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## Out of Equilibrium Majoranas in Interacting Kitaev Chains

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We employ a time-dependent real-space local density-of-states method to study the movement and fusion of Majorana zero modes in the 1D interacting Kitaev model, based on the time evolution of many-body states. We analyze the dynamics and both fusion channels of Majoranas using time-dependent potentials, either creating *Walls* or *Wells*. For fast moving Majoranas, we unveil non-equilibrium signatures of the "strong-zero mode" operator (quasi parity degeneracy in the full spectrum) and its breakdown in the presence of repulsive Coulomb interactions. Focusing on forming a full electron after fusion, we also discuss upper and lower limits on the Majorana speed needed to reduce non-adiabatic effects and to avoid poisoning due to decoherence.

Introduction. Majorana zero modes (MZMs) generate considerable interest because of potential applications in quantum information and computation [1-3]. MZMs obey non-Abelian exchange statistics. Because they are topologically protected from local perturbations and disorder, they are of value as possible qubits [4-6]. Signatures of MZMs are expected to develop in tunneling conductance experiments as zero bias peaks [7-9]. The simplest setup to realize Majoranas are quantum wires, where MZMs develop at the two edges [1, 14]. For ferromagnetic atomic chains with strong spin-orbit coupling placed over a superconductor, MZMs were indeed reported at the edges in spatially and spectral-resolved scanning tunneling experiments [10, 11]. In nanowires, most theoretical work neglect repulsion among particles. However, Coulomb repulsion plays an important role in 1D MZMs because it suppresses the pairing-induced bulk gap and can destroy topological protection [12, 13].

The movement of Majoranas and detection of fusion channels are important for quantum-information processing [14, 15]. MZMs behave as Ising non-Abelian anyons [4, 16] and obey the fusion rule [15]:  $\gamma \times \gamma = I + \psi$ , meaning two MZMs can fuse into the vacuum I or into an electron  $\psi$ . The fusion process requires a slow adiabatic movement of Majoranas, achieved by applying properly adjusted time-dependent local gates to the topological superconducting wire [14]. The rapid progress in quantum-wires with tunable local gates [14, 15, 17] provides a promising platform for creation, movement, and fusion of Majoranas [18].

Motivated by experimental progress in nanowires [19], here for the first time we employ a computationallyintensive time-dependent real-space local density-ofstates  $LDOS(\omega, j, t)$  method to observe the movement and fusion of Majoranas in the interacting Kitaev model. The uniqueness of our effort is that we can study Majorana movement at *any* speed by properly choosing the time dependence of gate voltages, namely we can access the non-equilibrium situation away from adiabaticity, difficult to reach by theoretical tools.  $LDOS(\omega, j, t)$ of moving Majoranas and fusion outcome can be measured in tunneling spectroscopy experiments based on gate-control nanowire devices [19, 20]. Compared to previous studies based on single particle states, here we use the exact-diagonalization method for the time evolution of the many-body states of *interacting* electrons in the 1D Kitaev model up to 16 sites. We address the outof-equilibrium properties and fusion rules of MZMs, via sequential application of time-dependent chemical potential gates. For fast moving non-interacting MZMs, using the time-dependent LDOS, we find the signature of a "strong-zero mode" operator in  $LDOS(\omega, j, t)$  [21, 22]. Remarkably, we found the total spectral weight at  $\omega = 0$ remains conserved (almost identical to the case of slow moving Majoranas). However, with interaction V, depending on its strength and switching time  $\tau$ , we find loss in spectral weight at  $\omega = 0$  and breakdown of the strong-zero mode properties. Furthermore, we provide the time scale to observe the fusion rules of interacting Majoranas. Although it is widely expected that for "adiabatic" movement of MZMs their fusion will lead to the formation of a fermion or vacuum states, only via calculations as presented here, that allows for any speed for the MZMs, is that we can establish how "slow" the movement must truly be in practice.

*Model and Method.* We consider the time-dependent interacting 1D Kitaev model for spinless fermions with open boundary conditions to anchor MZMs:

$$H(t) = -t_h \sum_{i=1}^{N-1} \left( c_i^{\dagger} c_{i+1} + H.c. \right) + V \sum_{i=1}^{N-1} \left( n_i n_{i+1} \right) + \Delta \sum_{i=1}^{N-1} \left( c_i c_{i+1} + H.c. \right) + \sum_i^N \mu_i(t) n_i, \qquad (1)$$

where  $n_i = c_i^{\dagger} c_i$  and  $c_i^{\dagger} (c_i)$  is the fermionic creation (annihilation) operator,  $t_h$  is the hopping amplitude, and  $\Delta$  is the *p*-wave pairing strength. The time dependent Hamiltonian H(t) (Eq.1) commutes with the parity operator  $P = e^{i\pi \sum_j n_j}$  [23, 24]. The time dependence is incorporated in the chemical potential  $\mu_i(t)$  as:

$$\mu_i(t) = 0 \ (i < i_0), \qquad \mu_i(t) = \mu \ (i > i_0),$$



FIG. 1. Schematic representation of the transfer of the right edge  $MZM^R$  to site  $i_0 - 1$  in a 1D Kitaev chain. The onsite chemical potential at any site can be tuned using timedependent local gates with quench rate  $1/\tau$ . Large (small)  $\tau$ corresponds to slow (fast) motion of  $MZM^R$ . Blue squares denote the topological region (with  $\mu_{i < i_o} = 0$ ) while red squares denotes the non-topological region (with  $\mu_{i > i_o} = \mu$ ) for this *Wall* case. For the *Well*, the red region has a negative  $\mu$ .

$$\mu_i(t) = \mu \frac{n\Delta t}{\tau} \ (i = i_0), \tag{2}$$

where  $1/\tau$  is the quenched rate,  $\Delta t = 0.001$  is the small time step we used, and n is the integer number of those steps, such that the on-site chemical potential  $\mu_i(t)$  at  $i = i_0$  increases approximately linearly from 0 to  $\mu$  in a time  $\tau$ (defined as the switching time of the local gate at site  $i = i_0$ ). The sequential application of onsite gates  $\mu_i(t)$  on the right half of the 1D chain, creates a moving *Wall* for  $\mu > 0$  (or moving *Well* for  $\mu < 0$ ), separating topological from non-topological regions at site  $i = i_0$ . Equating our number of sites with number of gates in a coarse-grained approach, this process leads to the movement of the right edge Majorana zero mode (MZM<sup>R</sup>) from the edge i = Nto site  $i_0 - 1$  in a finite time  $t = N_R \tau$ , with  $N_R$  the number of sites where the chemical potential reached its maximum value (here being  $|\mu| = 12$ ) at time t (Fig. 1).

To calculate the time-dependent local density-of-states (at zero temperature), we first time evolve the groundstate wave function  $|\psi(0)\rangle$  up to time  $t = N_R \tau$ , using the time-dependent Hamiltonian H(t) as:  $|\Psi(t)\rangle =$  $\mathcal{T}exp\Big(-i\int_0^t H(s)ds\Big)|\psi(0)\rangle$ , where  $\mathcal{T}$  is the time ordering operator [25]. Then, we calculate the double-time Green function G(t,t') [26], using the time-independent Hamiltonian  $H_f = H(t = t_f)$  at time  $t_f = N_R \tau$ :

$$G_j^{elec}(t,t') = \langle \Psi(t) | c_j^{\dagger} e^{iH_f t'} c_j e^{-iH_f t'} | \Psi(t) \rangle.$$
 (3)

The time-dependent  $LDOS_{elec}(\omega, j, t)$  for electrons is the Fourier transform with respect to t' of the local-Green function at site j:  $LDOS_{elec}(\omega, j, t) =$ 

$$= \frac{1}{\pi} Im \int_0^T dt' e^{i(\omega+i\eta)t'} iG_j^{elec}(t,t').$$
(4)

where we use T = 70 for the integration, and broadening  $\eta = 0.1$ . Similarly we obtained the  $LDOS_{hole}(\omega, j, t)$ for holes, using the Fourier transform of the Green function  $G_j^{hole}(t, t') = \langle \Psi(t) | c_j(t') c_j^{\dagger} | \Psi(t) \rangle$ . The total local density-of-states at site j is, thus,  $LDOS(\omega, j, t) =$  $LDOS_{hole}(\omega, j, t) + LDOS_{elec}(\omega, j, t)$ .



FIG. 2. Slow movement of Majoranas. (a) Local density-ofstates  $LDOS(j, \omega, t)$  vs. site j, at time t = 0 and for  $\omega = 0$ . The sharp peaks at sites j = 1 and j = 12 in  $LDOS(j, \omega = 0, t)$  represent Majorana edge modes for different values of V and  $\mu_j = 0$  (for 1 < j < 12). Inset:  $LDOS(\omega)$  vs.  $\omega$ at site j = 12 using broadening  $\eta = 0.1$ . Site-dependent  $LDOS(j, \omega, t)$  at  $\omega = 0$  and time  $t/\tau = 6$  for V = 0, 0.5, and 1.0 with  $\tau = 36$ , 60, and 72, respectively, for: (b) Positive  $\mu$  (*Wall*). Inset shows the site-dependent density  $\langle n(j, t) \rangle$  at  $t/\tau = 6$  for  $\mu_R = 12$ . (c) Negative  $\mu$  (*Well*). Inset shows the site-dependent density  $\langle n(j, t) \rangle$  at time  $t/\tau = 6$  for  $\mu_R = -12$ . L = 12 sites and  $t_h = \Delta = 1.0$  were used.

Slow movement of Majoranas. For fusion or braiding of MZMs, it is required to transfer the Majoranas slowly, close to the adiabatic limit [27, 28]. Figure 2(a) shows the real-space local density-of-states  $LDOS(\omega = 0, j, t = 0)$ vs. site j, with  $\mu_i = 0$  (for all sites), and at  $t_h = \Delta = 1$ . For small or zero V, these peaks are sharply localized at the end sites (i = 1 and 12), whereas for robust V the  $\omega = 0$  peaks are slightly delocalized over a few sites. In the inset, we show  $LDOS(\omega)$  at time t = 0 at the end site j = 12 and several V's. We find a sharp peak at  $\omega = 0$ , signaling a MZM mode at the end site. Integrating in  $\omega$  the LDOS( $\omega$ , j = 12) at V = 0.0 gives spectral weight 0.48, close to the analytically expected value 0.5 [29].

Next, with sequential application of the timedependent chemical potential  $\mu_j(t)$ , the right edge MZM<sup>R</sup> (at site j = 12) is moved to the middle site (j = 6) in a time  $t = N_R \tau$  (i.e.  $t/\tau = N_R = 6$  because we travel 6 sites). We study cases  $\tau = 36, 60$ , and 72, for interaction



FIG. 3. Slow Majorana fusion using  $\mu(t) > 0$  (*Wall*),  $\Delta = 1$ , at V = 0.0, 0.5, and 1.0, with  $\tau = 36, 60$ , and 72, respectively. Upper panels show Majoranas fusion for the initial state with parity P = -1. (a) Electron  $LDOS_{elec}(\omega)$  after moving a MZM from j = 12 to j = 1. (b) Hole  $LDOS_{hole}(\omega)$  at  $t/\tau = 11$  and site j = 1. (c) Charge density  $\langle n(j = 1, t) \rangle$  vs. time t, varying V. Lower panels show Majoranas fusion for the initial state with parity P = +1. (d) Electron  $LDOS_{elec}(\omega)$  at  $t/\tau = 11$  and site j = 1, (e) Hole  $LDOS_{hole}(\omega)$  after moving a MZM from j = 12 to j = 1. (f) Charge density  $\langle n(j = 1, t) \rangle$  vs. time t.

strengths V = 0.0, 0.5, and 1.0, respectively. In this case, $|\psi(t)\rangle$  remains close to the degenerate ground-state space (larger values of V require a slower-rate of increase in the onsite  $\mu_{i_0}(t)$ ). As shown in Fig. 2(b), for  $\mu = 12$  (i.e. when creating a potential *Wall*), the  $LDOS(\omega = 0, j, t)$ has peaks at sites j = 1 and j = 6 at time  $t/\tau = 6$ , indicating that a slow transfer of MZM<sup>R</sup> from j = 12 to j = 6 occurred. The average density  $\langle n(j,t) \rangle$  is close to zero for  $j \ge 7$ , while it is close to 0.5 for  $j \le 6$  (inset of Fig. 2(b)). Interestingly, at  $\mu_R = -12$  (when creating a potential *Well*), the effect of interaction increases. In the non-topological region  $(j \ge 7)$ , each site is occupied by one fermion, whereas in the topological region ( $j \leq$ 6) the mean occupancy is close to 0.5. With nonzero V, to minimize the Coulomb interaction between fermions at the topological to non-topological boundary, the fermions near the boundary become inhomogeneously distributed (Fig. 2(c), inset). This delocalizes  $MZM^R$  over more sites as V increases (Fig. 2(c)).

Slow fusion of Majoranas. For the fusion of Majoranas, we move the right edge MZM<sup>R</sup> slowly all the way to the left end (site j = 1) using sequential operations of  $\mu(t)$  in a time interval  $t = 11\tau$  (see caption Fig. 3). At t = 0, for V = 0,  $t_h = \Delta = 1$ , with  $\mu_i = 0$  (for all sites), the system has degenerate many-body ground states ( $|\psi_1\rangle$  and  $|\psi_2\rangle$ ). These degenerate ground states have different fermionic parity  $P = \pm 1$ . At t = 0, we



FIG. 4. Fast movement of Majoranas using a small quench rate  $\tau = 1$ ,  $\mu(t) > 0$  (*Wall*), and for the initial state with parity P = -1. (a) Site-dependent  $LDOS(j, \omega, t)$  at  $\omega = 0$ , for V = 0.0, 1.0, and 2.0, after moving the right MZM from site 12 to 8. Electron  $LDOS_{elec}(\omega)$  and Hole  $LDOS_{elec}(\omega)$ at site j = 8, for (b) V = 0.0, (c) V = 1.0, and (d) V = 2.0.

start the time evolution with those initial states  $|\psi_s\rangle$ (s = 1 or 2) up to  $t/\tau = 11$ , to confirm both fusion channels (*Electron* :  $\Psi$  and *Vacuum* : *I*). For positive chemical potential,  $\mu(t) > 0$  (*Wall*) and the initial states  $|\psi_s\rangle$  with parity P = -1, the electron  $LDOS_{elec}(\omega)$  at j = 1 shows a sharp peak close to  $\omega = 0$ , for V = 0, 0.5, and 1 (Fig. 3(a)). Meanwhile, the hole  $LDOS_{hole}(\omega)$  at j = 1, displays no peak (Fig. 3(b)). The time-dependent density  $\langle n(j = 1, t) \rangle$  at site j = 1 takes values one at  $t/\tau = 11$ , giving a clear indication of the formation of a single electron (spinless) at site j = 1 after Majoranas fusion (Fig. 3(c)). On the other hand, for the initial state  $|\psi_s\rangle$  with parity P = +1, the hole  $LDOS_{hole}(\omega)$ displays a sharp peak close to  $\omega = 0$ , for V = 0, 0.5, and 1 (Fig. 3(e)). The electron  $LDOS_{elec}(\omega)$  has no peak at  $t/\tau = 11$ , see Fig. 3(d). The density  $\langle n(j=1,t) \rangle$  at j=1approaches zero (Fig. 3(f)), confirming a vacuum state at  $t/\tau = 11$ . Because the MZM spreads over more than one site as V grows, density fluctuations occur at site j = 1as compared to V = 0 (Figs. 3(c,f)) (for the slow fusion of Majoranas, in the presence of a time-dependent potential Well see [40]).

Fast movement of Majoranas. Changing  $\mu(t)$  employing a faster rate (smaller  $\tau$ ), leads to a fast movement of MZMs<sup>R</sup> generating non-adiabatic effects [30, 31]. The faster change in  $\mu(t)$  results in a finite overlap of the time-evolving wave-function  $|\psi(t)\rangle$  with excited states of the instantaneous Hamiltonian H(t). Starting the time evolution with initial states  $|\psi_s\rangle$ , with parity P = -1 for  $\mu(t) > 0$  (Wall) and using all eigenvectors  $\{|n\rangle\}$  of the instantaneous Hamiltonian H(t) the electron LDOS at finite time  $t = N_R \tau$  can be written as:  $LDOS_{elec}(\omega, t) =$ 

$$\frac{-1}{\pi} Im\left(\sum_{m,n} \frac{\langle \Psi(t)|c_j^+|n\rangle\langle n|c_j|m\rangle\langle m|\Psi(t)\rangle}{e_n - e_m + \omega + i\eta}\right), \quad (5)$$

where  $\eta = 0.1$ , and the rest of the notation is standard (as reference for L = 12 the number of states is 4,096).

Figure 4(a) shows  $LDOS(\omega, j, t)$  after moving the MZM from j = 12 to j = 8 for different values of Coulomb interaction V and with fast quench rate  $\tau = 1$ . For V = 0, and for this faster change in  $\mu(t)$  the peak values of  $LDOS(\omega, j, t)$  almost remain the same as compared to slower changes in  $\mu(t)$  (Fig. 2(b)). On the other hand for the same faster moving MZMs but for a finite repulsion V, there is a significant reduction in the peaks magnitude of  $LDOS(\omega, j, t)$  (at sites j = 1 and j = 8) with increasing V. For larger values of V, the sitedependent  $LDOS(\omega, j, t)$  indicate finite overlap between left and right MZMs (Fig. 4(a)). Figures 4(b,c,d) contains the electron  $LDOS_{elec}(\omega)$  and hole  $LDOS_{hole}(\omega)$ vs.  $\omega$  at site j = 8, for different values of the Coulomb interaction V at time  $t/\tau = 4$ . Using full-diagonalization of H(t), we find for V = 0 and  $\Delta = 0.9$  [32] that all many-body eigenstates of H(t) come in pairs with opposite parity  $P = \pm 1$  and the states of each pair are almost degenerate (the parity degeneracy of all manybody states is compatible with a strong zero mode operator [21, 22]). The equal peak heights of  $LDOS_{elec}(\omega, t)$ and  $LDOS_{hole}(\omega, t)$  at  $\omega = 0$  in Fig. 4(b) (with spectral weight 0.5), can be associated with the presence of such strong-zero mode operator when in non-equilibrium. The spectral weight is dominated by only a few higher energy degenerate-pair states, all with comparable weight, contributing to  $\omega = e_n - e_m = 0$  (with |m - n| = 1) in the  $LDOS_{elec}(\omega, t)$  (Eq. 6) and  $LDOS_{hole}(\omega, t)$ .

This strong zero mode operator is immune to decoherence [33], potentially leading to topological qubits with infinite coherence time [34] This occurs because they are topologically protected due to global parity conservation and quasi-degenerate paired states in the full spectrum (involving opposite fermion parity) [21, 35]. This protection survives as long as the left and right MZMs do not overlap with each other. However, increasing V, the peaks of  $LDOS_{elec}(\omega, t)$  and  $LDOS_{hole}(\omega, t)$  start splitting and the peak values are no longer equal in magnitude (Fig. 4(c,d)). Furthermore, at large V the electron and hole parts of LDOS show peaks away from zero and these peaks are largely splitted. The split in  $LDOS_{elec}(\omega, t)$ and  $LDOS_{hole}(\omega, t)$  peaks is due to the breakdown of degeneracy of higher excited paired states increasing V.

Fast Fusion of Majoranas. In real Majorana nanowire setups, it is necessary to move the Majoranas with sufficient speed to be faster than the quasi-particle "poisoning" time [15, 36–38]. Here, we present the minimum required switching time of local gates for fast moving  $MZM^R$ , so that we obtain a full electron after the fusion of left and right MZMs. To fuse Majoranas, starting with the initial state having parity P = +1, we moved at various speeds the right MZM<sup>*R*</sup> all the way to the left end (site j = 1).



FIG. 5. Fast Majorana fusion using  $\mu(t) < 0$  (*Well*) at various  $\tau$ 's, for the initial state with P = +1. The right MZM is moved from site j = 16 to 1. (a)  $LDOS_{elec}(\omega)$  at site j = 1 and (b)  $LDOS_{hole}(\omega)$  at j = 1, for V = 1,  $\Delta = 0.6$ , and L = 16.

In Figs. 5(a,b), we show the fusion of Majoranas using a potential Well ( $\mu(t) < 0$ ) with  $\Delta = 0.6$ , V = 1, and for the system length L = 16. For these parameters, we believe our system is closer to realistic setups [39] for experiments [10, 11]. However, for negative chemical potential, after the formation of an electron at site j = 1 the repulsive nearest-neighbor V leads to the split in ground state energy (approximately of order V [40]), causing an energy shift in peak values of  $LDOS_{elec}(\omega)$  and  $LDOS_{hole}(\omega)$  increasing  $\tau$ . Also, increasing  $\tau$  (slower-motion of MZM<sup>R</sup>), the peak at  $\omega = V$  for  $LDOS_{elec}(\omega)$  starts increasing, while the  $LDOS_{hole}(\omega)$  peak value at  $\omega = V$  decreases. At  $\tau = 48$ , we obtain a sharp electron peak close to  $\omega = 1.0$ , whereas the  $LDOS_{hole}(\omega)$  peak vanishes to zero as  $\tau$  grows (Figs. 5(a,b)). Specifically, this shows the formation of a full electron at  $\tau = 48$  for  $\Delta = 0.6$ , V = 1, and L = 16.

In terms of SI units the switching time per gate, for V = 1 corresponds to  $\tau \hbar/\Delta \sim 0.17$ ns to 5.2ns (using  $\tau = 48$ , and  $\Delta = 180\mu$ eV or  $\Delta = 6\mu$ eV as in previous literature [18, 36]). Independently, the quasiparticle "poisoning" time in nano-wire systems has been estimated in a broad range 10ns to 10ms [36, 37]. Because in the worse case of 5.2ns, eight gates require a total time 41.6ns to move adiabatically the Majorana, and since this number is close to the poisoning time, we conclude that there should be a time range where moving Majoranas in chains can occur adiabatically before poisoning occurs for V = 1. As V increases, the situation deteriorates because we must use larger  $\tau$  to form a full electron after fusion (see for larger V in [40]).

*Conclusions.* We performed real-time dynamics and fusion of Majoranas in the interacting 1D Kitaev model using sequential application of time-dependent chemical potentials (gates). We show that the movement and fusion outcomes can be monitored using the time-

dependent local density-of-states, and should be observed in tunneling spectroscopy experiments [19]. We find that for non-interacting and fast moving Majoranas, the near degeneracy of MZMs exists even in many higher energy states and MZMs remain topologically protected. However, for the interacting case, with increasing V we find a decrease in spectral weight at  $\omega = 0$  in the timedependent local density-of-states Furthermore, we estimate the minimum required switching time of local gates to form a full electron after the fusion. Due to advancements in fabricating Majorana nanowires [19, 41] with long quasiparticle poisoning time [36, 37], and considering our estimations for the times needed for adiabatic movement, we believe proper Majorana movement could be realized in realistic gate-control nanowire devices [19, 20, 42].

Note Added. After completion of this work, we became aware of an experimental realization of the twosites Kitaev chain employing two quantum dots coupled through a short superconducting semiconducting hybrid (InSb nanowires) [43]. In this experiment, they were able to tune the hoping  $(t_h)$  and pairing term  $\Delta$  close to  $t_h = \Delta$ . Interestingly, using tunneling spectroscopy measurements they showed the signature of two localized Majorana modes on each quantum dot close to  $t_h = \Delta$ . This experimental realization of a minimal Kitaev chain opens the possibility of an experimental confirmation of our predictions about Majorana fusion (at  $t_h \sim \Delta$ ) using arrays of quantum dots.

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