



PAPER

Engineering non-equilibrium quantum phase transitions via causally gapped Hamiltonians

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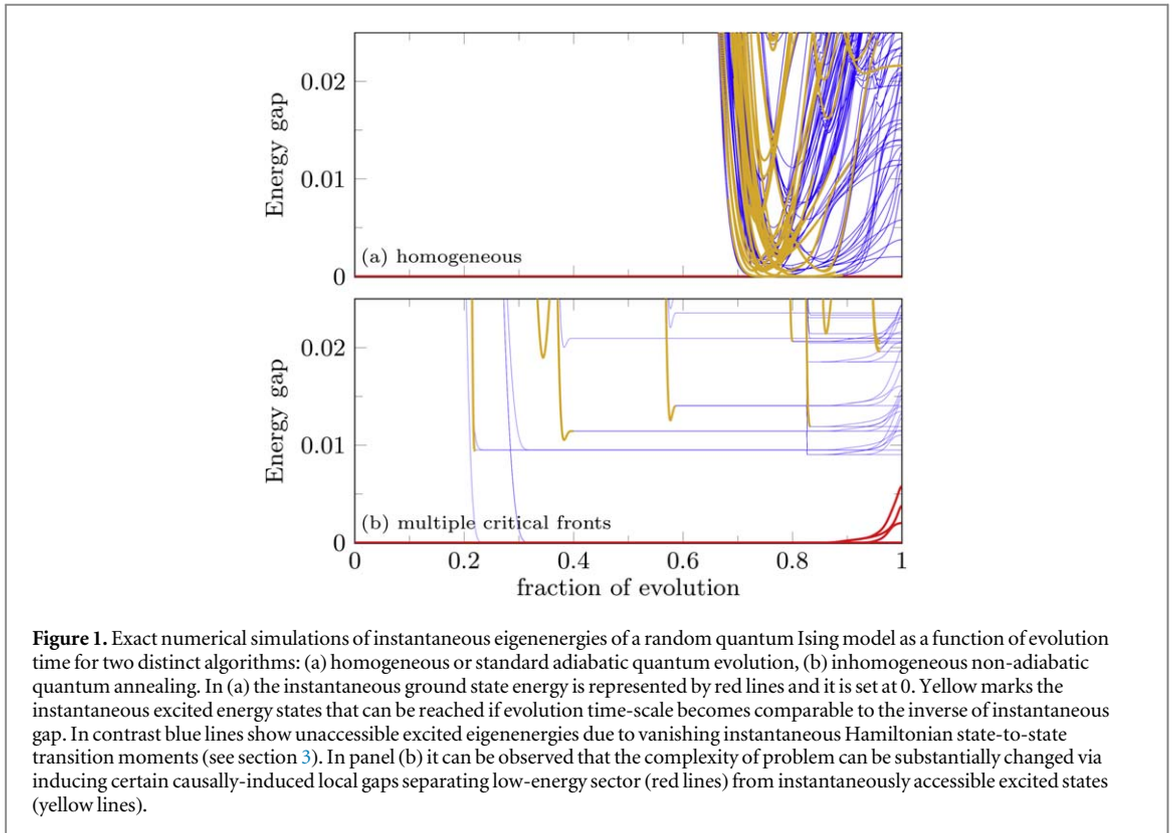
Masoud Mohseni¹, Johan Strumpfer² and Marek M Rams³ ¹ Google Quantum Artificial Intelligence Lab, Venice, CA 90291, United States of America² Google, San Francisco, CA 94109, United States of America³ Institute of Physics, Jagiellonian University, Łojasiewicza 11, 30-348 Kraków, PolandE-mail: mohseni@google.com**Keywords:** quantum phase transition, disordered spin chains, Kibble–Zurek mechanism, adiabatic protocol, random Ising model

Abstract

We introduce a phenomenological theory for many-body control of critical phenomena by engineering causally-induced gaps for quantum Hamiltonian systems. The core mechanisms are controlling information flow within and/or between clusters that are created near a quantum critical point. To this end, we construct inhomogeneous quantum phase transitions via designing spatiotemporal quantum fluctuations. We show how non-equilibrium evolution of disordered quantum systems can create new effective correlation length scales and effective dynamical critical exponents. In particular, we construct a class of causally-induced non-adiabatic quantum annealing transitions for strongly disordered quantum Ising chains leading to exponential suppression of topological defects beyond standard Kibble–Zurek predictions. Using exact numerical techniques for 1D quantum Hamiltonian systems, we demonstrate that our approach exponentially outperforms adiabatic quantum computing. Using strong-disorder renormalization group (SDRG), we demonstrate the universality of inhomogeneous quantum critical dynamics and exhibit the reconstructions of causal zones during SDRG flow. We derive a scaling relation for minimal causal gaps showing they narrow more slowly than any polynomial with increasing size of system, in contrast to stretched exponential scaling in standard adiabatic evolution. Furthermore, we demonstrate similar scaling behavior for random cluster-Ising Hamiltonians with higher order interactions.

Controlling non-equilibrium dynamics of quantum many-body systems is one of the main challenges in condensed matter physics and quantum control. Such complex quantum systems have very rich parameter space and unusual dynamical properties that makes them very hard to simulate and control as they are driven through critical regions [1]. The main difficulties arise from the fact that these systems generally contain high degree of disorders and effectively low dimensions such that they are not prone to exact analytical treatment or mean-field approximations. In principle their dynamics can be mapped to the dynamics of spin-glass systems that are driven/quenched by external control fields and could experience various first and second-order quantum phase transitions and Griffiths singularities [2, 3]. Quantum dynamics of such complex systems, except trivial cases, would be out-of-equilibrium when they are quenched in any finite time. However, such rich dynamical properties could lead to novel computational resources [4–7] provided that we obtain sufficient degree of control over their dynamics.

Adiabatic quantum computation (AQC) has been developed as a particular paradigm that utilize the continuous-time dynamics of driven many-body quantum systems for solving optimization tasks [4, 5]. In this model, the solution of a hard combinatorial optimization problem is encoded in the ground state of an interacting many-body system which can be prepared adiabatically from an initially trivial ground state, provided that time evolution is much longer than the inverse of minimum gap square [5]. One of the major challenges to AQC, that has been largely ignored in the quantum computing literature, is that for many realistic problems the analog quantum annealer will inevitably contain a significant amount of quenched disorder



smearing the corresponding quantum phase transitions for pure systems. Thus, the required time-scale for satisfying adiabatic limit could grow as a stretch exponential due to Griffiths singularity [3], even in the absence of any first order phase transitions. The Griffiths effects have pronounced consequences for finite-dimensional quantum systems, much stronger than in the classical counterparts. In fact, near-term quantum processors are best examples of low-dimensional quantum systems due to the inherent locality of physical interactions and geometrical constraints on the degree of connectivity [8]. After the embedding of a computational problem into quantum annealers, or their digital simulations [9], they will inevitably react to quantum fluctuations inhomogeneously at the physical level. Consequently, near-term quantum processors will typically experience locally inhomogeneous and smeared first and second order phase transitions, even if we drive them with an external field which is homogeneous in space. In particular, annealing schedules exhibit multiple vanishing gaps between ground state and first excited state, see figure 1(a), leading to exponentially long annealing time-scales. In practice, we always have a finite annealing time-scale that would inherently violate the adiabaticity condition, even for finite-size systems, leading to emergence of domain walls or topological defects that emerge at a relatively wide effective quantum critical region. This is in sharp contrast to a single, well-defined quantum critical point for pure system, where their density of defects can be estimated via Kibble–Zurek mechanism (KZM) in the thermodynamics limit [10–14]. As of today, there is no known way to guarantee the quality of solutions, given finite space–time physical resources, and there is no constructive or algorithmic way to improve performance for such analog quantum information processors within a given accuracy. These issues have lead us to the following fundamental questions: Is it possible to engineer quantum phase transition in disordered systems by inhomogeneous control fields to enforce spatially-induced gaps between low energy sector and higher energy states (see figure 1(b)).

Here, we present a general approach for controlling quantum critical dynamics. We introduce different classes of spatial and/or temporal inhomogeneous protocols to drive strongly disordered quantum spin chains across a quantum phase transition and minimize the residual energy of the final state. This is achieved by creating governing Hamiltonian with multiple critical fronts that can synchronize the local phase transitions in space and time. In each local region, the number of spins that simultaneously experience the critical dynamics is controlled by the length scale and shape of the inhomogeneity in which the magnetic field is modulated. Causality is introduced as the main control strategy to spatially coordinate symmetry breaking events among neighboring regions by finding the appropriate degree of inhomogeneities and the speed of critical fronts to reduce the number of topological defects. We explore the conditions for an optimal suppression of domain walls and show that we can beat the standard homogeneous KZM prediction for the density of the topological defects for strongly disordered transverse Ising problem in 1D. Moreover, we show that these phenomena can similarly

be observed for systems with k -local physical interactions. We demonstrate that inhomogeneous driving can be exponentially faster for such systems than conventional (homogenous) schemes such as adiabatic quantum annealing. Furthermore, we show that the universality of quantum critical phenomena holds for inhomogeneous quantum critical dynamics even in the presence of strong disorder.

The outline of this paper is as follows: in section 1, we review causal origins of topological defects in the context of KZM for pure and disorder systems. In section 2, we describe two general classes of inhomogeneous quantum annealing (IQA), type I and type II, for re-constructing phase transitions and present numerical results for strong-disorder 1D transverse Ising model. In this context, we show that AQC can be understood as a trivial form of either type I or type II IQA. In section 3, we first provide a phenomenological theory of the emerging local gaps and its connection to threshold velocities for critical fronts. We then derive an expression for distribution of local gaps as a function of inhomogeneity slope with a logarithmic correction on the system size. We demonstrate universality of critical fronts shapes via strong-disorder renormalization group (SDRG) techniques. We also discuss how the shape of inhomogeneity is related to its penetration depth into disordered phase. A generalization of our work for k -local Hamiltonian system is presented in section 4. A detailed treatment of our work as a generalization of KZM and discussions on lower- and upper-bounds for the shape of critical fronts is provided in a separate manuscript [15]. The generalization to spin-glass systems will be presented in another subsequent work [16, 17].

1. Causal origin of topological defects

We start by reviewing the KZM for pure systems (in absence of any disorders) which has been developed as the phenomenological theory to describe the breakdown of adiabaticity in critical systems [10–14]. The theory provides a rough estimate for the density of topological defects that arise when a quantum or classical many-body system is driven through a continuous critical point at a finite rate. The key observation is that in the vicinity of a quantum critical point a system at the thermodynamical limit effectively stops following the adiabatic evolution for any finite quench rate—no matter how slow it is driven. This results in emergence of universal KZ length scale which depends on the quench rate and manifests itself, among others, in the density of topological excitations.

The time dependent evolution of system can be expressed by a Hamiltonian as: $H(g) = gH_c + H_p$ where H_c is controllable Hamiltonian, $g(t)$ is a control parameter with value g_c at the critical point, and H_p is the Hamiltonian of interest or the ‘problem Hamiltonian’. Near a critical point the characteristic energy scale of the system behaves as $\Delta \sim 1/|\varepsilon|^{z\nu}$ at the thermodynamical limit. The system experiences a divergence of the equilibrium relaxation time, $\tau = \tau_0/|\varepsilon|^{z\nu}$, as well as a divergence of the equilibrium correlation length, $\xi = \xi_0/|\varepsilon|^\nu$, where $\varepsilon = (g_c - g)/g_c$ is the dimensionless distance to the critical point. The ν and $z\nu$ are the critical exponents that characterize the universality class of the phase transition. The derivation below assumes that the exponents are well defined, i.e. they do not depend on ε and describe pure power-law dependence, and that there are no other relevant long-distance scales in the problem.

The speed of information, or the speed of second sound, is on the order of the ratio of critical length-scale to the critical time-scale

$$v_s \sim \xi/\tau = (\xi_0/\tau_0)|\varepsilon|^{\nu(z-1)}. \quad (1)$$

A causal separation near a critical point for any pair of spins could emerge if their relative distance is much larger than length scale that the information can propagate with the corresponding second sound velocity, v_s , for given finite quenching time interval. Consequently, choices of broken symmetry for spins belong to two different causal zones are not necessarily related. This is the origin of topological defects formation. The Lieb–Robinson bound [18], which characterizes the maximum speed of information in quantum many-body systems with local interactions, provides an upper-bound for v_s . We note that v_s can achieve its Lieb–Robinson upper-bound when $z = 1$, such as the prototypical 1D transverse Ising model.

Within the vicinity of g_c the quenched external field can be linearized in the form $g(t) = g_c(1 - t/\tau_Q)$, such that $\varepsilon(t) = t/\tau_Q$, where τ_Q is the quench rate and the critical point is crossed at $t = 0$. The parameter regime close to the critical point in which the system is not able to adiabatically adjust to the slowly changing external field, and effectively, to zeroth order approximation not responding, is called frozen or impulse regime. The freezing occurs at a particular time scale \hat{t} in which the relaxation time $\tau(t)$ becomes approximately equal to quench rate $\varepsilon/\dot{\varepsilon}$. Thus, by setting $\tau(\hat{t}) = |\varepsilon(\hat{t})/\dot{\varepsilon}(\hat{t})|$ we arrive at

$$\hat{t} = (\tau_0 \tau_Q^{z\nu})^{\frac{1}{1+z\nu}}. \quad (2)$$

This equation gives the KZ time-scale relevant to describe the universal behavior of the system slowly quenched through the critical point. The corresponding length-scale is a power-law of the quench rate as well

$$\hat{\xi} = \xi[\varepsilon(\hat{t})] = \xi_0(\tau_Q/\tau_0)^{\frac{\nu}{1+\nu}}. \quad (3)$$

This length scale can be used to estimate the size of the domains in the broken symmetry phase. Consequently, the density of defects is expected to vanish as $d \sim \xi(\hat{t})^{-D}$, where D is the dimensionality of system and we assume that the defects are sufficiently robust and do not relax quickly during the subsequent evolution. This is the key predication of KZM. For example, in the well-studied case of 1D Ising model in absence of any disorder we have $\nu = z = 1$. The KZM prediction for the density of excitations reads $d \sim \xi(\hat{t})^{-1} \sim \tau_Q^{-1/2}$ in that case [19–21], which can indeed be verified analytically [20]. The above argument was later developed into full dynamical scaling hypothesis, which allows to obtain similar power-laws for other observables of interest [22–26].

Understanding causal effects in disordered systems near a critical point and any attempt for estimation of density of defects requires careful analysis and not much is known outside of specific cases. Experimentally, quenches from the superfluid to the Bose glass were reported [27], with the resulting residual energies vanishing very slowly with the increasing quench rate. Full theoretical understanding is still missing in this case. Theoretical investigations are mostly limited to the class of systems with the critical point in the universality class of so-called infinite-disorder fixed point. Here, we are interested in systems belonging to this class. We first consider the prototypical example of a random transverse Ising Hamiltonian for a chain of N spins, $\hat{H} = -\sum_{n=1}^N g(n)\sigma_n^x - \sum_{n=1}^{N-1} J_{n,n+1}\sigma_n^z\sigma_{n+1}^z$, with quenched (fixed) disorder in the nearest-neighbors couplings $J_{n,n+1}$. In this article we assume that they are drawn from the flat distribution over interval $[-1, 1]$. The unit of time is set by $\hbar = 1$. Using SDRG techniques, the equilibrium properties of this model were first evaluated by Daniel Fisher [28]. For a homogenous or uniform transverse field in the model, the distribution of disorders induces a critical point that can be evaluated by relation $g_c = \exp(\log(|J_{n,n+1}|))$. For uniform distribution of $J_{n,n+1} \in [-1, 1]$ this yields a critical value of $g_c = e^{-1} \simeq 0.367\ 879$. It should be pointed out that the critical point for similar systems in two-dimensions [29] and in presence of dissipation [30] are also known to belong to this universality class. We use numerical SDRG to demonstrate universality of our non-equilibrium protocols in the section 3. We also generalize our results to Ising model with certain k -local interactions in the section 4.

The presence of disorder, changes the universality class of the critical point of the Ising model from $\nu = z = 1$ to $\nu = 2$ and $z \rightarrow \infty$, and thus quantitatively and qualitatively modifies the dependence of correlation length and density of defects on the quench time-scale. Most importantly, using SDRG techniques, it was evaluated that as the system approaches the critical point the gap of random Ising model scales as $\Delta[\varepsilon] \simeq |\varepsilon|^{1/|\varepsilon|}$ [28], and consequently the critical exponent $z\nu = 1/|\varepsilon| + O(1)$ diverges as $\varepsilon \rightarrow 0$. For that reason the KZM derivation described earlier has to be modified to take this into account [31, 32]. The characteristic time-scale \hat{t} follows from the condition $|\hat{\varepsilon}(\hat{t})/\varepsilon(\hat{t})| = 1/(\tau_Q|\varepsilon(\hat{t})|) \approx \kappa|\varepsilon(\hat{t})|^{1/|\varepsilon(\hat{t})|}$, where κ is a constant factor on the order of one. The above relation can be solved in the limit of infinitely long annealing time, $\ln(\tau_Q) \gg 1$, yielding [31]

$$\hat{\xi} \sim \frac{\ln^2(\tau_Q/\kappa)}{\ln^2[\ln(\tau_Q/\kappa)]}. \quad (4)$$

The density of defects is then suppressed logarithmically with quench time $d \sim 1/\ln^2 \tau_Q$, which is quadratically faster than simulated annealing, where defects scale as $d \sim 1/\ln \tau_Q$ [33, 34]. The existence of these logarithmic scaling laws implies that one has to run exponentially long annealing times to reduce the residual energy of the final state. However, as we will show in the next section one can recover a polynomial scaling by driving the system with a spatially inhomogeneous transverse field.

2. Causal control of topological defects with multiple critical fronts

From carefully studying defect formation under homogeneous drive fields, one can see how a new way of suppressing or controlling topological defects can emerge by being aware of causal separation of subsystems due to the extremely small values of velocity for information propagation near a critical point according to equation (1). In other words, one can try causal synchronization of the local phase transitions by inhomogeneous driving fields, as far as the critical front do not move faster than a threshold velocity corresponding to the speed of information, see [35–38] for a quantum case and [39–44] for classical counterpart. Note that this is fundamentally different than the standard annealing paradigm which is guided by the inverse of a *global gap* of a quantum Hamiltonian system which provides an upper-bound for relaxation time scales according to the adiabatic theorem. In other words, adiabaticity provides a sufficient condition for annealing time and it is not necessary to get low-energy states or even the ground state of disordered Ising systems.

Here we provide a phenomenological description of causally-induced non-equilibrium quantum phase transitions. Specifically, we develop an algorithmic quantum annealing approach to create a causal sequence of locally gapped Hamiltonians. We note that for strongly disordered systems in low dimensions there is a

quantum Griffiths region that is spread in the disordered and ordered phases, i.e. on both sides of a critical point [28]. Within the Griffiths region the system undergoes effective local phase transitions that are space–time separated in nature even if the control fields are homogeneous. The key observation is that one can create situations in which the choices of symmetry-breaking events in a local neighborhood that have already experienced phase transitions earlier could influence the symmetry breaking events elsewhere, provided that the control fields have certain inhomogeneous spatiotemporal structures. These symmetry breaking events are perceived by the rest of the system, which is still in a disordered phase, as effective boundary conditions influencing their local fields.

In order to develop an algorithmic quantum annealer, here we construct a general class of IQA schedules. They are a function of a fixed total quench time or annealing time $T \sim \tau_Q$, proportional to the annealing rate of the homogeneous quench τ_Q introduced in the previous section. The performance of the algorithms are evaluated by computing the precision ϵ_Q of approximating the ground state. Here we mostly focus on the random instances of strongly-disorder spin chains, nevertheless our construction is general and can be applied to higher-dimensional systems [17]. There are two main reasons for such a choice. First, for 1D case we can simulate their dynamics exactly using a mapping to free-fermionic system, as e.g. in [31, 32, 37]. Also the critical behavior of such systems when driven via homogeneous external fields have been studied extensively, thus the new non-equilibrium physics of such systems when driven inhomogeneously can be better benchmarked and appreciated. The overall Hamiltonian for a system of N spins under an inhomogeneous driver field can be written as:

$$H(t) = -\sum_{n=1}^N g(n, t) \sigma_n^x - \sum_{\langle n,m \rangle} J_{nm} \sigma_n^z \sigma_m^z \quad (5)$$

and the quality of an output state is characterized by a normalized residual energy as: $\epsilon_Q = Q/N$, with $Q = \langle \psi(\tau_Q) | H_p | \psi(\tau_Q) \rangle - \langle \psi_{gs} | H_p | \psi_{gs} \rangle$, where $|\psi(\tau_Q)\rangle$ is the quantum state of the system at the final annealing time. $|\psi_{gs}\rangle$ is the ground state of the classical time-independent Hamiltonian, or the problem Hamiltonian, $H_p = -\sum_{\langle n,m \rangle} J_{nm} \sigma_n^z \sigma_m^z$, with eigenvalue $E_{gs} = \langle \psi_{gs} | H_p | \psi_{gs} \rangle$. We note that for pure systems, where $J_{nm} = J$, the normalized residual energy can be related to Kibble–Zurek correlation length $\hat{\xi}$ by $\epsilon_Q \sim |J| \hat{\xi}^{-D}$ where $\hat{\xi}^{-D}$ is the density of topological defects and D is the dimension of system.

Here we assume that the inhomogeneous drive field is a transverse field that can be locally modulated for every individual spin. This Hamiltonian can be realized with the near-term quantum annealing technologies currently being developed at the D-Wave Quantum Computing Systems and Google Quantum AI Lab.

As we describe in the next section, for any given instance of disorders $\{J_{nm}\}$ as we drive the system toward the quantum critical point, the system responds to quantum fluctuations within M distinct ‘clusters’, which are related to the emergence of rare local regions within the Griffiths phase. As we will show, the number and locations of clusters can be estimated via a simple preprocessing step that grows linearly with the size of the chain for 1D system. The generalization to higher dimensional system is presented in [17].

In each cluster we drive the many-body system by a transverse Ising Hamiltonian with some local structure. Thus, we drive these M clusters simultaneously into some space–time separated inhomogeneous transitions

$$g(n, t) = \sum_{l=0} h_l(n) \lambda_l(t) + \sum_{k=1}^M \omega_k g_k(|n - n_k| - v_k(n, t)t), \quad (6)$$

where each h_l is the time-independent global magnetic field which has a spatial structure and each $\lambda_l(t)$ is spatially uniform but it can generate nonlinear dependence to time. The terms $\sum_{k=1}^M \omega_k g_k(|n - n_k| - v_k(n, t)t)$ characterize various spatiotemporal dependencies of traveling quantum critical fronts, where $|n - n_k|$ denotes a distance measure of node n from some center node n_k per cluster where we trigger the quantum fluctuations. The center of these spatiotemporal inhomogeneities can be shifted linearly in time by $v_k(n, t)t$ with a spatiotemporal dependence for each k cluster. However, for simplicity, for rest of this work we concentrate on a constant critical front motion for each cluster; that is $v_k(n, t) = v_k$.

In the following section, we define the shape and two different kind of velocities for critical fronts and provide two simple examples of type I and II annealing, namely periodic inhomogeneous annealing, and multiple-critical-fronts inhomogeneity.

2.1. Shape and velocity of critical fronts

The inhomogeneity slope and its horizontal and vertical velocities of inhomogeneity can be characterized by a set of hyper-parameters $\{\alpha, v^h, v^v\}$ corresponding, respectively, to local slope of the instantaneous field in space and its spatial (horizontal) and temporal (vertical) velocities, that are defined by derivatives of $g(n, t)$ and $n(g_{fix}, t)$ as:

$$v^v(n, t) = -\partial g(n, t) / \partial t, \quad (7)$$

$$v_k^h(n, t) = \partial n(g_{\text{fix}}, t) / \partial t, \quad (8)$$

$$\alpha(n, t) = \partial g(n, t) / \partial n. \quad (9)$$

Thus, we can derive closed form expressions over these hyperparameters for different annealing schedules. To appreciate the generality of the shape of $g(n, t)$, we consider two concrete and qualitatively distinct classes for IQA with respect to possible temporal and/or spatial inhomogeneities.

2.2. Type I IQA: space and time separated inhomogeneity

In this class, we consider a general form of independent or separated space and time quantum fluctuations to drive the annealing dynamics

$$g(n, t) = \sum_{l=0} h_l(n) \lambda_l(t). \quad (10)$$

An example of this class will be a periodic spatial inhomogeneity (standing wave) combined with spatially-independent time-local inhomogeneity as:

$$g(n, t) = h_0 \lambda_0(t) + \lambda_1(t) \sum_{k=-\infty}^{\infty} a_k e^{i\pi k n},$$

where each term in the spatial contribution in the second term corresponds to an estimated cluster size. We provide a simple illustration of these periodic spatial inhomogeneities in the next section. We note that KZM—in the context of pure systems—was also extended to quenches that are homogeneous in space, but nonlinear (inhomogeneous) in time [25, 45]. Such inhomogeneity adjusts the quench rate to the distance from the critical. Consequently it allows to reduce the number of generated defects.

2.3. Type II IQA: spatiotemporal inhomogeneities

In this class, we build a sufficiently general example by creating a multiple-critical-fronts annealing schedule in M clusters where critical fronts in each cluster are moving with the speed $v_k(n)$ and each are governed by a separate activation function $\tanh[\theta_k(|n - n_k| - v_k(n)t)]$:

$$g(n, t) = g_c \left\{ 1 + \sum_{k=1}^M \omega_k \tanh[\theta_k(|n - n_k| - v_k(n)t)] \right\}, \quad (11)$$

where g_c is the critical value of transverse field. We note that there is no particular significance for our choice of activation function here. As an important special class of the above driver field, we linearize the activation function in each cluster near quantum critical point, that is $\tanh[\theta_k(n - v_k(n)t)] \simeq \theta_k(n - v_k(n)t)$, then for each cluster we get:

$$g_k(n, t) = g_c \{1 + \theta_k(n - v_k(n)t)\}. \quad (12)$$

In the first example of this type, we consider an inhomogeneity with constant v_k for each cluster of the form $g_k(n, t) = g_c \{1 + \tanh[\theta_k(n - v_k t)]\}$ which yields $n = \tanh^{-1}(g_{k-\text{fix}}/g_c - 1)/\theta_k + v_k t$.

$$v^v(n, t) = g_c [1 - \tanh^2[\theta_k(n - v_k t)]] \theta_k v_k = \alpha_k(n, t) v_k,$$

$$v_k^h(n, t) = v_k,$$

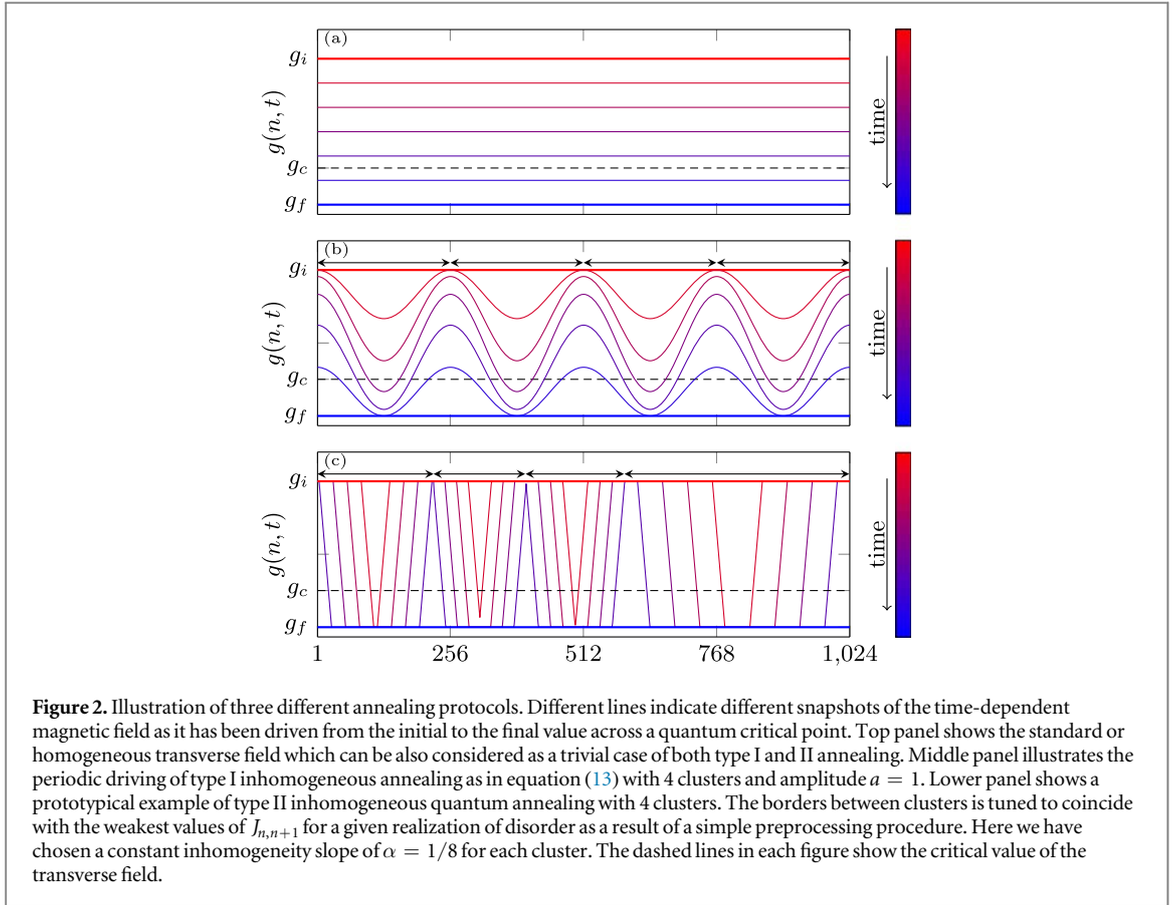
$$\alpha_k(n, t) = g_c [1 - \tanh^2[\theta_k(n - v_k t)]] \theta_k.$$

In the next example of this type, we consider linear approximation of activation function near critical point which yields $g_k(n, t) = g_c \{1 + \theta_k(n - v_k t)\}$ and $n = (g_{k-\text{fix}}/g_c - 1)/\theta_k + v_k t$. Thus we have:

$$v^v = g_c \theta_k v_k = \alpha_k v_k, v_k^h = v_k, \text{ and } \alpha_k = g_c \theta_k.$$

2.4. Standard AQC: absence of any inhomogeneity

We note that the standard or homogeneous quantum annealing schedule which has been extensively studied and numerically benchmarked for almost two decades can be considered as the extreme limit of either type-I or type-II of IQA. In the former case we have one spatially uniform transverse field, $g_0(n) = \text{const.}$, and linear velocity $\lambda_0(t) = (1 - t/\tau_Q)$, where τ_Q is the overall annealing time-scale. Thus we have the familiar form of homogeneous transverse field, which is linear in time, as: $g(n, t) = g_0(1 - t/\tau_Q)$. In order to see AQC as a extreme limit of type-II IQA, we must note that the homogeneous transfer field can be considered as a single critical front with a trivial flat shape with infinite velocity; that is $\theta \rightarrow 0$ and $v(n) \rightarrow \infty$, while $\theta v(n)$, which is basically the vertical velocity, will be equivalent to the inverse of annealing time and thus will be finite. The hyperparameters for homogenous annealing, respectively, become $v^v(n, t) = 1/\tau_Q$, $v_k^h(n, t) = \infty$, and $\alpha(n, t) = 0$.



2.5. Exponential suppression of defects

To illustrate the power of multi-front critical control, we numerically investigate two concrete forms of type I and II inhomogeneous annealing as described above and compare their performance against standard QA. All the simulations in this section are done using the Jordan–Wigner transformation that maps the Hamiltonian in equation (5) onto the system of free fermions where it can be solved numerically in a standard way. For details of these techniques, we refer the readers to the appendix B of [37]. For our numerical analysis here the cluster formation that we invoke is simple and has linear scaling with the system size for Ising chains. In the next section we use SDRG to examine construction and scaling of causal gaps.

Examples of the type I and II annealing schedules for one random instance of Ising chain are given in figure 2, in which different snap-shots of time-dependent transverse fields are plotted along the chain. In figure 2(a), we illustrate the trivial/homogeneous schedule. In figure 2(b) we explore the effects of periodic critical fronts by constructing the schedule

$$g(n, t) = g_i(1 - t/T) + a \cos(kn) \sin(\pi t/T), \quad (13)$$

where T is the total evolution time. Finally, in figure 2(c) we illustrate an example of multiple-critical-fronts strategy.

In the latter, in order to decide the position of the cluster, we employ a simple preprocessing. It is based on observation, that for strictly 1D geometry, the instantaneous local gap may be set by a single, very weak link. We discuss it in more detailed in section 3. Such weak link sets the local time-scale needed for adiabatic transition dividing the chain into two weakly interacting parts. We want to place the borders between the clusters at such links, as (i) they would require the longest time to align according to the weak coupling and (ii) the energy penalty for placing the defects there is the smallest. To that end, we look for the largest cluster—starting at the end of the chain or at the end of the previous cluster—where $\min_{\text{cluster}_k}(J_{n,n+1}) \cdot \kappa > v_k$. Here, v_k would be the velocity of the front for this (k th) cluster, and κ is a parameter fixing the exact value of the threshold (in practice $\kappa \approx 2$). This condition would allow for adiabatic transition as if the energy gap were set by single links only. If the condition is not satisfy the considered candidate for cluster is cut at its weakest link, creating a new smaller cluster where we check the condition again. The procedure is repeated until full chain is divided into clusters. As a result, for a total fixed annealing time, all velocity v_k are cluster dependent; allowing optimization of the computational parameters over the available time. For a cluster of size L_k sites the (vertical) velocity is given by $v_k^v = (|g_f - g_i| + \alpha L_k/2)/T$ for a given fixed total annealing time T . Each cluster is driven separately, and the

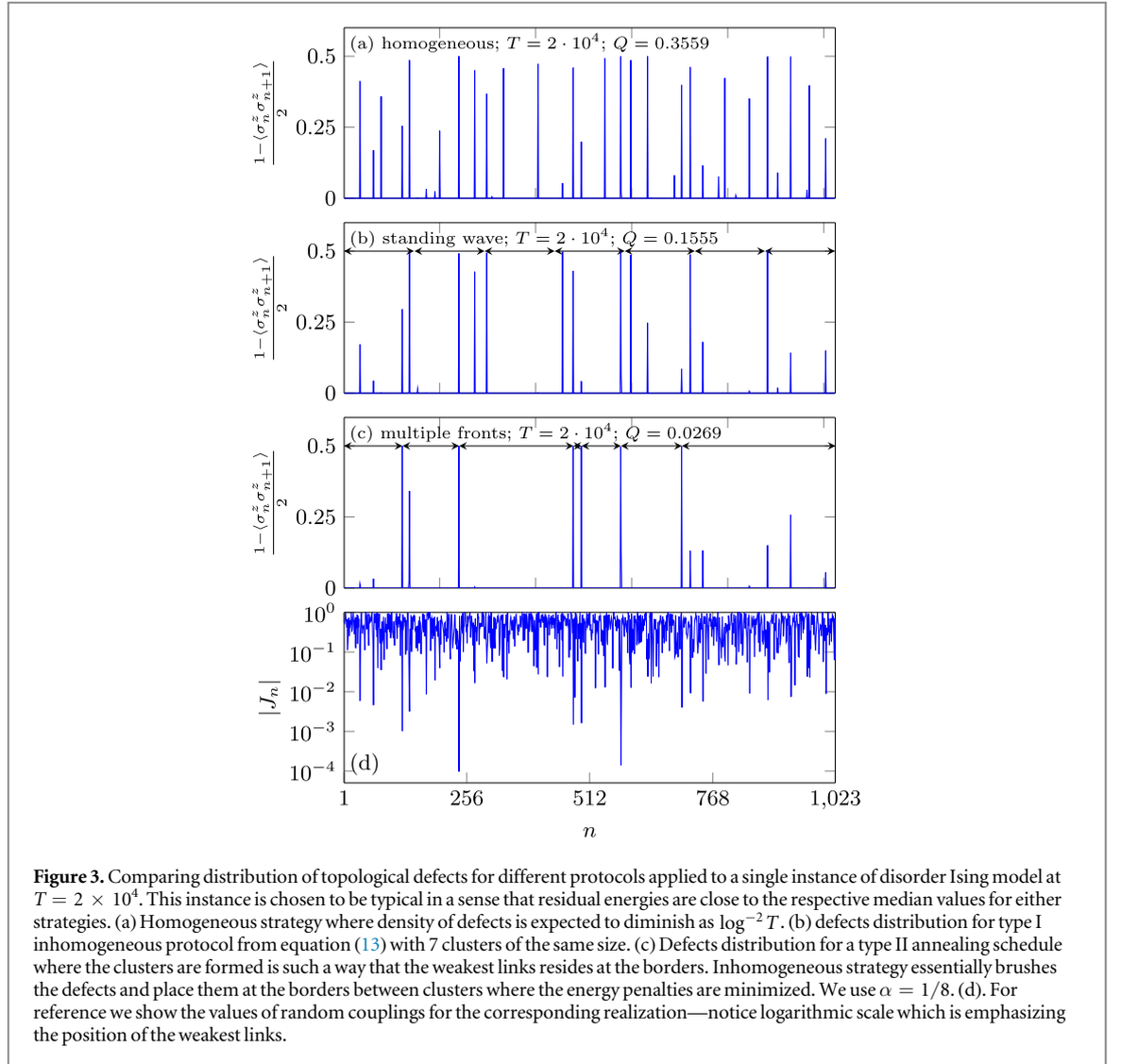


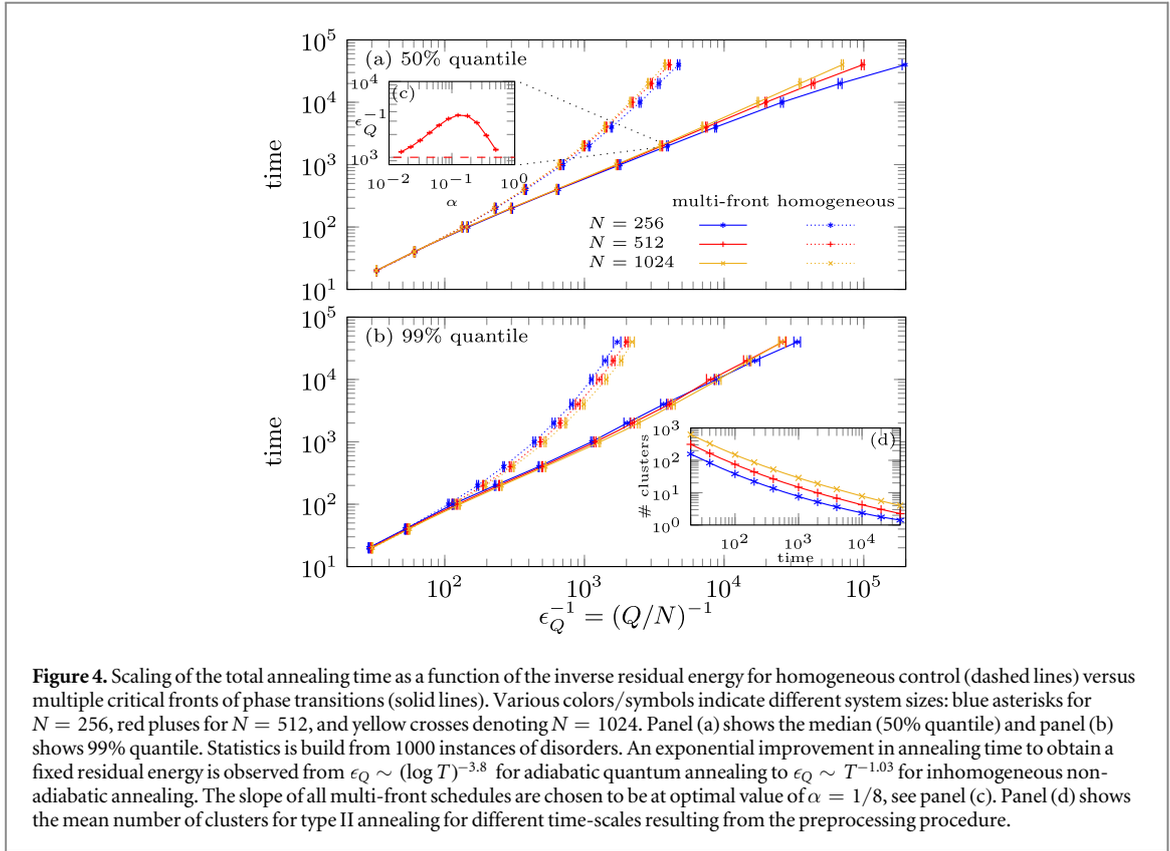
Figure 3. Comparing distribution of topological defects for different protocols applied to a single instance of disorder Ising model at $T = 2 \times 10^4$. This instance is chosen to be typical in a sense that residual energies are close to the respective median values for either strategies. (a) Homogeneous strategy where density of defects is expected to diminish as $\log^{-2} T$. (b) defects distribution for type I inhomogeneous protocol from equation (13) with 7 clusters of the same size. (c) Defects distribution for a type II annealing schedule where the clusters are formed in such a way that the weakest links resides at the borders. Inhomogeneous strategy essentially brushes the defects and place them at the borders between clusters where the energy penalties are minimized. We use $\alpha = 1/8$. (d). For reference we show the values of random couplings for the corresponding realization—notice logarithmic scale which is emphasizing the position of the weakest links.

inhomogeneous front is brushing from the middle of each cluster to both ends simultaneously. For each cluster spanning spins $n = 1, 2, \dots, L$ (counting from the beginning of the cluster) the magnetic field is constructed as:

$$g(n, t) = \begin{cases} g_i, & \left| n - \frac{L}{2} \right| - tv > \frac{g_i - g_f}{2\alpha}, \\ \frac{g_i + g_f}{2} + \alpha \left(\left| n - \frac{L}{2} \right| - tv \right), & \left| n - \frac{L}{2} \right| - tv \leq \frac{g_i - g_f}{2\alpha}, \\ g_f, & \left| n - \frac{L}{2} \right| - tv < \frac{g_f - g_i}{2\alpha}. \end{cases} \quad (14)$$

The probability distribution of topological defects for these protocols are presented in figure 3. The defect density is evaluated for a strongly disordered instance of a spin chain consist of up to about 1000 qubits with $J_{n,n+1}$ sampled randomly from $[-1, 1]$. Here, the inhomogeneous annealing can be regarded as a many-body quantum control strategy which can significantly reduce the number of topological defects by synchronizing the symmetry breaking events and can brush the remainder of defects into the weakest J_{ij} where they act as defect sinks. In other words, not only do these non-adiabatic paths suppress the emergence of domain walls, but also minimize the energy cost per defect by several order of magnitudes.

The scaling of annealing or quenching time as a function of inverse residual energy is plotted in figure 4 for 1000 random instances of 1D Ising chains ranging from 256 spins to 1024 spins. It can be observed that the annealing time is improved exponentially over standard AQC scheme for typical, 50% quantile, as well as harder instances, 99% quantile (each corresponds to the residual energy in which $x\%$ instances have smaller values). In the homogeneous protocol we have $\epsilon_Q \sim (\log T)^{-\gamma}$. We obtain $\gamma \approx 3.8 \pm 0.4$. It can be compared with $\gamma \approx 3.4$ which has been obtained by from Caneva *et al* [32] from smaller values of quench times T and for



slightly different protocol, where the magnetic field was also disordered. What is important here is that γ is larger than 2, i.e. the value of exponent governing the scaling of defect density. This results from defects being more likely to appear on the links with smallest $|J_{n,n+1}|$. For uniform distribution in $[-1, 1]$, $\gamma = 4$ would correspond to defects appearing only at the weakest links.

On the other hand, in type II protocol we observe $\epsilon_Q \sim T^{-1.03}$ (fit for $N = 1024$, $T > 100\%$ and 50% quantile). Multi-front protocol have been constructed by fixed $\alpha = 1/8$, where optimality of such choice is shown in panel (c) for $T = 1000$. Similar plots for other time-scales (not shown) suggest that in this system this value is optimal independently if $T \gg 1$. Panel (d) shows the mean number of clusters in type II annealing for different time-scales. For large times it should scale as $\sim N/T^{0.5}$ which results from the preprocessing procedure. This follows from the expected size of clusters which can be solved in given T for $J_{n,n+1}$ drawn from a uniform distribution in $[-1, 1]$.

3. Scaling relations for causal gaps

Here, we introduce the notation of causally gapped Hamiltonians that are created by time-dependent multiple critical fronts introduced in the previous section. In particular, we derive a scaling relation for the distributions of minimum causal gaps as the function of system size and inhomogeneity slope. In this work, we define the *causal gaps* as the inhomogeneously-induced instantaneous energy gap which becomes relevant when the critical front is driven below a threshold velocity allowing for information of symmetry-breaking events to propagate. The core idea is that there is an effective threshold velocity v_t^k that determines the suppression of topological defects formation within each cluster in disordered systems. If the velocity of the front in each cluster is much larger than this threshold speed, $v_k \gg v_t^k$ for all k , then the effect of the spatial variations of the control field become irrelevant in the sense that we will recover the standard critical dynamics created by homogeneous driving, which can be understood by the standard KZM. However, when we drive each critical front such that $v_k \ll v_t^k$, the length scale and shape of the critical front becomes highly relevant allowing to suppress creation of the topological defects in each cluster. The shape of the critical front determines the number of spins that simultaneously experience criticality creating an effective (finite-size) energy gap. Otherwise, the homogeneous system would be gapless at the critical point in the thermodynamical limit.

3.1. Causal gap for pure systems

In the absence of disorder and with sufficiently smooth critical front, that is $\alpha \ll 1$, one can invoke a variant of KZM to estimate when the inhomogeneity of the driving front is relevant. This question can be regarded from two perspectives [35, 36].

Firstly, starting from the limit of homogeneous driving, we note that the relevant speed of information at the critical front can be expressed as $\hat{v} \sim \hat{\xi}/\hat{t}$. Here, $\hat{\xi}$ and \hat{t} are the effective length scale and time scale that the system experience according to KZM given by the relations (2) and (3), respectively. This yields $\hat{v} = \tau_Q^{\nu(1-z)/(1+z\nu)}$. Next we consider the control parameter $\varepsilon(n, t)$ to be position dependent. We linearize the relation at the critical front for fixed position n_{fixed} as

$$\varepsilon(n, t) = \alpha(n_{\text{fixed}} - vt) = -\alpha vt + \text{const.} \quad (15)$$

This gives the local annealing rate $\tau_Q(n) = 1/(\alpha v)$. Causality implies that if $v \gg \hat{v}$, then the choice of symmetry breaking which happened earlier along the chain cannot influence what is happening later and we recover the independent defect formation assumed in the standard KZM. The self-consistency condition allows to express such threshold velocity as a function of our main control parameter α . This is obtained by inserting the above annealing rate into the expression for \hat{v} , which leads to

$$\hat{v}_t \sim \alpha^{\nu(z-1)/(1+\nu)}. \quad (16)$$

Alternatively, we could look at the instantaneous Hamiltonian resulting from inhomogeneous front in equation (15). We focus here on the instantaneous ground state of such system, which is interpolating—in space—between order and disorder phases. They are spatially separated by an effective critical regime which size, called a *penetration depth*, can be estimated as

$$\hat{\xi}_i \sim \alpha^{-\frac{\nu}{\nu+1}}. \quad (17)$$

It follows from a variant of KZM argument, so-called KZM in space [46–49], where one asks about characteristic distance from n_{fixed} up to which the system is able to locally adjust to $\varepsilon(n)$ changing in space as if it were locally homogeneous with local correlation length determined by $\varepsilon(n)$. Apart from the Ising model [46–49], the interplay between inhomogeneous external field and criticality was studied in the context of spin-1 Bose–Einstein [50], the XY model [51], and the XXZ model [52].

The finite size of the effective critical region in equation (17) allows to estimate the gap of the instantaneous Hamiltonian as

$$\hat{\Delta}_i \sim \alpha^{\frac{z\nu}{\nu+1}}. \quad (18)$$

By combining those characteristic scales we obtain a threshold velocity v_t which is again given by equation (16). The meaning of this threshold velocity is however different here. Namely, we can expect that if the velocity v in equation (15) is $v \ll v_t$, then the system would be able to follow its instantaneous ground state.

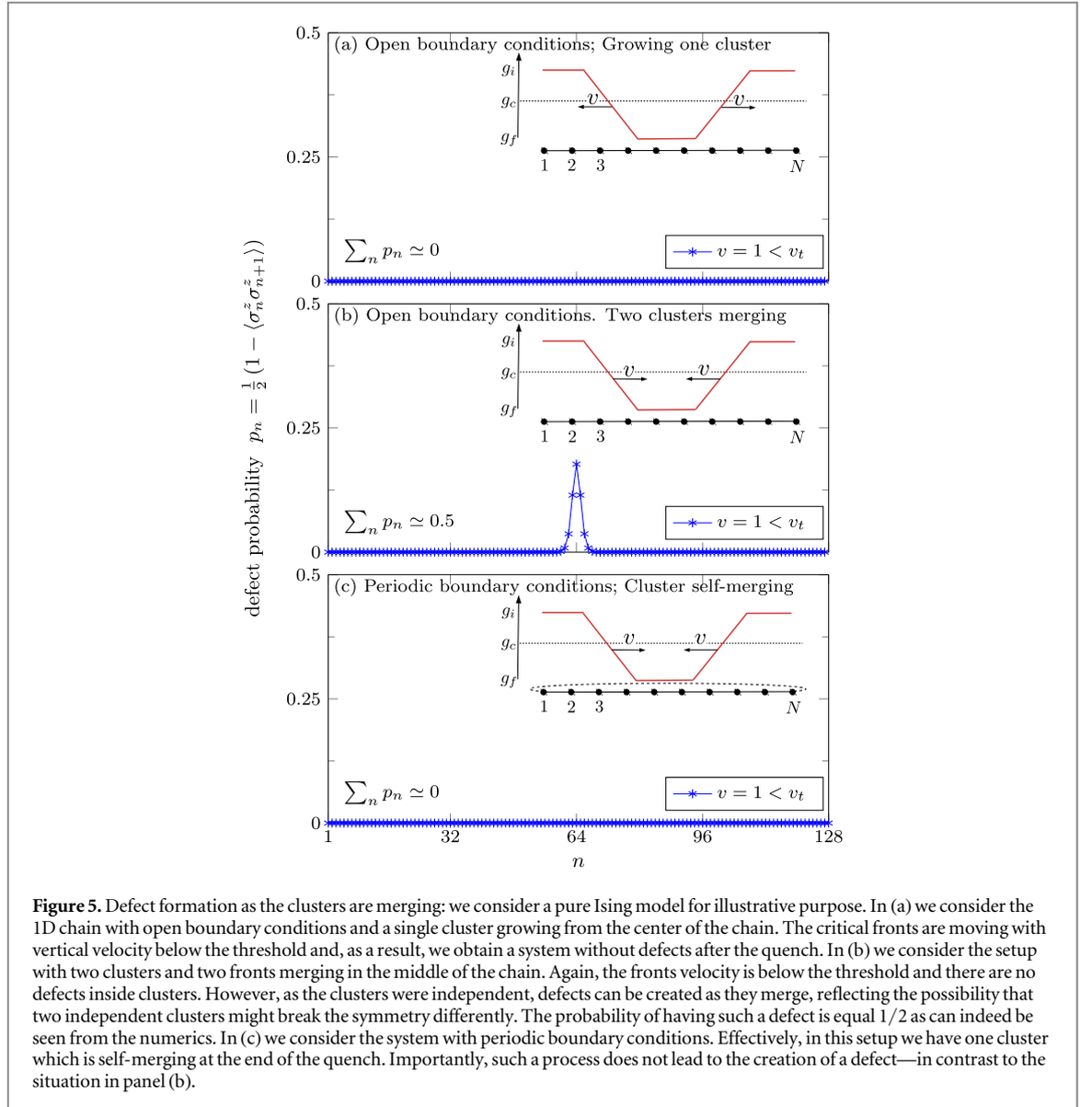
It should be noted that whenever $z = 1$ the threshold velocity becomes constant and we get a sharp suppression of topological defects whenever we drive the system with critical front that is slightly below v_t . The value of v_t becomes equal to 2 for a transverse Ising chain when all couplings are equal to one [35]. This suppression of defects and causal synchronization, however, could be affected when we drive the system with a multi-front strategy. For a simple pure system driven by two critical fronts, both moving with velocity smaller than v_t , in figure 5 we illustrate how the defects can be created when two clusters are merging together. This highlights the interplay of the causal effects and different boundary conditions on the defect suppression.

As we have discussed in section 1, the disordered systems have a completely different critical phenomena as $z\nu \rightarrow \infty$. In this case one has to modify equation (18) accordingly [37] by taking into account that at the critical point the gap is expected to vanish as a stretched exponential with the system size—effectively given in our case by equation (17). We elaborate on this in the next section in the context of multiple critical fronts driving strategy.

3.2. Universality of causal gaps via SDRG

In this section we use a combination of analytical and numerical SDRG techniques to show that the distribution of causal gaps are universal irrespective of the shape of inhomogeneities for strongly-disorder Ising systems. We also derive a scaling relation for the dependence of the minimal causally-induced gap on the actual system size and slope of inhomogeneity. We first present implementations of SDRG for disordered spin chains under various schedules.

The core concept of all RG techniques is to re-express the parameters which define a problem by coarse-graining some microscopic degrees of freedoms. In each step of the RG flow we arrive at effective Hamiltonian terms that have fewer and much simpler parameters acting on a lower energy and larger (macroscopic) length scale, such that certain physical or computational aspect of interest in the original problem remain unchanged. The procedure can be recursively repeated until the Hamiltonian is no longer changing which indicates that we

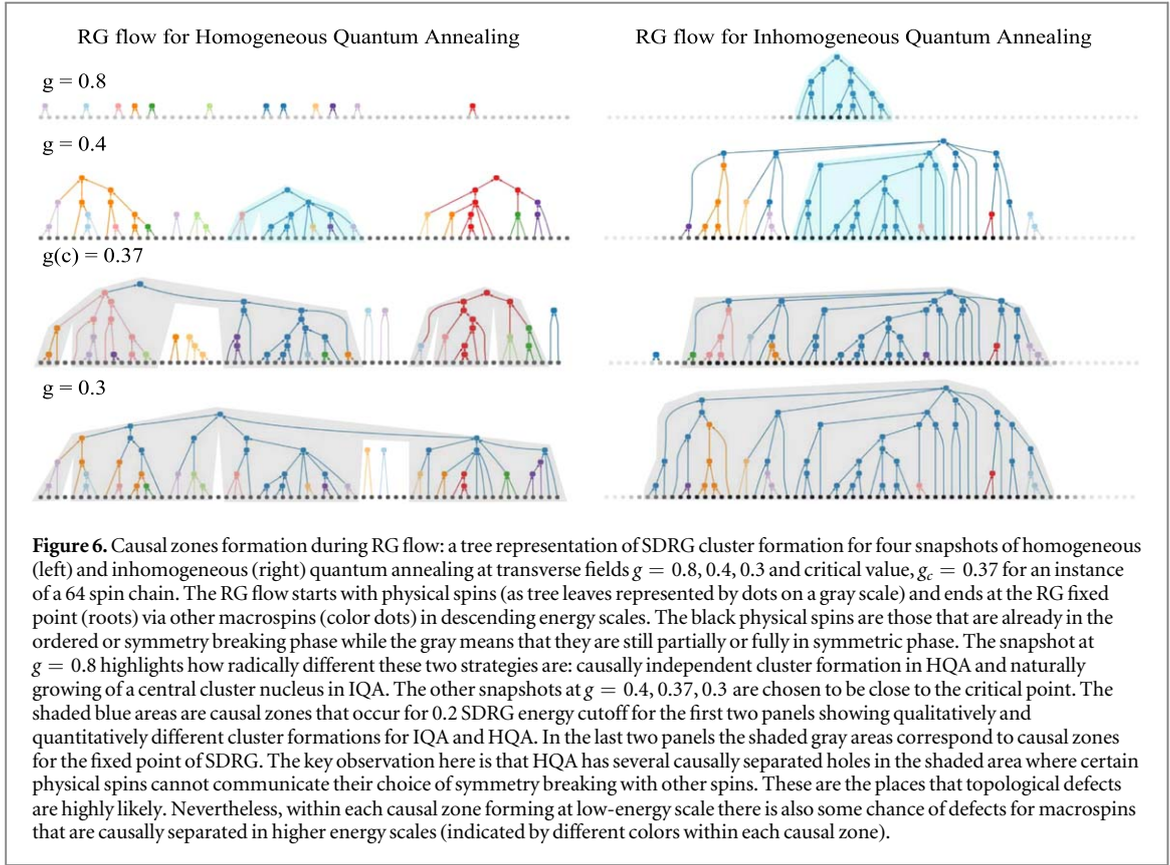


have arrived at the fixed point of the RG flow. In SDRG, that is specifically developed for disordered systems and has been generalized to higher-dimensional systems [53], the largest energy scale is systematically removed via two different operations: site decimation and bond decimation [28, 54], see [55] for a review.

A site decimation occurs whenever we have a site-dependent transverse field which is the largest energy scale within a local neighborhood of our system; that is $g_i > J_{ij} \forall j$. Site decimation means that we basically lock the spin i to the direction of its local transverse field. Such spin will be effectively decoupled from the rest of the system. Emerging new couplings are generated between all neighbors of the decimated spin i . These effective couplings can be computed within second order perturbation theory as $\tilde{J}_{jk} = \frac{J_{ij}J_{ik}}{g_i}$, if $J_{jk} = 0$ or otherwise

$\tilde{J}_{jk} = \max\left(J_{jk}, \frac{J_{ij}J_{ik}}{g_i}\right)$. A bond decimation is performed in similar fashion. A bond decimation occurs whenever we have two sites i and j interacting via J_{ij} that is the largest energy scale within a local neighborhood of our system; i.e. $J_{ij} \geq g_i$ and $J_{ij} \geq g_j$; and also $J_{ij} \geq J_{ik} \forall k$ and $J_{ij} \geq J_{jl} \forall l$. Bond decimation simply means that we lock the two sites i and j into a macrospin by projecting the combined pair into their local ground states. No additional bonds or coupling between any spins are generated in this case. All the spins that were previously coupled to at least one of the sites are now interacting with the combined cluster. For the spins that were coupled to both i and j we invoke a maximum selection rule. Effective transverse field at the emerging macrospins becomes $\tilde{g}_i = \frac{g_i g_j}{J_{ij}}$.

We apply the above RG procedure to our time dependent Hamiltonian by considering each time as a snapshot for different instances of static spin chains, see figure 6. The SDRG simulation confirms our assumption of causally independent clusters in the homogeneous strategy, in the low-energy or long wavelength



limit. In contrast, we observe inter-cluster causal dependence emerging in the multiple critical fronts strategy, see the SDRG visualizations of our protocols in figure 6.

Next, we discuss an upper-bound for a global threshold velocity such that the multiple critical fronts strategy can lead to suppression of excitation between the low energy manifold and the rest of excited states. If we have full parallel annealing in all clusters simultaneously, we essentially interrupt the causality of symmetry breaking events between different clusters as illustrated in figure 6. When the fronts are sufficiently separated, the RG flow makes the corresponding clusters exponentially decoupled. Consequently, when looking at the possible transitions, we can consider each front independently. The transition matrix elements for each cluster k :

$$\left\langle 0, t \left| \frac{d\hat{H}}{dn_k^f} \right| 1, t \right\rangle \frac{dn_k^f}{dt} = \Omega_k(n_k^f) v_k, \quad (19)$$

where n_k^f encodes the (time dependent) position of the front and $v_k = \frac{dn_k^f}{dt}$ is its vertical velocity. $\Omega_k(n_k^f)$ defines local instantaneous ground state to first excited state transition matrix elements for that front. For simplicity of analysis in following sections we choose a constant and uniform velocity for all the critical fronts in various clusters i.e. $v_k = v$. Due to causal independence of clusters in multiple criticality, the adiabatic condition for low-energy states (approximate solutions) can be characterized by maximum over of all possible local transition matrix over its local gap, $\Delta_k(n_k^f)$; that is $\frac{\Omega_k(n_k^f)}{\Delta_k^2(n_k^f)}$. Then the low-energy quasi-adiabatic dynamics is expected for

$$v \ll v_t = \min_k \frac{\Delta_k^2(n_k^f)}{\Omega_k(n_k^f)}. \quad (20)$$

In the following we are going to discuss the universality of the threshold velocity, reflected in its dependence on the shape of critical front characterized by the slope α .

Firstly, it is worth drawing the connection between the condition in equation (20) with the threshold velocity which was derived in the previous section. There, it was calculated as $\hat{v} = \hat{\xi}_i / \hat{t}_i$, where $\hat{\xi}_i$ and \hat{t}_i were the characteristic length scale (the penetration depth) and time scales (given by the inverse of the gap) related with the slope of inhomogeneity α . Similarly, the adiabatic condition is sometimes formulated as, $v \ll \Delta \cdot \Gamma$, where Δ is the energy gap, and Γ estimates the width of the region (in the driving parameter space) for which the gap is close to its minimal value, see e.g. [57]—in analogy to the avoided level crossing and the Landau–Zener problem. To resolve this seeming inconsistency (i.e. that the gap appearing in equation (20) is squared comparing to other expression), we note that $\Omega_k(n_k^f)$ and $\Delta_k(n_k^f)$ are not independent. We expect that $\Omega_k(n_k^f) / \Delta_k(n_k^f) \sim \xi_i^{-1}$, i.e. it

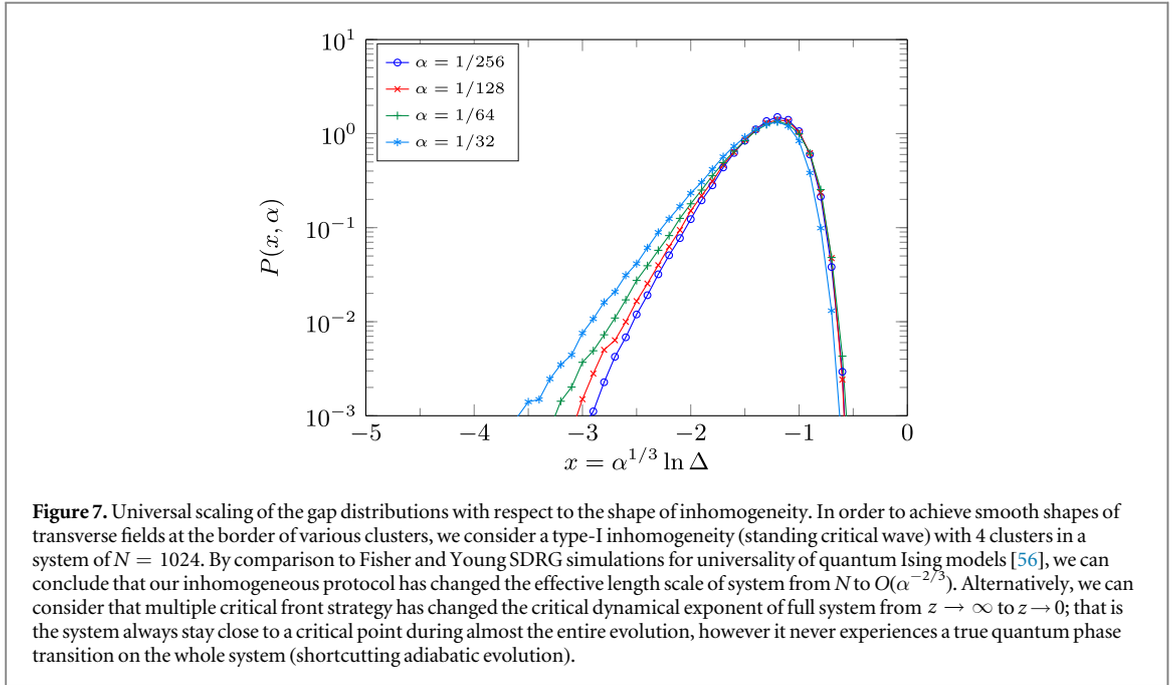


Figure 7. Universal scaling of the gap distributions with respect to the shape of inhomogeneity. In order to achieve smooth shapes of transverse fields at the border of various clusters, we consider a type-I inhomogeneity (standing critical wave) with 4 clusters in a system of $N = 1024$. By comparison to Fisher and Young SDRG simulations for universality of quantum Ising models [56], we can conclude that our inhomogeneous protocol has changed the effective length scale of system from N to $O(\alpha^{-2/3})$. Alternatively, we can consider that multiple critical front strategy has changed the critical dynamical exponent of full system from $z \rightarrow \infty$ to $z \rightarrow 0$; that is the system always stay close to a critical point during almost the entire evolution, however it never experiences a true quantum phase transition on the whole system (shortcutting adiabatic evolution).

is directly proportional to the size of the effective critical region. We show this in figure 7 and discuss further below. Importantly, this allows us to focus on the scaling of the gap.

The effective size of the critical region (for a given front) is given by equation (17). In our case $\nu = 2$, which gives $\xi_i \sim \alpha^{-2/3}$. The fronts are independent if their respective distances, or the sizes of the clusters, are $\gg \xi_i$. The typical gap related with such critical front is then expected to scale as a stretched exponential in ξ_i , $\Delta_k(n_k^f) \sim e^{-\text{const}\sqrt{\xi_i}} \sim e^{-\text{const}\alpha^{-1/3}}$ [37].

To be more precise, we consider $P(\Delta, \alpha)$ as a distribution of the minimal relevant instantaneous gap Δ , as each front is traveling within a cluster for a fixed α . We argue that we can observe the universality of local gap probability distributions by introducing a rescaled log-gap $x = \alpha^{1/3} \log \Delta$, with the expected $P(\Delta, \alpha)d\Delta = \tilde{P}(x)dx$, and universal distribution $\tilde{P}(x)$. In contrast, the universal scaling of the gap in the finite size homogeneous system at criticality is known to be described by universal distribution $P(\Delta, N)d\Delta = \tilde{P}(x)dx$ with $x = N^{-1/2} \log \Delta$ [56, 58]. Our ansatz is then a directly consequence of an assumption that the effective size of the critical region in our protocol is no longer given by N but instead is characterized by the penetration depth in equation (17).

We verify those scaling predictions in figure 7. We consider the system with two clusters, i.e. 4 critical fronts to highlights the independence of the front and that the scaling prediction naturally carry on to the case of multiple fronts. We calculate the minimal relevant gap (which would be related with one of the fronts), which we distinguish by finding the minimal energy eigenstate with the corresponding transition matrix element Ω above some threshold. We calculate the gap via numerical SDRG which ends at a system of few spins which is subsequently exactly diagonalized. While the SDRG procedure which we use is introducing some errors, we check that the results can be essentially reproduced by the numerically exact solutions based on free-fermionic picture for transverse-field Ising models. Using SDRG allows us to highlight that the fronts are independent as they become effectively decoupled during SDRG flow. We compute the distribution of Δ by sampling, for each disorder instance, from different equidistant front positions as they are moving though the chain. We disregard the beginning and the end of the protocol when the system is far from criticality and focus on the relevant intermediate part where we have independent critical fronts. In figure 7 we collect the statistics using 5000 disorder instances. Indeed, we can see that the peaks of the rescaled distribution collapse validating the scaling anzats. The tail corresponding to small energies, where the distributions do not properly collapse, is non-universal and results from occasional very weak links (as $J \in [-1, 1]$). They give rise to gaps of similar order (occurring again when the magnetic field acting on the neighboring spins is again almost switched off), which are characteristic for strictly 1D system. We elaborate more on this later in this section. The presence of such gaps is especially pronounced for larger values of α when the typical gap $\sim e^{-\text{const}\alpha^{-1/3}}$, which can be attributed to many-body effects, is larger.

The analysis above regards local, instantaneous gap related with the critical fronts traveling though the chain. To quantify the difficulty of the problem, we consider the minimal gap encounter during such quench. To that

end we focus on the distribution $P(\Delta, \alpha)d\Delta = \tilde{P}(x)dx$. Importantly $\tilde{P}(x)$, as calculated for the homogeneous critical case in [56, 58], has a Gaussian tail for $|x| \gg 1$,⁴ i.e.

$$\tilde{P}(x) \sim e^{-ax^2}. \quad (21)$$

Figure 7, especially in the limit of small α when gaps attributed to weak links are less relevant, indicates that the above holds also in our case⁵. Now, let the probability that Δ is smaller than some Δ_{\min}^q be

$$P(\Delta < \Delta_{\min}^q, \alpha) = \int_{-\infty}^{x_q} \tilde{P}(x)dx = \epsilon, \quad (22)$$

where $x_q = \log(\Delta_{\min}^q)\alpha^{1/3}$. In order to find the probability distribution for minimal gap we assume that we are sampling N times from $P(\Delta, \alpha)$ (or more precisely proportional to N times in the limit of large N). That way, the probability

$$P(\Delta_{\min} > \Delta_{\min}^q) = (1 - \epsilon)^N = q, \quad (23)$$

where Δ_{\min} is the minimum from the sample of N instantaneous gaps. This equation defines Δ_{\min}^q as a q -quantile for the global minimal gap. Now, we obtain Δ_{\min}^q from the above equation. This gives $\frac{1}{N} \log q = \log(1 - \epsilon) \simeq -\epsilon \sim \frac{1}{2ax_q} e^{-ax_q^2}$, which is obtained by expanding the error function for the Gaussian tail in equation (21), to the leading order in small x_q . Solving this equation in the leading order we obtain

$$\alpha^{1/3} \log \Delta_{\min}^q = x_q \approx -\sqrt{\frac{1}{2a} \log \frac{N}{2a \log q^{-1}}}. \quad (24)$$

This suggests that if we fix a quantile q , then in the asymptotic limit

$$\Delta_{\min}^q \sim \exp(-\text{const} \cdot \alpha^{-1/3} \cdot \sqrt{\log N}), \quad (25)$$

i.e. it is vanishing slower than any polynomial with increasing N . This has to be compared with homogeneous gap scaling as: $\Delta_{\min}^q \sim \exp(-\text{const} \cdot \sqrt{N})$, which vanishes as stretched exponential with N . We note here that similar analysis in case when $\tilde{P}(x)$ would have exponential tail for large negative x would give polynomial dependence for the minimal gap on N .

In order to fully analyze the tail of the gap distribution we also consider the situation that the minimum gap is enforced by very small local link rather than many-body gap of critical system of effective size ξ_i . This issue, which is essentially an artifact of 1D systems, can be largely avoided by a multi-front strategy where such links are placed in-between clusters which are driven quasi-adiabatically. To that end, let us assume that the links are drawn from uniform distribution $J_i \in [-1, 1]$. Probability that a single link is weaker than some ϵ is equal ϵ , or $P(|J_i| > \epsilon) = 1 - \epsilon$. Let us consider that q is the probability that all the links are stronger than this ϵ is $q = (1 - \epsilon)^N$. For small ϵ this yields

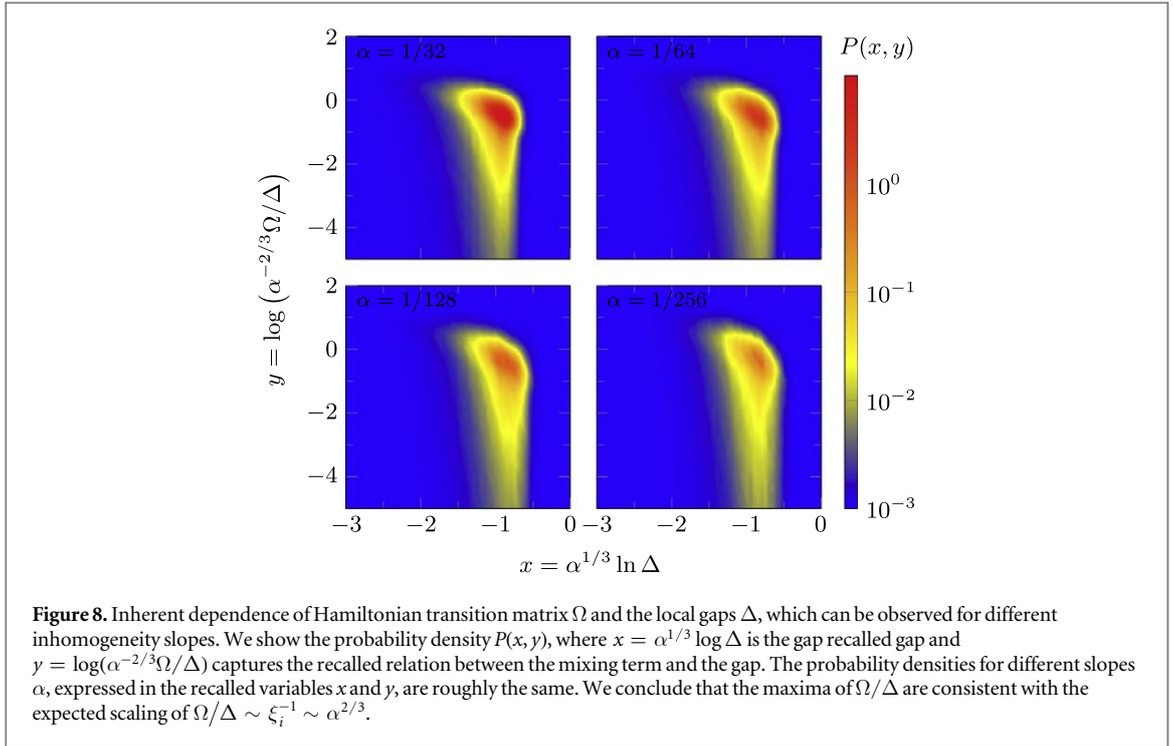
$$\epsilon \simeq -\frac{\log q}{N}.$$

The minimal gap related with such a weak link is of similar size, and such effects become relevant in 1D geometry. Finally, we should note that if we consider the distribution of the logarithm of the gap x , the uniform distribution and related weak links directly translate to the exponential tail of $P(x)$ —mentioned in the previous paragraph—which is indeed still visible in figure 7 for larger values of α .

Finally, in figure 8 we illustrate the relation between the transition matrix elements Ω and the gap Δ . To that end, for simplicity we consider protocol with single front and, as in figure 7, sample the values of minimal relevant gap and corresponding Ω as the front is traveling though the chain. The results are collected as probability distribution, where the statistics is collected from 5000 instances. In order to illustrate the universal behavior we employ rescaled variables. For the gap $x = \alpha^{1/3} \log(\Delta)$ as above, and $y = \log(\alpha^{-2/3}\Omega/\Delta)$ to reflect the expected relation $\Omega/\Delta \sim \xi_i^{-1} \sim \alpha^{2/3}$. We plot the obtained distributions of $P(x, y)$ in figure 8 for several different values of α . We observe that they are roughly similar in agreement with our prediction. We should note that local maxima of Ω/Δ —i.e. where it is most relevant—coincide with the local minima of energy gap. Apart from those point, Ω/Δ is quickly approaching zero, which is reflected by elongated shape of the distribution $P(x, y)$ in the direction of small (irrelevant) y .

⁴ This derivation corresponds to the smallest, single quasi-particle gap in the Ising model. Here due to conserved parity symmetry the relevant gap corresponds to two excited quasi-particle.

⁵ More precisely we expect the tail vanishing as $\tilde{P}(x) \sim e^{-a(x-x_0)^2}$, with some non-universal constant x_0 . We set $x_0 = 0$ for clarity of derivation as it is not relevant for the main conclusion.



4. Random cluster-Ising Hamiltonian

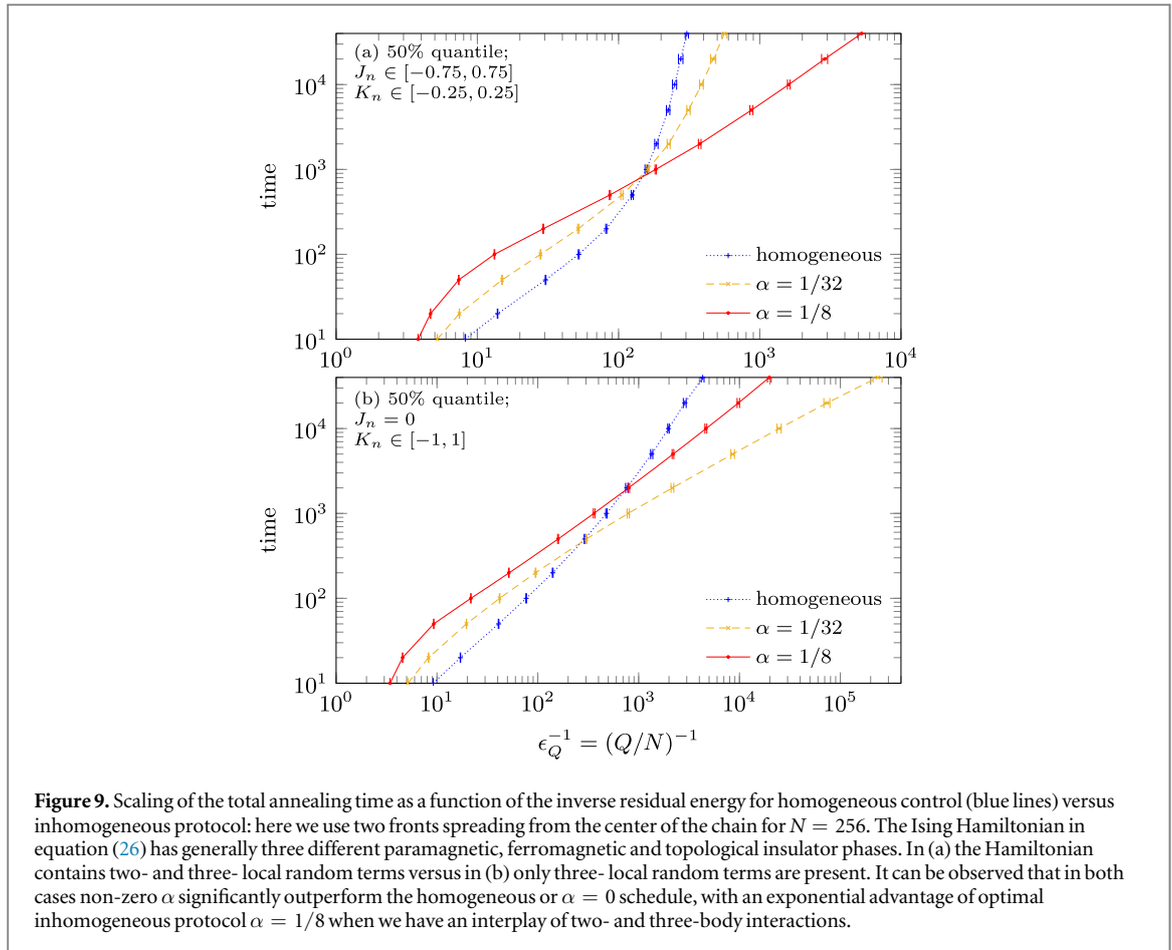
The numerical results in the previous sections were confined to 1D geometry where, in the final ground state, the neighboring spins are aligned according to corresponding $J_{n,n+1}$. In this section we show that the general approach discussed in this article does not hinge on the possibility to align nearest-neighbor interacting spins in strictly 1D Ising geometry. To that end we consider random cluster-Ising Hamiltonian as follows:

$$H(t) = - \sum_{n=1}^{N-1} J_n \sigma_n^z \sigma_{n+1}^z - \sum_{n=1}^{N-2} K_n \sigma_n^z \sigma_{n+1}^x \sigma_{n+2}^z - \sum_{n=1}^N g(n, t) \sigma_n^x \quad (26)$$

where the first two terms contains the problem Hamiltonian and the last term is the (inhomogeneous) driving term given by external transverse field. The quench dynamics generated by this Hamiltonian can be simulated analogously as for the Ising chain from the previous section as—using Jordan–Wigner transformation—it can be mapped onto a free fermionic system.

First, lets consider the pure and homogeneous model with $J_n = J > 0$, $K_n = K > 0$ and $g_n = g > 0$. Depending on the relative strength of those terms the model has 3 distinct phases, see e.g. [59–63]. It is convenient to set $K + J = 1$. When the magnetic field is dominating, $g > 1$, the system is in paramagnetic phase analogously to the Ising model. We are going to initialize the evolution in this phase. In the opposite limit, $g < 1$, the system is in ferromagnetic phase for $J + g > K$ and symmetry-protected topological order phase for $K > J + g$. For $J = g = 0$ it reduces to the cluster Hamiltonian [64]. The phases are separated by critical points with $z = 1$ or $z = 2$.

We consider random couplings J_n and K_n , which makes the final target state far from being trivial for a 256 spin system. We present the results of the quench in figure 9. We consider two different disorder distributions: in figure 9(a) J_n are dominating—that is $J_n \in [-0.75, 0.75]$ and $K_n \in [-0.25, 0.25]$, and in figure 9(b) all $J_n = 0$ and $K_n \in [-1, 1]$. For homogeneous driving the residual energy is vanishing logarithmically with the quench time in both cases, which is a similar behavior as for the random Ising model. It can be observed that there is crossover of the performance for sufficiently long times and the homogeneous protocol is considerably outperformed by an inhomogeneous driving fields with the optimal slope. The advantage for the case of random J_n could be exponential, see figure 9(a). In this case, the spatial inhomogeneity allows the system to reach the quality of solution (small residual energies) which are practically unattainable within the homogeneous approach. Here, we use a version with single cluster and two critical fronts spreading from the center of the chain. It should be noted that the optimal shapes (α) in those two cases are different. This optimal value of α can be found numerically for given distribution of disorders as a simple preprocessing, or hyper-parameter



characterization, similar to the spirit of finding the optimal annealing time or number of sweeps for simulated annealing or quantum Monte Carlo solvers.

5. Conclusion and future works

We have presented a model for engineering quantum phase transitions in disordered systems by manipulating information flow among clusters that are formed within a quantum critical region. We have shown that space-time inhomogeneities in the control fields could lead to reconstruction of causal zones (light cones), such that symmetry breaking events can be synchronized suppressing the density of topological defects and/or redistributing their spatial arrangements. We have used exact diagonalization techniques for 1D systems to show an exponential speedup of non-adiabatic IQA over standard adiabatic quantum computing, even in the presence of higher order interactions. By application of renormalization group techniques we have demonstrated that the effective causal gaps exhibit universality with respect to the shape of inhomogeneity. We have derived a scaling relation showing such effective gaps have sub-polynomial scaling with the system size, in contrast to stretch exponential for homogeneous control strategies. In a subsequent work [17], we will provide a detailed theoretical discussion of our work as a generalization of KZM for disordered systems including various bounds for the shape of critical fronts and threshold velocities under different assumptions. We will also discuss how our approach can be applied to low-dimensional spin-glass problem Hamiltonians [16, 17]. During the preparation of this manuscript an exponential speedup for IQA of p-spin model was reported showing ferromagnetic first-order phase transitions can be smeared with inhomogeneous control strategies [65].

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