Observation of critical phase transition in a generalized Aubry-André-Harper model on a superconducting quantum processor with tunable couplers

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Quantum simulation enables study of many-body systems in non-equilibrium by mapping to a controllable quantum system, providing a new tool for computational intractable problems. Here, using a programmable quantum processor with a chain of 10 superconducting qubits interacted through tunable couplers, we simulate the one-dimensional generalized Aubry-André-Harper model for three different phases, i.e., extended, localized and critical phases. The properties of phase transitions and many-body dynamics are studied in the presence of quasi-periodic modulations for both off-diagonal hopping coefficients and on-site potentials of the model controlled respectively by adjusting strength of couplings and qubit frequencies. We observe the spin transport for initial single- and multi-excitation states in different phases, and characterize phase transitions by experimentally measuring dynamics of participation entropies. Our experimental results demonstrate that the newly developed tunable coupling architecture of superconducting processor extends greatly the simulation realms for a wide variety of Hamiltonians, and may trigger further investigations on various quantum and topological phenomena.

Introduction.—Using controllable quantum systems, quantum simulation provides a powerful approach to study many-body physics, which might be challenging for a classical computer [1, 2]. In analogue quantum simulation, specific model Hamiltonians can be directly realized by engineering the platform Hamiltonians such that dynamics of real quantum systems can be studied in a controllable manner, such as in trapped ions [3–5], atoms in optical lattices [6–9], superconducting qubits [10–12], and nuclear spins [13, 14]. Particularly, superconducting quantum simulation can explore a wide regime from localization to weak and strong thermalization in non-equilibrium quantum many-body systems [15–23].

On the other hand, the 1D Aubry-André-Harper (AAH) model [24, 25], as a workhorse for studying localization and topological states, has attracted much attention both theoretically and experimentally [16, 26–33]. The original AAH model can be derived from a 2D quantum Hall system with nearest-neighbor hopping. When considering the next-nearest-neighbor hopping, one can deduce a generalization of the AAH model with both on-site and off-diagonal quasi-periodic modulations [34, 35]. The generalized AAH (GAAH) model shows different and interesting localization and topological properties, for instance, the critical phase featured by multifractal wave functions and the topological adiabatic pumping [36–41]. With the development of experimental technologies, the GAAH model has been realized in photonic crystals with on-site or off-diagonal modulation [31] and cold atoms systems in momentum space [42]. With flexible control and precise measurement of superconducting processor, the GAAH model may be simulated analogously, given off-diagonal quasi-periodic modulations can be implemented precisely.

In our experiment, taking advantage of newly developed tunable coupling architecture [43, 44], we simulate the GAAH model for a wide variety of parameters on a superconducting processor. By adjusting both qubits and couplers, we experimentally observe the dynamics of the extended, localized, and critical phase in the GAAH model, and investigate the phase transition from the perspective of non-equilibrium dynamics. We observe that in the critical phase, the spin can propagate over a range intermediate between that of the extended phase and of the localized phase, for both initial single- and multi-excitation states. In addition, we quantify how fast initial states spread over the Hilbert space for different phases by experimentally measuring the time evolution of participation entropies, and characterize the transition among the extended, localized, and critical phase by calculating averaged late-time participation entropies.

Model and set-up.—Our quantum processor consists of a chain array of $L = 10$ transmon superconducting qubits, and 9 tunable couplers with each placed between every two nearest-neighbor qubits, which enable an accurate control of couplings (see Fig. 1(a) and Supplementary Material for details). The effective Hamiltonian of the qubits system can be described by the Bose-Hubbard model,

$$\hat{H} = \sum_{j=1}^{9} (J_{j,j+1} \hat{a}^\dagger_{j} \hat{a}_{j+1} + \text{H.c.}) + \sum_{j=1}^{10} \hbar \omega_j \hat{n}_j - \frac{U}{2} \hat{n}_j (\hat{n}_j - 1),$$  (1)
where $\hat{a}_j^\dagger (\hat{a}_j)$ is the photon creation (annihilation) operator, $\hat{n}_j \equiv \hat{a}_j^\dagger \hat{a}_j$ is the number operator, $J_{j,j+1}$ is the tunable nearest-neighbor coupling strength, $h_j$ is the tunable local potential, and $U_j$ denotes the qubit anharmonicity serving as the on-site interaction. Since $U_j \gg J_{i,j}$ for our processor, an excess of energy is needed for having more than one photon at each site, so the system can be described by hard-core bosons, or equivalently spin-1/2 spins, with the conservation of the total photons (or spins) due to the $U(1)$ symmetry.

In our superconducting processor with tunable couplers, the coupling $J_{j,j+1}$ contains two parts: (i) direct coupling between nearest-neighbor qubits $J_{j,j+1}^0$ and (ii) superexchange interaction via the coupler in between $J_{j,j+1}^{SE} \propto 1/\Delta_{j,j+1}$ with $\Delta_{j,j+1}$ denoting frequency detuning between the $j$-th coupler and the two nearest-neighbor qubits (see Supplementary Material for details). Therefore, by applying fast voltages to the $Z$ control lines of the couplers, $J_{j,j+1}$ can be tuned individually from −30 to +4.8 MHz in our processor. In addition, by applying the fast Z pulse on each qubit, the local potential can also be arbitrarily tuned relative to resonant frequency $\sim 4.36$ GHz. In this way, we can finally realize a hard-core bosonic version of the GAAH model,

$$\frac{\hat{H}}{\hbar} = \lambda \sum_{j=1}^{9} \left( 1 + \mu \cos \left[ 2\pi \left( j + \frac{1}{2} \right) \alpha + \delta \right] \right) \hat{a}_j^\dagger \hat{a}_{j+1} + \text{H.c.} + \lambda \sum_{j=1}^{10} V \cos (2\pi j \alpha + \delta) \hat{a}_j^\dagger \hat{a}_j. \quad (2)$$

Here, as depicted in Fig. 1(b), we take a quasi-periodic modulation $J_{j,j+1} = \lambda (1 + \mu \cos [2\pi (j + 1/2) \alpha + \delta])$ and $h_j = \lambda V \cos (2\pi j \alpha + \delta)$, where $\mu$ and $V$ indicate the off-diagonal and on-site modulations amplitudes, respectively, and we choose $\lambda/(2\pi) = 4$ MHz throughout the work. Besides, $\alpha = (\sqrt{5} - 1)/2$ is the irrational frequency which takes the same value for on-site and off-diagonal modulations, and $\delta \in [-\pi, \pi)$ is an arbitrary global phase offset.

For different parameters $\mu$ and $V$, localization property of the eigenstates of Hamiltonian Eq. (2) can be characterized by the inverse participation ratio (IPR) [45]. In Fig. 1(c), we plot the localization phase diagram of Hamiltonian Eq. (2). The heatmap shows the eigenstate’s IPR $= \sum_{i} |\psi_{n,i}|^4$, where $\psi_{n,i} = \langle \psi_n | i \rangle$ is the wave function coefficient of the eigenstate $|\psi_n \rangle$ expressed in the computational basis $\{|i\rangle\}$. Here, we actually displays the negative logarithm of the IPR, in order to distinguish the three phases more clearly, and to associate with the participation entropy in the next section. Since mobility edges are absent in our model [39], (i) for $V < 2, \mu < 1$, all bulk eigenstates are extended, denoted as the extended phase, where the IPR vanishes and its negative logarithm is close to $\log L \sim 6.9$; (ii) for $V > 2 \max(1, \mu)$, all bulk eigenstates are localized with the IPR close to one and its negative logarithm close to zero, denoted as the localized phase; (iii) in the rest, the eigenstates are critical with intermediate IPRs, denoted as the critical phase.

**Observation of spin transport in the GAAH model.**—First, we show that the GAAH model can be simulated with our su-
perconducting qubits in a tunable coupling architecture. The experimental scheme is that three phases of the GAAH model manifest distinguishing localization properties by measuring the photon occupancy probabilities for specific initial states. We initially excite the leftmost qubit $Q_1$, i.e., the system is initialized as $|\psi(0)\rangle = |1000000000\rangle$, where $|0\rangle$ ($|1\rangle$) denotes the ground (excited) state of a qubit. Then we apply the fast Z pulse on each qubit and coupler, and the system will evolve under the Hamiltonian Eq. (2), satisfying Schrödinger equation $|\psi(t)\rangle = e^{-iHt}|\psi(0)\rangle$. We monitor its dynamics from $t = 0$ to 500 ns, by measuring the photon occupancy probabilities of each qubit $P_j(t) = \langle \psi(t) | a_j^\dagger a_j | \psi(t) \rangle$. For each time point, we perform 5000 repeated single-shot measurements.

The experimental results for the three phases are plotted in the left panel of Fig. 2(a-c), with a comparison of numerical simulations in the right panel of Fig. 2(a-c). As shown in Fig. 2(a), in the extended region, the spin transport is not blocked, and a lightcone-like propagation and reflection at the boundary are still visible when weak off-diagonal and on-site quasi-periodic disorder exists. As the opposite, for sufficiently large on-site disorder $V$, the spin is fully localized, and only the initially occupied site has a occupancy probability close to one at any time (Fig. 2(c)). In the critical region, the spin tends to oscillate around adjacent sites of the initially occupied site, and the propagation range is intermediate between the above two (Fig. 2(b)).

With the capability of precise simultaneous control and readout, we can also prepare initial product states to probe the largest Hilbert space (i.e., the half-filled sector), by exciting selected $M = N/2 = 5$ qubits and keeping the rest in their ground state. The experimental sequences are shown in Fig. 1(d). Here, we focus on an initial Néel state $|\psi(0)\rangle = |1010101010\rangle$, and measure the photon occupancy probabilities from $t = 0$ to 500 ns. The experimental and numerical results are plotted in Fig. 2(d-f). As shown in Fig. 2(d), in the extended region, the mean photon occupancy probabilities oscillates around 0.5 at long times for all ten qubits with a small fluctuation, showing a pattern of oscillation between odd and even sites back and forth. In Fig. 2(e), the photon occupancy probabilities for all qubits are also close to 0.5 in the critical region. However, different from the extended region, the photon occupancy probabilities in the critical region exhibit a certain degree of dependence on the initial configuration, that is, the photon occupancy probabilities at the initially occupied

FIG. 2. The time evolutions of qubit-resolved photon occupancy probabilities $P_j(t)$ for the system initialized in (a–c) $|\psi(0)\rangle = |1000000000\rangle$ , and (d–f) $|\psi(0)\rangle = |1010101010\rangle$, in (a, d) the extended phases with $\mu = 0.5$ and $V = 0.5$; (b, e) the critical phases with $\mu = 2.0$ and $V = 0.5$; (c, f) the localized phases with $\mu = 0.5$ and $V = 4.0$. The left panel of each figure shows experimental data, and the right panel shows numerical simulation. Experimental data are averaged over 5 realizations of random chosen phases $\delta$, while numerical simulation are averaged over 50 realizations of random chosen phases $\delta$ with decoherence taken into account.
The dynamical participation entropy is a characterization quantifying how fast $|\psi(t)|^2$ spreads over the Hilbert space. Here we select $2 \times M = 10$ initial states which is far from equilibrium, taking the form $|\psi_i(0)\rangle = |10\rangle \otimes |01\rangle \otimes |i\rangle$, and $|\psi_{M+i}(0)\rangle = G|\psi_i(0)\rangle$ with a global spin-flip operator $G = \prod_{j=1}^{L} \hat{\sigma}_j^z$, for $i = 1, \ldots, M$. The experimental data shown in Fig. 3 are averaged over these 10 initial states, and the measured multi-qubit probabilities $p_i$ are post-selected within the half-filled sector due to the $U(1)$ symmetry before calculating the participation entropy.

Figure 3(a) displays the time evolution of participation entropy for the system quenched into the three phases. After a fast initial relaxation, the participation entropy oscillates around some certain value, which varies in different phases. For both small $\mu$ and $V$, the system lies in the extended phase, where the late-time participation entropy keeps at a high value with a small oscillation (see the yellow points in Fig. 3(a)). For small $\mu$ and sufficiently large $V$, the participation entropy oscillates around a much smaller value (see the purple points), as we would expect in the localized phase. For relatively large $\mu$, the late-time participation entropy stands between the extended and localized cases (see the purple points), which behaves similarly to that with a small $\mu$ and intermediate $V$ around the extended to localized transition point (see the data of $V = 2$ in Fig. 3(b) for comparison).

Figure 3(b) and (c) show the time evolution of participation entropy with increasing $V$ for fixed $\mu = 0.5$, and increasing $\mu$ for fixed $V = 1$, respectively. As shown in Fig. 3(b), with the increase of $V$, the growth of participation entropy is suppressed significantly, reflecting a transition from the extended region to the localized region. In contrast, as $\mu$ increases to approach the theoretical transition point $\mu_c = 1$, the growth of participation entropy slows down compared to the extended phase. However, as $\mu$ continues to increase, the curve of time evolution of participation entropy rises slightly again. The late-time participation entropy in the critical phase reflect a multifractal behavior, and the multifractal analysis can be found in the Supplementary Material.

In addition, the averaged participation entropy at long times can be used as an experimentally accessible characterization of phase transition. Here, we experimentally measure the averaged late-time participation entropies $S_{\Omega}^{PE}$ along three paths I, II and III in the $\mu - V$ plane (see Fig. 4(a)), corresponding to extended to localized transition, extended to critical transition,
and localized to critical transition, respectively. Averages are taken over a time window from 350 to 450 ns (see the grey area in Fig. 3(a)), and among the 10 initial states as defined before. In Fig. 4(a), we also display the numerical results of the averaged late-time participation entropy calculated using Hamiltonian (2) in the same way as the experimental processing, for the whole \( \mu - V \) plane (\( 0 \leq V \leq 4, 0 \leq \mu \leq 2 \)) as a reference, which exhibits a similar phase diagram to IPR averaged over eigenstates in Fig. 1(c). Because the time window 350–450 ns is far less than the averaged \( T_2 \sim 22.3 \mu s \) (see Supplementary Material), and \( T_2 \) has little effect on the averaged late-time participation entropies, we ignore the effect of decoherence in the simulation here. The numerical simulation and experimental results are consistent well with each other.

Comparisons of the specific experimental data with numerical simulation of the three paths I, II and III are plotted in Fig. 4(b), (c) and (d), respectively. Different from the path I, where \( S_{PE}^2 \) decreases monotonically as \( \mu \) increases, for the path II, \( S_{PE}^2 \) first decreases to the minimum around the theoretical transition point \( \mu_c = 1.0 \), and then increases slightly but keeps almost unchanged for increasing \( \mu \) in the critical phase. Note that both the experimentally measured and numerically calculated minimum \( S_{PE}^2 \) are reached at \( \mu = 1.25 \) instead of \( \mu_c = 1.0 \) for \( L = 10 \), which we attribute to finite size effects. To see this, we also calculate the averaged late participation entropies for larger system sizes \( L = 14, 18, 22 \), which we rescale by a factor \( \frac{\log N_L}{\log N} \) with \( N_L = \left( \frac{L}{L/2} \right) \), i.e., \( S_{PE}^2(L) = \frac{\log N_L}{\log N} \cdot S_{PE}^2(L) \), for a direct comparison to the experimental system size \( L = 10 \). In the systems with \( L = 14, 18, 22 \), the minimum \( S_{PE}^2 \) are reached at \( \mu = 1.0 \). For the path III, i.e., the localized to critical transition, \( S_{PE}^2 \) decreases slightly for increasing \( \mu \leq 1 \), and then increases and finally reaches a intermediate value featuring a multifractal behavior, as \( \mu \) increases across the transition point \( \mu_c = 1.5 \) for fixed \( V = 3 \). It is worth pointing out that the slope at the transition point \( \mu_c = 1.5 \) increases with system size, and will diverge in the thermodynamic limit, as a signature of the localized to critical transition.

Conclusions.—We implement a simulation of the GAAH model for a wide range of parameters. The capability of individual control and multi-qubit simultaneous readout of our superconducting processor allows for observation of multi-qubit spin transport and measurement of participation entropies in the experiments revealing clearly phase transitions of the model. Our experiment paves the way to quantum simulation of rich many-body phases and may trigger further studies about dynamics of many-body systems.

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Supplementary Materials:
Observation of critical phase transition in a generalized Aubry-André-Harper model on a
superconducting quantum processor

QUBIT INFORMATION

Our experiment is performed on the superconducting quantum processor with tunable couplers, which is identical to the one in reference [44]. This device consists of 10 transmon qubits ($Q_1 \sim Q_{10}$) and 9 transmon-type couplers ($C_1 \sim C_9$). In this experiment all qubits are initialized at their idle frequencies $\omega_{\text{idle}}^j/2\pi (j = 1, 2, \cdots, 10)$ that spread in the range from 4.280 GHz to 4.900 GHz. The idle frequencies are carefully arranged to reduce unwanted interaction and crosstalk errors among qubits (or between qubits and couplers) during single-qubit operations. All relevant information about qubit characteristics are listed in Table S1. The anharmonicity of all qubits are around 200 MHz. The readout pulse for all qubits are 1.0 $\mu$s in length, and the readout frequency and readout power are optimized for a high visibility. The readout fidelity, which is denoted by $F_0$ and $F_1$, are listed in Table S1.

Z PULSE CROSSTALK CORRECTION

The $Z$ control line crosstalks are estimated by measuring the linear response with respect to the bias voltages. To achieve this, we excite the target and measure how much $Z$ pulse amplitude (zpa) of the target needs to compensate for the crosstalk from the source. It is routine to measure crosstalk between qubits, but specific means are required to measure crosstalk between qubits and couplers. Following Ref. [44], we measure the coupler-to-qubit $Z$ crosstalk and further add the measurement of qubit- and coupler-to-coupler $Z$ crosstalks. Pulse sequence used for such measurement of crosstalk to couplers is plotted in Fig. S1(a). Due to the AC Stark effect, the response of target zpa to source zpa is nonlinear. In order to obtain the classical $Z$ crosstalk that removes the AC Stark effect, we fit the data in the linear region where the source qubit (coupler) is far from the target coupler, see Fig. S1(b). We then obtain the total crosstalk matrix, as shown in Fig. S2. In our device, all crosstalks between qubits are reduced due to its coupler architecture.

MEASUREMENT OF EFFECTIVE COUPLING STRENGTHS

The system can be fully described by the generalized Aubry-André-Harper (GAAH) model with the Hamiltonian:

$$\hat{H} = \sum_{j=1}^{9} (J_{j,j+1} \hat{a}_j^\dagger \hat{a}_{j+1} + \text{H.c.}) + \sum_{j=1}^{10} h_j \hat{a}_j^\dagger \hat{a}_j. \tag{S1}$$

Here $J_{j,j+1} = \lambda \left( 1 + \mu \cos \left[ 2\pi \left( j + \frac{1}{2} \right) \alpha + \delta \right] \right)$ and $h_j = \lambda V \cos (2\pi j \alpha + \delta)$ represent the effective coupling of qubits and the frequency detuning relative to resonant frequency (± 4.36 GHz). In this paper, we set $\lambda/2\pi = 4 \text{ MHz}$ and $\alpha = (\sqrt{5} - 1)/2$. $\mu$ and $V$ represent modulations amplitudes of off-diagonal hopping and on-site incommensurate potential, respectively. We manipulate $\mu$ and $V$ by applying the fast voltages to the corresponding $Z$ control lines of the coupler and qubit respectively.

In our processor of coupler architecture, the effective coupling of qubits can be adjusted by tuning couplers frequencies. Mathematically, the effective coupling strength can be described by [43, 44]:

$$J_{j,j+1} = J_{j,j+1}^0 + \frac{J_{j,j+1}^0 J_{j+1,j+2}^0}{\Delta_{j,j+1}}, \tag{S2}$$

where $J_{j,j+1}^0$ is the direct coupling between $Q_j$ and $Q_{j+1}$, $J_{j}^0$ is the direct coupling between $Q_j$ and $C_j$, and $1/\Delta_{j,j+1} = \ldots$
correct the zpas that bias qubits frequencies. To eliminate the
coupling strengths can be measured precisely through the joint
probability as a function of qubit-qubit swapping time \( t \) and
coupler zpa \[44\]. Fig. S3(b) is the normalized Fourier trans-
formation corresponding to Fig. S3(a). Fig. S3(b) shows the
advantages of adjusting the effective coupling strength using
the coupler architecture. Varying coupler zpa not only makes
decoupling point possible, but also greatly improve the upper
limit of coupling strength. In the experiment, we first calcu-
late the coupling strength distribution satisfying Eq. (2) ac-

TABLE S1. Qubit characteristics. \( \omega_{\text{idle}} \) is the idle frequency of qubit where decoherence parameters (including energy relaxation time \( T_1 \) and
Ramsey Gaussian dephasing time \( T_2^\text{R} \)) are measured. \( \omega_{\text{max}}/2\pi \) and \( \omega/2\pi \) denote to qubits’ sweetpoint frequency and resonant frequency,
respectively. \( F_0 \) (\( F_1 \)) is the measurement probability of \( |0\rangle \) (\( |1\rangle \)) when the qubit is prepared in \( |0\rangle \) (\( |1\rangle \)), which is used to mitigate the readout
errors.

| \( \omega_{\text{idle}}/2\pi \) (GHz) | \( \omega_{\text{max}}/2\pi \) (GHz) | \( T_1 \) (\( F_0 \)) | \( T_2^\text{R} \) (\( F_1 \)) |
|-----------------|-----------------|---------------|---------------|
| 4.900           | 5.491           | 14.3          | 0.9           |
| 4.365           | 5.201           | 14.8          | 1.1           |
| 4.930           | 5.347           | 33.0          | 1.1           |
| 4.310           | 5.36            | 17.9          | 1.2           |
| 4.878           | 5.226           | 30.7          | 1.2           |
| 4.455           | 5.313           | 28.1          | 1.1           |
| 4.800           | 5.318           | 26.3          | 1.2           |
| 4.280           | 5.5            | 21.5          | 1.2           |
| 4.770           | 5.108           | 24.4          | 1.3           |
| 4.388           | 5.248           | 32.9          |               |

\[
\left( 1/(\omega_{Q_j} - \omega_{C_j}) + 1/(\omega_{Q_{j+1}} - \omega_{C_j}) \right)/2 \]
is related to frequency detuning between the \( j \)-th coupler and its two nearest neighbor qubits. As shown in Fig. S3(a), effective coupling strengths can be measured precisely through the joint probability as a function of qubit-qubit swapping time \( t \) and coupler zpa \[44\]. Fig. S3(b) is the normalized Fourier transformation corresponding to Fig. S3(a). Fig. S3(b) shows the advantages of adjusting the effective coupling strength using the coupler architecture. Varying coupler zpa not only makes decoupling point possible, but also greatly improve the upper limit of coupling strength. In the experiment, we first calculate the coupling strength distribution satisfying Eq. (2) according to Eq. (S2) and estimate the zpa of each coupler, and then fine-tune the zpa by performing two-qubit swapping experiment until the coupling strength is within the tolerance (\( \leq 0.1 \text{MHz} \)).

After all the coupling strengths are calibrated, we need to correct the zpas that bias qubits frequencies. To eliminate the zpa deviation caused by the AC Stark effect, we perform two specific Rabi oscillation experiments on the target qubit whose zpa needs to be determined. In details, we consider two staggered distributions of non-target qubits frequencies as shown in Fig. S4. Assuming that the zpa of target qubit measured by the first Rabi experiment with one of staggered frequency distribution is zpa \(_{1}\), and the second Rabi experiment corresponding to the reverse-staggered frequency distribution measures another zpa \(_{2}\), we can estimate the zpa of target qubit that bias it to working point as \((zpa_1 + zpa_2)/2\). Thus, the zpa deviation caused by the AC Stark effect that depends on the frequency detuning can be cancelled approximately. We calibrate all the qubits zpas one by one in this way, and finally bias the frequencies of all the qubits to the specified distribution.

**EFFECTS OF DECOHERENCE**

In order to increase the adjustment range of coupling strength and reduce the distortion caused by high zpa, we choose a low resonant frequency (\( \approx 4.36 \text{GHz} \), deviates about 800 MHz from the sweet point) and decrease the idle frequencies \( \omega_{\text{idle}} \) accordingly. However, as a consequence, the decoherence of qubits is exacerbated by low frequency noise, leading to the decrease of dephasing time \( T_2^\text{R} \). Besides, the existence of energy relaxation time \( T_1 \) may also contribute to the dissipation. Such dissipation dynamics can be described by the Lindblad master equation

\[
\frac{d\hat{\rho}(t)}{dt} = -\frac{i}{\hbar} [\hat{H}, \hat{\rho}(t)] + \sum_{n=1}^{L} \left( \hat{K}_n \hat{\rho}(t) \hat{K}_n^{\dagger} - \frac{1}{2} \left( \hat{K}_n \hat{K}_n^{\dagger} \hat{\rho}(t) + \hat{\rho}(t) \hat{K}_n^{\dagger} \hat{K}_n \right) \right),
\]

(S3)

with \( \hat{\rho}(t) \) being the time-dependent density matrix of \( L \) qubits and \( \hat{K}_n \) (\( \hat{K}_n^{\dagger} \)) as Lindblad operators. The first term of Eq. (S3) represents the unitary evolutions of the system and the other terms imply the dissipation of the system due to interaction with the environment. Here we consider the dephasing and energy relaxation effects, and the corresponding Lindblad operators are written as \( \hat{K}_n = (1 - 2\hat{a}_n^{\dagger} \hat{a}_n)/\sqrt{2T_2} \) and \( \hat{K}_n^{\dagger} = \hat{a}_n/\sqrt{T_1} \), respectively.

The numerical results of the dynamics of participation entropies with decoherence effects, shown in Fig. 2–3 in the main text, and Fig. S5 in Supplementary Material, are obtained by solving Eq. (S3) with the averaged \( T_1 \sim 22.3 \mu s \) and \( T_2^\text{R} \sim 4.0 \mu s \). Here, the time-dependent participation en-

![FIG. S2. Z line crosstalk matrix. All the element values between qubit pairs or coupler pairs is at a low level (<2%) compared with high crosstalk between qubits and couplers.](image-url)
tropies can be calculated by the output density matrix \( \hat{\rho}(t) \), corresponding to \( p_i(t) = \langle i | \hat{\rho}(t) | i \rangle \), with \(| i \rangle \) being the computational basis in \( \mathcal{N} \)-dimension Hilbert space. Furthermore, post-selection is also employed to mitigate the effect of energy relaxation in the experimental data processing. Taking the above factors into account, the numerical results match with the experimental data quite well, showing the characteristics of phase transition despite the inevitable decoherence.

### FIRST-ORDER PARTICIPATION ENTROPY

In the limit \( q \to 1 \), we can obtain the dynamical first-order participation entropy:

\[
S_{1PE}^{PE}(t) = \lim_{q \to 1} \frac{1}{1-q} \log \sum_{i} p_i(t)^q - \frac{N}{\sum_{i} p_i(t)} \log p_i(t).
\]  

(S4)

The dynamical behavior of first-order participation entropy is very similar to the second-order participation entropy we discussed above, but with a different \( q \)-dependent fractal dimension \( D_q \) in terms of scaling behavior. Here, we also present the dynamics of the first-order participation entropy in Fig. S5, and the averaged late-time first-order participation entropy for different parameters \( \mu \) and \( V \) in Fig. S6. Comparing with Fig. S5 and Fig. 3 in the main text, we can find that the growth trend over time of first-order participation entropy is the same as that of the second-order, but with slightly higher values than those of the second-order. Thus, the averaged late-time first-order participation entropy can also be used to characterize phase transitions as the second-order, as shown in Fig. S6.

### SCALING BEHAVIOR OF PARTICIPATION ENTROPY

For many-body states, the participation entropy \( S_{q}^{PE} \) can characterize the localization in the Hilbert space. In the thermodynamic limit \( \mathcal{N} \to \infty \), \( S_{q}^{PE} \) is (i) \( \log N \) for a perfectly delocalized state, (ii) a const for a localized state, (iii) \( D_q \log N \) (where \( D_q < 1 \)) for a state with a fractal dimension \( D_q \), which occurs at localization transition or in the critical phase [36, 49].

To see this, we display the scaling behavior of the averaged late-time participation entropy \( \overline{S_{q}^{PE}} \) in Fig. S7. Since the numerical simulation match with the experimental data quite well, here we use numerical results of larger system sizes to fit with the form \( \overline{S_{q}^{PE}} = a_2 \log(\mathcal{N}) + b_2 \). We choose three pairs of parameters \( (\mu = 0.5, V = 1.0), (\mu = 2.0, V = 1.0) \) and \( (\mu = 1.0, V = 3.0) \) for the extended, critical, and localized phases, respectively. In the extended phase, \( a_2 \approx 0.928 \), which is very close to one for a perfectly delocalized state. In the critical phase, multifractal behavior of many-body states is revealed by the fractal dimension \( a_2 \approx 0.502 \). Besides, in the localized phase, \( a_2 \approx 0.243 \) is much smaller than those in the extended and critical phases.
FIG. S5. (a) The dynamics of first-order participation entropy for the system quenched into the three phases. The yellow, red, and purple points are experimental data for the extended ($\mu = 0.5, V = 1.0$), critical ($\mu = 2.0, V = 1.0$), and localized ($\mu = 0.5, V = 4.0$) phases, respectively. Lines with the same color are numerical simulation under the same parameters with decoherence taken into account. (b) The dynamics of first-order participation entropy with fixed $\mu = 0.5$ and $V = 1, 2$ and 4 (extended to localized transition). (c) The dynamics of first-order participation entropy with fixed $V = 1$ and varying $\mu = 0.5, 1$ and 2 (extended to critical transition).

FIG. S6. Comparisons between experimental data and numerical simulation (a) with fixed $\mu = 0.5$ and varying $V$ from extended to localized transition, (b) with fixed $V = 1$ and varying $\mu$ from extended to critical transition, and (c) with fixed $V = 3$ and varying $\mu$ from localized to critical transition. Points with statistical error bars are experimental data, and solid lines are numerical simulation using the Hamiltonian 2 in the main text.
FIG. S7. The averaged late-time participation entropy as a function of \( \log N \). We choose \( \mu = 0.5, V = 1.0 \) for the extended phase, \( \mu = 2.0, V = 1.0 \) for the critical phase, and \( \mu = 1.0, V = 3.0 \) for the localized phase. The results of the fit are drawn with dashed lines.