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Strong pairing in mixed-dimensional bilayer antiferromagnetic Mott insulators

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Interacting many-body systems in reduced-dimensional settings, such as ladders and few-layer systems, are characterized by enhanced quantum fluctuations. Recently, two-dimensional bilayer systems have sparked considerable interest because they can host unusual phases, including unconventional superconductivity. Here we present a theoretical proposal for realizing high-temperature pairing of fermions in a class of bilayer Hubbard models. We introduce a general and highly efficient pairing mechanism for mobile charge carriers in doped antiferromagnetic Mott insulators. The pairing is caused by the energy that one charge gains when it follows the path created by another charge. We show that this mechanism leads to the formation of highly mobile but tightly bound pairs in the case of mixed-dimensional Fermi-Hubbard bilayer systems. This setting is closely related to the Fermi-Hubbard model believed to capture the physics of copper oxides, and can be realized in currently available ultracold atom experiments.

ne of the fundamental results in the theory of strongly interacting electron systems is the Kohn-Luttinger theorem, which states that electron pairing is inevitable in systems with purely repulsive interactions. Following the original paper by Kohn and Luttinger in 19651, it was commonly assumed that their pairing mechanism gives rise to critical temperatures that are too small to be observable in realistic experimental systems. The discovery of high-temperature superconductivity in cuprates² triggered renewed interest in the study of pairing in systems with repulsive interactions. Even today, one key question in the field of cuprate superconductors is whether electron pairing in these materials can be explained by purely repulsive Coulomb interactions. The lack of theoretical consensus on this question motivated the development of the field of quantum simulation using ultracold atoms: the Fermi-Hubbard model, commonly believed to capture the main properties of cuprate superconductors, is naturally realized using fermions in optical lattices³⁻⁵. During the past few years, considerable experimental progress has been made in analysing the properties of this model, including the observation of long-range antiferromagnetic order⁶, magnetic polaron properties⁷, the crossover from polarons to Fermi liquids⁸ and bad metallic transport⁹. However, the demonstration of pairing has so far been out of reach as temperatures substantially below the superexchange would be required.

This Article has two goals. First we identify a microscopic bilayer model that features repulsive interactions between fermions, yet exhibits pairing at temperatures comparable to the superexchange, realizable by state-of-the art quantum simulators (Fig. 1). A special characteristic of this model is its mixed-dimensional (mixD) character: it features interlayer exchange interactions, but no interlayer single-particle tunnelling (Fig. 1a,b). Such systems can be realized experimentally using ultracold atoms by applying a potential gradient in bilayer and ladder models, which have been recently realized in cold atom experiments (12-15).

The second goal of our paper is to reveal a general mechanism for pairing in doped antiferromagnetic Mott insulators (AFMI). It is based on the idea that a single hole can be understood as a bound state of chargon and spinon¹⁶⁻²⁰, carrying the respective charge and spin quantum numbers of the underlying AFMI²¹⁻²⁵, which are connected by a geometric string of displaced spins. The antiferromagnetic correlations (which may also be short-range) of the parent state are assumed to induce a linear string tension²⁶. Within the same framework, a bound state of two holes corresponds to a bound state of two chargons. We demonstrate that the interplay of the potential energy, arising from the string tension, and kinetic energy of the partons leads to large binding energies for mesonic pairs of chargons. Our work goes beyond earlier analysis^{27–31} by including adverse effects that suppress pairing in a generic system. Effects detrimental to pairing will be suppressed in the mixD bilayer model we propose, featuring overwhelmingly attractive emerging

In the present work we focused on the case without interlayer tunnelling, $t_{\perp} = 0$, which enabled an effective theoretical description. We considered strong spin–charge coupling, where the intralayer tunnelling t_{\parallel} exceeds the relevant superexchange J_{\perp} ($t_{\parallel} \gg J_{\perp}$; Fig. 1). We demonstrate that the deeply bound pairs are highly mobile. This constitutes a notable exception from the common expectation that bipolarons should carry a heavy effective mass. The light mass of bipolarons is also significant, because it allows for high condensation temperatures T_c and has important implications for the ground state at finite doping: a light mass renders the localization of pairs unlikely. Therefore, we expect density wave states, such as stripes, to be less competitive with the superfluid states of pairs. In the opposite limit, $t_{\parallel} \ll J_{\perp}$, we also expect pairing, but in this case the pairs are heavy and thus T_c is low.

The microscopic mixD bilayer model we studied hosts a Bose–Einstein condensate (BEC) regime with mobile but tightly bound

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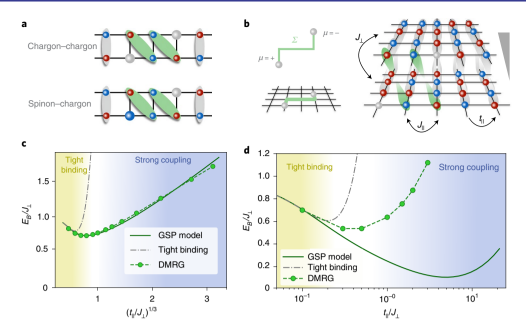


Fig. 1 | **String-based chargon pairing in mixD bilayers.** We introduce an efficient pairing mechanism for distinguishable chargons with flavours $\mu=\pm$ connected by a string Σ . **a,b**, The string pairing mechanism can be realized in a model of d-dimensional bilayers with spin-1/2 particles and strong repulsive interactions for d=1 (**a**) and d=2 (**b**). We consider mixD systems, where the hopping t_\perp between the layers is suppressed by a gradient Δ , while the superexchange J_\perp is kept intact and $J_\perp\gg J_{||}$. The strong interlayer superexchange leads to the formation of rung singlets (dimers indicated by grey ellipses with red and blue spheres), depicted here for d=1 (**a**) and d=2 (**b**). The motion of doped holes (grey spheres) via t_\parallel within different layers $\mu=\pm$ tilts the singlets along their paths, which corresponds to the formation of Σ . The potential energy associated with the string increases with its length as more singlets are tilted. Grey (green) shading corresponds to vertical (tilted) singlets. **c**,**d**, E_B obtained by comparing the energies of spinon-chargon and chargon-chargon mesons in the GSP model, and the ground-state energy for a system with one versus two holes in DMRG as described in the text. We assumed that $J_\parallel/J_\perp=0.01$ and varies t_\parallel/J_\perp in equation (4). In the DMRG, weak interlayer tunnelling of strength $t_\perp/J_\perp=0.01$ was added to ensure convergence. The shading indicates the tight binding and strong coupling regimes. **c**, E_B for a 40×2 ladder (d=1). **d**, E_B for a bilayer system on a 12×4 cylinder.

pairs at low doping, and a Bardeen–Cooper–Schrieffer (BCS) regime at high doping⁵. Hence our findings may have further implications for our understanding of the phenomenology of high-temperature superconductivity: when the interlayer couplings J_{\perp} are switched off, the bilayer systems we consider reduce to independent copies of the two-dimensional (2D) Fermi–Hubbard model. The latter is widely believed to capture the essential physics of the cuprate superconductors. Although experiments indicate that the cuprates do not realize a BEC phase with tightly bound Cooper pairs, they may be close in parameter space to models that exhibit a full BEC-to-BCS crossover. This would help to explain the pseudogap observed in cuprates, which is reminiscent of the pseudogap that arises at strong coupling in a BEC-to-BCS crossover, as realized in the mixD bilayer model.

String-based chargon pairing

The main result of this Article is the identification of a general string-based pairing mechanism for doped holes in AFMIs (Fig. 1). We begin by introducing a general formulation of the model, which we dub the dimer parton model, in an abstract way not tied to any microscopic Hamiltonian. We present the assumptions that went into the construction of the model and summarize its key predictions, and provide technical details in the Methods. In the following section we propose a microscopic bilayer model (Fig. 1a,b), and demonstrate that it realizes the abstract string scenario we envisioned. In particular, this allows us to understand the surprisingly high binding energies that we found for mobile (that is, strongly coupled) holes in 2D mixD bilayers shown in Fig. 1d.

In our general dimer parton model, we described excitations of AFMIs by two different types of parton, spinons and chargons, which carry the spin and charge quantum number, respectively. The partons also carry a flavour index $\mu = \pm$, allowing us to work with

distinguishable chargons without a mutual hard-core constraint, which we predicted to pair up at strong couplings. The two different flavours $\mu=\pm$ could, for example, be internal degrees of freedom or different layers in a bilayer system (Fig. 1). In the dimer parton model, we can treat arbitrary lattice geometries, but we assumed a homogeneous coordination number z for all sites; for example z=4 for the bilayer system shown in Fig. 1b.

Furthermore, we considered rigid strings Σ on the underlying lattice, which connect the partons and fluctuate only through the motion of the latter. We assumed a linear string tension σ_0 ; that is, a string is associated with an energy cost $V(\ell_{\Sigma}) = \sigma_0 \ell_{\Sigma}$ proportional to its length ℓ_{Σ} , which reflects how the short-range antiferromagnetic correlations of the underlying AFMI are disturbed by the parton motion. In the microscopic mixD bilayer models discussed below (Fig. 1a,b), this string tension corresponds to the cost of breaking up rung singlets via the parton motion.

Since we considered rigid strings with a linear string tension, the partons are always confined in our model, forming mesonic states. The binding energy for two charges was hence obtained by comparing the ground state energy of a spinon-chargon (labelled sc) pair to the energy per chargon of a chargon-chargon (labelled cc) pair

$$E_{\rm B} = 2E_{\rm sc} - E_{\rm cc}.\tag{1}$$

At strong couplings (that is, when the hopping amplitude $t\gg\sigma_0$ of the holes exceeds the string tension $\sigma_0\propto J$ determined by the super-exchange J), we predict a binding energy with a characteristic scaling

$$E_{\rm B} = \alpha \underbrace{(2 - 2^{1/3})}_{-0.740} t^{1/3} \sigma_0^{2/3} + \mathcal{O}(J), \tag{2}$$

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where $\alpha > 0$ is a non-universal constant proportional to $(z-1)^{1/6}$ (see Methods for details).

Since a combination of t and $\sigma_0 \propto J$ appears in equation (2), we predicted a remarkably strong asymptotic $E_{\rm B}$. The appearance of t in this expression highlights the key aspect of the underlying binding mechanism, where two chargons share one string and gain equal amounts of superexchange and kinetic energy: the motion of one chargon frustrates the underlying AFMI along the string Σ . In the case of two chargons, the second chargon can retrace the string created by the first, thus lowering the energy cost due to frustration in the spin sector. $E_{\rm B}$ becomes positive because binding to a light second chargon with mass $\propto 1/t$ is favoured over binding two chargons individually to heavy spinons with mass $\propto 1/J$.

In a 2D square lattice $\alpha \approx 2$ and using a string tension of $\sigma_0 \approx J = 4t^2/U$, where U are on-site interactions, we expect a binding energy on the order of $|E_B| \simeq t^{1/3} J^{2/3} \simeq 2.5 t (t/U)^{2/3}$. This suggests that binding energies of a significant fraction of a typical tunnel coupling t may be possible, at least in principle, potentially exceeding room temperature.

As a second main result of the dimer parton model we estimated the effective mass of the chargon pair at strong couplings

$$M_{\rm cc}^{-1} = 4t\sqrt{z - 1/z}. (3)$$

Despite being tightly bound, the pair is highly mobile—contrary to common expectations for bipolarons.

We anticipate that the general string-based pairing mechanism plays a role in different microscopic models of AFMIs. Below, we discuss a specific realization in the mixD bilayer systems shown in Fig. 1.

Microscopic model

We considered a bilayer of d-dimensional sheets with spin-1/2 particles $\hat{\mathbf{c}}_{\mathbf{j},\mu,\sigma}$, which can be either bosons or fermions. Here, $\mu,\sigma=\pm$ are layer and spin indices, respectively, and \mathbf{j} is a d-dimensional site index. In the following, we considered fermions. The particles were assumed to be strongly repulsive, allowing us to work in a subspace without double occupancies. The Hamiltonian we considered includes nearest-neighbour hopping t within the layers and spin-exchange couplings J_{\parallel},J_{\perp} :

$$\hat{\mathcal{H}} = -t_{\parallel} \hat{\mathcal{P}} \sum_{\langle \mathbf{i}, \mathbf{j} \rangle} \sum_{\mu, \sigma = \pm} \left(\hat{\mathbf{c}}_{\mathbf{i}, \mu, \sigma}^{\dagger} \hat{\mathbf{c}}_{\mathbf{j}, \mu, \sigma} + \text{h.c.} \right) \hat{\mathcal{P}} +
+ J_{\parallel} \sum_{\mu} \sum_{\langle \mathbf{i} \mathbf{j} \rangle} \left(\hat{\mathbf{S}}_{\mathbf{i} \mu} \cdot \hat{\mathbf{S}}_{\mathbf{j}, \mu} - \frac{1}{4} \hat{\mathbf{n}}_{\mathbf{i}, \mu} \hat{\mathbf{n}}_{\mathbf{j}, \mu} \right) +
+ J_{\perp} \sum_{\mathbf{i}} \left(\hat{\mathbf{S}}_{\mathbf{i}, +} \cdot \hat{\mathbf{S}}_{\mathbf{i}, -} - \frac{1}{4} \hat{\mathbf{n}}_{\mathbf{i}, +} \hat{\mathbf{n}}_{\mathbf{i}, -} \right),$$
(4)

where $\hat{\mathcal{P}}$ projects to the subspace with maximum single occupancy per site; $\hat{\mathbf{S}}_{\mathbf{j},\mu}$ and $\hat{\mathbf{n}}_{\mathbf{j},\mu}$ denote the on-site spin and density operators, respectively, and h.c. denotes the Hermitian conjugate. The sum in the second line includes nearest-neighbour bonds within the layers, whereasthethirdline corresponds to bonds between the layers (Fig. 1b).

Experimentally, the model in equation (4) can be realized in d=1,2 starting from a Hubbard Hamiltonian⁵ with on-site interactions U. A strong interlayer gradient Δ can be used to obtain a mixD setting¹⁰ along the third direction where the two layers $\mu=\pm$ are physically realized. Note that the antiferromagnetic coupling $J_{\perp}>0$ can be realized both for fermions and bosons via an appropriate choice of the gradient Δ and the Hubbard interaction U^{33-36} . In the following we assume that the interlayer spin-exchange is dominant, $J_{\perp}\gg |J_{\parallel}|$, which can be achieved by choosing $U\gg t_{\parallel}$.

As a central result of our Article, we predict strong binding energies of holes, on the order of $|E_{\rm B}| \gtrsim J_{\rm L}$, in the strong coupling regime $t_{\parallel} \gg J_{\perp}$ where charges are highly mobile. In Fig. 1c,d we show

binding energies obtained from density-matrix renormalization group (DMRG) simulations³⁷ in mixD ladders (d=1) and bilayers (d=2) for J_{\parallel} =0. The strong binding observed numerically can be explained by the string-based pairing mechanism introduced above.

The undoped ground state of equation (4) at half filling and for $|J_{\parallel}| \ll J_{\perp}$ corresponds to a product of rung singlets

$$|\Psi_0\rangle = 2^{-V/2} \prod_{\mathbf{j}} \left(\left| \uparrow \atop \uparrow \right\rangle_{\mathbf{j}} - \left| \uparrow \atop \downarrow \right\rangle_{\mathbf{j}} \right),$$
 (5)

where $V = L^d$ is the volume. Excitations correspond to the break up of singlets and dope the system with holes. The motion of a hole introduces tilted singlets along its path, directly imprinting the string Σ between two partons into the spin background. As illustrated in Fig. 1a,b, the mixD bilayer model naturally realizes spinon–chargon and chargon–chargon bound states.

Effective parton models

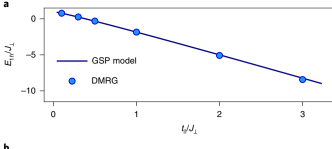
Now we discuss how the microscopic model, equation (4), can be related to effective parton models (see the Supplementary Information for a detailed derivation). We considered systems with two distinguishable partons, n = 1, 2, where the label n summarizes the set of properties layer μ , parton type (spinon or chargon) and spin σ . To obtain distinguishable partons, at least one of these properties has to be different between n = 1 and n = 2. We introduce the notation $|\mathbf{x}_1, \mathbf{x}_2, \Sigma\rangle$, where Σ is a string of tilted singlets connecting \mathbf{x}_2 to \mathbf{x}_1 , and \mathbf{x}_n are the positions of the partons.

Below, we discuss two parton models, the dimer parton model and the refined Gram–Schmidt parton (GSP) model. They can be related to the model in equation (4) within the following approximations (Supplementary Information):

- (1) The frozen-spin approximation³⁸ neglects fluctuations of spins along Σ ; this is justified by a separation of time scales when $t_0 \gg J_1$.
- (2) We neglect the effects of self-crossings of strings, which becomes exactly valid for large coordination numbers $z\gg 1$ (or in ladders with d=1). Even for a d=2 square lattice bilayer, such effects are expected to be quantitatively small due to their small relative share of the string Hilbert space³⁹.
- (3) In the dimer parton model, following the spirit of the Rokhsar–Kivelson quantum dimer model⁴⁰, we assumed that spin configurations $|\mathbf{x}_1, \mathbf{x}_2, \Sigma\rangle$ with parton positions $\mathbf{x}_{1,2}$ form an approximately orthonormal basis, $\langle \mathbf{x}_1', \mathbf{x}_2', \Sigma' | \mathbf{x}_1, \mathbf{x}_2, \Sigma\rangle \approx \delta_{\mathbf{x}_2',\mathbf{x}_2} \delta_{\mathbf{x}_1',\mathbf{x}_1} \delta_{\Sigma',\Sigma}$ where δ denotes the Kronecker delta, which is justified by the small overlaps of configurations with non-identical singlet configurations. This leads to an effective dimer parton Hamiltonian with nearest-neighbour hopping of the partons under simultaneous adaption of the string states, and a string potential V_{Σ} .

The third approximation differs between the dimer parton and the GSP models. In the GSP model, the assumption that string states form an orthonormal basis is partially dropped. To this end, we performed a Gram–Schmidt orthogonalization of the string states defined in the t–J Hilbert space and accounted for the non-orthogonal nature of spinon–chargon states, up to loop effects (Supplementary Information). This led to an accurate prediction of the spinon dispersion. For chargon–chargon states, the GSP and dimer parton models are equivalent.

In the parton models, for simplicity, we assumed a linear string potential, $V^{\rm sc}_{\Sigma} = \sigma_0 \, \ell_{\Sigma}$, where ℓ_{Σ} is the length of string Σ . For the tilted singlets in the mixD bilayer setting, equation (4), the linear string tension is $\sigma_0 = 3/4J_{\perp}$, corresponding to the energy difference between a singlet and a random spin configuration. In the chargon-chargon case, the string potential has an additional on-site attraction, $V^{\rm cc}_{\Sigma} = V^{\rm cc}_{\Sigma} - \delta_{\Sigma,0}J_{\perp}/4$, which leads to slightly stronger binding than that predicted by equation (2).



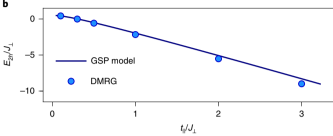


Fig. 2 | MixD bilayers in 2D. We studied ground state energies in a mixD bilayer system with d=2 using the GSP model and DMRG for a bilayer system on a 12×4 cylinder. **a,b**, Energy per hole as a function of t_{\parallel}/J_{\perp} for one hole (**a**) and two holes (**b**).

Two holes are distinguishable if they move within separate layers $\mu_1 = -\mu_2$, and they can thus occupy the same position \mathbf{x} in the lattice. Note that the motion of a second hole in the opposite layer and along the same path can completely remove the geometric string Σ of displaced spins.

Numerical results

In Fig. 1c we show the $E_{\rm B}$ predicted by the GSP model for the mixD ladder (d=1). Plotting the results over $(t_{\parallel}/J_{\perp})^{1/3}$ shows that $E_{\rm B}$ at strong couplings, $t_{\parallel} > J_{\perp}$, scales as $|E_{\rm B}| \simeq t_{\parallel}^{1/3} J_{\perp}^{2/3}$ as predicted by the dimer parton model. In the opposite tight binding regime, $t_{\parallel} \ll J_{\perp}$, we found excellent agreement with our perturbative result from ref. 5 (see the dash-dotted line in Fig. 1c). For our DMRG simulations, we defined the binding energy as $E_{\rm B} = 2E_1 - (E_0 + E_2)$, where E_N is the ground state energy of the state with N holes. Our DMRG results are in excellent agreement with our prediction from the GSP model for couplings t_{\parallel}/J_{\perp} ranging over several orders of magnitude, from 0.1 to >30.

In Fig. 1d, we show our results for the mixD bilayer system (d=2). From the dimer parton model we again expected strong pairing when $t_{\parallel} \gg J_{\perp}$. This prediction was confirmed by our DMRG results in $2 \times 12 \times 4$ systems, where the first direction denotes the two layer indices and we assumed periodic (open) boundary conditions along the short (long) axes of the cylinder. For $t_{\parallel}/J_{\perp} = 3$ we found remarkably strong $E_{\rm B}$ below J_{\perp} . For small $t_{\parallel}/J_{\perp} \ll 1$, the perturbative tight binding result⁵ was obtained.

We also compared predictions made by the GSP model with DMRG results for d=2. In Fig. 2 we show the energies per hole in the spinon–chargon and chargon–chargon cases, respectively, and obtained very good agreement. However, a small deviation of a few per cent could be observed at strong couplings in the two-hole case, where the GSP and dimer parton models were identical. We expect that loop effects, ignored in the parton models, led to the slightly lower energy per hole found by DMRG. Since $E_{\rm B}$, shown in Fig. 1d, was obtained as the small difference between the large one-and two-hole energies, see equation (2), the deviation of the GSP model from DMRG seems sizable at strong couplings. We emphasize, however, that GSP and DMRG consistently predicted positive

 $E_{\rm B}$ throughout, and a crossover from tightly bound to strongly fluctuating extended pairs of holes.

Signatures of parton formation in one- and two-hole spectra

To further corroborate the parton structure of the one- and two-hole states we found at strong coupling and demonstrate the high degree of mobility of the paired states, we investigated their spectral properties. We performed time-dependent DMRG simulations⁴¹⁻⁴³ in the mixD ladder (d=1) and bilayer (d=2) to extract the following spectral functions A

$$A_{1/2}(\mathbf{k},\omega) = \sum_{n} \delta(\omega - E_n^{(1/2)} + E_0) \left| \left\langle \psi_n^{(1/2)} \right| \hat{\mathbf{C}}_{(1/2),\mathbf{k}} |\psi_0\rangle \right|^2, \quad (6)$$

with frequency ω , wavevector \mathbf{k} and where $|\psi_0\rangle$ (E_0) is the ground state (energy) without holes, and $|\psi_n^{(1/2)}\rangle$ ($E_n^{(1/2)}$) are the eigenstates (energies) with one (two) holes. The excitations are created through the operators $\hat{\mathbf{C}}_{(1/2),\mathbf{k}}$, which are defined as the Fourier transforms of $\hat{\mathbf{C}}_{\mathbf{l},\mathbf{i}} = \hat{\mathbf{c}}_{\mathbf{i},+,\uparrow}$ and $\hat{\mathbf{C}}_{2,\mathbf{i}} = \hat{\mathbf{c}}_{\mathbf{i},+,\uparrow}\hat{\mathbf{c}}_{\mathbf{i},-,\downarrow}$ (see insets in Fig. 3a,b).

Our results in Fig. 3a,b for the ladder (d=1) reveal a series of narrow lines that are visible up to high energies, in both the one-hole (Fig. 3a) and two-hole (Fig. 3b) spectra. They correspond to long-lived vibrational excitations of the spinon–chargon and chargon–chargon mesons, respectively. This intuition was confirmed by the excellent agreement we obtained with energies predicted by the GSP model. In the Supplementary Information, we show that the parton model also captures the spectral weights. The existence of discrete internal excitations, as revealed in Fig. 3, is a hallmark of emerging mesonic states in doped AFMIs. In addition to the vibrational states visible in Fig. 3, we also predicted odd-parity spinon–chargon and chargon–chargon states from the parton model, whose spectral weights are negligible in Fig. 3 (Supplementary Information).

For the two-hole case (Fig. 3b), we observed a strong centre-of-mass dispersion. This indicated that the deeply bound pair was highly mobile, confirming a key prediction of the parton model equation (3). Additionally, at $k_x = \pi$ the relative motion of the partons is suppressed by quantum interference, and only the lowest mesonic state has non-vanishing spectral weight.

The parameter set we used in Fig. 3a,b assumed that $J_{\parallel}/J_{\perp}=0.31$ and corresponded to a situation that could be realistically realized in a Ferm–Hubbard ladder with a potential gradient between the two legs⁵. Hence, the emerging spinon–chargon and chargon–chargon meson states we predicted in our model should be experimentally accessible with ultracold atoms in optical lattices with current technology.

In Fig. 3c we show two-hole spectral cuts at rotationally invariant momenta $\mathbf{k} = 0 = (0,0)$ and $\boldsymbol{\pi} = (\pi,\pi)$ for the mixD bilayer where d=2; we considered $t_{\parallel}/J_{\perp} = 2$ in the strong coupling regime. Here, our underlying time-dependent DMRG simulations on a $2 \times 40 \times 4$ cylinder were more challenging and we reached shorter times than in d=1. This resulted in significant Fourier broadening, and prevented us from identifying individual vibrational excitations at high energies. Nevertheless, at $\mathbf{k} = 0$ a first peak was visible in the DMRG data. Comparison with a broadened GSP model spectrum suggested that this peak probably corresponds to the first vibrational meson state.

At $\mathbf{k} = \pi$, the two-hole spectrum in Fig. 3c collapsed to a single peak. As in d=1 at $k=\pi$, this can be understood within the dimer parton model from destructive quantum interference that leads to a cancellation of the chargon's relative motion. The energy of the tightly bound two-hole state at $\mathbf{k} = \pi$ was accurately predicted by the GSP model.

As suggested by the large energy difference of the chargon-chargon state between k=0 and $k=\pi$, on the order of $6t_{\parallel}$ in Fig. 3c, we

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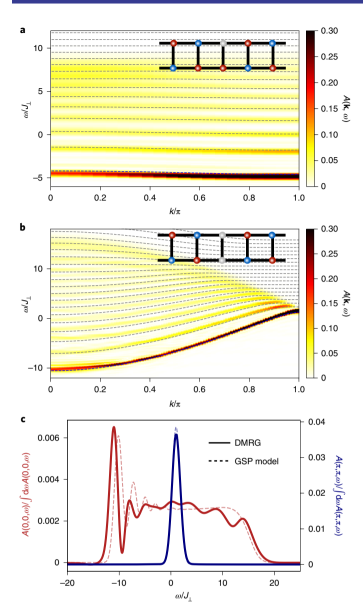


Fig. 3 | Spectral function in mixed dimensions. a-c, Spectra for one hole (**a**) and two holes (**b**) in a ladder (d=1) and two holes in a mixD bilayer (d=2) (**c**). In **a** and **b**, we assumed that t_{\parallel}/J_{\perp} = 3.4 and J_{\parallel}/J_{\perp} = 0.31; and in **c** that t_{\parallel}/J_{\perp} = 2 and J_{\parallel}/J_{\perp} = 0.01. In our time-dependent DMRG (colour map) we added weak tunnelling t_{\perp}/J_{\perp} = 0.01 to simplify convergence. The dashed lines in **a** and **b** correspond to eigen energies of the GSP model (Supplementary Information), where only states with even inversion symmetry are shown. The observed discrete spectral lines correspond to vibrational excitations of emerging spinon-chargon and chargon-chargon mesons in the system. The dashed lines in **c** describe GSP spectra broadened by the Fourier resolution of our time-dependent DMRG simulations (solid lines). Red (blue) lines correspond to momenta **k**=0 (**k**= π).

found that the deeply bound state at $\mathbf{k}=0$ had a light effective mass. A fit $k_x^2/(2M_{cc})$ to the lowest spectral peak $E_{cc}(k_x\mathbf{e}_x)-E_{cc}(0)$ yielded $M_{cc}=2.44/t_{\parallel}$ at $t_{\parallel}/J_{\perp}=2$. This compares reasonably with our estimate $M_{cc}\geq 0.63/t_{\parallel}$ from the dimer parton model; we used an improved version of equation (3) (Supplementary Information). This result should be contrasted with the much heavier $M_{cc}=25/J_{\perp}$ in the tight binding regime⁵, at $t_{\parallel}/J_{\perp}=0.1$ with a comparable $E_{\rm B}$ on the order of $|E_{\rm B}|\approx 3/4J_{\perp}$.

Finite-temperature phase diagram

At finite temperatures, the emergence of two kinds of mesons can lead to rich physics. Although the ground states we found are always paired, we expected a mixture of spinon–chargon and chargon–chargon pairs in chemical equilibrium when T > 0. Because a gas of N spinon–chargon pairs has more entropy than a gas of N/2 chargon–chargon pairs at high T, we expected a crossover from spinon–chargon to chargon–chargon domination around some temperature $T_{\rm mix}$.

In $d \ge 2$ dimensions, we expected an even richer phase diagram. The chargon–chargon pairs can form a (quasi-) condensate below the respective temperature ($T_{\rm BKT}$) $T_{\rm c}$ in (d=2) d>2 dimensions at finite doping and before spinon–chargon pairs dominate beyond $T_{\rm mix}$. Furthermore, at a critical temperature T^* on the order of J_{\perp} we expect a thermal deconfinement transition of the mesons to take place when $d \ge 2$ (ref. ⁴⁴).

Discussion and outlook

We introduced a general string-based pairing mechanism for distinguishable chargons in AFMIs, and demonstrated that the resulting binding energy scales as $|E_{\rm B}| \simeq t^{1/3} J^{2/3}$, or $|E_{\rm B}| \simeq t(t/U)^{2/3}$, when $t\gg J$. The highest binding energies are expected when the chargon–chargon mesons become extended, with an average string length scaling as $\ell \simeq (t/J)^{1/3}$ (ref. ²⁷; see also the Supplementary Information). We confirmed the validity of our analytical arguments using DMRG simulations of bilayer toy models that can be realized experimentally using, for example, ultracold atoms in optical lattices ⁵. Another interesting situation corresponds to cuprate materials with a multilayer structure and an effective chemical potential offset between the layers ^{45,46}: starting from half filling, we expect that a strong pump pulse may allow the creation of metastable pairs of doublons and holes in opposite layers. These may then form the mesonic bound states that we propose, and could be probed by a subsequent laser pulse.

The analysis we performed here was limited to individual mesons, but our prediction of deeply bound and highly mobile chargon pairs for $t\gg J$ has important implications at finite doping. In such a regime, we expect a quasi-condensate with power-law correlations below a critical BKT temperature $T_{\rm BKT}$ for the mixD bilayer (d=2). Owing to the high mobility of the chargon–chargon pairs, a sizable $T_{\rm BKT}$ at a fraction of t and robustness against localization are expected.

Another interesting direction would be the investigation of an emergent SO(5) symmetry in the bilayer models, see refs. 47,48 for studies using a related model.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41567-022-01561-8.

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NATURE PHYSICS ARTICLES

Methods

String-based chargon pairing. Here we introduce the general string-based pairing mechanism for an idealized parton model of AFMIs in d dimensions (the dimer parton model). We considered two flavours of chargons $\hat{\mathbf{h}}_{\mathbf{x},\mu}$ with $\mu=\pm$, which allowed us to work with distinguishable chargons; $\mathbf{x}\in\mathbb{Z}^d$ denotes a lattice vector. The two different flavours $\mu=\pm$ could, for example, be internal degrees of freedom or different layers in a bilayer system (Fig. 1). In the dimer parton model, we can treat arbitrary lattice geometries (in principle), but we assumed a homogeneous coordination number z for all sites; for example z=4 for the bilayer system shown in Fig. 1b. We note that the coordination number considered here is from the point of view of the hopping charges, which are confined to the 2D planes and therefore have the same coordination number as in a square lattice. We also introduced 2×2 flavours of spinons, $\hat{\mathbf{f}}_{\mathbf{x},\mu,\sigma}$, where $\sigma=\pm$ denotes the spin index.

Furthermore, we considered rigid strings Σ on the underlying lattice, which connect the partons and fluctuate only through the motion of the latter. In our idealized scenario, we assumed that such geometric strings retain a full memory of the parton trajectories up to self-retracing paths, but including loops. These strings can also be viewed as a manifestation of the gauge fluctuations expected from any parton representation of an AFMI.

We worked in a regime of low parton density and considered only isolated pairs of two distinguishable partons, n=1,2. Hence, their quantum statistics played no role in the following. The label n summarizes the set of μ , parton type (spinon or chargon) and spin σ in the case of a spinon. To obtain distinguishable partons, at least one of these properties has to be different between n=1 and n=2. We can write the orthonormal basis states of the dimer parton model as $|\mathbf{x}_1, \mathbf{x}_2, \Sigma\rangle$, where Σ connects \mathbf{x}_2 to \mathbf{x}_1 ; that is:

$$\langle \mathbf{y}_1, \mathbf{y}_2, \Sigma' | \mathbf{x}_1, \mathbf{x}_2, \Sigma \rangle = \delta_{\mathbf{y}_1, \mathbf{x}_1} \delta_{\mathbf{y}_2, \mathbf{x}_2} \delta_{\Sigma', \Sigma}.$$
 (7)

The effective Hamiltonian of the two partons written in first quantization reads

$$\hat{\mathcal{H}}_{2p} = -\sum_{n=1}^{2} \frac{1}{2m_{n}} \sum_{\langle \mathbf{y}, \mathbf{x} \rangle, \Sigma} \left(\left| \mathbf{y}_{n}, \Sigma'_{\mathbf{x}_{n}, \mathbf{y}_{n}, \Sigma} \right\rangle \langle \mathbf{x}_{n}, \Sigma | + \text{h.c.} \right)
+ \sum_{\mathbf{x}} |\Sigma\rangle \langle \Sigma| V_{\Sigma}.$$
(8)

The first line describes nearest-neighbour hopping of the partons, with amplitudes $1/2m_n$, and simultaneous adaption of the string state from Σ to $\Sigma'_{\mathbf{x}_n,\mathbf{y}_n}$: hopping along the existing string leads to a retraction of the string along $\mathbf{y}_n - \mathbf{x}_n$, otherwise testing is extended by the same element. Extensions to next-nearest-neighbour hopping can also be considered. The second line in equation (8) describes a general string potential V_Σ that we assume to be independent of the parton flavours.

In the following we consider a linear string potential

$$V_{\Sigma} = \sigma_0 \,\ell_{\Sigma}. \tag{9}$$

In a weakly doped AFMI we may assume that $\sigma_0 \simeq J$ is proportional to J (refs. ^{21,25}). Furthermore, we assumed that the parton masses strongly depended on flavour in the following way

$$(2t)^{-1} = m_h \ll m_f \simeq J^{-1}. \tag{10}$$

That is the chargon mass m_h is significantly lighter than the spinon mass m_P since $t\gg t^2/U\simeq I$ in AFMIs.

Our goal in the following is to demonstrate that two distinguishable chargons (that is, with different μ) form a strongly fluctuating pair with a large $E_{\rm B}$ when $t\gg J$. Since we considered rigid strings with a linear string tension, the partons were always confined in our model. Hence $E_{\rm B}$ for two charges was obtained by comparing mesonic states constituted by a spinon–chargon and a chargon–chargon pair, respectively:

$$E_{\rm B} = 2E_{\rm sc} - E_{\rm cc}. \tag{11}$$

First, we considered spinon–chargon mesons. Because $1/m_i \simeq J$, we could treat the spinon motion perturbatively and start from a localized spinon. In this case, all charge fluctuations are described by nearest–neighbour tunnelling between adjacent string configurations, and equation (8) becomes a single–particle hopping problem on the Bethe lattice $\hat{\mathcal{H}}_{2p}|_{sc} = -t\sum_{\langle \varSigma',\varSigma\rangle} \left(|\varSigma'\rangle \langle \varSigma| + \text{h.c.} \right) + \sigma_0 \sum_{\varSigma} \ell_{\varSigma} |\varSigma\rangle \langle \varSigma| + \mathcal{O}(J)$ (Fig. 1a). Its spectrum is composed of decoupled rotational and vibrational sectors \mathbb{S}^3 . The ground state has no rotational excitations, and its energy in the limit $t\gg J$ follows the well-known universal form \mathbb{S}^3 .

$$E_{\rm sc} = -2t\sqrt{z-1} + \alpha t^{1/3} \sigma_0^{2/3} + \mathcal{O}(J), \tag{12}$$

where $\alpha > 0$ is a non-universal constant proportional to $(z-1)^{1/6}$. Note that contributions from the spinon motion are contained in corrections $\mathcal{O}(J)$.

Next we considered chargon-chargon mesons composed of a pair, $\mu=\pm$, of two distinguishable chargons. We then needed to include the dynamics of both partons in equation (8). Since $\hat{\mathcal{H}}_{2p}$ is translationally invariant, we can transform to the

co-moving frame of the first parton \mathbf{x}_1 by a Lee–Low–Pines transformation 49 . For a given centre-of-mass quasimomentum \mathbf{k} from the Brillouin zone corresponding to the underlying lattice, one again obtains a hopping problem on the Bethe lattice. Although the contributions from the string potential and from \mathbf{x}_2 were identical to the spinon–chargon case, the motion of the first chargon contributes additional \mathbf{k} -dependent terms that tend to frustrate the motion of the pair for $\mathbf{k} \neq \mathbf{0}$ (ref. 28). Around the dispersion minimum at $\mathbf{k} = \mathbf{0}$ we obtained

$$\hat{\mathcal{H}}_{2p}|_{cc} = -2t \sum_{\langle \Sigma', \Sigma \rangle} \left(\left| \Sigma' \right\rangle \langle \Sigma \right| + \text{h.c.} \right) + \sigma_0 \sum_{\Sigma} \ell_{\Sigma} \left| \Sigma \right\rangle \langle \Sigma \right|, \quad (13)$$

up to corrections of order $\mathcal{O}(\mathbf{k}^2)$ (Supplementary Information). The only difference from the spinon–chargon problem is that $t \to 2t$ has been replaced by twice the chargon tunnelling. This replacement can be easily understood by noting that the relative motion of the two chargons involves the reduced mass, $1/m_{\rm red} = 2/m_{\rm h}$; that is, $t_{\rm red} = 2t$. Thus we obtained the energy of the chargon–chargon meson by using the same scaling result as in the spinon–chargon case, equation (12), and merely replacing $t \to 2t$. It is important to note that this leaves the non-universal constant α unchanged, and we obtained

$$E_{cc} = -4t\sqrt{z-1} + \alpha (2t)^{1/3} \sigma_0^{2/3} + \mathcal{O}(J, \mathbf{k}^2). \tag{14}$$

Now we were in a position to calculate $E_{\rm B}$, equation (11), of two holes in the limit $t\gg J$. While the kinetic zero-point energies $\propto -t(z-1)^{1/2}$ cancel, the leading-order string binding energies $\propto t^{1/3}\sigma_0^{2/3}$ yield

$$E_{\rm B} = -\alpha \underbrace{(2 - 2^{1/3})}_{= 0.740...} t^{1/3} \sigma_0^{2/3} + \mathcal{O}(J). \tag{15}$$

This is a remarkably strong $E_{\rm B}$, which depends on a combination of t and $\sigma_0 \simeq J$. The appearance of t in this expression highlights the underlying binding mechanism, where two chargons share one string, gaining equal amounts of potential and kinetic energy.

Finally, we estimated the effective mass of the chargon pair on a hypercubic lattice. We made a translationally invariant ansatz for the chargon–chargon meson in the co-moving frame of the first chargon, and expand the variational energy up to order \mathbf{k}^2 . As shown in the Supplementary Information, for $t\gg J$ this yields a centre-of-mass dispersion $\mathbf{k}^2/2M_{cr}$ of the pair, with

$$M_{\rm cc}^{-1} = 4t\sqrt{z - 1/z}. (16)$$

Despite being tightly bound, the pair is highly mobile—contrary to common expectations for bipolarons.

Data availability

The datasets generated and/or analysed during the current study are available from the corresponding author on reasonable request.

Code availability

The data analysed in the current study has been obtained using the open-source tenpy package; this DMRG code is available via GitHub at https://github.com/tenpy/tenpy and the documentation can be found at https://tenpy.github.io/#.

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Author contributions

All authors contributed to the writing of the manuscript. A.B. performed all numerical DMRG simulations. L.H., F.G. and A.B. derived the effective parton model. A.B. and F.G. proposed the study of string-based pairing in mixed dimensions and all authors contributed to the conceptual ideas.

Competing interests

The authors declare no competing interests.

Additional information

 $\label{thm:continuous} \textbf{Supplementary information} \ The online version contains supplementary material available at $$https://doi.org/10.1038/s41567-022-01561-8.$

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