A moiré superlattice is formed when two or more van der Waals layers are stacked with small differences in lattice constant or orientation. When the isolated layers are semiconductors or semimetals, the electronic properties of the bilayer are accurately described by continuum models that have the periodicity of the moiré superlattice, thereby realizing moiré materials - artificial two-dimensional crystals in which the lattice constant is on the moiré pattern scale. One of the most attractive aspects of moiré materials is that the longer periodicity allows the number of electrons per effective atom to be tuned through large ranges with electrical gates. When the moiré minibands are flat, electronic correlations are strong and can lead to new physics. In magic angle twisted bilayer graphene, for example, strong correlations are manifested by insulating states surrounded by superconducting domes [14], and quantum anomalous Hall ferromagnets [5, 6].

In this article we report on an exact diagonalization study of the moiré superlattices formed in transition metal dichalcogenide (TMD) heterobilayers in which correlated insulators and Wigner crystal states have already been observed [14]. In heterobilayer systems, which have different two-dimensional semiconductors on opposite sides of the junction, there is an interval of energy near the band extremum within which carriers are localized in one of the two layers. For example, for WSe$_2$ heterobilayers formed with either MoSe$_2$ or MoS$_2$, the carriers at the top of the valence band are localized in the WSe$_2$ layer, but experience a periodic potential due to the moiré pattern. Spin-valley locking in WSe$_2$ then leads to low-energy physics described by a Hubbard-like model in which spinful electrons experience a periodic potential whose extrema form a triangular lattice [14, 15]. We limit our attention to the case of one-electron per moiré period and focus on the metal to insulator phase transition (MIT) [16, 17] that is expected to occur when interactions become strong compared to moiré miniband widths.

The bilayer is described by a continuum model that depends on moiré potential depth $V_m$ and on the moiré period $a_M$ (or equivalently the twist angle), which determine the kinetic energy scale $W_M$ and interaction energy scale $U_M$. Our main results are summarized by the phase diagram in Fig. 1 which is controlled by the dimensionless parameters $V_m/W_M$ and $U_M/W_M$. We find that the metal-insulator phase transition points can be readily identified by calculating the charge gap $\Delta_c$ vs. $U_M/W_M$.
for fixed $V_M/W_M$, as shown in Fig. 1(a). Repeating these calculations at different modulation strengths $V_m$ yields the phase diagram shown in Fig. 1(b). In order to emphasize the importance of a non-mean-field theory treatment of the metal-insulator phase transition, we have included an estimate for transition line obtained from the Hartree-Fock method applied to a groundstate without broken translational symmetry. The insulating state is favored, as expected, at large $V_m$ and $U_M$ but its stability is overestimated by the Hartree-Fock calculation. Momentum-state occupation-number distribution functions $\langle \Psi_{GS} | n_k | \Psi_{GS} \rangle$ where $| \Psi_{GS} \rangle$ is the many-body ground state, plotted as insets in Fig. 1(b), clearly distinguish the two states by the presence or absence of a Fermi surface (schematically represented as a green circle). These numerical results clearly indicate that a MIT occurs at half-filling in moiré materials, demonstrating that they are an attractive platform for searches for superconductivity in doped Mott-insulators, and spin-liquid states on the insulating side of metal-insulator phase transitions. Below we first explain the technical details of our calculations and then discuss their implications.

II. MOIRÉ BAND MODEL

The moiré Hamiltonian of twisted TMD heterobilayer valence bands is [14]

$$\hat{H} = -\frac{\hbar^2}{2m^*} \mathbf{k}^2 + \Delta(\mathbf{r}),$$  \hspace{1cm} (1)

where $\Delta(\mathbf{r})$ is an external potential with moiré periodicity. Experimental [15] and theoretical [19, 20] values for the effective mass of valence band monolayer WSe$_2$ vary; here we take $m^* = 0.35 m_0$. In the dominant harmonic approximation $\Delta(\mathbf{r}) = 2V_m \sum_j \cos(\mathbf{b}_j \cdot \mathbf{r} + \psi)$, where $\mathbf{b}_j = k_0 (\cos(2\pi j/3), \sin(2\pi j/3))$ and $k_0 = 4\pi/(\sqrt{3}a_M)$. In this approximation the moiré modulation potential is completely characterized by strength ($V_m$) and shape ($\psi$) parameters. The potential strength $V_m$ depends on heterobilayer and, when strain effects are accounted for, also on twist angle. The shape parameter $\psi$ controls the relative depth of potential extrema locations and, as shown in Ref. [19], strongly influences the strength of particle-hole asymmetry relative to the half-filled moiré band case considered in this work. For concreteness we choose the value $\psi = -94^\circ$, estimated from ab initio calculations for WSe$_2$/MoSe$_2$ in Ref. [14]. For this $\psi$ the valence band potential has a single maximum centered at the AA positions of the moiré superlattice.

An example of the moiré minibands obtained by diagonalizing the Hamiltonian in Eq. (1) in a plane wave basis is shown in Fig. 2(a). Clearly, the topmost valence moiré band is well-separated and flat in this case. The width of the topmost moiré band, and the energy gap to the second moiré band are plotted as a function of twist angle and shape parameter in Fig. 2(b) and 2(c) respectively. The width increases with twist angle but is almost $\psi$-independent except near $\psi = -60^\circ$ and $\psi = -180^\circ$, where the bands broaden. This property is explained by Fig. 2(c). Outside of the blue regions, the topmost moiré miniband is not spectrally isolated, and any mapping to a one-band Hubbard model is inaccurate. At both $\psi = -60^\circ$ and $\psi = -180^\circ$, the shape parameter value imposed by emergent symmetries in the case of $\Gamma$-valley TMD homobilayers [21], the moiré potential has two identical maxima that sit on honeycomb lattice sites, and it is therefore necessary to retain at least two bands to model the low-energy physics. As $\psi$ is varied there is a smooth crossover between triangular and honeycomb lattice limits, with intermediate values of $\psi$ providing a realization [15, 22] of charge-transfer insulator physics. In this work we limit our attention to the one-band Hubbard model case.

Using a harmonic oscillator approximation near the highest extremum, we estimate that the moiré triangular lattice hopping parameter $t \sim \exp(-\kappa V_m^{1/2}/\theta)$, where $\kappa$ is a constant, and that the on-site Coulomb interaction $U_0 \sim e^2/(e\alpha_W) \sim (e^2 \sqrt{\theta V_m^{1/4}})/\epsilon$, where $\alpha_W$ is the width of the flat band Wannier function and $\epsilon$ is the static dielectric constant. By varying the values of $\theta$, $V_m$ and $\epsilon$, the ratios between interaction strength and hopping can be tuned. Experimentally, the potential depth can be varied in situ by applying pressure [13, 23] or gate-controlled displacement fields [24], while the dielectric environment can be modified by varying the carrier density of surrounding graphene gates [23] and their separation from the active layer.
III. MANY-HOLE HAMILTONIAN

Since our goal is to investigate the electronic properties of moiré materials, we simplify the many-body problem by projecting the continuum Hamiltonian to the Hilbert space of the topmost moiré miniband:

$$H = \sum_{k,\sigma} \epsilon_{k,\sigma} c_{k,\sigma}^\dagger c_{k,\sigma} + \frac{1}{2} \sum_{k,l,m,n} \sum_{\sigma,\sigma'} V_{klmn} c_{k,\sigma}^\dagger c_{l,\sigma'}^\dagger c_{n,\sigma'} c_{m,\sigma},$$

(2)

where $c_{k,\sigma}^\dagger$ creates (destroys) a hole with momentum $k$ in valley $\sigma$, $k, l, m, n$ are momentum labels, $\epsilon_{k,\sigma}$ is a flat valence band single particle energy, and $V_{klmn}$ is a two-particle matrix element

$$V_{klmn} = \langle k, \sigma | l, \sigma' | V | m, \sigma, n, \sigma' \rangle.$$ 

(3)

The Coulomb long-range interaction is given by $V = e^2/\epsilon r_1 - r_2$ and $e^{-1}$ is an interaction strength parameter related to the two-dimensional system’s three-dimensional dielectric environment. The $V_{klmn}$ matrix elements are sensitive to the tails of the flat band wavefunctions at positions away from their maxima in the moiré unit cell. The size of these tails is sensitive to the confinement potential at lattice sites, which is weaker in the moiré material case than in atomic lattices. For small twist angles the interaction physics is expected to be accurately described by a model with only on-site interactions. For larger angles, however, longer range Coulomb interaction and non-local terms become more important (see Supplemental Material for further comment [26]).

IV. METAL-INSULATOR TRANSITION

Our analysis is based on exact diagonalizations of Eq. [2] with periodic boundary conditions applied to different finite system sizes, limiting the number of momentum points in the discretized first Brillouin zone to $N$. We note that for half-filling $N$ is also the number of particles in the spinful system. Typical results are illustrated in Fig. 4(a) where we plot the lowest 1700 many-body energies relative to the ground state as a function of $e^{-1}$ for $N = 9$. We see a set of $2^N$ low-energy states separated by a Hubbard gap to higher states at strong interactions. This identifies a parameter range of insulating states where the many-body physics can be described by a spin model.

The picture of localized spins breaks down with decreasing interaction strength and a transition to a metallic phase is expected. Fig. 4(b) shows the energy gap to the first many-body excited state for system sizes $N = 9, 12$ and $N = 16$. For strong interactions, the total spin of the system is size dependent with $S = 0$ for $N = 12$ and $N = 16$, and $S = 3/2$ for $N = 9$. The spectra for $N = 9$ and $N = 12$ show level crossings around $e^{-1} \sim 0.045$, signaling a possible spin liquid intermediate phase with minimum total spin between a Fermi liquid and the strong interaction limit, as predicted for related models [27–29]. There is a level crossing in the gray-shaded region ($e^{-1} \sim 0.0175 – 0.025$) in all three geometries, that we identify with the MIT. We have estimated the ratio between onsite Hubbard interaction $U_0$ and the nearest neighbor hopping integral $t$ for the shaded region, calculated from our model using Wannierization, obtaining $U_0/t \sim 7.9 – 9.9$. This estimate is consistent with previous studies of the triangular Hubbard model [27–29]. For moiré materials the precise value of $U_0/t$ is dependent on $\theta$, $V_m$ and $\epsilon$, because of longer range hopping and non-standard interaction terms.

To examine the MIT more directly we evaluate the charge gap $\Delta_c$, i.e., the difference between the energy to add a particle and the energy to remove a particle from a given ground state, to see if it remains finite in the thermodynamic limit. The charge gap shown in Fig. 5(a) is defined as $\Delta_c \equiv \lim_{N \to \infty} \Delta_c(N)$, where

$$\Delta_c(N) = E_0(N + 1) + E_0(N - 1) - 2 E_0(N).$$

(4)

The values of $\Delta_c$ for each potential strength $V_m$ shown in
Fig. 4 were obtained from extrapolations of $N = 4, 9, 16$ results to $N = \infty$ (see Supplemental Material for further details [26]). The values obtained for the charge gaps in the insulating region of our phase diagram are in the order of tens of meV, in agreement with results reported in Refs 7 and 8. As noted earlier, these charge gap calculations show clear metal-insulator phase transitions at positions that can be accurately estimated. The insulating state is favored, as expected, at large potential strengths and at smaller twist angles, which decrease $W_M$ and produce a rapid decrease in band width at a fixed $V_m$.

Metal-insulator transitions are more interesting when continuous but are usually first-order [29, 31]. In magnetically frustrated systems such as the organic compounds [27, 32] metal-insulator transitions are often only weakly first-order. Some theoretical work [33, 34] suggests that spin liquids with a spinon Fermi surface could undergo first-order. Some theoretical work [33, 34] suggests that spin liquids with a spinon Fermi surface could undergo first-order. Some theoretical work [33, 34] suggests that spin liquids with a spinon Fermi surface could undergo first-order.

![Graph](image)

FIG. 4. (a) Expectation value of single-particle kinetic energy relative to the flat-band average $\bar{T}$ vs. $\epsilon^{-1}$ for several system sizes. The gray shading indicates the approximate position of the MIT as estimated by the charge gap calculations. (b) Numerical first derivative of (a), that reveals peaks for $N = 9$ and $N = 12$ at both phase transitions. In these calculations $\theta = 2.5^\circ$ and $V_m = 11$ meV, corresponding to $V_m/W_M = 2.86$. The top axis indicates the values of $U_M/W_M$ corresponding to a given $\epsilon^{-1}$.

and $N = 12$ in the shaded area, while for $N = 16$ the derivative seems to be continuous. This evidence, combined with the apparently continuous vanishing of the charge gap $\Delta_c$ vs. $U_M/W_M$ in Fig. 4(a), clearly shows that the MIT in moiré materials is not a simple strongly first order phase change.

V. DISCUSSION

The theory of metal-insulator transitions in two or more dimensions continues to be a challenge, partly because of the absence of a clear order parameter. In the case of triangular lattice systems, magnetic frustration in the insulating state adds an additional complication. A standard way to approach this problem theoretically is to study generalized single band Hubbard models in particular lattice geometries. Some layered organic compounds are believed to be described by a triangular lattice Hubbard model with on site interactions and nearest-neighbor hopping. In those systems an intermediate spin-liquid state seems to appear [30, 32, 35, 36] in the vicinity of pressure-induced MITs. Previous numerical studies of the frustrated Hubbard model motivated by these experiments do identify the expected insulating (120°-Néel state) and the Fermi liquid states in the strong and weak on-site interaction limits [27, 29, 31, 37]. Between those phases an insulating phase without apparent magnetic order appears in agreement with experiment, separated from the metal by a first-order transition [28, 29, 35]. Our calculations suggest that there is also a delicate intermediate state close to the MIT line in triangular lattice moiré materials and that the transition occurs under experimentally realizable conditions. It is clear from our numerical study that the moiré material metal-insulator transition is not strongly first order, in agreement with known properties of organic compound triangular lattice systems, in which magnetic frustration plays an important role.

The principal difference between moiré materials and atomic crystals is that the potential that attracts particles to lattice sites is bounded in the former case, and unbounded Coulomb ion-core attraction in the latter. In some cases [15, 22] the moiré potential can have two minima, and even two-identical minima per moiré unit cell [21]. One signal of this behavior is a relatively small splitting between the two topmost moiré minibands. In these cases the moiré insulator is more like a charge-transfer insulator than like a Mott-Hubbard insulator, and the minimal model for the description of its MIT includes at least two-bands. The boundary between Mott and charge-transfer insulators is set by the band structure shape parameter $\psi$, as we show in Fig. 2.

Moiré materials are of special interest because of the possibility they present for in situ, tuning of relevant parameters. Most important among these is the possibility of using gates to alter the carrier density and to measure the chemical potential as a function of carrier...
density \[10\] [11]. For metal-insulator phase transitions, the implication is that the charge gap at half-filling is directly measurable. Because the band-width in all heterobilayer moiré materials is very sensitive to twist angle, this knob can be used to prepare samples that are in the neighborhood of the metal-insulator transition. In situ tuning through the metal-insulator phase transition can then be achieved using gates, or pressure, or by changing gate screening properties. The prospects for unambiguous experimental determination of the order of the metal-insulator phase transition using transport \[12\] [13] and chemical potential measurements, and of the presence or absence of a spin-liquid state are excellent, and would set the stage for careful studies of weakly doped Mott insulators.

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**Note added** – Two experimental studies of the metal-insulator phase transition in moiré superlattices have appeared on the arXiv since our original submission \[14\] [15].

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for (a) $\theta = 0.5$ and (b) $\theta = 2.5$ with $V_m = 11$ meV in a Brillouin zone mesh of 225 momentum points. For $\theta = 0.5$, the histograms have substantial overlap, showing a peak near $|\langle k, l | V | m, n \rangle| = 2.0$ meV. If we considered approximating the moiré material by an on-site only $U_0$ Hubbard model, a single peak would be present near $|\langle k, l | V | m, n \rangle| = 1.8$ meV. The Wannier-estimation of the nearest-neighbor interaction $U_1$ is an order of magnitude smaller than the estimated $U_0$, but for $\theta = 0.5$ it spreads the distribution into several peaks and partially explains the differences compared to the moiré material model. We conclude that at small twist angles the moiré materials Hamiltonian is faithfully represented by a simple Hubbard model, with on-site and perhaps near-neighbor interactions. For larger angles, i.e. $\theta = 2.5$, the discrepancy between the models is clear, suggesting that non-local interaction terms that are normally neglected in lattice models start to play an important role. A detailed analysis of the role of long range Coulomb elements and non-local terms in moiré materials is a subject of future work.

**EXTRAPOLATIONS TO THE THERMODYNAMIC LIMIT AND COMPARISON WITH HARTREE-FOCK METHOD**

We compare extrapolations to larger system sizes for quantities obtained from exact diagonalization with the corresponding extrapolations for quantities obtained from Hartree-Fock method calculations. For the Hartree-Fock calculations we assume that translational symmetry is not broken, i.e. we assume a ferromagnetic or paramagnetic ground state, which allows access to larger system sizes. In Fig. 6(a) we show that Hartree-Fock method ground state energy per particle results for system sizes $N = 4, 9, 16, 36, 81$ and 144 lie in a line when plotted as function of $N^{-1/2}$, as do the exact diagonalization results for system sizes $N = 4, 9$ and 16 discussed in the main text. The thermodynamic limit Hartree-Fock energy extrapolated from results for $N = 4, 9, 16$ equals the result extrapolated from calculations at larger sys-
tem sizes to within less than 1 meV per unit cell. We expect that the exact-diagonalization ground state energy results discussed in the main text have a similar accuracy. Fig. 6(b) compares charge-gap $\Delta_c \sim N^{-3/2}$ extrapolations based on ferromagnetic Hartree-Fock and exact diagonalization calculations. We see that the results follow the expected power laws accurately. In Fig. 6(c) we show how the charge gaps obtained from Hartree-Fock compare to the exact diagonalization charge gaps as a function of interaction strength. As pointed out in the MS, Hartree-Fock mean-field method tends to overestimate the stability of the insulating phase, which is manifested in larger values for $\Delta_c$ and also in metal-insulator transition points at smaller values of $U_M/W_M$. This discrepancy between charge gaps justifies the necessity of using a non-perturbative method like exact diagonalization in order to study physics near the metal-insulator phase transition.

FIG. 6. System-size extrapolations for exact diagonalization and Hartree-Fock methods. (a) Groundstate energies per particle as a function of $N^{-1/2}$ for half-filling at two values of $e^{-1}$, yellow (blue) points show results from exact diagonalization (Hartree-Fock) for different system sizes. Solid (dashed) lines correspond to extrapolations to the thermodynamic limit from ED (HF) data, for $V_m=11$ meV and $\theta = 2.5^\circ$. (b) Extrapolations for the charge gaps calculated from ED (solid lines) and Hartree-Fock (dashed lines) for the same parameters as in (a). (c) Extrapolated charge gaps vs $U_M/W_M$, for the two values of the normalized moiré potential depth $V_m/W_M$ presented in Fig. 1(a) (lines with points) and the corresponding charge gaps in a ferromagnetic groundstate calculated from Hartree-Fock (dashed lines).