Observation of Pauli Crystals

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The Pauli exclusion principle is a fundamental law underpinning the structure of matter. Because of their antisymmetric wave function, no two fermions can occupy the same quantum state. Here, we report on the direct observation of the Pauli principle in a continuous system of up to six particles in the ground state of a two-dimensional harmonic oscillator. To this end, we sample the full many-body wave function by applying a single atom resolved imaging scheme in momentum space. We find so-called Pauli crystals as a manifestation of higher order correlations. In contrast to true crystalline phases, these unique high-order density correlations emerge even without any interactions present. Our work lays the foundation for future studies of correlations in strongly interacting systems of many fermions.

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Correlated fermions lie at the heart of many open questions concerning quantum matter that remain unresolved to this day. Knowledge of the type and origin of correlations, especially of higher orders, is an essential cornerstone in the endeavor of solving these complex many-body systems [1–3]. Strong correlations are, however, not exclusive to interacting particles as was demonstrated already in 1956 by the famous experiment of Hanbury Brown and Twiss [4]. Bosons tend to occupy the same quantum state while multiple occupation of a single state is forbidden for fermions due to the Pauli exclusion principle.

In a degenerate gas of neutral fermions, Pauli exclusion reveals itself through suppression of collisions [5,6] and an effective Fermi pressure [7]. Antibunching has been observed directly in time-of-flight measurements [8,9] or via the suppression of density fluctuations [10,11]. Quantum gas microscopy advances have led to the first single atom resolved observation of Pauli blocking in the band insulating regime of a lattice potential [12].

Here, we extend single atom resolved measurements of fermionic correlations to continuous systems. We study ultracold, fermionic atoms that are confined to a two-dimensional harmonic oscillator potential. Even in the absence of interactions, the Pauli exclusion leads to local high-order density correlations between the atoms that go beyond a simple Fermi hole. The geometric patterns that we observe are clearly distinct from those in interaction driven systems and they have been termed Pauli crystals [13].

Pauli crystals only emerge at very low temperatures where the particles become quantum degenerate and their Fermi energy dominates over temperature and trap imperfections. This requires charge-neutral noninteracting systems and control on very low absolute energy scales [14]. The structures act as a starting point for the study of correlations in continuous systems with single atom resolution.

The experimental observation of Pauli crystals relies on two essential capabilities: the preparation of 6 noninteracting fermions in a well-defined quantum state and the detection of 6-body correlations in the relative positions or momenta of these particles. We perform our experiments with a balanced mixture of two hyperfine states of ⁶Li confined by the superposition of an optical tweezer and a single site of a one-dimensional optical lattice in the vertical direction [see Fig. 1(a)]. The large ratio between axial [ωₐ = 2π × 6560(6) Hz] and radial [ωᵣ = 2π × 983(5) Hz] trap frequencies allows us to work in a quasi-2D regime for sufficiently small temperature T and particle number N. In this limit all the atoms occupy the motional ground state in the axial direction and the dynamics are limited to the harmonic confinement in the radial direction.

The nth energy level of a symmetric two-dimensional harmonic oscillator is (n+1)-fold degenerate, leading to three lowest closed-shell configurations filled with 1, 3, and 6 fermions per spin state respectively [see Fig. 1(a)]. A spilling technique that was initially pioneered for one-dimensional systems [15] and that we extended to two dimensions [16] allows us to reach these configurations filled with two spin components. The preparation fidelities are 92(7)% for the 3 + 3 (N⇑ + N⇓) and 56(3)% for the 6 + 6 ground states, respectively. We work with a two component mixture instead of a single component gas since attractive interactions during the spilling sequence improve the preparation fidelity. To create a noninteracting mixture, we make use of a Feshbach resonance [17] and adiabatically ramp the magnetic offset field B to a zero crossing of the scattering length a₃D at 568 G once the system is initialized in the ground state. For all the measurements presented in the following, the atoms of only one of the two spin components are imaged.

We extract momentum correlations from our system by first mapping the initial momenta of the particles onto their
position by a TOF expansion for a quarter trap period in a single lattice site with \( \omega_{\text{TOF}} = 2\pi \times 20.7(5) \text{ Hz} \). This is followed by the single atom detection fluorescence imaging scheme discussed in detail in Ref. [19]. By collecting on average 20 photons per atom on a single photon counting camera, this method allows us to detect atoms of a single spin component in free space with fidelities on the order of 95% [see Fig. 1(b)]. Each image obtained in this way represents a single sample \( \frac{\hat{p}}{\sqrt{\hbar m \omega}} \) of the \textit{in situ} momentum distribution for every particle \( i = 1, \ldots, N \) of one of the spin components [see Fig. 1(c)].

The harmonic confinement plays an important role for our measurements. We prepare our atoms in eigenstates of the harmonic oscillator. The real space wave functions are therefore given by Slater determinants that contain superpositions of Hermite polynomials [20]. These wave functions are invariant under continuous Fourier transforms and therefore also invariant under our TOF expansion. The expansion simply corresponds to a magnification of the in-situ wave function by a factor of 50. This leads to an effective imaging resolution of approximately 200 nm. The natural scale of the harmonic oscillator states is given by \( p_0 = \sqrt{\hbar m \omega} \), or \( l_0 = \hbar / p_0 = 1.31(1) \ \mu \text{m} \).

Our measurements have been performed by preparing the system at the lowest accessible temperatures with two \( (N = 3) \) or three \( (N = 6) \) harmonic oscillator shells filled and collecting 9994 and 19291 TOF images, respectively. Only images where the correct number of atoms have been detected are investigated further, leading to postselection rates of 25% and 28%. We process these measurements as suggested by Ref. [13] to reveal correlations between the fermionic particles [see Fig. 1(c)]. In the first step, we subtract the respective center of mass momentum \( \vec{p} \) from each set of momenta \( p_i \). We find that the width of the center of mass momentum scales with the inverse square root of the total mass (i.e., \( m_{\text{tot}} = N m \), where \( m \) is the mass of one \(^6\text{Li} \) atom) as expected [20].

A histogram of the remaining relative momenta yields the one-particle momentum distribution [see Fig. 2(a)]. The latter expresses the probability to find one particle with momentum \( p_x - \vec{p}_x \) and \( p_y - \vec{p}_y \) when integrating over all possible momenta of all other particles. We stress that the one-particle momentum density can also be obtained from average density images without single particle resolution and does not reveal any higher order correlations. Our data, however, contain more information: we know the full configuration of all particles in every single realization of the experiment.

To extract correlations, a second processing step is necessary. Because of the radial symmetry of our system, the angle distribution of all particles \( N(\phi) \) is homogeneous (see Fig. 3). The rotational symmetry is only broken in each experimental realization by the measurement itself and the particles align with respect to a random axis that is different for each set of momenta \( p_i \). We rotate each set independently to a new coordinate system \( \vec{\hat{p}}_i \) by an angle that minimizes the distance to a chosen target configuration [13, 20]. Strong correlations in both the \( N = 3 \) and the \( N = 6 \) particle state become apparent immediately in the \( N \) particle momentum configuration distributions that we obtain in this way [see Figs. 2(b,c)].

The observation of these so-called Pauli crystals confirms theoretical predictions [13]. To rule out an artifact of our analysis, we compared to images that were obtained after shuffling the atom momenta between different experimental runs [20]. This is especially important since it has been shown that other distance measures can cause a bias towards the target configuration [21]. In addition, we calculate the angular density-density correlation function \( C_2(0, \phi) \) that expresses the probability of finding a second particle at an angle \( \phi_2 = \phi \) when one particle is fixed at \( \phi_1 = 0 \). The result for \( N = 6 \) is shown in Fig. 3 and clearly shows the presence of four peaks, as expected for the Pauli crystal and in agreement with a Monte Carlo simulation [20].

We find that the observed configuration distributions exhibit weaker modulation than the simulations we perform for systems at zero temperature. We quantify this effect...
through the contrast $C(p)$, which we define as one minus the ratio of the minima and maxima of a fit to the configuration distribution at a fixed radial momentum $p = 2p_0$ [see Figs. 4 (a,d)]. Apart from technical limitations like the point spread function of our imaging setup or fluctuations of trap potentials, we identify the finite temperature of our initial state as the main cause of this reduction.

To study the effect of finite temperature in more detail, we “melt” the $N = 6$ crystal by increasing the mean energy of the initial state [see Figs. 4(a)–4(c)]. To this end, we modulate the confining potential at twice the trap frequency $2\omega_r$ with variable amplitudes and take around 3000 images
at each setting. Trap imperfections like anharmonicity and anisotropy in combination with small potential drifts cause the system to dephase on timescales much faster than the modulation time of $t = 50$ ms and the excitation is therefore not coherent.

The total energy of the system is extracted from the momentum measurements and averaged over each data set. We find that the energy of the lowest temperature initial state, without any applied modulation, is $E = 13.1 \hbar \omega_{\text{r}}$. This value is about 5% below the expected $N = 6$ ground state energy of $E_g = 14 \hbar \omega_{\text{r}}$. Our measurement of the energy entails systematic uncertainties like the error on the frequency measurement of the expansion potential $\omega_{\text{TOF}}$ (≈2%) or the magnification of our imaging setup that both enter quadratically. Taken together, these uncertainties may account for the observed shift to lower energies, which is systematic for all data points.

A comparison of the relative change in energy and contrast clearly displays the effect of the modulation [see Fig. 4(e)]. We find that the contrast reduces with increasing mean energy. A linear fit to the contrast leads to a slope of $dC/dE_{\text{exp}} = -0.075(13)/\hbar \omega_{\text{r}}$. We compare this value to the slope of $-0.048(3)/\hbar \omega_{\text{r}}$ that we extract from a simulation using thermal states as described in Ref. [14] [see Fig. 4(f)]. While our finite, noninteracting system is not expected to thermalize after the modulation, the number of excited states that might contribute to the density matrix at a given excitation energy is very large. The dimension of the Hilbert space of excitations from the $N = 6$ particle ground state to the next few higher shells is already on the order of a few hundred thousands. Trap imperfections like anharmonicity and anisotropy lead to coupling between the different degrees of freedom. Together with small potential drifts this motivates our comparison of the measured final state to a thermal mixture. In addition to deviations from a thermal state, the discrepancies we find may be due to the systematic uncertainties in determining the total kinetic energy or due to additional excitations in the axial direction. The thermalization dynamics that may occur in the presence of interactions are an exciting topic for future studies.

In conclusion, we find that the finite temperature of our experiments is one factor that contributes to the reduced contrast of the measured Pauli crystals. The ability to melt the Pauli crystal clearly shows that the observed correlations originate from the fermionic nature of our initial state. Neither fidelity nor resolution of our imaging technique depend on the initial state energy.

We have observed that Pauli’s principle leads to the formation of striking geometric configurations of fermions confined in a trap, even in the absence of any interactions. The structure is not apparent in the density distribution directly but only reveals itself in correlations between relative positions or momenta. Each single experimental realization still fluctuates and can deviate significantly from the most probable configuration.

Many interacting mesoscopic systems, like ions [22], Rydberg atoms [23] or dipolar gases [24] show self-ordering and crystalline structures akin to what is observed here. While this motivates the term Pauli crystal, we stress that in our case translational symmetry is not broken and no long-range order is present. The ground state is a coherent superposition of all possible configurations and the rotational symmetry is only broken through the actual measurement.

We generically expect this kind of order driven by quantum statistics to be present in few-fermion systems of fixed particle number. Our simulations show that similar structures appear, for example, also in box potentials. While the exact trapping potential is not important, the order will be pronounced as long as the interparticle spacing is not much smaller than the size of the system. As the system size is increased, we expect the structures to decrease in contrast until they vanish for a homogeneous, infinite Fermi gas.

Our measurements demonstrate that the correlation environment of individual particles can now be accessed in continuum systems. This unique capability will be extremely useful for future studies of correlations in strongly interacting systems. Our single atom imaging scheme can be made sensitive to a second spin state [19], which may directly reveal pairing correlations near the few-body precursor of a phase transition that we have recently observed [16]. Scaling up the system size will enable us to shed further light on many open questions concerning two-dimensional Fermi gases like the nature of its normal phase [25] or to study the emergence of Cooper pairing [1]. The detection of momentum correlations represents one milestone on the path towards understanding many of these complex fermionic many-body systems.

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