

Fingerprints of Preformed Pairs in Two-Electron Angle-Resolved Photoemission Spectroscopy

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We use variational exact diagonalization (VED) to calculate the two-electron removal spectral weight for the Hubbard-Holstein model, starting from the ground state with two electrons on a one-dimensional chain. We argue that this spectral weight provides a valuable proxy for the intensity of 2eARPES processes. Our results show that when contrasted to the presumably larger signal due to two electrons ejected from two different pairs, the presumably weaker signal due to two electrons ejected from the same pair (i) is segregated in energy, appearing at a lower binding energy, and (ii) has a very characteristic momentum dependence, with a different symmetry than that of the signal corresponding to two electrons emitted from two different pairs. We verify that these fingerprints appear for pairs with different symmetries, and prove that they arise as a direct consequence of momentum and energy conservation, therefore they are generic for any model with electron-boson coupling that can lead to formation of electron pairs. Experimental observation of these fingerprints will confirm the existence of pairs. Moreover, the momentum dependence map allows one to distinguish whether the pairs are coherent (superconducting) or not. Finally, we argue that these considerations generalize to finite but low electron concentrations, finite temperatures, and higher dimensions.

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Introduction—Angle-resolved photoemission spectroscopy (ARPES) is a well-established and extremely successful experimental technique that provides direct access to the quasiparticle band structure and Fermi surfaces of a variety of interesting materials [1]. Particularly useful for strongly correlated electron systems is the fact that ARPES also enables access to the quasiparticle self-energy. This quantity can be compared against self-energies corresponding to various correlated model Hamiltonians, allowing theorists to gain insights into the most relevant physics of the system of interest.

While access to these single-particle properties is extremely valuable, it is not sufficient to fully characterize the state of a correlated system. For example, direct signatures of pair formation, e.g., in the context of superconductivity, is seen with ARPES if the superconducting gap opened in the single-particle density of states is within the energy resolution of the ARPES system. However, generically ARPES cannot identify whether the gap is due to superconductivity or some other order; additional measurements are needed to settle that [2].

This becomes problematic when trying to identify new correlated states that are not well understood theoretically. Of primary interest are liquids of preformed pairs, hypothesized to appear in systems where boson exchange favors pairing of the electrons, however these pairs are not

coherent (the system is not superconducting) because the electron density is too low or the temperature is too high. We recently argued that signatures of such a state could be seen in ARPES [3], however it is recognized that one of the most direct diagnostics for pairing correlations is provided by two-electron coincidence ARPES (2eARPES); other possible correlation-focussed spectroscopies are reviewed in Ref. [4]. Indeed, analysis of the probability of coincidence detection of a pair of ejected electrons upon absorption of one photon has allowed the observation of the exchange-correlation hole surrounding an electron in a correlated (but unpaired) metal [5], while recent theoretical work argued that 2eARPES can be used to identify the pairing state in Bardeen-Cooper-Schrieffer (BCS) superconductors whose Cooper pairs have a nonzero center-of-mass momentum [6], and to extract dynamic electron interactions in other ordered phases, such as a charge density wave [7]. Exact calculation of two-photons in, two-electrons out 2eARPES intensities has revealed clear signatures of preformed pair formation for an eight-site cluster attractive Hubbard model at half filling [8].

In this Letter, we use the formalism developed in Refs. [8–10] to calculate 2eARPES intensities. First, we argue that when compared to the presumably larger signal due to two electrons ejected from two different pairs, the signal due to two electrons ejected from the same pair is

always (i) segregated in energy, and (ii) has a characteristic momentum-dependence. These fingerprints are a direct consequence of momentum and energy conservation, therefore they must appear in any model hosting preformed pairs. Their observation in 2eARPES will directly confirm the presence of pairs, allow for an estimate of their real-space radius, and also indicate whether the pairs are coherent or not. We then verify this by calculating numerically the 2eARPES intensity of a preformed pair (a bipolaron) in the 1D Hubbard-Holstein model. Varying the electron-phonon coupling strength and the on-site Coulomb repulsion, we can contrast the results for ground states (GSs) consisting of two unbound polarons (strong repulsion, weak el-ph coupling) from those for GS with a bipolaron (weak repulsion, strong el-ph coupling). We confirm that the two fingerprints are observed both for s - and p -wave pairs.

Model and method—We study an N -site 1D chain with electrons that experience Hubbard repulsion and are also Holstein coupled to dispersive optical phonons, as described by the Hamiltonian,

$$\begin{aligned}
 H = & -t_{\text{el}} \sum_{j,\sigma} \left(c_{j,\sigma}^\dagger c_{j+1,\sigma} + \text{H.c.} \right) + g \sum_j \hat{n}_j \left(a_j^\dagger + a_j \right) \\
 & + t_{\text{ph}} \sum_j \left(a_{j+1}^\dagger a_j + \text{H.c.} \right) + \omega_0 \sum_j a_j^\dagger a_j \\
 & + U \sum_j n_{j\uparrow} n_{j\downarrow},
 \end{aligned} \quad (1)$$

where $c_{j\sigma}^\dagger$ and $a_j^\dagger$ are electron and phonon creation operators at site j , respectively, $\hat{n}_j = \sum_\sigma c_{j\sigma}^\dagger c_{j\sigma}$ is the density operator, and U is the Hubbard repulsion. The free-electron dispersion $\epsilon_k = -2t_{\text{el}} \cos(k)$ is controlled by the nearest-neighbor hopping amplitude taken as the energy unit $t_{\text{el}} \equiv 1$, while the optical phonon energy $\Omega_q = \omega_0 + 2t_{\text{ph}} \cos(q)$ is such that $\omega_0 > |2t_{\text{ph}}|$. We set the lattice constant $a = 1$, also $\hbar = 1$. The electron-phonon coupling strength is hereafter characterized by the dimensionless $\lambda = g^2 / [2t_{\text{el}} \sqrt{\omega_0^2 - 4t_{\text{ph}}^2}]$ [11].

We study this Hamiltonian for $N_e = 2$ electrons on a finite chain with periodic boundary conditions employing variational exact diagonalization (VED) [12–17]. We typically use $N_h \sim 18$ iterations to generate the variational space by repeated application of H ; we verified that the results are converged and correspond to the thermodynamic limit $N \rightarrow \infty$.

2eARPES intensity—We calculate the spectral weight $A_2(\omega, k_1, k_2) = (1/\pi) \text{Im} \mathcal{G}_{k_1 k_2}(\omega)$ of the two-particle propagator

$$\mathcal{G}_{k_1 k_2}(\omega) = \sum_n \frac{\left| \left\langle \psi_{-k_1 - k_2}^{(n, N_e - 2)} \right| c_{k_1 \sigma_1} c_{k_2 \sigma_2} \left| \psi_0^{(GS, N_e)} \right\rangle \right|^2}{\omega - i\eta + E_{-k_1 - k_2}^{(n, N_e - 2)} - E_0^{(GS, N_e)}}. \quad (2)$$

Here, $\mathcal{H}|\psi_k^{(n, N_e)}\rangle = E_k^{(n, N_e)}|\psi_k^{(n, N_e)}\rangle$ are the eigenstates and eigenenergies with N_e particles and total momentum k ; n labels other quantum numbers. The ground-state GS corresponds to $n = 0$ and has total momentum 0. For results in the singlet sector we set $\sigma_1 = -\sigma_2$ and verify that double occupancy has a nonvanishing probability. For the triplet sector, we set $\sigma_1 = \sigma_2$. In this case double occupancy is forbidden by the Pauli principle, and the value of U becomes irrelevant.

This two-particle propagator can be linked directly to the quantity defined in Eq. (23) of Ref. [8],

$$\begin{aligned}
 D_{\mathbf{k}_1 \mathbf{k}_2}^{(0)}(\omega) = & \int_{-\infty}^{\infty} dt e^{-i\omega t} \int_{-\infty}^{\infty} d\tau \left\langle c_{k_1 \sigma_1}^\dagger(t) c_{k_2 \sigma_2}^\dagger(t + \tau) \right. \\
 & \left. \times c_{k_2 \sigma_2}(\tau) c_{k_1 \sigma_1}(0) \right\rangle.
 \end{aligned} \quad (3)$$

Up to a factor containing photoexcitation matrix elements, which is ignored for simplicity, $D_{\mathbf{k}_1 \mathbf{k}_2}^{(0)}(\omega)$ was shown by Devereaux *et al.* [8] to represent the 2eARPES coincidence detection rate integrated over all possible differences $\Delta\omega = \omega_1 - \omega_2$ at a fixed total $\omega = \omega_1 + \omega_2$, where $\omega_i = \omega_{ph} - (\hbar^2 \mathbf{k}_i^2 / 2m) - W$ is the energy imparted to the system upon absorption of photon i and emission of the photoelectron with momentum $\mathbf{k}_i$. We denote $\mathbf{k}_i = k_i \mathbf{x} + \mathbf{k}_{i,\perp}$, i.e., the conserved (quasi)momentum along the chain is called k_i (note that $k_i \neq |\mathbf{k}_i|$); this is why the electron operators associated with the states involved in the photoemission processes have (quasi) momenta k_1, k_2 . Finally, W is the work function and the expectation value $\langle \dots \rangle$ is over the GS with N_e particles. Physically, τ is the time difference between the emission of the first and the second electron of the detected 2e ARPES pair [8].

For reasons that will be justified *a posteriori*, in the following we focus on the simpler case $\tau \approx 0$, when Eq. (3) simplifies straightforwardly to $D_{\mathbf{k}_1 \mathbf{k}_2}^{(0)}(\omega) \propto A_2(\omega, k_1, k_2)$ (the dependence on the transverse momenta of the photo-electrons is through the total energy ω imparted to the system).

Next, we analyze Eq. (2) to infer the generic properties of the expected 2eARPES intensity for a ground-state of preformed pairs that are very weakly interacting with one another. In Ref. [3] we showed that such a liquid of s -like singlet bipolarons is the GS of the 1D Hubbard-Holstein model at low carrier concentrations, if the coupling λ is sufficiently large and the repulsion U is sufficiently small. Here, the preformed pairs are bipolarons of energy $E_{BP}(K = 0) = 2E_P(0) - \Delta$ (the bipolaron GS momentum is $K = 0$), where Δ is the binding energy and $E_P(k)$ is the energy of a single polaron with momentum k (the single polaron GS is also at $k = 0$). Interestingly, this model also allows for the binding of p -like, triplet bipolarons if $t_{ph} < 0$. As shown in Ref. [16], dispersive phonons mediate

an effective nearest-neighbor interaction $\Delta E = 2t_{ph}g^2/\omega_0^2$, which becomes attractive if $t_{ph} < 0$ and suffices to bind triplet pairs when ΔE is sufficiently negative, see Supplemental Material for more details [18].

Equation (2) shows that 2eARPES spectral weight is expected at the energy $\omega = E_0^{(GS,N_e)} - E_{-k_1-k_2}^{(\alpha,N_e-2)}$. First we consider the case where the two electrons originate in the same bipolaron. Then $E_0^{(GS,N_e)} \approx E_{BP}(0) + E_0^{(GS,N_e-2)}$ due to the weak bipolaron interactions. The lowest binding energy feature corresponds to $E_{-k_1-k_2}^{(\alpha,N_e-2)} = E_0^{(GS,N_e-2)}$, i.e., the bipolaron is removed and no phonons are left behind. This process is only possible if $k_1 + k_2 = 0$ and has energy $\omega = E_{BP}(0) = 2\mu$ [3], confirming that this bipolaron liquid has no gap to pair excitations, $\omega - 2\mu \leq 0$. The next feature appears at $\omega - 2\mu = -\Omega_{-k_1-k_2}$ if the 2eARPES process removes the bipolaron but leaves behind one phonon carrying the total momentum imparted to the system; this is followed by a two-phonon continuum when two phonons that share the total momentum $-k_1 - k_2$ are left behind, etc.

This process (both electrons ejected from the same pair) competes with the much more probable process of the two electrons being ejected from two different pairs: the latter's probability scales like $N_p(N_p - 1)$ where $N_p = N_e/2$ is the number of pairs, while the former's scales like N_p . This is why we assume throughout that the latter's contribution to the 2eARPES intensity is presumably larger than the former's.

Consider having two electrons ejected from two pairs, starting from $E_0^{(GS,N_e)} \approx 2E_{BP}(0) + E_0^{(GS,N_e-4)}$. The lowest energy eigenstate upon removing an electron from a bipolaron corresponds to formation of a polaron, followed by higher energy replicas when phonons are also left behind. It follows that $E_{-k_1-k_2}^{(\alpha,N_e-2)} \geq E_P(-k_1) + E_P(-k_2) + E_0^{(GS,N_e-4)}$ (we ignore weak interactions between the two polarons and the remaining bipolarons, which is reasonable at very low densities). If the two polarons are far apart, they cannot instantaneously bind into a new bipolaron, thus the lowest binding energy feature is at $\omega \leq 2E_{BP}(0) - E_P(-k_1) - E_P(-k_2) \leq 2E_{BP}(0) - 2E_P(0) = 2\mu - \Delta$, i.e., at energy Δ below the lowest binding energy feature for two electrons ejected from the same pair. This energy separation allows the presumably weaker 2eARPES signal originated from the same pair to be distinguished from the presumably stronger signal from two electrons emitted from two pairs.

This prediction is reminiscent of the results presented in Ref. [6] for a BCS superconductor, which also shows the intensity due to both electrons coming from the same Cooper pair to be energetically separated from that for electrons emitted from different Cooper pairs. Here, we confirm that this energy separation between the two signals also appears for a liquid of incoherent preformed pairs, and that it is an exact result, not a mean-field or BCS approximation.

Results—We now confirm these general expectations using VED for $N_e = 2$, both for singlet s - and triplet p -symmetry pairs, by calculating $A_2(\omega, k_1, k_2)$ from Eq. (2). We find that 50 Lanczos steps typically sufficed to obtain accurately the bipolaron GS wave function and energy. We truncated the sum over the $N_e - 2 = 0$ eigenstates in Eq. (2) at $n \leq n_{\max} = 200$ and employed the Gram-Schmidt reorthogonalization procedure to ensure their orthogonality. An important test for the accuracy of the numerical results is the sum rule $\sum_{k_1, k_2} \int_{-\infty}^{+\infty} d\omega A_2(\omega, k_1, k_2) = 1$ (in the singlet sector) and 2 (in the triplet sector). For all results shown here, this sum rule is satisfied to four significant digits.

Figure 1 shows contour plots of $A_2(\omega, k, -k)$ obtained with VED. We begin with panel (a), which shows the results in the singlet sector, when $U = 0$. Because of the strong el-ph coupling $\lambda = 1$, the $N_e = 2$ GS is a strongly bound bipolaron, hence $A_2(\omega, k, -k)$ gives the 2e ARPES

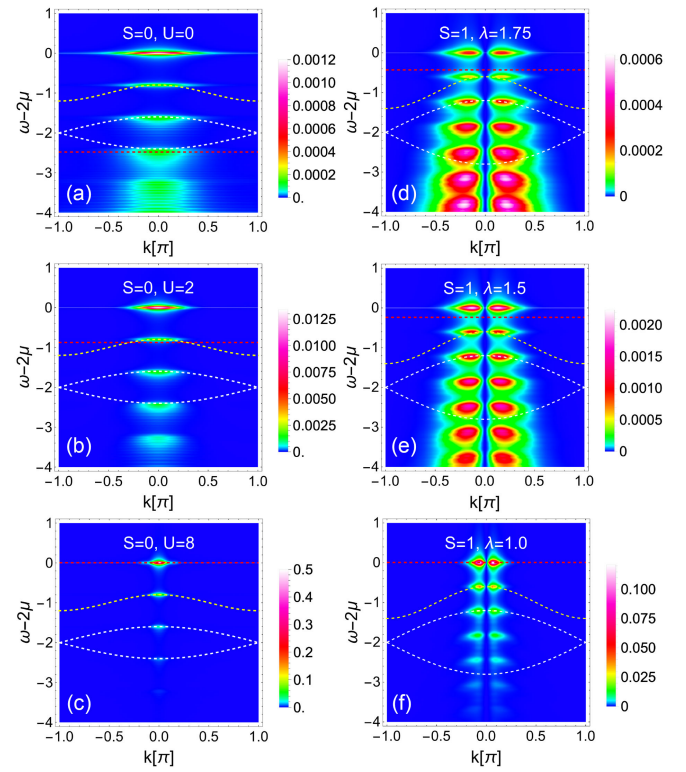


FIG. 1. Contour plots of $A_2(\omega, -k, k)$ for $N_e = 2$ and $\omega_0 = 1$. Panels (a)–(c) are in the singlet channel with $\lambda = 1$, $t_{ph} = -0.1$, and $U = 0, 2, 8$, respectively. Panels (d)–(f) are in the triplet channel with $t_{ph} = -0.2$ and $\lambda = 1.75, 1.5, 1$, respectively. In all panels, the upper yellow dashed line shows the single-phonon sideband at $\omega_{1ph}(k) = -\Omega_k$, and the two lower white dashed lines enclose the shifted continuum representing two-phonon excitations: $\omega_{2ph}^{\pm}(k) = -2\omega_0 \pm 4t_{ph} \cos(k/2)$. The red dashed line at $\omega - 2\mu = -\Delta$ marks the upper edge of the 2eARPES signal from electrons coming from different pairs (not shown). A thin white line indicates $\omega - 2\mu = 0$. We used $N_h = 18$ and a system size $N = 64$.

weight when both electrons are ejected from the same pair, with opposite momentum. As discussed above, the lowest binding energy feature appears indeed at $\omega - 2\mu = 0$ when no phonons are left behind. Conservation of momentum then requires $k_1 + k_2 = 0$, explaining our choice $k_1 = -k_2 = k$ (we verified that this feature is invisible if $k_1 + k_2 \neq 0$).

The next lowest binding energy feature has one phonon in the final state, with momentum $q = -k_1 - k_2 \rightarrow 0$ if $k_1 = -k_2 = k$. Indeed, its energy shift agrees with the phonon energy, shown by the yellow dashed line, at $q = 0$. We checked that this feature is visible at other values of $k_1 + k_2$ and follows the phonon dispersion (see Supplemental Material [18]). The next lowest binding energy feature has two phonons with momenta $q_1 + q_2 = -k_1 - k_2$ in the final state, forming a continuum with weight expected in between the two lower white dashed lines, in agreement with the numerical results. The observation of these higher binding energy features allows the direct measurement of the phonon dispersion. It may also be possible to extract information about the nature of the electron-phonon coupling from their weight [19]. However, we remind the reader that in a system with a finite concentration of carriers, all features with energy $\omega - 2\mu < -\Delta$ (horizontal red dashed line) are covered by the presumably much larger spectral weight from processes where the two electrons are ejected from different pairs, see Supplemental Material for more details [18]. In panel (a) both the one- and the two-phonon features are above this threshold, however as U is increased and Δ decreases, eventually only the feature at $\omega - 2\mu = 0$ will be resolved [panel (b)]. Finally, for a large U [panel (c)], the $N_e = 2$ GS corresponds to two unbound polarons and there is no “single pair” signal.

Similar results are obtained for $A_2(\omega, k, -k)$ corresponding to emission from a triplet p -symmetry pair, as shown in panels (d)–(f) of Fig. 1. Here U is irrelevant, and we can control the binding energy through varying $|\Delta E|$, see discussion above. The obvious difference between the triplet and singlet pairs 2eARPES intensity is the “node” obtained at $k = 0$ for the former, as required by Pauli’s principle.

Next, we analyze the evolution of the lowest binding energy feature at $\omega - 2\mu = 0$ with the binding energy, and argue that its existence (and thus, the existence of preformed pairs) can be confirmed even when the energy resolution is comparable to or even worse than the gap size Δ . Comparison of this feature in panels (a) and (b) of Fig. 1 shows that it becomes narrower in k space as Δ decreases. This is easy to understand if we consider the bipolaron GS,

$$\begin{aligned}
 |\psi_0^{(GS,2)}\rangle = & \sum_k \alpha_k c_{k,\uparrow}^\dagger c_{-k,\downarrow}^\dagger |\emptyset\rangle + \sum_{k,q} \alpha_{k,q} c_{k,\uparrow}^\dagger c_{-k-q,\downarrow}^\dagger b_q^\dagger |\emptyset\rangle \\
 & + \sum_{k,q_1,q_2} \alpha_{k,q_1,q_2} c_{k,\uparrow}^\dagger c_{-k-q_1-q_2,\downarrow}^\dagger b_{q_1}^\dagger b_{q_2}^\dagger |\emptyset\rangle + \dots
 \end{aligned}$$

The spectral weight of the lowest binding energy feature (due to ejection of the two electrons and no phonons left behind) is proportional to $|\alpha_k|^2$. The Fourier transform of α_k defines the amplitude of probability to find the two bound electrons at a relative distance δ apart. For more strongly bound pairs, this peaks at a smaller δ , implying a larger spread in k —precisely what is observed by comparing panels (a) and (b) for the singlet pair, and (d) and (e) for the triplet pair.

Another way to illustrate this is to show the momentum resolved 2eARPES weight $\gamma_{pair}(k_1, k_2) = A_2(\omega = 2\mu, k_1, k_2)$ due to this feature only, which can be obtained by integrating over a narrow energy range. Figure 2 shows $\gamma_{pair}(k_1, k_2)$ for parameters identical to those in Fig. 1, illustrating beautifully the inverse relationship between the spread of this feature along the $k_1 + k_2 = 0$ line, and the binding energy of the pair.

These results verify that the 2eARPES signal at $\omega = 2\mu$ that originates when the electrons are emitted from the same pair appears only when $k_1 = -k_2$. If the detectors are at equal distance from the sample and select photoelectrons with equal momenta $|\mathbf{k}_1| = |\mathbf{k}_2|$, then the times of flight are equal. Coincidence detection of such photoelectrons means that they must have been emitted at the same time, i.e.,

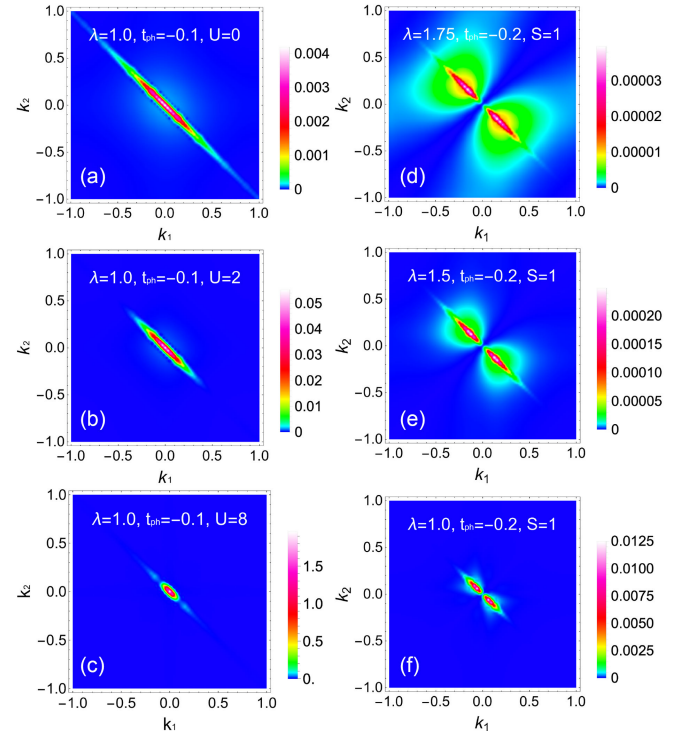


FIG. 2. Momentum-resolved spectral weight of the lowest binding energy feature $\gamma_{pair}(k_1, k_2) = A_2(\omega = 2\mu, k_1, k_2)$. In order to increase the number of k -points, we used $N_h = 16$ and $N = 64$. Panels (a)–(c) are in the singlet channel, while panels (d)–(f) are in the triplet channel, for parameters identical to those in Fig. 1.

$\tau = 0$, explaining why we made this choice when simplifying Eq. (3). However, the conservation of momentum of the system that underlies the $k_1 + k_2 = 0$ condition is always valid, hence we expect a similar conclusion even if the full Eq. (3) is evaluated.

This asymmetry of the single-pair signal in the (k_1, k_2) plane explains how the existence of preformed pairs could be inferred even if the binding energy Δ was smaller than the energy resolution of the apparatus, so that this signal was covered by the presumably larger signal from electrons emitted from different pairs. Based on Eq. (2), we can estimate the latter to be proportional to the convolution $\int d\omega_1 \int d\omega_2 A_1(\omega_1, k_1) A_1(\omega_2, k_2) \delta(\omega - \omega_1 - \omega_2)$, where the (ARPES) single particle spectral weights are,

$$A_1(\omega, k) = \frac{1}{\pi} \text{Im} \sum_n \frac{|\langle \psi_{-k}^{(n, N_e-1)} | c_{k\sigma} | \psi_0^{(GS, N_e)} \rangle|^2}{\omega - i\eta + E_{-k}^{(n, N_e-1)} - E_0^{(GS, N_e)}}. \quad (4)$$

Let $\beta(k_1, k_2)$ be the result when this signal from two different pairs is integrated for a small energy range below $\omega = 2\mu - \Delta$. Because $A_1(\omega, k) = A_1(\omega, -k)$, it follows immediately that $\beta(k_1, k_2) = \beta(-k_1, k_2) = \beta(k_1, -k_2) = \beta(-k_1, -k_2)$, i.e., this contour plot has C_4 symmetry in the (k_1, k_2) space, unlike the strong C_2 symmetry of $\gamma_{\text{pair}}(k_1, k_2)$ [plots of $\beta(k_1, k_2)$ are shown in Supplemental Material [18]]. As a result, if the total energy-integrated signal at the top of the spectrum has more weight along $k_1 + k_2 = 0$ than along $k_1 = k_2$, this confirms the existence of preformed pairs.

While the numerical results presented here are at $T = 0$ and for a single pair on a 1D chain, we believe that these two fingerprints pointing to the existence of preformed pairs are present at finite T for finite densities in any dimension.

At finite temperatures, a fraction of the pairs are thermally dissociated. As noted above, 2eARPES signal from unbound quasiparticles starts at $\omega - 2\mu = 0$, so it is always superimposed over the 2eARPES signal coming from electrons ejected from the same pair. Nevertheless, its C_4 symmetry (and different evolution with temperature) should make it distinguishable from the pair signal.

At finite concentrations and low- T , the system is superconducting if a macroscopic fraction of the pairs condense in the $K = 0$ state. In this case, the 2eARPES signal coming from electrons ejected from the same pair is proportional to that shown in Figs. 1 and 2 for a single pair with $K = 0$, in agreement with the findings of Ref. [8].

Another option is an incoherent liquid of preformed pairs, where each pair momentum K has a microscopic occupation number $n_K/N \rightarrow 0$. This occurs at $T = 0$ for our 1D model [3], but more generally might be expected as an intermediate state between a superconductor and the normal state, if superconductivity is lost due to phase fluctuations (not to unpairing, like in BCS). The occupied pair K momenta are set by k_F , as demonstrated in Ref. [3]

for the “Bose sea” GS of our model. The 2eARPES signal from electrons ejected from a pair with momentum $K \neq 0$ is similar to those shown in Fig. 2 but shifted to $k_1 + k_2 = K$. Thus, the very asymmetric C_2 pattern remains, however it acquires a finite “transverse” width proportional to k_F , as further discussed in the End Matter. (In Figs. 1, 2, the widths are set by the broadening η). All these considerations carry over in higher D in the plane parallel to the sample surface, $\mathbf{k}_{1,\parallel} + \mathbf{k}_{2,\parallel} = \mathbf{K}_{\parallel}$.

Summary—To conclude, we identified two fingerprints that allow the separation of the presumably small contribution of processes where both electrons are emitted from the same pair, from the total 2eARPES intensity. These fingerprints are direct consequences of the conservation of momentum and energy, therefore we are confident that they will also appear in more sophisticated theoretical descriptions of 2eARPES spectroscopy and in more comprehensive numerical methods, such as diagrammatic Monte Carlo, which can calculate 2eARPES intensities for finite concentrations in the thermodynamic limit of Hamiltonian (1) in any dimension [20–23]. Experimental observation of these fingerprints will confirm the existence of electron pairs in the system, and whether they are coherent (superconducting) or not.

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Data availability—The data that support the findings of this article are not publicly available upon publication because it is not technically feasible and/or the cost of preparing, depositing, and hosting the data would be prohibitive within the terms of this research project. The data are available from the authors upon reasonable request.

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End Matter

Figure 3 shows the evolution of the momentum-resolved spectral weight of the lowest-binding energy feature $\gamma_{pair}(k_1, k_2) = A_2(\omega = 2\mu, k_1, k_2)$ (integrated over a narrow energy range) with increasing electron density $n = N_e/N = 0.03, 0.16$, and 0.28 (from top to bottom panels). The left column panels are for the singlet channel with $\lambda = 1, t_{ph} = -0.1, U = 2$, while the right column panels are in the triplet channel with $\lambda = 1.75, t_{ph} = -0.2$. For these values, the GS are liquids of incoherent pairs (coherence is not possible in 1D because of quantum fluctuations).

These results were obtained by summing individual 2eARPES intensities obtained with VED from Eq. (2) for individual pairs with momenta $|K| \leq n\pi = k_F$. This “Bose sea” of noninteracting pairs was shown in Ref. [3] to reproduce accurately results obtained with density matrix renormalization group at finite concentrations.

The results of Fig. 3 confirm that even for a finite-density incoherent liquid of pairs, the strong C_2 symmetry is maintained. This is a direct consequence of momentum conservation, as the two electrons ejected from a pair with momentum K must obey $k_1 + k_2 = K$. The “transverse” broadening then gives a direct measure of the occupied pair momenta K .

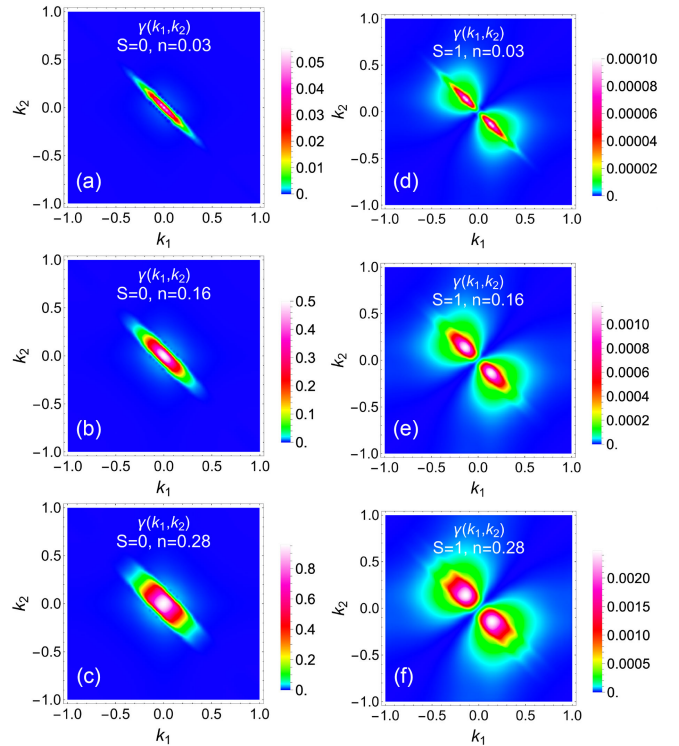


FIG. 3. Momentum-resolved spectral weight of the lowest-binding energy feature $\gamma_{pair}(k_1, k_2) = A_2(\omega = 2\mu, k_1, k_2)$ for densities $n = N_e/N = 0.03, 0.16, 0.28$ in the top, middle, and bottom panels, respectively. Panels (a)–(c) are in the singlet channel with $\lambda = 1, t_{ph} = -0.1, U = 2$, while panels (d)–(f) are in the triplet channel with $\lambda = 1.75, t_{ph} = -0.2$.