus, although this density-dependent hopping contribution is small in comparison 
with the on-site interaction $U$, it can be neglected only if also small in comparison with the 
conventional tunneling $J$. Considering two neighboring sites $i$ and $j$, the density-induced 
tunneling operator takes the form

$$\hat{J}_{BC} = -J_{BC} \hat{b}_j^\dagger (\hat{n}_i + \hat{n}_j) \hat{b}_j \quad \text{with} \quad J_{BC} = -U_{iii}$$

and $J_{BC} > 0$. Due to the structure of this operator, we can combine it with the conventional 
tunneling $J$ to an effective hopping

$$\hat{J}_{BC}^\text{eff} = -[J + J_{BC}(\hat{n}_i + \hat{n}_j - 1)] \hat{b}_j^\dagger \hat{b}_j.$$ 

As a consequence, the total tunneling increase is of the order of several tens of per cent for 
standard $^{87}$Rb conditions (see figure 4, dashed line) and even more for stronger interactions 
and deeper lattices. Note that density–density off-site interactions ($U_{ijij}$) [6] and correlated pair 
tunneling ($U_{ijij}$) are considerably weaker than bond-charge interactions (see appendix C).

The nature of bond-charge interactions can be intuitively illustrated as an effective potential. In equation (5), the term $\hat{n}_i + \hat{n}_j - 1$ corresponds to the density $\rho_{BC}(\mathbf{r}) = n_i |w_i(r)|^2 + (n_j - 1)|w_j(r)|^2$ on sites $i$ and $j$ excluding the hopping particle. By inserting explicit 
expressions for the integral $J_{BC}$ and the tunneling $J$, we can write the effective hopping 
operator (5) as

$$\hat{J}_{BC}^\text{eff} = \int \text{d}^3r \, w_j^*(r) \left( \frac{p_j^2}{2m} + V(r) + g\rho_{BC}(r) \right) w_j \hat{b}_j^\dagger \hat{b}_j.$$ 

We can now identify the expression $V(r) + g\rho_{BC}(r)$ as an effective tunneling potential, which 
is illustrated in figure 4(c). Since the density of the repulsively interacting particles is localized

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to the centers of the lattice sites, the effective potential corresponds to a shallower lattice, i.e. increased tunneling. Note that the effective potential can be used to determine the total tunneling via band structure calculations (see appendix E).

4. Multi-orbital tunneling

While the extension to the Hubbard model above is still within the lowest-band approximation, the following discussion covers the influence of higher orbitals. The calculation of the effective tunneling can be performed by taking two adjacent sites of the lattices into account (figure 1(d)). Writing initial and final states as a product of wave functions of the two sites allows us to express the renormalized tunneling in terms of single-site expectation values. Consequently, the renormalized parameters depend only on the solution of the single-site problem. The presented scheme allows us to derive the effective Hamiltonian (1) independently of the applied method for treating the on-site many-particle problem. Here, the orbital occupations are computed by exact diagonalization of \( n \) particles on a single site using a finite many-particle basis with a high-energy cut-off. The short-range interaction between the atoms is modeled as a box-shaped interaction potential (see appendix A for details). The single-particle orbitals are represented by the Wannier functions \( w^{(\alpha)}(r) \) with a 3D band index \( \alpha = (\alpha_x, \alpha_y, \alpha_z) \).

In the following, we consider multi-orbital hopping processes, where one atom tunnels from the left site (L) with initially \( n_L \) particles to the right site (R) with initially \( n_R \) particles (figure 1(d)). As a direct consequence, the total tunneling \( J_{n_L,n_R} \) becomes intrinsically occupation dependent. The multi-orbital hopping operator \( \hat{J}_\text{MO} = \sum_\alpha J_\alpha \hat{b}_R^{(\alpha \dagger)} \hat{b}_L^{(\alpha)} \) sums over all

\[ \alpha = (\alpha_x, \alpha_y, \alpha_z), \]

\[ J_\alpha = J^{\alpha}_\text{MO} + J^{\alpha}_\text{MO} + J^{\alpha}_\text{MO} + \cdots, \]

where the latter two scale with the prefactor \( \nu = n_i + n_j - 1 \). The deviations from the bare Hubbard tunneling \( J_0 \) are plotted as a function of (a) the lattice depth \( V_0 \) and (b) the interaction strength \( \alpha_s \). (c) The single-band bond-charge interaction gives rise to an effective tunneling potential (see text). (d) Qualitatively, the multi-orbital tunneling scales with the population of higher orbitals (shading) where the tunneling \( J_\alpha \) (arrows) is substantially enhanced in higher bands.

Figure 4. Contributions to the effective tunneling \( J_{n_i,n_j}^{\text{tot}} \) with \( n_i = n_j = 3 \) by multi-orbital tunneling \( J_{n_i,n_j}^{\text{MO}} \), bond-charge interactions \( J_{BC} \) and multi-orbital bond-charge interactions \( J_{BC,MO} \), where the latter two scale with the prefactor \( \nu = n_i + n_j - 1 \). The deviations from the bare Hubbard tunneling \( J_0 \) are plotted as a function of (a) the lattice depth \( V_0 \) and (b) the interaction strength \( \alpha_s \). (c) The single-band bond-charge interaction gives rise to an effective tunneling potential (see text). (d) Qualitatively, the multi-orbital tunneling scales with the population of higher orbitals (shading) where the tunneling \( J_\alpha \) (arrows) is substantially enhanced in higher bands.
possible multi-orbital processes
\[
J_α = - \int \mathbf{d}^3r u_R^{(α)σ} \left( \frac{\mathbf{p}^2}{2m} + V(\mathbf{r}) \right) w_L^{(α)}(\mathbf{r}),
\]
(7)
where tunneling takes place only between equal orbitals due to orthogonality relations. It is possible to reduce this complex multi-orbital operator to an effective hopping operator
\[
\hat{J}_{\text{eff}} = -\hat{b}_i^\dagger \hat{b}_j J_{\text{eff}}^{\text{MO}}_{j,i}.
\]
(8)
The occupation-dependent tunneling matrix elements are given by
\[
J_{\text{eff}}^{\text{MO}} = \frac{1}{\sqrt{n_L(n_R+1)}} \sum_α J_α \langle \Psi(n_R+1) \vert \hat{b}^{(α)\dagger} \vert \Psi(n_R) \rangle \langle \Psi(n_L-1) \vert \hat{b}^{(α)} \vert \Psi(n_L) \rangle,
\]
(9)
where \(\Psi(n)\) is the single-site wave function. This can be evaluated using the coefficients obtained by exact diagonalization as elaborated in appendix B.

The tunneling in higher bands \(J_α\) is drastically enhanced compared with the lowest band tunneling \(J_0\). Therefore, the contributions of higher-orbital processes are significant and grow exponentially with the lattice depth, although the population of higher bands is usually smaller than 1%. As an example, the matrix element \(J_{\text{eff}}^{\text{MO}}\) for three bosons per site is plotted in figure 4 (green lines). In general, the effective tunneling is increased except for the elements \(J_{\text{eff}}^{\text{MO}} = J_{\text{eff}}^{\text{MO}}\) where the tunneling is slightly reduced.

In an intuitive picture, the many-particle calculation can be reduced to the occupation of single-particle orbitals \(\langle \hat{n}^{(α)} \rangle\). The fraction of particles occupying higher orbitals \(α\) tunnels via the respective tunneling matrix elements \(J_α\), as indicated in figure 4(d). In this approximation, the effective tunneling can be estimated by the weighted sum (see appendix B)
\[
J_{\text{eff}}^{\text{MO}} \approx \frac{1}{\sqrt{n_L(n_R+1)}} \sum_α J_α \sqrt{\langle \hat{n}^{(α)}_L \rangle \langle \hat{n}^{(α)}_R \rangle}.
\]
(10)
This simplified approach is in qualitative agreement with the fully correlated calculation above.

5. Multi-orbital bond-charge interaction

It proves necessary to apply the same multi-orbital treatment of the normal tunneling developed in the last section to the bond-charge-assisted hopping. By analogy, we can define an effective multi-orbital bond-charge hopping
\[
\hat{J}_{\text{eff}}^{\text{BC,MO}} = -\hat{b}_i^\dagger (\hat{n}_i + \hat{n}_j) \hat{b}_j J_{\text{eff}}^{\text{BC,MO}}_{j,i},
\]
(11)
with an occupation-dependent parameter. It depends on the single-site matrix elements \(\langle \Psi(n-1) \vert \hat{b}^{(α)} \vert \Psi(n) \rangle\) and \(\langle \Psi(n-1) \vert \hat{b}^{(β)} \hat{b}^{(γ)} \hat{b}^{(δ)} \vert \Psi(n) \rangle\) as discussed in appendix D. The results shown in figure 4 for the multi-orbital bond-charge interaction (solid red line) differ substantially from the single-orbital case (dashed line) for intermediate and deep lattices. Processes involving higher bands start to dominate for deep lattices, rendering the lowest-band calculation invalid and can even turn the sign of the bond-charge contributions negative. Processes of the type \(J_{n,0}^{\text{BC,MO}}\) are only weakly influenced by multi-orbital corrections.
Figure 5. Occupation-dependent total tunneling $J_{n_i,n_j}^{\text{tot}}$ and on-site interactions $U_{n_i,n_j}^{\text{MO}}$ as a function of (a) the lattice depth $V_0$ and (b) the interaction strength $a_s$. The empirical fit $\tilde{U}_2$ is plotted as a dashed black line. Note that $J_{n_i,n_j}$ and $J_{n_i+1,n_j-1}$ describe time-reversed hopping processes and thus are equivalent. For an explanation of the error bars see text.

6. Effective tunneling

The total tunneling consists of normal and bond-charge-assisted tunneling, both effectively including higher orbital processes. The total tunneling

$$J_{n_i,n_j}^{\text{tot}} = J_{n_i,n_j}^{\text{MO}} + (n_i+n_j-1)J_{n_i,n_j}^{\text{BC,MO}}$$

and the individual contributions are shown in figures 4(a) and (b). The occupation-dependent enhancement of the tunneling is depicted in figure 5. The deviation from the Hubbard tunneling $J$ can easily reach 30% for three particles per site at moderate $a_s$ and $V_0$. The data in this figure are calculated using a box-shaped interaction potential (see appendix A) with a width $W = 5 \text{ nm}$ and a lattice constant $a = 377 \text{ nm}$. It is important to note that the results are almost independent of $W$, provided that $W$ is much smaller than the lattice constant. The error bars in figure 5 show the results for $W = 25 \text{ nm}$ and $W \to 0$.

Whereas the normal tunneling is increased by the multi-orbital renormalization, the density-induced tunneling is reduced. Coincidentally, both multi-orbital corrections compensate for each other and the overall deviations can be approximated surprisingly well by the single-band bond-charge tunneling for moderate particle numbers (dashed line in figure 4). This justifies the applicability of the model Hamiltonian (2), where tunneling is restricted to the lowest band, i.e. $J_{n_i,n_j}^{\text{tot}} \approx J + (n_i+n_j-1)J_{\text{BC}}$.

7. Occupation-number-dependent on-site interaction

In a similar fashion as the inclusion of higher orbitals alters the tunneling, also the on-site interaction $U$ is modified. The orbital degree of freedom decreases the on-site
interaction [34–36] as the particles tend to avoid each other. The results for \( U_n^{MO} \), representing the eigenvalue of the single-site exact diagonalization [35], are depicted for occupation numbers \( n = 2–4 \) in figure 5. Note that for strong interactions, it may be necessary to use realistic interaction potentials. It is convenient to express \( U_2^{MO} \) within equation (2) by a fitted empirical function \( \tilde{U}/U = \lambda_1 + (\lambda_2 + \lambda_3 a_s/a) \sqrt{V_0/E_R + \lambda_4 e^{S_M/K(a_s/a)}} \), which describes the on-site interaction for two particles but also serves as an approximation for three and four particles.

8. Mean-field/Gutzwiller phase diagrams

The SF–MI phase diagram (figure 2) for the extended Hamiltonian (1) is derived within mean-field theory [2, 4]. By introducing the SF order parameter \( \psi = \langle \hat{b}_i \rangle = \langle \hat{b}^\dagger_i \rangle \) and by neglecting the fluctuations of quadratic order, the lattice sites are decoupled (see appendix F). Within second-order perturbation theory for the Mott state with \( n \) particles, we find an analytical expression for the SF–MI transition point

\[
\left| | n \rangle | \tilde{J}_{BC}(n-1) + 1 |^2 + (n+1) | \tilde{J}_{BC} n + 1 |^2 \right| E_0(n) - E_0(n-1) + 1 = 0
\]

with \( \tilde{J}_{BC} = J_{BC,MO}/J_{n,n}^{MO} \) and the unperturbed energy

\[
E_0(n) = (1/2) U_n^{MO} (n-1) - \mu)n/n \tilde{J}_{BC}^{MO}.
\]

Due to the decoupling in the mean-field approach, only tunneling processes with \( n_i = n_j = n \) can be accounted for. Thus, this approach cannot fully implement the occupation-number-dependent Hamiltonian (1). This can be achieved by using the GW approach, where the energy functional is minimized with respect to the coefficients \( f_n \) of the trial wave function \( |G\rangle = \prod_i \sum_n f_n |n_i\rangle \) (see appendix G). The GW results are depicted in figure 2 (red lines) and show nearly perfect agreement with the mean-field calculation (blue lines). The mean-field solution (13) can also be applied to the simplified occupation-independent model (2) using the lowest-band tunneling parameters \( J \) and \( \tilde{J}_{BC} = J_{BC}/J \) as well as the fitted on-site interaction \( \tilde{U} \).

9. Conclusions

We have presented an occupation-dependent model Hamiltonian incorporating multi-orbitally renormalized tunneling, density-induced tunneling and on-site interaction. The effectively increased tunneling and reduced on-site interaction cause a substantial modification of the Bose–Hubbard phase diagram for bosonic atoms. In addition, we have discussed a simplified occupation-independent model that includes the lowest-band bond-charge tunneling and reproduces the phase diagram well. In general, we have derived a multi-orbital renormalization procedure for one- and two-particle processes. It is capable of describing strongly correlated optical lattice systems accurately that cannot be calculated correctly by means of variational mean-field treatments [25–28]. Furthermore, it is important to include both bond-charge tunneling and multi-orbital corrections as extensions of the Hubbard Hamiltonian. Several effects such as the reduction of the bond-charge tunneling cannot be described with an effective

\[ \text{The parameters are } \lambda_1 = 0.91, \lambda_2 = -0.012 E_R^{-1}, \lambda_3 = -0.80(E_R a_s)^{-1}, \lambda_4 = 0.14 \text{ and } \lambda_5 = -34 a_s^{-1}. \]

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wave function. The presented results can be transferred to fermionic systems, quantum gas mixtures, low-dimensional systems, long-range interactions and different lattice geometries. In these systems, the occupation-dependent tunneling may also lead to novel quantum phases. Related work on defining a basis with renormalized operators can be found in [45].

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Appendix A. Multi-orbital interactions

The treatment of multi-band Hamiltonians for lattices represents a challenging and still open problem posed by its complexity. A common approach is the restriction to the lowest band, which neglects interaction-induced admixtures of higher orbitals. In order to account for orbital degrees of freedom, we focus on two neighboring sites of the lattice, where we apply a multi-orbital diagonalization, the so-called configuration interaction method, to each site individually (see figure 1(d), main text). In this approach, off-site interactions are neglected as the occupation of higher orbitals is mainly caused by the local on-site interaction. The 3D Wannier functions

$$w^{(a)}_{L/R}(r) = w^{(a)}(r - r_{L/R})$$ (A.1) constitute the single-particle orbitals, where $r_{L/R} = (\pm \frac{1}{2}a, 0, 0)$ is the coordinate of the left and the right site, respectively, $a$ is the lattice spacing and

$$\alpha = (\alpha_x, \alpha_y, \alpha_z)$$ (A.2)

is the 3D band index. The maximally localized Wannier functions are obtained by means of band-structure calculations for a separable cubic lattice potential $V(r) = V_0 \sum_i \cos^2(\pi r_i / a)$ with the lattice depth $V_0$. To include correlations of particles fully, a many-particle Fock basis $|N\rangle = |n_0, n_1, \ldots \rangle$ with $n$ particles is used, where $n_i$ is the number of particles in orbital $i$. The Hamiltonian for a single lattice site reads

$$\hat{H}_{\text{site}} = \sum_a \epsilon^{(a)} \hat{n}^{(a)} + \frac{1}{2} \sum_{a \beta \gamma \delta} U^{(a \beta \gamma \delta)} \hat{b}^{(a)\dagger} \hat{b}^{(\beta)\dagger} \hat{b}^{(\gamma)} \hat{b}^{(\delta)},$$ (A.3)

where $\hat{n}^{(a)} = \hat{b}^{(a)\dagger} \hat{b}^{(a)}$, $\hat{b}^{(a)\dagger}$ creates and $\hat{b}^{(a)}$ annihilates a particle in the Wannier orbital $a$ with single-particle energies $\epsilon^{(a)}$. In general, the ground state of the single-site problem with $n$ particles is given as a superposition of Fock states

$$|\Psi(n)\rangle = \sum_N c_N(n) |N(n)\rangle$$ (A.4)

with real coefficients $c_N(n)$. Since the Hamiltonian (A.3) preserves parity the ground state consists of many-particle states with even parity. The lowest eigenvalue of the diagonalized Hamiltonian matrix directly corresponds to the occupation-dependent on-site interaction $U_{n}^{\text{MO}}$ as discussed in the main text.
We model the short-range interaction between the atoms as a box-shaped interaction potential

$$V(r - r') = \frac{\tilde{g}}{(2W)^3} \prod_{i=1}^{3} \theta (W - |r_i - r'_i|)$$  \hspace{1cm} (A.5)

with the Heaviside step function $\theta$ and a variable box width $W$. The influence of the finite interaction range depends only on the ratio $W/a$ and increases with the lattice depth since the wave functions are contracted. However, for $W \ll a$ the results (see figure 5, main text) are almost independent of $W$. The interaction integrals in equation (A.3) are given by

$$U^{(\alpha\beta\gamma\delta)}(\tilde{g}, W) = \int d^3r \int d^3r' \, w^{(\alpha)*}(r) \, w^{(\beta)*}(r') \, V(r - r') \, w^{(\gamma)}(r') \, w^{(\delta)}(r),$$  \hspace{1cm} (A.6)

where $\tilde{g}$ is fixed by the condition $U^{(0000)}(\tilde{g}, W) = g \int d^3r |w(r)|^4$ with $g = \frac{4\pi h^2}{m} a_s$ and s-wave scattering length $a_s$. Note that the $\delta$-interaction potential can be understood as the limiting case of the box interaction potential. However, it behaves differently if including an infinite number of orbitals requiring a regularization procedure as described in [33] (for numerical diagonalization see [46]). For real interaction potentials, the mathematical subtlety of the $\delta$-interaction is not present and under certain experimental conditions [35] decay processes in higher orbitals may even effectively lead to a finite Hilbert space. Here, for the numerical diagonalization, nine bands are accounted for in each spatial direction restricted to the energetically lowest $6 \times 10^3 \times n^2$ many-particle states.

**Appendix B. Multi-orbital tunneling**

In the following, we consider hopping processes, where one atom tunnels from the left site (L) with initially $n_L$ particles to the right site (R) with initially $n_R$ particles (see figure B.1(d), main text). The tunneling operator accounting for all possible orbital hopping processes reads

$$\hat{J}^{\text{MO}} = \sum_{\alpha\beta} J_{\alpha\beta} \hat{b}_{\text{R}}^{(\alpha)*} \hat{b}_{\text{L}}^{(\beta)},$$

with tunneling matrix elements

$$J_{\alpha\beta} = - \int d^3r \, w^{(\alpha)*}_{\text{R}}(r) \left( \frac{p^2}{2m} + V(r) \right) w^{(\beta)}_{\text{L}}(r).$$  \hspace{1cm} (B.1)

Because of orthogonality relations, only matrix element with $\alpha = \beta$ have nonzero contributions. Thus, the tunneling between the left and the right site simplifies to

$$\hat{J}^{\text{MO}} = \sum_{\alpha} J_{\alpha} \hat{b}_{\text{R}}^{(\alpha)*} \hat{b}_{\text{L}}^{(\alpha)}$$  \hspace{1cm} (B.2)

with matrix elements $J_{\alpha} = J_{\alpha\alpha}$, which only depend on the band index $\alpha_x$ in the $x$-direction. The respective tunneling matrix elements $J_{\alpha}$ for the band $\alpha_x$ are plotted in figure B.1, which are positive for even bands and negative for odd bands. The tunneling amplitude is strongly enhanced for higher orbitals due to the considerably larger overlap of the Wannier functions.

As the occupation of higher orbitals is decoupled from off-site interactions and tunneling in our model, it is possible to reduce the complex multi-band hopping to an effective hopping $J_{n_L,n_R}^{\text{MO}}$. It describes the transition from an initial state $|\Psi_I\rangle$ to the final state $|\Psi_F\rangle$ with

$$|\Psi_I\rangle = |\Psi(n_L)\rangle_{\text{L}} \otimes |\Psi(n_R)\rangle_{\text{R}},$$

$$|\Psi_F\rangle = |\Psi(n_L - 1)\rangle_{\text{L}} \otimes |\Psi(n_R + 1)\rangle_{\text{R}}$$  \hspace{1cm} (B.3)
Figure B.1. Single-particle tunneling matrix elements $J_\alpha$ for the lowest five bands as a function of the lattice depth $V_0$ and in units of the recoil energy $E_R = \hbar^2/8ma^2$.

(see figure B.1(d)). As discussed in the main text, the effective multi-orbital tunneling can be defined as

$$J_{\text{eff}}^{\text{MO}} = -\hat{b}_{\text{R}}^{\dagger}J_{\text{nL},\text{nR}}^{\text{MO}}$$  \hspace{1cm} (B.4)$$

with

$$J_{\text{nL},\text{nR}}^{\text{MO}} = \frac{\langle \Psi_{1} | J^{\text{MO}} | \Psi_{1} \rangle}{\sqrt{n_{\text{L}}(n_{\text{R}} + 1)}}. \hspace{1cm} (B.5)$$

As the initial and final states (B.3) are product states, the expression separates into two terms for the left- and the right-hand sites

$$\langle \Psi_{1} | J^{\text{MO}} | \Psi_{1} \rangle = \sum_{\alpha} J_{\alpha} \langle \Psi(n_{\text{R}} + 1) | \hat{b}^{(\alpha)^{\dagger}} | \Psi(n_{\text{R}}) \rangle \langle \Psi(n_{\text{L}} - 1) | \hat{b}^{(\alpha)} | \Psi(n_{\text{L}}) \rangle_{\text{L}}. \hspace{1cm} (B.6)$$

Using the many-particle ground state (A.4) obtained by exact diagonalization, we can define

$$j_{n}^{(\alpha)} = \langle \Psi(n - 1) | \hat{b}^{(\alpha)^{\dagger}} | \Psi(n) \rangle = \sum_{N/N'} c_{N}(n - 1) c_{N}(n) \langle N'(n - 1) | \hat{b}^{(\alpha)} | N(n) \rangle. \hspace{1cm} (B.7)$$

Note that since the coefficients $c_{N}$ can be chosen to be real, the conjugated term reads

$$\langle \Psi(n - 1) | \hat{b}^{(\alpha)^{\dagger}} | \Psi(n) \rangle = j_{n-1}^{(\alpha)} = j_{n-1}^{(\alpha)}.$$ The effective tunneling can thus be written as

$$J_{\text{nL},\text{nR}}^{\text{MO}} = \frac{1}{\sqrt{n_{\text{L}}(n_{\text{R}} + 1)}} \sum_{\alpha} j_{\alpha} j_{n_{\text{L}}}^{(\alpha)} j_{n_{\text{R}}+1}^{(\alpha)}. \hspace{1cm} (B.8)$$

Due to parity conservation of the Hamiltonian (A.3), the ground states at the left and the right site are superpositions of Fock states $|N(n)\rangle$ with even parity. Tunneling in an odd orbital would alter the parity of both sites, resulting in final states with vanishing coefficients. Consequently,
tunneling can only occur in even orbitals $\alpha$ with positive tunneling matrix elements $J_\alpha$. It turns out that the effective tunneling is increased by the tunneling in higher bands with the exception of the processes $J_{n,0}^{\text{MO}} = J_{1,0}^{\text{MO}} - 1$. Here, the lack of interaction in the left or the right site prohibits tunneling via higher bands. The interaction at one site, however, depopulates the Fock state $|n, 0, 0, \ldots\rangle$, which results in slightly lowered tunneling energy. The process $J_{1,0}^{\text{MO}}$, where no interactions are present, is equivalent to the uncorrelated tunneling $J = J_0$.

Intuitively, the mean-field wave functions of the left and right sites

\[
\psi_{\text{MF}}^L(r) = \sum_\alpha c_n^\alpha \psi_{\alpha}^L(r),
\]

and

\[
\psi_{\text{MF}}^R(r) = \sum_\alpha c_n^{\alpha+1} \psi_{\alpha}^R(r),
\]

can be used to calculate the multi-orbital tunneling \cite{28}. The coefficients can be obtained from the orbital occupation numbers $c_n^\alpha = \sqrt{\langle \hat{n}_n^{\alpha} \rangle} / \sqrt{n}$, with the expectation value $\langle \hat{n}_n^{\alpha} \rangle = \langle \Psi(n) | \hat{n}_n^{\alpha} | \Psi(n) \rangle$. This allows us to estimate the multi-orbital by

\[
J_{\mu, n}^{\text{MO}} \approx \int d^3r \psi_{\text{MF}}^\dagger R(r) \left( \frac{p^2}{2m} + V(r) \right) \psi_{\text{MF}}^\dagger L(r)
\]

\[
= \sum_\alpha c_n^\alpha c_n^{\alpha+1} J_\alpha.
\]

### Appendix C. Off-site interactions

The full lowest-band interaction

\[
\hat{H}_{\text{int}} = \frac{1}{2} \sum_{ijkl} U_{ijkl} \hat{b}_i^\dagger \hat{b}_j^\dagger \hat{b}_k \hat{b}_l
\]

with interaction integrals \eqref{eq:interaction_integrals} is commonly restricted to the on-site interaction. However, off-site interactions between neighboring sites can reach the same order of magnitude as the tunneling. Expanding the lowest-band Hamiltonian accordingly leads to three distinct physical processes, namely bond-charge interaction (density-dependent hopping), correlated pair tunneling and density–density interaction \cite{16–19,37}. While the latter process has to be compared with the on-site interaction, the other two constitute tunneling processes. The full lowest-band Hamiltonian with nearest-neighbor interaction reads

\[
\hat{H} = -J \sum_{\langle i, j \rangle} \hat{b}_i^\dagger \hat{b}_j - J_{\text{BC}} \sum_{\langle i, j \rangle} \hat{b}_i^\dagger (\hat{n}_i + \hat{n}_j) \hat{b}_j + J_{\text{pair}} \sum_{\langle i, j \rangle} \hat{b}_i^\dagger \hat{b}_j^2 + \frac{1}{2} U \sum_i \hat{n}_i (\hat{n}_i - 1) + V \sum_{\langle i, j \rangle} \hat{n}_i \hat{n}_j
\]

with matrix elements $U = U_{i\bar{i}i\bar{i}}$, $J_{\text{BC}} = -U_{i\bar{i}ij}$, $J_{\text{pair}} = U_{i\bar{i}jj}$ and $V = U_{ijij}$.

In figure C.1 these parameters are plotted as a function of the lattice depth. The off-site density–density interaction is very small compared to the on-site interaction and can consequently be neglected. This also applies for the correlated pair tunneling, which is even negligible when compared with the single-particle tunneling matrix element $J$. The bond-charge tunneling matrix element, however, reaches 10% of the conventional tunneling amplitude for
Figure C.1. Lowest-band parameters for on-site interaction $U$, tunneling $J$, bond-charge tunneling $J_{BC}$, correlated pair-tunneling $J_{\text{pair}}$ and density–density interaction $V$.

intermediate and deep lattices. In addition, it scales with the total particle number and can thus be a significant contribution to the total tunneling. Note that the multi-orbital renormalization can influence the individual parameters very strongly.

Appendix D. Multi-orbital bond-charge tunneling

In analogy to the multi-orbital tunneling, the orbital degree of freedom also affects the bond-charge interaction. The single-band bond-charge or density-induced tunneling is introduced in the main text (see equation (3) and figure 1(b)). As discussed before, we can restrict the calculation to two neighboring sites and write the multi-orbital bond-charge operator (see figure 1(c), main text) as

$$\hat{J}_{BC, MO} = \frac{1}{2} \sum_{\alpha \beta \gamma \delta} \hat{b}^{(\alpha \dagger)}_R (J_{\alpha \beta \gamma \delta}^{BC} \hat{b}^{(\beta \dagger)}_L \hat{b}^{(\gamma)}_L + P \left(J_{\beta \gamma \alpha \delta}^{BC} \hat{b}^{(\beta \dagger)}_R \hat{b}^{(\gamma)}_R \hat{b}^{(\delta)}_L \right) \hat{b}^{(\delta)}_L \right)$$

with

$$J_{\alpha \beta \gamma \delta}^{BC} = -\int d^3r \int d^3r' w^{(\alpha)}_R(r) w^{(\beta)}_L(r') V(r-r') w^{(\gamma)}_L(r') w^{(\delta)}_L(r)$$

$$= -P \int d^3r \int d^3r' w^{(\alpha)}_L(r) w^{(\beta)}_R(r') V(r-r') w^{(\gamma)}_R(r') w^{(\delta)}_R(r),$$

where the Wannier functions are chosen to be real and the sign $P = (-1)^{\alpha + \beta + \gamma + \delta}$ is parity dependent. Note that for a $\delta$-interaction potential, we have $J_{\alpha \beta \gamma \delta}^{BC} = J_{\alpha \beta \gamma \delta}^{BC}$. The effective multi-orbital bond-charge tunneling reads

$$\hat{J}_{BC, MO}^{\text{eff}} = -\hat{b}^\dagger_R (\hat{n}_R + \hat{n}_L) \hat{b}_L J_{\alpha \beta \gamma \delta}^{BC, MO} \hat{b}^{(\delta)}_L$$
with occupation-number-dependent matrix elements

\[ J^{BC,MO}_{n_L,n_R} = \frac{\langle \Psi_f | J^{BC,MO} | \Psi_i \rangle}{\sqrt{n_L(n_R + 1)(n_L + n_R - 1)}}. \]  

(D.4)

Using that \( \Psi_i \) and \( \Psi_f \) are product states as well as the definitions (B.7) and

\[ j_n^{(\beta \gamma \delta)} = \langle \Psi(n - 1) | \hat{b}^{(\beta \gamma \delta)} \hat{b}^{\dagger (\gamma \delta)} | \Psi(n) \rangle, \]  

(D.5)

the effective matrix elements can be computed by

\[ J^{BC,MO}_{n_L,n_R} = \sum_{a \beta \gamma \delta} \frac{1}{2} \left( J^{BC,MO}_{a \beta \gamma \delta} + J^{BC,MO}_{a \beta \delta \gamma} \right) \frac{\left[ J^{(a)}_{n_L} j^{(\beta \gamma \delta)}_{n_R+1} + j^{(a)}_{n_R+1} j^{(\beta \gamma \delta)}_{n_L} \right]}{\sqrt{n_L(n_R + 1)(n_L + n_R - 1)}}. \]  

(E.6)

Note that \( \langle \Psi(n - 1) | \hat{b}^{(\beta \gamma \delta)} \hat{b}^{\dagger (\gamma \delta)} | \Psi(n) \rangle = j_{n-1}^{(\beta \gamma \delta)} \) and that because of the same arguments as in the last section, the orbitals \( \alpha \) and \( \beta + \gamma + \delta \) need to be even; thus the sign \( P \) in equation (D.1) vanishes.

### Appendix E. Effective potential analogy

Considering bond-charge tunneling between neighboring sites \( i \) and \( j \) with the restriction to the lowest band, the bond-charge hopping simplifies to

\[ \hat{J}_{BC} = \hat{b}_{i}^{\dagger} \left( U_{iiij} \hat{b}_{i}^{\dagger} \hat{b}_{i} + U_{ijji} \hat{b}_{j}^{\dagger} \hat{b}_{j} \right) \hat{b}_{j} \]  

(see equation (4), main text), where \( \hat{b}_{i}^{\dagger} = \hat{b}_{i}^{(0 \gamma \delta)} \) and \( \hat{b}_{i} = \hat{b}_{i}^{(0)} \) are lowest-band creation and annihilation operators, respectively. The interaction integral reads

\[ U_{ijkl} = g \int d^{3}r w_{i}^{*}(r) w_{j}^{*}(r) w_{l}(r) w_{j}(r) \]  

(E.2)

with the lowest-band Wannier functions \( w_{i}(r) = w_{i}^{(0)}(r - r_{i}) \). By inserting the integral expressions (E.2) in equation (E.1) and using the commutation relations for \( \hat{b}_{i}^{\dagger} \) and \( \hat{b}_{i} \), the bond-charge operator becomes

\[ \hat{J}_{BC} = \hat{b}_{i}^{\dagger} \hat{b}_{j} g \left( \int d^{3}r w_{i}^{*}(r) \hat{n}_{i} | w_{i}(r) |^{2} w_{j}(r) + \int d^{3}r w_{j}^{*}(r) (\hat{n}_{j} - 1) | w_{j}(r) |^{2} w_{j}(r) \right). \]  

(E.3)

In order to combine bond-charge hopping and conventional tunneling \( \hat{J} = -J \hat{b}_{i}^{\dagger} \hat{b}_{j} \) with

\[ J = - \int d^{3}r w_{i}^{*}(r) \left( \frac{p^{2}}{2m} + V(r) \right) w_{j}(r), \]  

we define a reduced density \( \hat{\rho}_{ij} = \hat{n}_{i} | w_{i}(r) |^{2} + (\hat{n}_{j} - 1) | w_{j}(r) |^{2} \) on neighboring sites \( i \) and \( j \) excluding the hopping particle. Omitting the hopping particle corresponds to the exclusion of self-interactions. As a consequence, the total tunneling can be written as

\[ \hat{J}_{BC}^{eff} = \hat{J} + \hat{J}_{BC} \]  

(E.5)

\[ = \int d^{3}r w_{i}^{*}(r) \left( \frac{p^{2}}{2m} + V(r) + g \hat{\rho}_{ij} \right) w_{j}(r) \hat{b}_{i}^{\dagger} \hat{b}_{j}. \]

In this integral, \( \hat{\rho}_{ij} \) can be replaced by \( \hat{\rho}_{ij} \rightarrow \rho(r) - | w_{j}(r) |^{2} \) with the density \( \rho(r) = \sum_{i} n_{i} | w_{i}(r) |^{2} \) of all particles in the lattice, since the Wannier functions are localized. Finally, we can identify
the effect of the bond-charge interaction as a normal tunneling within the effective potential $V_{\text{eff}} = V(\mathbf{r}) + g(\rho(\mathbf{r}) - |w_i|^2)$ experienced by the hopping particle (see figure 4(c), main text). However, since $U_{iiij} = U_{ijji}$, one may also define $\hat{\rho}_{ij} = (\hat{n}_i - \frac{1}{2})|w_i|^2 + (\hat{n}_j - \frac{1}{2})|w_j|^2$. In this case, the effective potential for a homogeneous filling $n \geq 1$ can be written as $V(\mathbf{r}) + g \sum_i (n - \frac{1}{2})|w_i|^2$. This form of the potential can be used to perform a band structure calculation in the effective potential [21]. Note, however, that in general the effective potential is not separable into its spatial directions. Furthermore, in a band structure calculation the Wannier functions are adapted to the effective potential whereas in equation (E.5) the Wannier functions of the bare lattice potential, are used.

Appendix F. The mean-field approach

The full Hamiltonian including the bond-charge interaction as well as the multi-orbital corrections to the tunneling, bond-charge and on-site interactions reads

$$\hat{H} = - \sum_{\langle i, j \rangle} \tilde{b}^\dagger_i \tilde{b}_j J_{\text{MO}}^{\text{BC},\langle i, j \rangle} - \sum_{\langle i, j \rangle} \tilde{b}^\dagger_i (\tilde{n}_i + \tilde{n}_j) \tilde{b}_j J_{\text{MO}}^{\text{MO},\langle i, j \rangle} + \frac{1}{2} \sum_i U_{\text{MO}}^{\text{MO}} \tilde{n}_i (\tilde{n}_i - 1) - \mu \sum_i \tilde{n}_i$$  \hspace{1cm} (F.1)

(see equation (1), main text), where $\langle i, j \rangle$ denotes nearest-neighbor tunneling and $\mu$ is the chemical potential. The hopping in the renormalized multi-orbital basis is described by $\tilde{b}^\dagger_i \tilde{b}_j$, while $\tilde{n}_i$ corresponds to the number of particles on site $i$. In order to apply mean-field theory (see [2, 4]), we introduce the SF order parameter $\psi = \langle \tilde{b}_i \rangle = \langle \tilde{b}^\dagger_i \rangle$, where $\psi \neq 0$ corresponds to the SF phase and $\psi = 0$ defines the Mott insulator (MI) with a fixed number of atoms per lattice site. The decoupling of the lattice sites is achieved by neglecting the fluctuations between $\tilde{b}^\dagger_i$ and $\tilde{b}_j$ of quadratic order, i.e.

$$\tilde{b}_i \tilde{b}_j \approx \tilde{b}_i^\dagger \tilde{b}_j - (\tilde{b}_i^\dagger - \langle \tilde{b}_i^\dagger \rangle)(\tilde{b}_j - \langle \tilde{b}_j \rangle) = \psi (\tilde{b}_i^\dagger + \tilde{b}_j) - \psi^2.$$  \hspace{1cm} (F.2)

However, the occupation-number-dependent tunneling parameter still couples the sites and it is necessary to approximate

$$\sum_{\langle i, j \rangle} (\psi (\tilde{b}_i^\dagger + \tilde{b}_j) - \psi^2) J_{\text{MO}}^{\text{MO},\langle i, j \rangle} \approx z \sum_i (\psi (\tilde{b}_i^\dagger + \tilde{b}_i) - \psi^2) J_{\text{MO}}^{\text{MO},\langle i, i \rangle},$$  \hspace{1cm} (F.3)

which corresponds to substituting $J_{\text{MO}}^{\text{MO},\langle i, j \rangle}$ with $J_{\text{MO}}^{\text{MO},\langle i, i \rangle}$. This is necessary, as the fluctuations are described by a single parameter $\psi$, which has no internal structure that would allow for the implementation of an occupation-dependent tunneling amplitude. This can be circumvented by using the GW approach, where the information on the population of the individual Fock states is preserved (see appendix G). However, we find that this approximation causes only small deviations. Analogously, we neglect quadratic fluctuations in the bond-charge term between neighboring lattice sites. Disregarding terms of the order $O(\psi^3)$ we find that

$$\tilde{b}_i^\dagger \tilde{n}_i \tilde{b}_j + \tilde{b}_i^\dagger \tilde{n}_j \tilde{b}_j \approx \psi (\tilde{b}_i^\dagger \tilde{n}_i + \tilde{n}_j \tilde{b}_j).$$  \hspace{1cm} (F.4)

We perform second-order perturbation theory in $\psi$ of a Mott lobe with $n$ particles per site and thus restrict the tunneling to the symmetric terms $J_{\text{MO}}^{\text{MO}}$ and $J_{\text{BC},\text{MO}}^{\text{MO}}$. This results in a decoupled single-site Hamiltonian $\hat{H}_i^{\text{eff}}/z J_{\text{MO}}^{\text{MO},\langle n, n \rangle} = \hat{H}_0 + \psi \hat{V}$ with

$$\hat{H}_0 = \frac{1}{2} \tilde{U}_n \tilde{n} (\tilde{n} - 1) - \tilde{\mu} \tilde{n} + \psi^2,$$

$$\hat{V} = -(\tilde{b}_i^\dagger + \tilde{b}_i) - (\tilde{b}_i^\dagger \tilde{n} + \tilde{n} \tilde{b}_i) J_{\text{MO}}^{\text{BC},\langle n, n \rangle},$$  \hspace{1cm} (F.5)

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z = 6 the number of nearest neighbors, and $(\tilde{U}_{\bar{n}}, J_{n,n}^{BC}, \tilde{\mu}) = (U_{\bar{n}}^{MO}, J_{n,n}^{BC,MO}, \mu)/z J_{n,n}^{MO}$. The unperturbed energy for a Mott state $|n\rangle = \frac{1}{\sqrt{n!}} \tilde{n}^{|n|} |0\rangle$ with $n$ particles is given by

$$E_0(n) = \langle n | \hat{H}_0 | n \rangle = \frac{1}{2} \tilde{U}_n n(n-1) - \tilde{\mu} n.$$  \hspace{1cm} (F.6)

The perturbation series up to third order in $\psi$ reads

$$E(\Psi) = E_0(n) + E_2(n) \Psi^2 + O(\Psi^4)$$  \hspace{1cm} (F.7)

with the second-order correction

$$E_2(n) = \frac{n | J_{n,n}^{BC} (n-1) + 1 |^2}{E_0(n) - E_0(n-1) + E_0(n) - E_0(n+1)} + 1.$$  \hspace{1cm} (F.8)

The boundary of the SF–MI phase transition is given by the Landau criterion $E_2(n) = 0$ for second-order phase transitions, which can be solved for $\mu$ (see figure 1, main text).

**Appendix G. The Gutzwiller approach**

In the mean-field approach elaborated above, only the tunneling matrix elements $J_{n,n}^{MO}$ and $J_{n,n}^{BC,MO}$ are taken into account due to the decoupling approximation. Using a GW approach [3, 47], the full occupation-dependent Hamiltonian (F.1) can be evaluated without this restriction. In this approach, the trial wave function

$$|G_i\rangle = \sum_{n=0}^{\infty} f_n |n\rangle_i$$  \hspace{1cm} (G.1)

with coefficients $f_n$ is used, which assumes that the wave function at each site $i$ can be written as a sum of local Fock states. In a homogeneous lattice with $M$ sites, all sites are equivalent and we can write $|G\rangle = \prod_{i=1}^{M} |G_i\rangle$. The Hamiltonian (F.1) can be split into four parts, namely

$$\hat{H} = \hat{J}_{\text{eff}} + \hat{J}_{\text{eff}}^{BC} + \hat{U} + \hat{\mu}.$$  \hspace{1cm} (G.2)

The expectation values for this Hamiltonian, depending exclusively on the expectation values for two adjacent sites $i$ and $j$, are given by

$$\langle \hat{J}_{\text{eff}} \rangle = -z M \langle G_i G_j | \tilde{b}^-_i \tilde{b}^-_j J_{n_i,n_j}^{MO} | G_i G_j \rangle$$

$$= -z M \sum_{n_i,n_j} f_{n_i+1} f_{n_j+1} f_{n_i} f_{n_j} \sqrt{n_i + 1} \sqrt{n_j + 1} J_{n_i+1,n_j}^{MO},$$  \hspace{1cm} (G.3)

$$\langle \hat{J}_{\text{eff}}^{BC} \rangle = -z M \langle G_i G_j | \tilde{b}^-_i (n_i + 1) \tilde{b}^-_j J_{n_i,n_j}^{BC,MO} | G_i G_j \rangle$$

$$= -z M \sum_{n_i,n_j} f_{n_i+1} f_{n_j+1} f_{n_i} f_{n_j} (n_i + 1) \sqrt{n_i + 1} \sqrt{n_j + 1} J_{n_i+1,n_j}^{BC,MO},$$  \hspace{1cm} (G.4)

$$\langle \hat{U} \rangle = \frac{1}{2} M \sum_n U_n n(n-1) |f_n|^2,$$  \hspace{1cm} (G.5)

$$\langle \hat{\mu} \rangle = -M \sum_n \mu n |f_n|^2.$$  \hspace{1cm} (G.6)
with \( z = 6 \) nearest neighbors. The system is in an MI state when all fluctuations vanish, i.e. when \( |G⟩ = ∏ |n⟩ \) is the energetically lowest state. The resulting GW phase diagram is plotted together with the mean-field results in figure 1 in the main text. Both the approaches, GW and mean-field, are in almost perfect agreement.

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