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# **Quantum resources in quantum technologies:**

## **From quantum optimization to many-body physics**

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*To Thee, the Constant through all variables, —*

*Whom I pursue, Who art both the Way and the Destiny;*

*This thesis is dedicated unto Thee.*

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cense.



## Abstract

Significant progress has been made in quantum technologies over the past decade. Despite this success, a central question remains: to what extent quantum resources are being exploited. This question is crucial not only for benchmarking quantum systems but also to unlock their full potential. This thesis analyses the role of quantum resources spanning quantum optimization algorithms to quantum many-body systems. First, we show how squeezing establishes a natural connection between the Quantum Approximate Optimization Algorithm and quantum metrology, revealing the role of quantum correlations and providing a benchmarking method for quantum-optimization devices. Further, we study multipartite entanglement in quantum optimization and demonstrate its presence on quantum hardware. However, entanglement alone is insufficient for a quantum advantage, as stabilizer states—though highly entangled—are classically simulable. We therefore study the role of nonstabilizerness in quantum optimization. Building on such resources, we further examine how they manifest in disordered systems. We show the emergence of non-Markovianity in disorder-averaged dynamics. Moreover, we examine complexity-resources across chaotic to integrable regimes in random-matrix models, highlighting the need for a multifaceted approach in quantum simulation. These investigations provide a deeper understanding of quantum technologies, from optimization to simulation, and lay the foundation for future developments toward quantum advantage.



## Zusammenfassung

In den letzten zehn Jahren wurden bei den Quantentechnologien erhebliche Fortschritte erzielt. Trotz dieses Erfolges bleibt eine zentrale Frage bestehen: In welchem Umfang werden die Quantenressourcen genutzt? Diese Frage ist nicht nur für das Benchmarking von Quantensystemen von entscheidender Bedeutung, sondern auch für die Erschließung ihres vollen Potenzials. In dieser Arbeit wird die Rolle von Quantenressourcen analysiert, die von Quantenoptimierungsalgorithmen bis hin zu Quanten-Vielteilchensystemen reichen. Zunächst zeigen wir, wie Squeezing eine natürliche Verbindung zwischen dem Quantum Approximate Optimization Algorithm und der Quantenmetrologie herstellt, indem wir die Rolle von Quantenkorrelationen aufdecken und eine Benchmarking-Methode für Quantenoptimierungsgeräte bereitstellen. Darüber hinaus untersuchen wir die mehrteilige Verschränkung in der Quantenoptimierung und demonstrieren ihr Vorhandensein auf Quantenhardware. Verschränkung allein reicht jedoch nicht aus, um einen Quantenvorteil zu erzielen, da Stabilisatorzustände — obwohl sie stark verschränkt sind — klassisch simulierbar sind. Wir untersuchen daher die Rolle der Nicht-Stabilisierung in der Quantenoptimierung. Aufbauend auf diesen Ressourcen untersuchen wir, wie sie sich in ungeordneten Systemen manifestieren. Wir zeigen das Auftreten von Nicht-Markowianität in unordnungsgemittelter Dynamik. Darüber hinaus untersuchen wir Komplexitätsressourcen in chaotischen bis hin zu integrierbaren Regimen in Zufallsmatrixmodellen und unterstreichen damit die Notwendigkeit eines vielschichtigen Ansatzes in der Quantensimulation. Diese Untersuchungen ermöglichen ein tieferes Verständnis der Quantentechnologien, von der Optimierung bis zur Simulation, und legen den Grundstein für zukünftige Entwicklungen in Richtung Quantenvorteil.





## Sommario

Negli ultimi dieci anni sono stati compiuti notevoli progressi nelle tecnologie quantistiche. Nonostante questi successi, rimane una domanda centrale: in che misura le risorse quantistiche vengono effettivamente sfruttate? Tale quesito è cruciale non solo per il benchmarking dei sistemi quantistici, ma anche per sbloccarne appieno il potenziale. Questa tesi analizza il ruolo delle risorse quantistiche in contesti che spaziano dagli algoritmi di ottimizzazione quantistica ai sistemi a molti corpi. In primo luogo, mostriamo come lo squeezing stabilisca una connessione naturale tra l'algoritmo di ottimizzazione approssimata quantistica (QAOA) e la metrologia quantistica, rivelando il ruolo delle correlazioni quantistiche e fornendo al contempo un metodo di benchmarking per i dispositivi di ottimizzazione quantistica. Successivamente, studiamo l'entanglement multipartito nell'ottimizzazione quantistica e ne dimostriamo la presenza nell'hardware quantistico. Tuttavia, l'entanglement da solo non è sufficiente a garantire un vantaggio quantistico: gli stati stabilizzatori, infatti, pur essendo altamente entangled, sono simulabili in modo classico. Analizziamo quindi il ruolo della non-stabilizzabilizzazione nell'ottimizzazione quantistica. Partendo da queste risorse, ci spingiamo oltre a esaminarne le manifestazioni nei sistemi disordinati. Mostriamo l'emergere della non-markovianità nelle dinamiche mediate sul disordine e analizziamo le risorse di complessità attraverso regimi caotici e integrabili in modelli a matrici aleatorie, mettendo in luce la necessità di un approccio multiforme alla simulazione quantistica. Queste indagini forniscono una comprensione più approfondita delle tecnologie quantistiche, dall'ottimizzazione alla simulazione, e pongono le basi per futuri sviluppi verso il pieno raggiungimento del vantaggio quantistico.



## সারাংশ

গত দশকে কোয়ান্টাম প্রযুক্তিতে অনেক উল্লেখযোগ্য অগ্রগতি হয়েছে। এত সাফল্য সত্ত্বেও, একটি মূল প্রশ্ন রয়েই গেছে: সত্যিকারের কতটা কাজে লাগান হচ্ছে কোয়ান্টাম সম্পদকে? এই প্রশ্নটি কেবল কোয়ান্টাম সিস্টেমের বেঞ্চমার্কিংয়ের জন্যই নয়, বরং তাদের পূর্ণ সম্ভাবনা উন্মোচনের জন্যও গুরুত্বপূর্ণ। এই থিসিসটি কোয়ান্টাম অপ্টিমাইজেশন অ্যালগরিদম থেকে শুরু করে কোয়ান্টাম বহু-বস্তু-সম্বন্ধীয় পদার্থবিদ্যা পর্যন্ত বিভিন্ন ব্যপারে কোয়ান্টাম সম্পদের ভূমিকা বিশ্লেষণ করেছে। প্রথমে, আমরা দেখাই কিভাবে squeezing, কোয়ান্টাম আনুমানিক অপ্টিমাইজেশন অ্যালগরিদম এবং কোয়ান্টাম মেট্রোলজির মধ্যে একটি স্বাভাবিক সংযোগ স্থাপন করে— একদিকে কোয়ান্টাম correlation সম্পর্কে ধারণা দেয়, আবার অন্যদিকে কোয়ান্টাম-অপ্টিমাইজেশন যন্ত্রের জন্য একটি বেঞ্চমার্কিং পদ্ধতিও আমাদেরকে দেয়। এরপরে, আমরা কোয়ান্টাম অপ্টিমাইজেশনে multipartite entanglement- এর ভূমিকা অধ্যয়ন করি এবং কোয়ান্টাম হার্ডওয়্যারে এর উপস্থিতি প্রদর্শন করি। তবে, কোয়ান্টাম প্রযুক্তির শ্রেষ্ঠত্ব প্রদর্শনের জন্য কেবল entanglement থাকলেই চলে না, কারণ এক ধরনের অবস্থা আছে, স্টেবিলাইজার অবস্থা নামের, যারা অত্যন্ত entangled হওয়া সত্ত্বেও, খুবই সহজেই সিমুলেশন করা যায়। তাই আমরা কোয়ান্টাম অপ্টিমাইজেশনে non-stabilizerness-এর ভূমিকাও অধ্যয়ন করি। এই ধরনের সম্পদের উপর ভিত্তি করে, আমরা আরও পরীক্ষা করি যে তারা বিশৃঙ্খল অবস্থায় কীভাবে প্রকাশ পায়। আমরা এই বিশৃঙ্খল সিস্টেমের মধ্যে থেকে সময়ের সাথে কিভাবে non-Markovianity-এর (অতীতকে মনে রাখা) উত্থান হতে পারে সেটা দেখাই। এরপরে, আমরা বিভিন্ন random-ম্যাট্রিক্স মডেলে বিশৃঙ্খল থেকে সংহত অবস্থা পর্যন্ত জটিলতা-সম্পর্কিত সম্পদদের পরীক্ষা করি, যা থেকে বোঝা যায় যে কোয়ান্টাম সিমুলেশনে বহুমুখী পদ্ধতির প্রয়োজন আছেই। এই তদন্তগুলি অপ্টিমাইজেশন থেকে সিমুলেশন পর্যন্ত কোয়ান্টাম প্রযুক্তি সম্পর্কে গভীর ধারণা প্রদান করে এবং ভবিষ্যতে কিভাবে এই প্রযুক্তি নতুন সম্ভাবনা এনে দিতে পারে, সেই উন্নয়নের ভিত্তিপ্রস্তর স্থাপন করে।



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## Chapter 1

# Introduction

In the last decades, quantum technologies have rapidly evolved, transforming quantum computation and simulation from a theoretical pursuit into an experimental reality [1–8]. This transformation has been propelled by continual progress in qubit coherence, scalable architectures, and the emergence of quantum error correction techniques across multiple hardware platforms, such as superconducting qubits [9–12], trapped ions [13–15], neutral atoms [16, 17], and photonic systems [18]. This technological momentum is matched by global enthusiasm: governments and private industries alike are investing heavily in quantum research, motivated by its potential to revolutionize fields such as cryptography [19], materials discovery [20], drug design [21], and optimization [22]. At the same time, academic research is broadening the scope of quantum simulation, aiming to tackle problems that lie beyond the reach of classical computation. For example, in condensed matter physics, quantum simulators are being explored to simulate non-equilibrium dynamics and quantum phase transitions [23–26]; in high-energy physics, to study real-time dynamics and non-perturbative effects in lattice gauge theories [27–33]; in quantum chemistry, to model molecular interactions and reaction mechanisms [34, 35], or in the realms of quantum gravity and quantum chaos, to understand holographic dualities and information scrambling [36–38]. Together, these developments illustrate not only the versatility of quantum technologies but also the diverse and growing landscape of quantum applications, from fundamental physics to real-world impact.

Despite such remarkable progress, skepticism about quantum technologies persists, driven by challenges such as noisy hardware, limited coherence time, the significant overhead of error correction, or the lack of broadly applicable quantum algorithms that can outperform classical methods. At the heart of this skepticism lies a key question: *Do quantum effects truly drive the observed results, or are they merely artifacts of noise, offering no clear advantage over classical simulation?* In response, the quantum community has developed rigorous benchmarking techniques [39–41] to address this ambiguity. Identifying and isolating truly quantum signatures is critical for validating devices [42, 43] and assessing their real-world utility [44, 45]. Beyond verification, active efforts focus on developing new algorithms, error mitigation strategies [46], and resource theories [47] to better harness and understand quantum resources. This deeper understanding clarifies the realistic potential of quantum technologies, helps set achievable milestones, and inspires confidence in the field’s progress toward transformative applications.

To understand which quantum resources are essential and how to harness them effectively, it helps to revisit the key milestones that have shaped the development of quantum technology and our understanding of its fundamental phenomena. Quan-

tum mechanics emerged in the early 20th century as a framework to describe microscopic phenomena that classical theories could not explain. Pioneering work by scientists such as Planck, Einstein, Heisenberg, Schrödinger, Dirac, Born, Bohr, von Neumann [48–59], and provoking key experiments such as Davisson-Germar’s and later, Jönsson’s double-slit interference experiments [60, 61], fundamentally reshaped physics and enabled transformative technologies like transistors, lasers, and atomic clocks. Riding on the concepts developed during this first quantum revolution, the second revolution arrived to transform the quantum theory of fundamental curiosity into practical engineering and technology. By recognizing the limitations of classical computers for simulating quantum phenomena, visionaries like Manin and Feynman proposed the idea of quantum simulators [62, 63]. As Feynman famously stated, “Nature isn’t classical, dammit, and if you want to simulate it, you’d better make it quantum” [64]. Concurrently, quantum information theory introduced protocols such as the BB84 protocol for cryptography [65], and Shor’s algorithm [66] for factoring large numbers. Experimental platforms have advanced steadily to realize these theoretical concepts in practice [67–70]. Later, the development of quantum error correction and fault-tolerance schemes [71, 72] shifted the focus from entanglement to magic states as the central resource for robust quantum computation. Alongside these developments, the tensor network methods [73, 74] have also advanced for the simulation of many-body quantum systems, complementing the growth of quantum technologies by enabling classical simulations that can benchmark experimental efforts in understanding complex quantum phenomena.

In the presence of such a variety of theoretical frameworks, numerical techniques, and experimental platforms, with their respective strengths and limitations, focusing on a single approach or perspective is inadequate to show quantum advantage [45, 75, 76]. For example, Google’s 2019 quantum supremacy claim [11] demonstrated that their 53-qubit Sycamore processor could sample from a random quantum circuit in about 200 seconds—a task believed to take classical supercomputers thousands of years. However, the problem was artificial and specifically chosen to favor quantum devices. And yet, subsequent classical algorithms [77–80] were able to show a reduced simulation time, challenging the original supremacy claim. Another notable case is that Farhi *et al.* introduced the Quantum Approximate Optimization Algorithm (QAOA) as a promising method for problems like MaxCut [81]. Initial results suggested potential quantum advantage, but subsequent improvements in classical algorithms challenged these claims [82, 83], highlighting the evolving competition between quantum and classical approaches. These examples serve as a clear reminder that reaching quantum advantage is not straightforward; it calls for broad exploration and inventive use of quantum resources.

Throughout this thesis, we investigate the multifaceted nature of quantum resources [47] both theoretically and experimentally with a central goal to understand how they are harnessed in contexts ranging from quantum optimization to many-body physics. Quantum resources provide a rigorous framework for distinguishing between what is considered easy and what is considered hard (expensive), based on the identification of free and resourceful operations within a given resource theory. This unified perspective enables us to identify which features of a quantum state or process can be exploited to gain an advantage over classical alternatives. In this quest, the classical paradigm serves as a free resource; by identifying what is tractable within classical computation or simulation, we can define appropriate resource theories and

---

corresponding measures that highlight the power of quantum systems. For instance, classical systems can easily represent product states, but struggle with superposition, leading to the formulation of entanglement as a quantum resource [84]. Similarly, stabilizer states can be efficiently simulated classically, whereas non-stabilizer states cannot, thereby motivating the resource theory of nonstabilizerness [85]. Thus, quantum resources can be viewed as a unifying framework to understand quantumness that helps to assess the current capabilities of quantum systems and their potential to surpass classical approaches, which is discussed in Chapter 2.

Within quantum computation, quantum optimization [22] stands out as particularly crucial. Combinatorial optimization problems [86] are widespread across science and industry, so any quantum improvement over state-of-the-art classical algorithms can have a profound impact. This makes quantum optimization one of the most promising pathways for realizing the practical value of quantum technologies. We discuss in detail the complexity of optimization problems and how quantum algorithms may tackle such issues in Chapter 3. Given the limited capability of current quantum devices [87], and the remarkable ability of existing classical computers, hybrid algorithms are emerging as a potential test-ground for quantum advantage, for example, the Quantum Approximate Optimization Algorithm (QAOA) [81]. However, despite its successful applications, QAOA depends heavily on classical optimization routines. This raises an important question: *to what extent do such hybrid algorithms truly harness quantum resources?* Although one may think that practical performance is all that matters, understanding the role of quantum resources is crucial for achieving genuine quantum advantage, especially in predicting its capability on large-scale problems that exceed currently accessible benchmarks. Without leveraging the aspects of quantum mechanics that make the computation classically intractable, the algorithm may ultimately fail to deliver a true quantum advantage. In a more quantum information way, it is about asking a fundamental question: *when a quantum algorithm outperforms its classical counterpart, what quantum resource underpins that advantage? Is it entanglement, nonstabilizerness, or some other feature of quantumness? And more importantly, does our current quantum hardware possess the capacity to harness these resources? If not, how can such a quantum effect be introduced into the hardware?* This is some of the underlying quests of this thesis for Part II (Chapter 4-7), where we explore resources in quantum optimization.

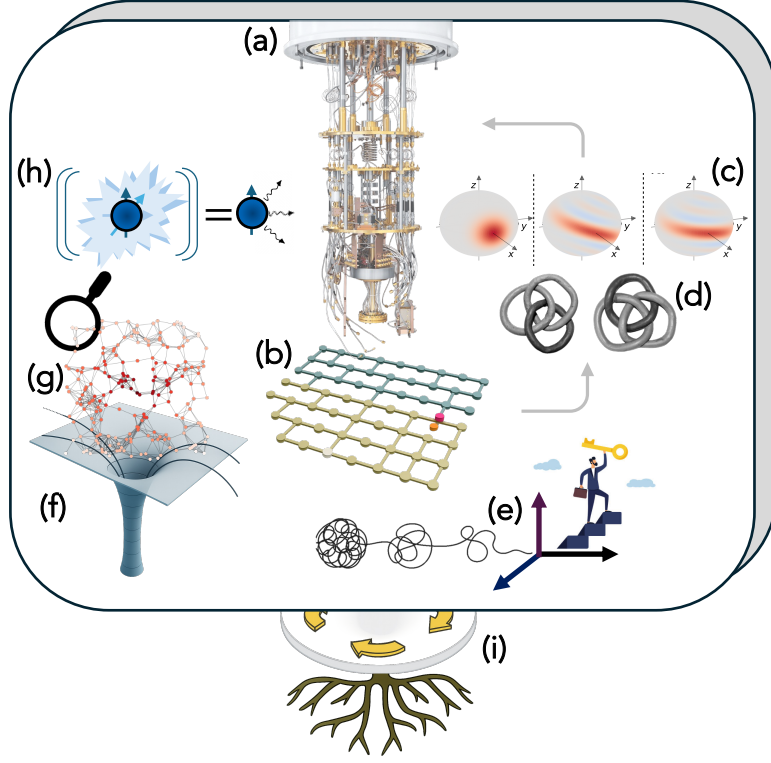
In Chapter 4, we establish a connection between quantum optimization and quantum metrology via squeezing to understand the role of quantum correlations in QAOA. We also introduce a hardware benchmarking technique based on spin-squeezing. To probe the necessity of multipartite entanglement—known to be essential for quantum computational speed-up [88]—we examine genuine multipartite entanglement in quantum annealing and uncover its correlation with success probability in Chapter 5. We further validate the presence of such entanglement experimentally on IBM’s 156-qubit quantum computer in Chapter 6. Recognizing that entanglement alone does not capture all aspects of quantum advantage, we turn our attention to nonstabilizerness in Chapter 7, studying its role in both QAOA and quantum annealing and its relationship to algorithmic fidelity.

In the realm of quantum technologies, quantum simulation [89, 90] forms another fundamental block. Its goal is to simulate highly complex physical systems, providing a powerful alternative to classical approaches. This is particularly relevant in regimes where traditional classical methods, such as tensor networks [91] or Monte

Carlo simulations [92], become inapplicable. This can also be thought of as the native and most natural application of quantum computers, where we aim to use a quantum computer to mimic the rules that describe physical microscopic quantum systems [75]. Analog quantum simulators have already yielded significant insights into many-body physics [16, 24, 93], showcasing the potential of quantum devices to explore otherwise intractable models. One of the aims of quantum simulation is to push these boundaries further, tackling increasingly complex problems. In this case, our goal is to assess the extent to which quantum resources are being utilized in quantum simulation of complex systems, particularly in the presence of disorder. In Part III (Chapter 8-9), we investigate how quantum resources manifest in disordered systems, probing whether such scenarios truly demand quantum computational power or if we are merely applying a quantum solution to problems that may not fundamentally require it.

In Chapter 8, we examine the effect of disorder in quantum dynamics, a ubiquitous feature arising from imperfections in fabrication, control, or the environment of the quantum device. We develop an analytical framework that enables the exact calculation of disorder-averaged dynamics for a broad class of quantum systems governed by periodic Hamiltonians. In Chapter 9, we explore whether different complexity markers, such as entanglement entropy, fractal dimension, and stabilizer Rényi entropy, respond uniformly when transitioning between chaotic and integrable regimes, or if there exist intermediate phases where these markers diverge, labeling the same quantum state as resourceful in one context and free in another.

The chapters in this thesis were developed as independent projects, each addressing specific questions within the field of quantum technologies. Despite this, a unifying theme emerges when viewed through the lens of quantum resources. By identifying conceptual similarities—such as the role of entanglement, nonstabilizerness, and other resource measures in optimization, as well as in disordered systems—we bring together diverse results under a common framework. These connections are visually represented in the cover illustration (Fig. 1.1), with further details provided in the caption.



**Figure 1.1. Cover illustration:** Visual representation of the thesis: (a) A superconducting quantum computer (image taken from [94]) with (b) qubit layouts representing Chapter 6, where the quantum circuit is being run (motivated from Fig. 6.16, [95]), to understand the resource requirements of the hardware part of quantum optimization, either (c) via squeezing (see Chapter 4) or magic (see Chapter 7) generated in the process of QAOA, or (d) multipartite entanglement in quantum annealing (see Chapter 5) (image taken from [96]). (e) A recurring theme throughout the thesis is the exploration of how various quantum resources contribute to optimization success (right side) and relate to the complexity of chaotic systems (left side) [95]. (f) An artist’s impression of the spacetime around a black hole, showing the strong curvature due to the high gravitational mass [95]. (g) The Sachdev-Ye-Kitaev model at low-temperature known to share similar chaotic behavior as a black hole [95, 97], is shown as a cartoon model of randomly interacting particles (Chapter 9), which can also represent the spin-glass model (Chapter 7) (image taken from [98]). (h) Among all these particles, consider a single particle under disorder, and in Chapter 8, we observe how a closed disorder system behaves like an open system. (i) All such analyses on resources, benchmarking, and complexity in quantum technology, from quantum optimization to quantum many-body physics, are rooted in the quantum informational notion of quantum resources (Chapter 2) (figure motivated from [99]).





# **Part I**

## **Background**



## Chapter 2

# Quantum resources

### *unification of quantumness*

---

Quantum mechanics provides a framework to describe physical phenomena that often defy classical explanations. Concepts such as entanglement have no classical analogs and only become meaningful in the quantum regime. In this light, it becomes both natural and necessary to ask: what exactly makes quantum systems different and possibly resourceful? Identifying and characterizing the aspects of physical systems that enable such distinction leads to the notion of quantum resources. This perspective not only deepens our understanding of quantum phenomena but also guides the development of applications by framing them within the unifying structure of quantum resource theories. This process can sometimes reveal insights beyond what could have been gained by studying the phenomena individually and allows us to exploit such properties as resources in quantum technological applications.

For example, quantum entanglement was once a topic of philosophical debate during the first quantum revolution. However, once recognized as a valuable resource, it became the essential driving force behind numerous quantum information tasks, e.g., quantum simulation, quantum computation, quantum metrology, quantum communication, quantum cryptography, etc., in the second quantum revolution.

Given the success of the resource theory of entanglement, it is natural to explore other physical phenomena that can also be recognized as valuable resources. Beyond well-understood bipartite entanglement, this thesis will consider multipartite entanglement, quantum magic or nonstabilizerness (in the context of quantum computation), quantum Fisher information, and non-Markovianity, all of which will be discussed in detail in this chapter. The classic review by Chitambar and Gour [47] serves as our primary reference.

## 2.1 General structure of resource theory

Basic economic principles suggest that an object gains value when it is not readily accessible [47]. For instance, diamonds are deemed valuable precisely because they are rare in nature. In this way, value is inherently a relative concept. A resource theory builds on this idea by identifying which operations are considered free or restrained in a given context and then examining what can be achieved under those constraints. Any object that cannot be created using only free operations is then classified as a resource.

Each quantum resource theory is structured upon two objects: free states and free operations [47]. Free states are those quantum states that can be ‘easily’ created within the defined framework, and free operations represent quantum manipulations that can be effortlessly carried out. Thus, it does not encompass all quantum mechan-

ically allowed physical processes, and only certain states can be prepared freely. Any states that are not free are then called a ‘resource’ state. In this way, by identifying the free states and free operations, resource theory also classifies every quantum state as either free or a resource. To compare with a real-life example, see Fig. 2.1.

### Quantum resource theory

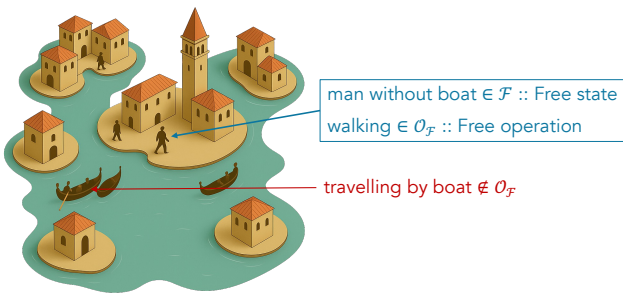
**Definition 1.** The pair  $\mathcal{R} = (\mathcal{F}, \mathcal{O}_{\mathcal{F}})$ , consisting of free states ( $\mathcal{F}$ ) and a set of completely positive and trace-preserving (CPTP) operations<sup>a</sup> ( $\mathcal{O}_{\mathcal{F}}$ ), known as free operation, is called a *quantum resource theory* (QRT) if the free operations map any free state to another free state,

$$\Lambda : \rho \in \mathcal{F} \mapsto \tilde{\rho} \in \mathcal{F}, \quad \forall \Lambda \in \mathcal{O}_{\mathcal{F}}. \quad (2.1)$$

<sup>a</sup>See Sec. 2.8.1 for the definition of CPTP operations.

Definition 1 is considered the golden rule of QRT. However, in practice, one has to understand the details that are hidden behind such an intuitive definition. For example, the state  $\rho$  and  $\tilde{\rho}$  may belong to two different systems. In general,  $\mathcal{O}_{\mathcal{F}}$  maps two physical system  $A$  and  $B$ , with corresponding Hilbert spaces  $\mathcal{H}_A$  and  $\mathcal{H}_B$ , i.e.,  $\mathcal{O}_{\mathcal{F}}(A \rightarrow B) \equiv \mathcal{O}_{\mathcal{F}}(\mathcal{H}_A \rightarrow \mathcal{H}_B) \subset \mathcal{Q}(A \rightarrow B)$ , where  $\mathcal{Q}$  denotes the set of all quantum channels from  $A$  to  $B$ . From the def. 1, one can ensure that (1)  $\mathcal{O}_{\mathcal{F}}(A \rightarrow A)$  contains the identity map  $\text{id}^A$ , i.e., doing nothing is free, and (2) if  $\Phi(A \rightarrow B) \in \mathcal{O}_{\mathcal{F}}$ , and  $\Lambda(B \rightarrow C) \in \mathcal{O}_{\mathcal{F}}$ , then  $\Lambda \circ \Phi(A \rightarrow C) \in \mathcal{O}_{\mathcal{F}}$ , i.e., the free operations can be performed in composition as many times as one wants without any cost. Thus, there is no way free operations can create a resource state.

In a QRT, the set  $\mathcal{F}(\mathcal{H}) \subset \mathcal{S}(\mathcal{H})$  are thus *free states*, and  $\mathcal{S}(\mathcal{H}) \setminus \mathcal{F}(\mathcal{H})$  are called *resource states* or *static resources*, where  $\mathcal{S}(\mathcal{H})$  are the set of valid density matrices on  $\mathcal{H}$ . Similarly, as  $\mathcal{O}_{\mathcal{F}}$  is called *free operation*, and the rest of the CPTP maps are called *dynamical resources*.



**Figure 2.1.** Island of Freedom, and Canal of Cost. A man can walk freely within his domain, operating without constraint. But beyond his island lies potential he cannot reach unaided. To cross the water, he must acquire a resource: a boat, and perform a resource-bound operation: rowing. Here, the resource theory  $\mathcal{R} = (\text{man without boat}, \text{walking})$ .

In the experimental scenario, free operations are considered more fundamental, despite the equal footing of both free states and free operations in the definition of a QRT. Because, for an experimentalist, even the free states are not provided for granted, and hence they must be able to prepare the initial free states using free operations. Mathematically also, the free states are an induced mapping of free operations, i.e.,  $\mathcal{F}(\mathcal{H}) := \mathcal{O}_{\mathcal{F}}(\mathbb{C} \rightarrow \mathcal{H})$ . However, in the theoretical sense, multiple free operations can be considered given the same free states, and thus, in a QRT, one needs the pair to define a concrete theory. Although the definition of a QRT does not include the definition of a resource state, it is indirectly implied through it. There are

several ways to have a resourceful operation. For example, a free operation together with a resource state ( $\sigma$ ) can become a resourceful operation (dynamical resource) on a free state, i.e., given  $\sigma \notin \mathcal{F}(B)$ , there may exist free operation  $\Phi \in \mathcal{O}_F(AB)$ , such that  $\Phi(\rho \otimes \sigma) = \Lambda(\rho)$ , where  $\Lambda \notin \mathcal{O}_F(A)$ , although  $\rho \in \mathcal{F}$ . One well-known example is quantum teleportation ( $\Lambda$ ) with entangled state ( $\sigma$ ) [100].

Often in quantum applications, one expects to amplify the quantum effects by considering a bigger system size. However, the isomorphism  $\mathbb{C}^2 \otimes \mathbb{C}^2 \cong \mathbb{C}^4$  does not translate into the same set of free states or operations. For example, a system of 2 qubits, although it has the same Hilbert space dimension (i.e., 4) as 1 ququart, can have a different QRT compared to a 1 ququart. On the other hand, within the tensor-product structure, one can relabel the Hilbert space, i.e., the density matrices acting on  $\mathcal{H}_A \otimes \mathcal{H}_B$  represent the same physical states as the density matrices acting on  $\mathcal{H}_B \otimes \mathcal{H}_A$ . As most operations in quantum mechanics can be represented by the tensor-product structure, it is crucial to understand when a QRT can also follow this structure.

### Tensor-product structure

**Definition 2.** A QRT  $\mathcal{R} = (\mathcal{F}, \mathcal{O}_F)$  admits a *tensor-product structure* if

1.  $\mathcal{O}_F$  is ‘completely free’, i.e., it remains free when acted on just one part of any joint system:  $\Phi \in \mathcal{O}_F(A \rightarrow B) \Rightarrow \text{id}^C \otimes \Phi \in \mathcal{O}_F(CA \rightarrow CB)$ , where  $\text{id}^C$  is the identity map on  $C$ .
2. Appending free states is a free operation:  $\Phi_\sigma(\rho) := \rho \otimes \sigma \in \mathcal{O}_F \quad \forall \rho \in \mathcal{F}$ .
3. Discarding a system is a free operation:  $\mathcal{O}_F(\mathcal{H} \rightarrow \mathbb{R})$  is not empty.

**Corollary 1.** The tensor-product structure has several important consequences:

- (1)  $\Rightarrow$  If  $\Phi, \Phi' \in \mathcal{O}_F$ , then  $\Phi \otimes \Phi' = (\text{id} \otimes \Phi') \circ (\Phi \otimes \text{id}) \in \mathcal{O}_F$ .
- (3)  $\Rightarrow$  Trace of a system is a free map.
- (1) + (3)  $\Rightarrow$  Partial trace  $\text{Tr} \otimes \text{id}$  is free.
- (2) + partial trace  $\Rightarrow$  If  $\rho, \sigma, \rho \otimes \sigma \in \mathcal{F}$ , then  $\rho$  can be converted to  $\sigma$  by free operation, and vice-versa.
- Inter-convertibility + (1)  $\Rightarrow$  If  $\rho, \sigma \in \mathcal{F}$ , then  $\rho \otimes \sigma \in \mathcal{F}$ . This implies  $\mathcal{F}(A) \otimes \mathcal{F}(B) \subset \mathcal{F}(AB)$ .
- Partial trace is free  $\Rightarrow$  If  $\rho^{AB} \in \mathcal{F}$ , then  $\rho^A, \rho^B \in \mathcal{F}$ .

Most physically motivated QRTs follow a tensor-product structure, such as entanglement, coherence, etc. However, Bell non-locality may not admit such a structure, for example, even if  $\rho, \sigma \in \mathcal{F}$ ,  $\rho \otimes \sigma$  may not [101].

## 2.2 Types of resource theories

In addition to the definition of QRTs, these theories may also possess extra mathematical structures. Theories that share a similar structure can utilize common mathemati-

cal tools and physical insights. We briefly mention some of these here for completeness, as the thesis explores some examples of them in more detail later.

**Convex resource theories.** A QRT is called *convex*, if the free operation  $\mathcal{O}_{\mathcal{F}}$  is convex, i.e.,

$$\forall \Phi, \Lambda \in \mathcal{O}_{\mathcal{F}} \Rightarrow p\Phi + (1-p)\Lambda \in \mathcal{O}_{\mathcal{F}}, \quad p \in [0, 1]. \quad (2.2)$$

This also implies that the set of free states is convex. Example- QRT of entanglement, coherence, asymmetry. However, sometimes convex resource theories can be defined through states, i.e., if the free states are convex.

**Non-convex resource theories.** An example is the QRT of non-Gaussianity in continuous variable quantum systems, where the sets of Gaussian states and Gaussian operations are not convex [102].

## 2.3 Quantifying through resource measures

Besides the significant effort involved in generating resource states or implementing resource operations, it is equally important to verify whether the quantum system exhibits the desired quantum features. A key challenge often lies in quantifying the available resources, which is crucial for drawing meaningful conclusions about the system. In this context, QRT offers various methods for constructing such measures. We will discuss some of these approaches in this section, as they provide the foundation for the measures used in our research analysis.

### 2.3.1 Axiomatic approach

We first discuss some necessary and desirable properties that the resource measures should satisfy. A true measure should be able to quantify the resource of a density operator irrespective of its Hilbert space dimension in terms of a non-negative real value, and thus, the measure  $f : \mathcal{S}(\mathcal{H}) \rightarrow \mathbb{R}_{\geq 0}$ . However, such system-size independence can be relaxed, and often one can work on a fixed dimension and rescale it to compare different dimensions, such as the density of some measure. The two essential requirements are:

**Vanishing for free states.** Free states has ‘no resource’:

$$\rho \in \mathcal{F} \Rightarrow f(\rho) = 0. \quad (2.3)$$

The converse is called *faithfulness*, i.e.,  $f(\rho) = 0$  implies  $\rho$  is free. Although it is appealing to be considered in the axiom, a certain resource state  $\sigma \notin \mathcal{F}$  may have no operational advantage over a free state, and one would like to attribute  $f(\sigma) = 0$ , thus not considered as essential. For example, distillable entanglement vanishes for all bound entangled states [103].

**Monotonicity.**  $f$  cannot increase under free operation:

$$f(\rho) \geq f(\Phi(\rho)), \quad \forall \rho \in \mathcal{S}(\mathcal{H}), \quad \Phi \in \mathcal{O}_{\mathcal{F}}. \quad (2.4)$$

This is a direct implication of Def. 1. Besides the free operation, sometimes one would like to consider that quantum measurement also does not increase the measures in some QRTs. For example, in QRTs of entanglement and magic, quantum measurement in the form of quantum-classical (QC) maps  $\Phi(\cdot) = \sum_i \Phi_i(\cdot) \otimes |i\rangle \langle i|$  is physically allowed. Such a function  $f$  is called

$$\text{convex linear} \Leftrightarrow f\left(\sum_i p_i \sigma_i \otimes |i\rangle \langle i|\right) = \sum_i p_i f(\sigma_i \otimes |i\rangle \langle i|), \quad \forall \sigma^{QC} = \sum_i p_i \sigma_i \otimes |i\rangle \langle i|. \quad (2.5)$$

The von Neumann entropy [Eq. (2.35)] has the property. Given the  $f$  is convex linear, then monotonicity [Eq.(2.4)] would further implies,

$$f(\rho) \geq \sum_i p_i f(\sigma_i \otimes |i\rangle \langle i|), \quad (2.6)$$

i.e., non-increasing on average under any flagged-outcome quantum measurement (projective measurement). Now, if appending and discarding classical flags is also a free operation (e.g., tensor product structure) in the QRT, then  $f(\rho_i) = f(\rho_i \otimes |i\rangle \langle i|)$ , and then Eq. (2.6) would become

$$f(\rho) \geq \sum_i p_i f(\sigma_i), \quad \forall \rho \mapsto \sum_i \Phi_i(\rho) \otimes |i\rangle \langle i|. \quad (2.7)$$

By rewriting the same equation, one finds [104],

$$\text{Strong monotonicity : } f(\rho) \geq \sum_i \text{Tr}[\Phi_i(\rho)] f\left(\frac{\Phi_i(\rho)}{p_i}\right). \quad (2.8)$$

Such *strong monotonicity* conditions imply  $f$  is non-increasing even after post-selection or multiple flagged outcomes.

**Convexity.**  $f$  is convex under mixing states,

$$f\left(\sum_i p_i \rho_i\right) \leq \sum_i p_i f(\rho_i), \quad (2.9)$$

i.e., by mixing states, one cannot increase the amount of resource. Note, this makes sense, as mixing in this sense means losing information. However, such property is not essential, but rather convenient when computing some functions within QRTs.

**Sub-additivity.**  $f$  is sub-additive

$$f(\rho \otimes \sigma) \leq f(\rho) + f(\sigma) \quad \forall \rho, \sigma, \quad (2.10)$$

i.e., two states together cannot have more resources than the sum of their resources. Although this feels intuitive to include as an essential condition, some measures may not follow this, for example, non-locality [101]. When the equality holds,  $f$  is called

additive. Such a property simplifies calculation and provides reasoning, for example, having  $N$  states has the same resources as having 1 state. In Chapter 7, it helps us to infer results as stabilizer Rényi entropy has this property (sec. 7.5.1). Although most measures do not have additivity, by regularization, one can construct functions that are additive on multiple copies of the same state,

$$f^\infty(\rho) = \lim_{n \rightarrow \infty} \frac{1}{n} f(\rho^{\otimes n}), \quad (2.11)$$

provided the limit exists. One sufficient condition for a limit's existence is a weaker form of sub-additivity:  $f(\rho^{\otimes(m+n)}) \leq f(\rho^{\otimes m}) + f(\rho^{\otimes n})$ ,  $\forall \rho, m, n$  [105]. If  $f$  is resource monotone,  $f^\infty$  is also.

**Asymptotic continuity.**  $f$  is asymptotically continuous if [106]

$$|f(\rho) - f(\sigma)| \leq K\epsilon \log[\dim(\mathcal{H})] + c(\epsilon), \quad \epsilon = \frac{1}{2} \|\rho - \sigma\|_1; \quad \forall \rho, \sigma \in \mathcal{S}(\mathcal{H}), \quad (2.12)$$

where  $K$  is some constant, and  $c(\epsilon) \rightarrow 0$  as  $\epsilon \rightarrow 0$ . The trace norm is defined as  $\|M\|_1 = \text{Tr}\sqrt{M^\dagger M}$ . Physically, it implies that if one state is perturbatively close to another state, then one would naturally anticipate their resource content to be similarly close to each other. Of course, the difference will increase as the system size increases, hence  $\log[\dim(\mathcal{H})]$ . For example, von Neumann entropy  $S_{\text{vN}}$  is an asymptotically continuous function, as ensured by the Fannes-Audenaert inequality [107, 108],

$$|S_{\text{vN}}(\rho) - S_{\text{vN}}(\sigma)| \leq \epsilon \log[\dim(\mathcal{H}) - 1] + h(\epsilon), \quad (2.13)$$

where  $h(x) = -x \log x - (1-x) \log 1-x$ , also known as binary entropy function. Such a property is often useful when proving bounds on the measures using perturbation theory.

### 2.3.2 General distance-based constructions

One can define resource measures by quantifying ‘how far’ the state is from the set of free states. Here, instead of using standard metric measures for distance, one may use monotonicity as a defining quality. A distance function  $d : \mathcal{S}(\mathcal{H} \otimes \mathcal{H}) \rightarrow \mathbb{R}_{\geq 0}$  is called

$$\text{contractive under all CPTP maps } \Phi \Leftrightarrow d(\rho, \sigma) \geq d(\Phi(\rho), \Phi(\sigma)), \quad \forall \rho, \sigma \in \mathcal{S}(\mathcal{H}). \quad (2.14)$$

Then, one has to minimize the distance from all free states to get a reliable measure<sup>1</sup>,

$$\mathcal{R}_d(\rho) = \inf_{\sigma \in \mathcal{F}(\mathcal{H})} d(\rho, \sigma). \quad (2.15)$$

It is obvious, that  $\mathcal{R}_d = 0, \forall \rho \in \mathcal{F}$ , and monotonicity is also satisfied as

$$\mathcal{R}_d(\Phi(\rho)) = \inf_{\tau \in \mathcal{F}} d(\Phi(\rho), \tau) \leq \inf_{\sigma \in \mathcal{F}} d(\Phi(\rho), \Phi(\sigma)) \leq \inf_{\sigma \in \mathcal{F}} d(\rho, \sigma) = \mathcal{R}_d(\rho). \quad (2.16)$$

---

<sup>1</sup>Note that we do not use min, rather we use inf, because it is possible that the minimum may not be found within the set. However, when the set  $\mathcal{F}(\mathcal{H})$  is closed, each infimum is attained by some free state  $\sigma$ .



The first inequality implies that measuring the distance from a state  $\Phi(\sigma)$  cannot yield a smaller value than measuring the distance from a free state  $\tau \in \mathcal{F}$ , and the second inequality uses  $d$  as contractive under  $\Phi$  [Eq.(2.14)]. Thus, it becomes clear how one can use a contractive distance function to define the resource measure. Quantum Fisher Information (QFI) is an example of such a contractive measure (sec. 2.6.3).

### 2.3.3 Entropic measures

Most entropic measures are some generalization of the relative Rényi entropies from classical information theory [109]. The *quantum relative Rényi entropy* is defined by [110]

$$D_\alpha(\rho||\sigma) = \frac{1}{1-\alpha} \log [\text{Tr}(\rho^\alpha \sigma^{1-\alpha})], \quad 0 < \alpha < 1. \quad (2.17)$$

Often, we deal with only the limiting case,

$$D_0(\rho||\sigma) = -\log [\text{Tr}(\Pi_\rho \sigma)], \quad \text{where } \Pi_\rho \text{ projects onto } \text{supp}(\rho), \quad (2.18)$$

$$\lim_{\alpha \rightarrow 1} D_\alpha(\rho||\sigma) \rightarrow S_{\text{rel}}(\rho||\sigma) := -\text{Tr}[\rho(\log \sigma - \log \rho)], \quad (2.19)$$

where  $S_{\text{rel}}(\rho||\sigma)$  is the quantum relative entropy [111]. The well-known Rényi entropy can be obtained by taking  $\sigma = \mathbb{I}_{\mathcal{H}}$ :

$$S_\alpha(\rho) := -D_\alpha(\rho||\mathbb{I}_{\mathcal{H}}) = \frac{1}{1-\alpha} \log [\text{Tr}(\rho^\alpha)]. \quad (2.20)$$

At  $\alpha = 1$ , we can recover von Neumann entanglement entropy,

$$S_{\text{vN}}(\rho) = -D_1(\rho||\mathbb{I}_{\mathcal{H}}) = -\text{Tr}[\rho \log \rho]. \quad (2.21)$$

One crucial property of  $D_\alpha(\rho||\sigma)$  is that they are contractive under CPTP maps, i.e.,

$$D_\alpha(\rho||\sigma) \geq D_\alpha(\Phi(\rho), \Phi(\sigma)), \quad \alpha \in [0, 1]. \quad (2.22)$$

This is also known as a data processing inequality [110]. Now, using this contractive property, we can define resource measures for any QRT,

$$\mathcal{R}_\alpha(\rho) := \inf_{\sigma \in \mathcal{F}} D_\alpha(\rho||\sigma), \quad \text{for } \alpha \in [0, 1]. \quad (2.23)$$

Similarly, the *relative entropy of a resource* is defined as

$$\mathcal{R}_{\text{rel}}(\rho) = \inf_{\sigma \in \mathcal{F}} S_{\text{rel}}(\rho||\sigma). \quad (2.24)$$

Interestingly,  $\mathcal{R}_{\text{rel}}$  is asymptotically continuous [Eq. (2.12)] when the maximally mixed state is free in the QRT [112].

### 2.3.4 Geometric measures

Relative entropy measures, while useful as resource monotones, are not proper metrics since they do not satisfy the triangle inequality. In some cases, one may be interested in using a measure with geometric interpretation, i.e., far from free states means more resources. It is indeed a special scenario under general distance-based constructions

(Sec. 2.3.2), which are contractive under CPTP. One such example is *trace distance of resource*,

$$\mathcal{R}_{\text{Tr}}(\rho) = \inf_{\sigma \in \mathcal{F}} D_{\text{Tr}}(\rho, \sigma) = \inf_{\sigma \in \mathcal{F}} \frac{1}{2} \|\rho - \sigma\|_1. \quad (2.25)$$

Operationally, the trace distance quantifies how distinguishable two states are: for a system prepared in either state  $\rho_0$  or  $\rho_1$  with probabilities  $p_0$  and  $p_1$ , the minimum error in identifying the correct state decreases as the distinguishability between the states increases, and the probability is given by  $\frac{1}{2}(1 + \|p_0\rho_0 - p_1\rho_1\|_1)$  (*Holevo-Helström theorem*) [113, 114].

Similarly, using the Bures metric,  $D_B(\rho, \sigma) = \sqrt{2}\sqrt{1 - F(\rho, \sigma)}$ , one can also define the Bures distance of resource measure as its infimum,

$$\mathcal{R}_B(\rho) = \inf_{\sigma \in \mathcal{F}} D_B(\rho, \sigma). \quad (2.26)$$

Here,  $F(\rho, \sigma)$  is the Uhlmann fidelity between two states, defined as  $(\text{Tr} \sqrt{\sqrt{\rho}\sigma\sqrt{\rho}})^2$ , and for the pure states  $F(|\psi\rangle, |\phi\rangle) = |\langle\psi|\phi\rangle|^2$ .

As both the trace-distance and Bures metric are true metrics, these are called geometric measures. However, one can directly minimize the fidelity  $F(\rho, \sigma)^2$  as a geometric measure, which historically happened in entanglement [115, 116], and then generalized as the geometric measure. In an arbitrary QRT,

$$\mathcal{R}_G = \inf_{\sigma \in \mathcal{F}} (1 - F(|\psi\rangle\langle\psi|, \sigma))^2 = \frac{1}{4} \mathcal{R}_B(|\psi\rangle)^4. \quad (2.27)$$

If  $\mathcal{F}(\mathcal{H})$  is a convex set with pure states being extreme points, then the infimum is always attained by a pure state,  $\sigma = |\phi\rangle\langle\phi|$ . In the entanglement theory, it is also the case that minimization over product states is enough (see Sec. 2.6.4). In Sec 2.6.4, we use such a geometric measure [Eq. (2.66)] to measure multipartite entanglement. Although  $\mathcal{R}_G$  does not have an operational meaning like trace distance, it is still valuable for deriving bounds, such as

$$1 - \sqrt{1 - \mathcal{R}_G(\rho)} \leq \mathcal{R}_{\text{Tr}}(\rho) \leq \sqrt{\mathcal{R}_G(\rho)}, \quad (2.28)$$

using the following inequality:  $1 - F(\rho, \sigma) \leq D_{\text{Tr}}(\rho, \sigma) \leq \sqrt{1 - F(\rho, \sigma)^2}$  [117].

## 2.4 Entanglement as a resource

In classical physics, it is easy to correlate distant systems by sharing information and acting on it locally. For example, Alice flips a coin and sends the result to Bob. If it is a head, he turns on a red light; if tails, a blue one. Repeating this many times creates a shared pattern based on a probability distribution. This approach is called Local Classical Systems, Operations, and Classical Communication (LCSOCC).

Quantum mechanics allows for more powerful correlations. One key framework is Local Operations and Classical Communication (LOCC), where local systems are quantum rather than classical. So, every LOCC involves a local measurement operation, followed by a broadcast of the measurement outcome, which can be written as Kraus-operator notation (see Sec. 2.8.1):

$$\Lambda(\cdot) = \sum_k \left( \bigotimes_{i=1}^N M_{k,i}^{A_i} \right) (\cdot) \left( \bigotimes_{i=1}^N M_{k,i}^{A_i} \right)^\dagger, \quad (2.29)$$

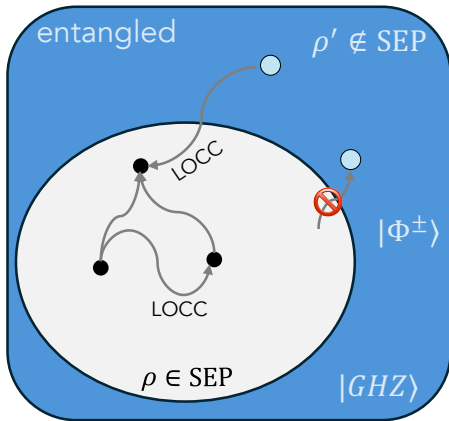
where  $M_{k,i}^{A_i}$  acts on  $A_i$  for a  $N$ -partite state space  $\mathcal{H} = \bigotimes_{k=1}^N \mathcal{H}^{A_k}$ . The QRT of entanglement can be defined using such LOCC as free operation [84, 118]. Any states obtained by LOCC would impose a tensor product structure (LO) and probabilistic mixture (CC) of the following form

$$\rho = \sum_{k=1}^n p_k \rho_{1,k}^{A_1} \otimes \rho_{2,k}^{A_2} \otimes \cdots \otimes \rho_{N,k}^{A_N}, \quad (2.30)$$

where  $\rho_{i,k}^{A_i}$  is an arbitrary state for party  $A_i$ . Any state having such a form is called a *separable* state, which forms the set of free states called SEP, and LOCC keeps SEP invariant. As every convex combination of separable states is also separable, the QRT is convex. Any state not belonging to SEP is called *entangled*. Thus, operationally, entanglement can be defined as a resource in this QRT (see Fig. 2.2).

### Quantum Entanglement

**Definition 3.** Entanglement is a characteristic of a composite physical system that cannot be created or enhanced through local (quantum) operations and classical communication (LOCC).



**Figure 2.2.** The set of quantum states contains the subset of separable states SEP. The LOCC operations map states within the set of separable states (SEP) to other states that also remain within SEP. Entangled states lie outside this set, for example, the Bell state, the GHZ state, etc. LOCC can bring an entangled state (outside) to a separable state (inside). However, it cannot generate entanglement.

Deciding whether a state is in SEP or not is difficult; in fact, the problem is **NP-hard** [119] (see sec. 3.2.1 for complexity classes). Instead, traditionally, research has been pushed along identifying separability criteria, which are necessary but not sufficient conditions for a state to be separable [120]. It might seem that fully understanding the set of free operations would resolve the problem. However, determining whether a given map  $\Lambda$  belongs to the class of LOCC operations remains an open question. Note that the structure described by Eq. (2.29) for LOCC is not the only structure for the separability condition. For example, there exist separable maps that cannot be realized via LOCC, and such maps were initially introduced to illustrate the phenomenon known as *nonlocality without entanglement* [121]. Hence, to obtain interesting QRTs, one needs to consider free operations more powerful than LOCC.

## 2.5 Bipartite entanglement

Bipartite systems serve as a fundamental framework for studying entanglement. Historically, the EPR paradox [122] and the subsequent discovery of Bell inequalities [123] are also primarily concerned with such systems. Moreover, quantum technological applications such as dense coding [124], quantum key distribution [125], and quantum teleportation [100] rely heavily on bipartite entanglement as a key resource for enabling secure and efficient quantum communication.

Consider a bipartite system  $AB$ , admitting a tensor product structure

$$\mathcal{H}_{AB} = \mathcal{H}_A \otimes \mathcal{H}_B, \quad (2.31)$$

where each subsystem is a qudit. For a pure state,

$$|\psi\rangle = \sum_{i,j} c_{ij} |i\rangle_A \otimes |j\rangle_B, \quad |i\rangle_A \in \mathcal{S}(\mathcal{H}_A), |j\rangle_B \in \mathcal{S}(\mathcal{H}_B). \quad (2.32)$$

The coefficient matrix  $c_{ij}$  can be simplified with a singular value decomposition (SVD), yielding a common Schmidt basis,

$$|\psi\rangle = \sum_k \lambda_k |k\rangle_A \otimes |k\rangle_B, \quad (2.33)$$

where  $\lambda_k$  are singular values, also known as Schmidt values in the context of quantum information, where SVD is also called the Schmidt decomposition of a state. Moreover, as the Schmidt values are the probability amplitude of the state, they fulfill the normalization condition, i.e.,  $\sum_k \lambda_k^2 = 1$ .

Separable states have only one non-zero Schmidt value, hence  $\lambda_0 = 1$ , i.e.,  $|\psi\rangle = |\psi\rangle_A \otimes |\psi\rangle_B$ . Such states can be obtained using LOCC, and belong to the set of free states  $\text{SEP}(\mathcal{H})$ . Moreover, they do not contain any local correlation, and hence, one can claim that there is no loss of information when a separable state is restricted to one of the subsystems.

For an arbitrary state, one may not have the full information by looking at a subsystem. Thus, we can differentiate between a separable state and a non-separable state by quantifying the information content of the restricted subsystem state. For example, consider measuring  $\hat{O}_A$  only on subsystem  $A$ :  $\hat{O}_A \otimes \hat{\mathbb{I}}_B$  on  $|\psi\rangle$ ,

$$\langle\psi| \hat{O}_A \otimes \hat{\mathbb{I}}_B |\psi\rangle = \sum_k \lambda_k \langle k|_A \hat{O}_A |k\rangle_A = \text{Tr} \left( \underbrace{\sum_k \lambda_k |k\rangle_A \langle k|_A}_{:=\rho_A} \hat{O}_A \right). \quad (2.34)$$

Thus, the restriction to a subsystem makes the pure state  $|\psi\rangle$  a mixed state  $\rho_A$ . The *mixedness* of  $\rho_A$ , can be measured by its entropy, known as von Neumann entanglement entropy,

$$S_{\text{vN}}(\rho_A) = - \sum_k p_k \log p_k. \quad (2.35)$$

Interestingly, one can see that the entanglement entropy depends on  $\lambda_k$  — probability amplitude shared between two subsystems, and thus indeed it contains the information of the indissociability of two subsystems, i.e., how correlated two systems are, and how

one can lose information if dissociated. Note that the operation  $\rho \rightarrow \rho_A$ , i.e., acting only on one part of the joint system, and then discarding the subsystem, is still a free operation (see Definition. 2.1), thus it does not change the resource contents. For a product state, as  $\lambda_k = 1$ ,  $S_{\text{vN}} = 0$  for any bipartitions.

The Bell states are the four maximally entangled two-qubit states and form an orthonormal basis for the two-qubit Hilbert space:

$$|\Phi^\pm\rangle = \frac{1}{\sqrt{2}}(|00\rangle \pm |11\rangle), \quad |\Psi^\pm\rangle = \frac{1}{\sqrt{2}}(|01\rangle \pm |10\rangle). \quad (2.36)$$

Each Bell state has equal absolute value of Schmidt coefficients  $(\frac{1}{\sqrt{2}}, \frac{1}{\sqrt{2}})$ , indicating maximal entanglement.

Von Neumann entanglement entropy follows some of the additional properties mentioned in Sec. 2.3.1. For example, subadditivity. For a system with two subsystems  $A$  and  $B$ ,

$$S_{\text{vN}}(\rho_{AB}) \leq S_{\text{vN}}(\rho_A) + S_{\text{vN}}(\rho_B) \quad (2.37)$$

We utilize the fact that the quantum relative entropy [Eq. (2.19)] is non-negative for states. Consider  $\sigma_{AB} = \rho_A \otimes \rho_B$ , then:

$$\begin{aligned} S_{\text{rel}}(\rho_{AB} \| \rho_A \otimes \rho_B) &= \text{Tr}[\rho_{AB} (\log \rho_{AB} - \log(\rho_A \otimes \rho_B))] \\ &= \text{Tr}[\rho_{AB} \log \rho_{AB}] - \text{Tr}[\rho_{AB} (\log \rho_A \otimes I_B + I_A \otimes \log \rho_B)] \\ &= -S_{\text{vN}}(\rho_{AB}) + S_{\text{vN}}(\rho_A) + S_{\text{vN}}(\rho_B). \end{aligned}$$

Now, using  $S_{\text{rel}} \geq 0$ , we can obtain Eq. (2.37).

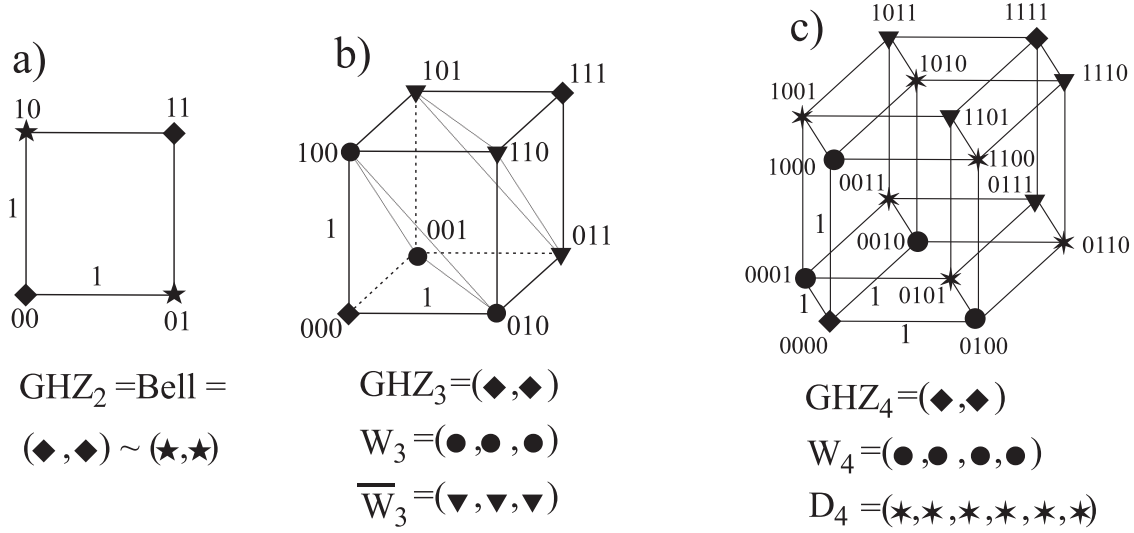
## 2.6 Multipartite entanglement

The concept of bipartite entanglement extends naturally to multipartite systems. For instance, in a system composed of three subsystems,  $\mathcal{H}_{ABC} = \mathcal{H}_A \otimes \mathcal{H}_B \otimes \mathcal{H}_C$ , a state is said to be (fully) separable if it can be written as a product of individual states:  $|\psi_A\rangle \otimes |\psi_B\rangle \otimes |\psi_C\rangle$ . Any state that does not take this form is considered entangled. At first glance, this might appear to be a straightforward generalization. However, it raises an important question: *Is there a fundamental difference between entanglement in multipartite systems and the well-understood bipartite case?* The answer is yes that multipartite entanglement exhibits qualitatively richer and more complex behavior than bipartite entanglement [96]. Even three qubits show fundamentally new types of entanglement beyond Bell states [126].

### 2.6.1 Example of multipartite entangled states: GHZ, W, Dicke

To understand the possible entangled states with multiple qubits, we will follow a diagrammatic approach for the sake of visualization. Let's consider the  $2^N$  computational basis states as the corners of the hypercube (see Fig. 2.3). In this way, every

edge is traversed by a Hamming distance<sup>2</sup> of 1. For the 2-qubit case (square), any superposition of two states forming an edge of the square is separable, but equal-weight superpositions of states represented by two corners on a diagonal give maximally entangled Bell states [Eq. (2.36)]. A similar picture is easily generalized to the case of three qubits. Here we can see that superpositions of two states at Hamming distance one, belonging to the same edge of the cube are separable, and superpositions of states at a Hamming distance of two display bipartite entanglement.



**Figure 2.3.** Distinguished pure states for systems of (a) two, (b) three, and (c) four qubits can be represented geometrically. In the case of two qubits, the Bell states correspond to pairs of opposite corners along a diagonal of a square. For three qubits, the Greenberger–Horne–Zeilinger (GHZ) state,  $|\text{GHZ}_3\rangle = |000\rangle + |111\rangle$ , is represented by two opposite corners along a main diagonal of a cube. Similarly, for four qubits, the GHZ state  $|\text{GHZ}_4\rangle = |0000\rangle + |1111\rangle$  corresponds to two opposite corners of a hypercube. The states  $W_3$  and  $\bar{W}_3$ , which are locally equivalent, are represented by two parallel triangles in the three-qubit case (b), while in the four-qubit case (c), the locally equivalent  $W_4$  and  $\bar{W}_4$  states correspond to two parallel tetrahedra. Additionally, the state labeled  $D_4$  represents a four-qubit Dicke state (Taken from [96]).

Now consider the following state, which can be decomposed into an entangled state and a product state.

$$|\psi_{AB|C}\rangle = \frac{1}{\sqrt{2}}(|000\rangle + |110\rangle) = |\Phi_{AB}^+\rangle \otimes |0_C\rangle. \quad (2.38)$$

States that cannot be decomposed in this manner are said to possess genuine multipartite entanglement. One can rightfully guess that a superposition of two states corresponding to maximally distant corners (Hamming distance equals the number of qubits) may be highly entangled. This is indeed the case, and they are called Greenberger–Horne–Zeilinger (GHZ) states [127]:

$$|\text{GHZ}\rangle_N = \frac{1}{\sqrt{2}}(|0\rangle^{\otimes N} + |1\rangle^{\otimes N}). \quad (2.39)$$

<sup>2</sup>The Hamming distance between two binary strings of equal length is the number of positions at which the corresponding bits differ. It measures how many substitutions are needed to change one string into the other.

The three-qubit GHZ state was created in 1999 [128].

Another genuinely multipartite entangled state is the  $W$  state, formed by triangles for the 3-qubit case, or tetrahedra for the 4-qubit case (see Fig. 2.3), where each basis state has only one qubit in state 1 :

$$|W_3\rangle = \frac{1}{\sqrt{3}}(|001\rangle + |010\rangle + |100\rangle), \quad (2.40)$$

$$|W_4\rangle = \frac{1}{2}(|0001\rangle + |0010\rangle + |0100\rangle + |1000\rangle). \quad (2.41)$$

One can also observe that the superposition of permutationally invariant states can form entangled states, which are known as Dicke states. For example,

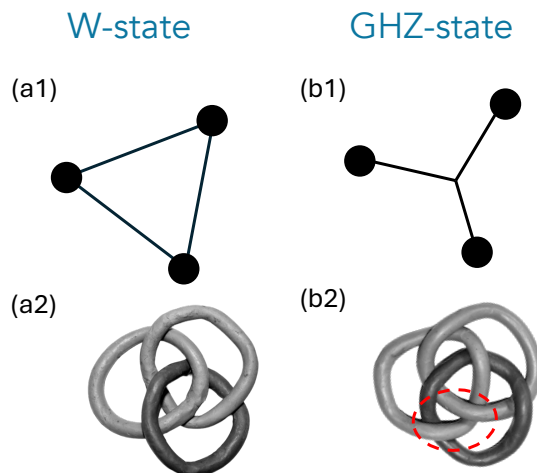
$$|D_4\rangle = \frac{1}{\sqrt{6}}(|1001\rangle + |1010\rangle + |1100\rangle + |0101\rangle + |0110\rangle + |0011\rangle). \quad (2.42)$$

More generically,

$$D_k^n = \binom{n}{k}^{-1/2} \sum_i P_i \left( |1\rangle^{\otimes k} \otimes |0\rangle^{\otimes (n-k)} \right). \quad (2.43)$$

### Perseverance (fragility) of the entangled states

Given a different kind of multipartite entangled states, one may wonder if there is something different between them. One can perform a perseverance (fragility) analysis in the sense that, by tracing out any subsystem, we can determine whether it remains entangled (not fragile) or becomes a separable state (fragile) [See Fig. 2.4]. GHZ states are fragile, as they become separable after tracing out, which also means that all the entanglement is of a global nature. Interestingly, this property holds if and only if the state is already in Schmidt decomposable form [129]. On the other hand, the  $W$  state is more robust than the GHZ state, because after tracing out, an entangled mixed state remains, and it cannot be written as a superposition of less than three separable states [130]. Such fragility does not say if one state is more entangled than the other; instead, it shows that they are entangled in different ways. For instance,  $W$  and GHZ states cannot be transformed into each other by LOCC [130].



**Figure 2.4.** A schematic comparison of  $W$  and GHZ states is illustrated using strings and knots, where you can envision the fragility of the state. For the  $W$  state (a1, a2), tracing out one subsystem leaves the remaining two subsystems still entangled. In contrast, for the GHZ state, which can be represented by three objects connected by a single thread or by three rings, removing any one ring unlinks the others (b1, b2) [Adapted from [96]].



As more qubits are added, classification becomes increasingly difficult and relies on detailed knowledge of the states, making it hard to link to observable features. Instead, here we use a hierarchy of separability criteria that offers less structural detail but is more practical and scalable.

## 2.6.2 Partitions of many-body systems

Consider a quantum system composed of  $N$ -parties,

$$|\psi\rangle = \sum_{i_k} c_{i_1, \dots, i_N} |i\rangle_1 \otimes \dots \otimes |i\rangle_N, \quad (2.44)$$

where the state belongs to the Hilbert space  $\mathcal{H} = \otimes_{k=1}^N \mathcal{H}_k$ . Such a multipartite system is quite different from the bipartite case. As there are multiple subsystem choices, and entanglement explicitly relies on the subsystem, we require a systematic formalism to treat all possible partitions on equal footing. To understand the multipartite entanglement [131], the main question we ask is whether it is possible to cluster the  $N$  parties into  $n$  groups, for  $n \leq N$ , such that  $|\psi\rangle$  is a product state with respect to the partition,

$$\textbf{\textit{n-separable:}} \quad |\psi\rangle = |\phi_1\rangle \otimes |\phi_2\rangle \otimes \dots \otimes |\phi_n\rangle. \quad (2.45)$$

If this is possible, then we can call  $|\psi\rangle$  as *n-separable*. If it is  $N$ -separable, then it is a product state with respect to all subsystems, and hence we call such a state *fully separable*. On the other extreme, if the state is not even biseparable, then it is *genuine N-partite entangled*. However, some states are in neither of the above classes; instead, they are expressed as some superposition of various classes of entangled or separable states. In that case, we ask if a state  $|\psi\rangle$  is producible by  $k$ -party entangled states, which let us define  $k$ -producible states. For instance, if we can write

$$\textbf{\textit{k-producible:}} \quad |\psi\rangle = |\phi_1\rangle \otimes |\phi_2\rangle \otimes \dots \otimes |\phi_m\rangle, \quad \forall |\phi_i\rangle \text{ containing maximally } k\text{-parties}, \quad (2.46)$$

we can call it  $k$ -producible, and it has to fulfill that  $k \cdot m \geq N$ <sup>3</sup>. In this way,  $k$ -producible implies it is enough to generate specific  $k$ -party entanglement to prepare  $|\psi\rangle$ . In the same way, we can say the converse statement that a state has *k-party entanglement*, if it is not producible by  $(k-1)$ -party entanglement, as it requires at least  $k$ -party entanglement. Formally, we can define,

$$\textbf{\textit{k-party entangled:}} \quad k\text{-producible, but not } (k-1)\text{-producible}. \quad (2.47)$$

For example, if a 6-qudit system can be written like

$$\underbrace{|\psi\rangle}_{\{1,2,3,4,5,6\}} = \underbrace{|\psi\rangle_{R_1}}_{\{1,2,3\}} \otimes \underbrace{|\psi\rangle_{R_2}}_{\{4\}} \otimes \underbrace{|\psi\rangle_{R_3}}_{\{5,6\}}, \quad (2.48)$$

it is called 3-producible, where the partition  $P = \{R_1, R_2, R_3\}$  is with  $R_1 = \{1, 2, 3\}$ ,  $R_2 = \{4\}$ ,  $R_3 = \{5, 6\}$ . If the  $|\psi\rangle_{R_1}$  cannot be represented by further partition, then we can say  $|\psi\rangle$  is 3-party entangled. Thus, the notion of multipartite entanglement stands on the idea of  $k$ -producible states.

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<sup>3</sup>Note, this provides a way to relate  $m$ -separability and  $k$ -producibility.



Note that defining  $n$ -separability does not reveal some structures of the entanglement. For example, for large  $N$ ,  $n$ -separability cannot distinguish whether the two partitions are of equal size or just one partition has one qubit and the other partition has a large, genuinely multipartite entangled state. Moreover, to tell about  $n$ -separability, one needs to know what  $N$  is. On the other hand, the notion of  $k$ -producibility, by design, can say ‘how many parties are entangled’. Moreover, deciding whether a state is  $k$ -producible does not require knowledge of  $N$ , for example, if  $k \ll N$  subsystems, one can still conclude if the system has  $k$ -party entanglement.

### 2.6.3 Multipartite entanglement through Fisher information

Entanglement plays a vital role in phase estimation, and hence in quantum metrology [132], where the task is to estimate a parameter shift  $\theta$  as precisely as possible. If one uses a probe state of  $N$  classically correlated particles, it is possible to obtain a parameter uncertainty of  $\Delta\theta \sim 1/\sqrt{N}$ , whereas if an  $N$ -particle entangled state is used, the uncertainty can be further reduced to  $1/N$ . Thus, parameter uncertainty can be related to the number of particles being genuinely entangled.

**Phase estimation.** In a general phase estimation scenario, a probe state  $\rho$  is perturbed under  $O$ ,

$$\rho \xrightarrow{O} \rho_\theta = e^{-i\theta O} \rho e^{i\theta O}, \quad (2.49)$$

where the operator  $O$  is known, and  $\theta$  is unknown, and the goal is to estimate  $\theta$  by studying the relation between  $\rho_\theta$  and  $O$ . An estimator  $\theta_{\text{est}}(\{\mu_i\}_m)$  is used depending on the results obtained from  $m$ -independent repeated measurements:  $\{\mu_i\}_m = \{\mu_1, \dots, \mu_m\}$ , with mean  $\langle \theta_{\text{est}} \rangle$ , and variance  $\Delta^2 \theta_{\text{est}} = \langle \theta_{\text{est}}^2 \rangle - \langle \theta_{\text{est}} \rangle^2$ . This variance quantifies the precision of the estimator and provides a figure of merit for its quality. In case of an unbiased estimator, one can recover the true phase,  $\langle \theta_{\text{rmest}} \rangle = \theta$ , and in this case, the minimal standard deviation is bounded by something called Fisher information, according to the Cramér–Rao inequality [133, 134]

$$\text{Cramér–Rao inequality: } \Delta^2 \geq 1/F(\theta), \quad (2.50)$$

where  $F(\theta)$  is called Fisher information as we defined below.

**Fisher information (classical).** The expectation value  $\langle \theta_{\text{est}} \rangle$  is dependent on the probability distribution obtained after repeated measurement,

$$\langle \theta_{\text{est}} \rangle = \sum_{\mu} p(\mu|\theta) \theta_{\text{est}}(\mu). \quad (2.51)$$

Assuming an unbiased estimator, the quality of the estimation depends on the quality of the measurement scheme itself, which is related to the probability distribution  $p(\mu|\theta)$ . The Fisher information [135, 136] precisely defines a function that measures the sensitivity of  $p(\mu|\theta)$  to changes of the parameter:

$$F(\theta) = \Delta^2 \underbrace{[\partial_\theta \ln p(\mu|\theta)]}_{=-s(\mu|\theta)} = \sum_{\mu} p(\mu|\theta) \left( \partial_\theta \ln p(\mu|\theta) \right)^2. \quad (2.52)$$

Here  $s(\mu|\theta)$  is the score of the distribution and measures the change in the likelihood function with respect to  $\theta$ . As its mean is zero, the variance defines the simplest non-trivial measure of the properties of the score.

**Quantum Fisher information.** In the quantum case, the probability distribution is made of outcomes of positive operator-valued measurements (POVM), denoted as  $\{\Pi_i\}$ , such that

$$\sum_{\mu} \Pi_{\mu} = \mathbb{I}, \quad p(\mu|\theta) = \text{Tr}(\rho(\theta)\Pi_{\mu}). \quad (2.53)$$

By removing the explicit dependence on POVM, i.e., by optimizing over all possible POVMs, one obtains quantum Fisher information (QFI) as

$$F_Q(\rho_{\theta}) = \max_{\{\Pi_{\mu}\}} F(\theta). \quad (2.54)$$

In this case, we can obtain [137, 138]

$$\text{quantum Cramér–Rao bound: } \Delta^2 \geq 1/F(\theta) \geq 1/mF_Q(\rho_{\theta}). \quad (2.55)$$

where  $m$  is the number of independent repetitions. However, this way of defining QFI associates the classical outcomes of quantum experiments with precision. One can see the explicit connection to  $\rho_{\theta}$  through the original definition of QFI, defined as a geometrical measure of statistical distinguishability in the space of states [137, 138].

We first define the symmetric logarithmic derivative (SLD)  $L(\theta)$  implicitly through

$$\partial_{\theta}\rho(\theta) = \frac{1}{2}\{\rho(\theta), L(\theta)\} = \frac{1}{2}(\rho(\theta)L(\theta) + L(\theta)\rho(\theta)), \quad (2.56)$$

which is connected to the probability distribution as

$$\partial_{\theta}p(\mu|\theta) = \text{ReTr}(\rho(\theta)\Pi_{\mu}L(\theta)). \quad (2.57)$$

Now we can use the definition of classical Fisher information [Eq. (2.52)],

$$\begin{aligned} F(\theta) &= \sum_{\mu} \frac{(\partial_{\theta}p(\mu|\theta))^2}{p(\mu|\theta)} = \sum_{\mu} \frac{(\text{ReTr}(\rho(\theta)\Pi_{\mu}L(\theta)))^2}{\text{Tr}(\rho(\theta)\Pi_{\mu})} \\ &\leq \sum_{\mu} \frac{\left| \text{Tr}(\sqrt{\rho}\sqrt{\Pi_{\mu}}\sqrt{\Pi_{\mu}}L(\theta)\sqrt{\rho}) \right|^2}{\text{Tr}(\rho(\theta)\Pi_{\mu})} \end{aligned} \quad (2.58)$$

$$(\text{Schwartz inequality}) \leq \text{Tr}(\rho(\theta)L(\theta)^2). \quad (2.59)$$

One can show that there is a POVM for which the above bound is attained,

$$F_Q[\rho(\theta)] = \text{Tr}(\rho(\theta)L(\theta)^2). \quad (2.60)$$

By inserting the particular form of  $\rho(\theta)$  of Eq. (2.49), we obtain,

$$\text{QFI : } F_Q[\rho, O] = 2 \sum_{\mu, \nu} \frac{\rho_{\mu} - \rho_{\nu}}{\rho_{\mu} + \rho_{\nu}} (\rho_{\mu} - \rho_{\nu}) |\langle \mu | O | \nu \rangle|^2, \quad (2.61)$$

where  $\{|\mu\rangle\}$  basis diagonalizes  $\rho$ . For a pure state, it simplifies to the variance,

$$F_Q[|\psi\rangle\langle\psi|, O] = 4 \sum_{\mu, \psi} |\langle\mu|O|\psi\rangle|^2 = 4(\langle\psi|O^2|\psi\rangle - \langle\psi|O|\psi\rangle^2). \quad (2.62)$$

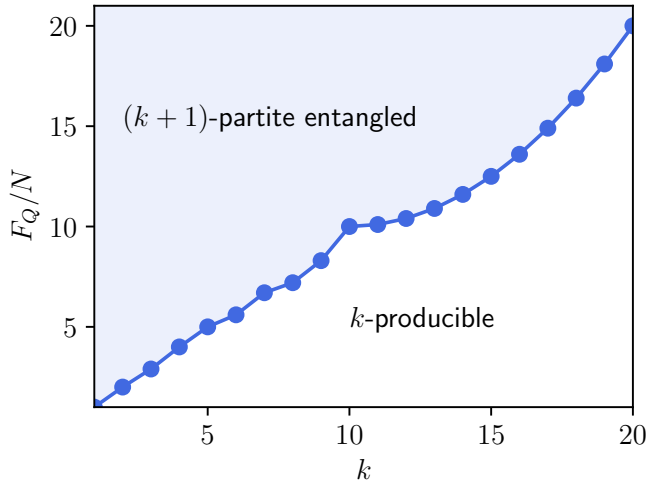
**Relation between  $k$ -party entanglement and  $F_Q$ .** The relation between QFI and  $k$ -partite entanglement can be understood by computing the maximum QFI of a  $k$ -producible state, where one can consider the operator  $\hat{O}$  as some linear operator of Pauli matrices, for example,

$$\hat{O}_{\text{lin}} = \frac{1}{2} \sum_{l=1}^N \vec{n}_l \cdot \vec{\sigma}_l = \frac{1}{2} \sum_l \alpha_l \sigma_x^{(l)} + \beta_l \sigma_y^{(l)} + \gamma_l \sigma_z^{(l)}. \quad (2.63)$$

For  $k$ -producible states and an arbitrary linear operator  $\hat{O}_{\text{lin}}$ , the QFI is bounded by [139, 140]

$$F_Q^{k+1} \text{ criterion: } F_Q[\rho_{k\text{-prod}}, O_{\text{lin}}] \leq sk^2 + r^2, \quad (2.64)$$

where  $s = \lfloor \frac{N}{k} \rfloor$  and residue  $r = N - sk$ . Hence, a violation of the bound implies  $(k+1)$ -partite entanglement. Thus, utilizing the inequality, one can estimate that at least  $k$ -partite entanglement is present in the state (see Fig. 2.5).

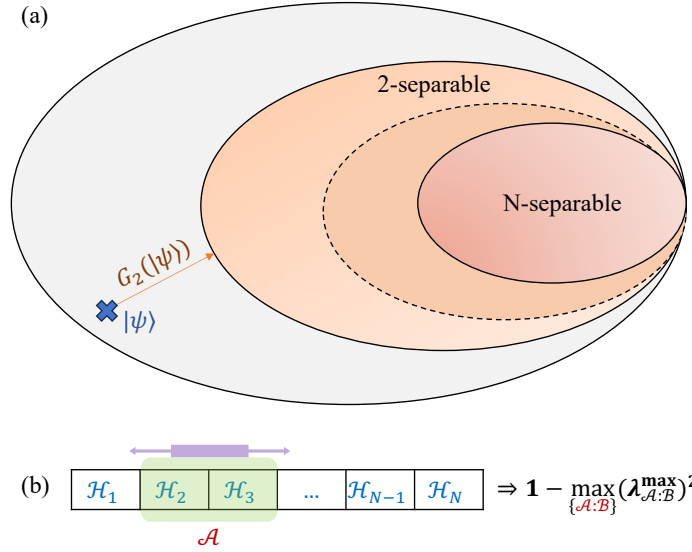


**Figure 2.5.**  $F_Q^{k+1}$  criterion. The solid line denotes the bound, which provides the bound for  $k+1$ -partite entanglement. Larger  $k$ -party entanglement requires higher QFI density. Note that to obtain the same  $k$ -partite entanglement, a larger  $N$  would require a higher  $F_Q$ .

## 2.6.4 Generalized geometric measure of entanglement

The Generalized Geometric Measure of entanglement (GGM) of entanglement generalizes the notion of geometric measure of entanglement and quantifies genuine multipartite entanglement [115, 116, 141–143]. Intuitively, the GGM measures the shortest distance between a quantum state and the set of product states [see Fig. 2.6(a)]. Formally, we can define a hierarchy of geometric measures  $G_k$  with  $2 \leq k \leq N$  of a  $N$ -party pure quantum state  $|\psi\rangle$  as the minimum distance between  $|\psi\rangle$  and the set  $\mathcal{S}_k$  of  $k$ -separable states  $|\pi\rangle = |\psi_1\rangle \otimes |\psi_2\rangle \otimes \cdots \otimes |\psi_k\rangle$ , given by

$$G_k(|\psi\rangle) = \min_{|\pi\rangle \in \mathcal{S}_k} (1 - |\langle\pi|\psi\rangle|^2) = 1 - \max_{|\pi\rangle \in \mathcal{S}_k} |\langle\pi|\psi\rangle|^2. \quad (2.65)$$



**Figure 2.6.** (a) Pictorial representation of the GGM: The GGM of a state is the minimum distance from all separable states, which coincides with the minimum distance from the 2-partite product states. (b) For pure states, instead of needing to minimize the distance of a given pure state to all 2-separable states, a much more efficient way exists to compute the GGM. Namely, it can be obtained from the maximum Schmidt coefficient across all possible bipartitions of the system. Sketched is a system described as a tensor product of local Hilbert spaces  $\mathcal{H}_j$ , bi-partitioned into sets  $\mathcal{A}$  and  $\mathcal{B}$ .

This distance measure resembles the Fubini–Study [144] and the Bures metrics [145]. The original geometric measure of entanglement  $G_{k=N}$  measures the distance between  $|\psi\rangle$  and fully separable states. At the other extreme,  $G_{k=2}$  is zero if  $|\psi\rangle$  can be written as a product state across any *single* bipartition of the Hilbert space. Even if that is the case, bipartite entanglement can still be found in other bipartitions. Since  $\mathcal{S}_k \subset \mathcal{S}_2$  for all  $k$  [Fig. 2.6(a)], the GGM measures the distance between  $|\psi\rangle$  and the union of all forms of  $k$ -separable states ranging from fully separable to biseparable states. The  $G_2$  thus captures the amount of genuine multipartite entanglement present in the system.

One may think that the inclusion of all kinds of biseparable states makes the measure hard to compute. However, one can show that for pure states the GGM takes the simple form [see Fig. 2.6(b)]

$$G_2(|\psi\rangle) = 1 - \max_{|\pi\rangle_{\mathcal{A}:\mathcal{B}} \in \mathcal{S}_2} |\langle \pi | \psi \rangle|^2 = 1 - \max_{\mathcal{A}:\mathcal{B}} (\lambda_{\mathcal{A}:\mathcal{B}}^{\max})^2. \quad (2.66)$$

Here,  $|\psi\rangle$  belongs to the Hilbert space  $\mathcal{H}_{A_1} \otimes \mathcal{H}_{A_2} \otimes \cdots \otimes \mathcal{H}_{A_N}$ . The bipartition  $\mathcal{A} : \mathcal{B}$  satisfies  $\mathcal{A} \cup \mathcal{B} = \{1, 2, \dots, N\}$ , and  $\mathcal{A} \cap \mathcal{B} = \emptyset$  [142, 143]. In other words, maximizing  $|\langle \pi | \psi \rangle|$  across the bipartition  $\mathcal{A} : \mathcal{B}$  is equivalent to finding the maximum Schmidt coefficient,  $\lambda_{\mathcal{A}:\mathcal{B}}^{\max}$ , for that bipartition. Finally, to compute the GGM, one has to find the global maximum of the Schmidt coefficient over all bipartitions  $\mathcal{A} : \mathcal{B}$  of the system<sup>4</sup>. Although in general the number of partitions grows exponentially with the system size, one can often use symmetry or bounds to estimate the GGM effectively. Moreover, in the context of analyzing the ground state of local many-body Hamiltonians, it is often observed that the main contribution to GGM comes from

<sup>4</sup>In the absence of simplifications such as symmetries, we need to calculate over the following number of bipartitions—  $\binom{N}{1} + \cdots + \binom{N}{\frac{N-1}{2}} = 2^{N-1} - 1$  for odd  $N$  and  $\binom{N}{1} + \cdots + \binom{N}{\frac{N}{2}-1} + \binom{N}{\frac{N}{2}} = 2^{N-1} - 1 + \frac{1}{2} \binom{N}{\frac{N}{2}}$  for even  $N$ .

the bipartitions of the system into contiguous blocks. Hence, focusing on a smaller number of blocks would be sufficient to obtain the exact value of GGM in these cases, and one can expect a similar approach to work in some cases of quantum optimization as well. One can show that the theoretical maximum value of GGM is obtained when, for a given bipartition, all the Schmidt coefficients become equal  $\frac{1}{\sqrt{d}}$ , and thus we have  $G_2^{\max}(|\psi\rangle) = 1 - 1/d$ , where  $d$  is the local Hilbert space dimension, also known as qudit dimension. We can understand this intuitively as follows. Consider an arbitrary bipartition  $\mathcal{A} : \mathcal{B}$  where  $|\mathcal{A}| = m$  and  $|\mathcal{B}| = N - m$ . For this bipartition, there can be at most  $d^m$  non-zero Schmidt coefficients, denoted as  $\{\lambda_{\mathcal{A}:\mathcal{B}}^1, \lambda_{\mathcal{A}:\mathcal{B}}^2, \dots, \lambda_{\mathcal{A}:\mathcal{B}}^{k=d^m}\}$ . To maximize the GGM, we need to arrange these Schmidt coefficients so that  $1 - (\lambda_{\mathcal{A}:\mathcal{B}}^1)^2$  is as large as possible. This is achieved when all  $\lambda_{\mathcal{A}:\mathcal{B}}^k$  are equal, i.e.,  $\lambda_{\mathcal{A}:\mathcal{B}}^k = \frac{1}{\sqrt{d^m}}$  for all  $k$ . Therefore, among all possible bipartitions, the maximum of  $1 - (\lambda_{\mathcal{A}:\mathcal{B}}^1)^2$  is obtained in the case where  $m = 1$ , and the maximally possible GGM value is  $G_2 = 1 - \frac{1}{d}$ . Extending the GGM to mixed states is non-trivial, though it can still be calculated by exploiting symmetries [141, 146, 147].

## 2.7 Nonstabilizerness

One of the main questions in quantum information theory is to understand to what extent quantum computers can offer advantages over classical computers. Entanglement is considered one of the underlying resources that can provide such advantages, as an entangled state of  $N$  qudits typically requires storing on the order of  $\mathcal{O}(d^N)$  complex amplitudes for its classical representation. Thus, one expects a quantum algorithm utilizing entanglement to be complex enough not to have a fast counterpart in the classical algorithm. However, this is not always the case. Stabilizer states can exhibit large entanglement yet remain efficiently simulable classically via Clifford circuits, as guaranteed by the Gottesman–Knill theorem [148–151]. The existence of such states calls for a different resource-theoretic formalism where free operations are quantum-computational processes that can be efficiently simulated using a classical computer. In other words, although entanglement is considered one of the primary necessities for quantum advantages, it is certainly not enough to acquire something beyond classical simulation.

### 2.7.1 Classical simulation of quantum computation

In order to distinguish the power of quantum computation from classical computation, the notion of classical simulation of quantum computation provides a good handle. An efficient classical simulation of quantum computation can be precisely defined [88, 152].

Consider a uniform family of quantum circuits  $\mathcal{U}_N$  acting on the  $N$ -qubit input state  $|\mathbf{0}\rangle \equiv |0\rangle^{\otimes N}$ , and then followed by a measurement of one qubit in the computational basis. This has two possible outcome  $\alpha \in \{0, 1\}$ , and the probability of its outcome being  $\alpha$  is

$$\pi(\alpha) = \langle \mathbf{0} | \mathcal{U}_N^\dagger [|\alpha\rangle \langle \alpha| \otimes I] \mathcal{U} | \mathbf{0} \rangle. \quad (2.67)$$

**Strong simulation.** If it is possible to evaluate  $\pi(0)$  up to  $M$  digits in  $\text{poly}(N, M)$  time on a classical computer, we say that the above quantum computation can be efficiently simulated classically in the strong sense.

**Weak simulation.** If it is possible to *sample once* from a probability distribution which is not necessarily exactly  $\{\pi(\alpha)\}$ , but sufficiently close to it, in  $\text{poly}(N)$  time on a classical computer, it is called weak simulation.

Most of the understanding on simulation of quantum computations is related to *strong* classical simulation, e.g., the Gottesman-Knill theorem [149, 150], matchgate circuits [152], etc. Also, most of the quantum circuits become hard when asked for strong simulation, more precisely  $\#\mathbf{P}$ -hard<sup>5</sup> problem, a set much bigger than the class of decision problems solvable by a quantum computer in polynomial time (**BQP**). On the other hand, weak simulation provides a more appropriate way to understand the difference between classical and quantum computation.

## 2.7.2 Stabilizer formalism and Clifford group

Some circuits do not provide any speedups, and such systems are studied in literature [153, 154], mainly via the help of stabilizer formalism. In fact, one of the ways to understand the Gottesman-Knill theorem is indeed through this formalism.

### Stabilizer formalism

The basic idea is to fix a state  $|\psi\rangle$  with the set of operators  $O_1, O_2, \dots, O_k$  such that

- $|\psi\rangle$  is +1 eigenvector for all  $O_i$ , i.e.,  $O_i |\psi\rangle = +1 |\psi\rangle$ ,  $\forall O_i$ ,
- $|\psi\rangle$  is unique up to a complex phase.

These conditions allow for a full characterization of the state by only specifying the list of operators. Such operators  $O_i$  forms a set  $\mathcal{A}_N \subset U_N(d)$ , called *stabilizing set*, and any such state  $|\psi\rangle$  is a *stabilizer state*. Here, we consider the state as an  $N$ -qudit state with local dimension  $d$ .

From here, one can introduce *generalized Clifford group* as the normalizer of the set of operators,  $O_i \in \mathcal{A}_N$ <sup>6</sup> as follows [155]

$$\text{generalized Clifford group: } \mathcal{C}_N(\mathcal{A}_N) = \{U \mid UOU^\dagger \in \mathcal{A}_N, \forall O \in \mathcal{A}_N\}. \quad (2.68)$$

Such a structure ensures that a stabilizer state  $|\psi\rangle$  remains a stabilizer state after the action of any Clifford unitary,  $O |\psi\rangle$ ,  $O \in \mathcal{C}_N$ .

A classic example of such stabilization is given by the Pauli group (see Fig. 2.7 for single-qubit Clifford gates), i.e.,  $\mathcal{A}_N = \mathcal{P}_N$ . The generalized Pauli matrices generate

<sup>5</sup>A function  $f : \{0, 1\}^* \rightarrow \mathbb{N}$  is called  $\#\mathbf{P}$ -hard if, for every function  $g \in \#\mathbf{P}$ , there exists a polynomial-time Turing reduction from  $g$  to  $f$ . That is,  $f$  is at least as hard as any problem in  $\#\mathbf{P}$ .

<sup>6</sup>The **centralizer** of a subset  $A \subseteq G$ , denoted  $C_G(A)$ , is the set of all elements in  $G$  that commute with every element of  $A$ .

$$C_G(A) = \{g \in G \mid ga = ag \text{ for all } a \in A\}$$

The **normalizer** of a subset  $A \subseteq G$ , denoted  $N_G(A)$ , is the set of all elements in  $G$  that conjugate  $A$  to itself.

$$N_G(A) = \{g \in G \mid gAg^{-1} = A\}$$

Whereas centralizer assures commutativity, normalizer assures stability under conjugation, and they follow a subset structure as:

$$C_G(A) \subseteq N_G(A) \subseteq G$$

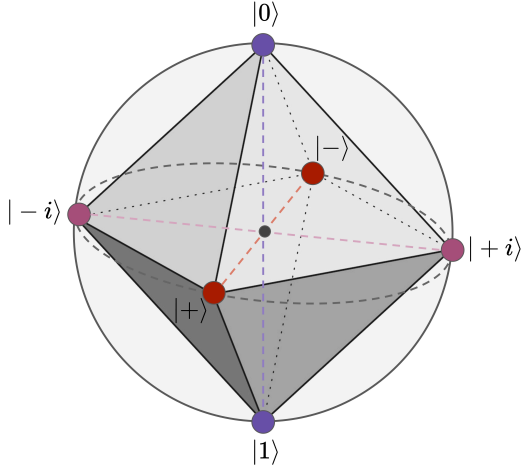
the Pauli group, also known as the Heisenberg-Weyl group, which defines quantum kinematics and the states of the quantum register, and it is a subgroup of the unitary group  $U(N)$ . The generalized Pauli (Heisenberg-Weyl) operators are defined as products of  $X$  and  $Z$  [85, 156],

$$T_{\vec{v}} = \omega^{v_1} X^{v_2} Z^{v_3}; \quad v_j \in \{0, 1, 2, \dots, d-1\}, \quad (2.69)$$

where  $Z$  and  $X$  are defined as the phase and shift operators,

$$Z = \sum_{j=0}^{d-1} \omega^j |j\rangle\langle j|, \quad X = \sum_{j=0}^{d-1} |j+1\rangle\langle j|; \quad \omega = e^{2\pi i/d}. \quad (2.70)$$

Note that there are  $d^3$  Paulis, which can be labelled by  $(v_1, v_2, v_3)$ , where  $v_j \in \mathbb{Z}$  has a physical interpretation as the position of the basis states  $|j\rangle$ .



**Figure 2.7.** The octahedron in the Bloch sphere defines the states accessible via single-qubit Clifford gates (Taken from PennyLane demonstration [157]).

We consider a system of  $N$  qudits with Hilbert space  $\mathcal{H} = \otimes_{i=1}^N \mathcal{H}_i$ , where  $\mathcal{H}_i$  correspond to local Hilbert space for each qudits. The  $N$ -qudit Pauli group  $\mathcal{P}_N$  encompasses all the possible Pauli strings with overall phases  $\pm 1$  or  $\pm i$ . The  $N$ -qudit Pauli group  $\mathcal{P}_N$  is defined as the set of  $\{T_{\vec{v}}\}$

$$T_{\vec{v}} := T_{\vec{v}(1)} \otimes \dots \otimes T_{\vec{v}(N)}; \quad \theta \in [0, 2\pi]. \quad (2.71)$$

The well-known *Clifford group* is the largest subgroup of the unitary group that acts on the Pauli/Heisenberg-Weyl group by conjugation and normalizes it; thus, the Clifford group ( $\mathcal{C}_N$ ) is also called the normalizer of the Heisenberg-Weyl group ( $\mathcal{P}_N$ ):

$$\mathcal{C}_N(\mathcal{P}_N) = \{U \text{ such that } UPU^\dagger \in \mathcal{P}_N \text{ for all } P \in \mathcal{P}_N\}. \quad (2.72)$$

The Clifford group can be generated by

- Hadamard gate  $\frac{1}{\sqrt{2}} \begin{pmatrix} 1 & 1 \\ 1 & -1 \end{pmatrix}$ ,
- $\pi/2$  phase gate  $\begin{pmatrix} 1 & 0 \\ 0 & i \end{pmatrix}$ ,



$$\bullet \text{ CNOT gate } \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 \\ 0 & 0 & 1 & 0 \end{pmatrix}.$$

The pure stabilizer states are defined as the states formed by the action of Clifford unitaries on  $|0\rangle$ ,

$$\textbf{Pure stabilizer states: } \{S_i\} = \{U|0\rangle \text{ such that } U \in \mathcal{C}_N\}, \quad (2.73)$$

and the convex hull of this set defined the full set of stabilizer states:

$$\textbf{Stabilizer states: } \text{STAB}(\mathcal{H}) = \left\{ \sigma \in L(\mathcal{H}) \text{ such that } \sigma = \sum_i p_i S_i \right\}, \quad \sum_i p_i = 1. \quad (2.74)$$

A circuit made of only Clifford unitaries is called a Clifford circuit, and is efficiently classically simulable via the Gottesman–Knill theorem [149].

### Gottesman–Knill theorem

**Theorem 1.** *Every uniform family of Clifford circuits, applied to the input state  $|0\rangle$  and followed by measurement on one qubit in the computational basis, can be efficiently simulated classically in the strong sense.*

We define *stabilizer operations* to be any combination of computational basis preparation, Clifford rotations, and computational basis measurements. Using stabilizer formalism, one can prove the above theorem. However, it is not necessary to go through the formalism in order to understand why the Clifford group is easier to simulate [158].

The Gottesman-Knill theorem is  $\oplus\mathbf{L}$  complete<sup>7</sup>. Only a few classical computing structures allow for the simulation of arbitrary Clifford circuits. For example, a classical circuit model comprising only NOT and CNOT gates should be enough. The strong simulation then asks if the state of the first qubit can be learned with certainty. Interesting, such a problem can be mapped, under a logarithmic-space reduction, to a problem of simulating a classical poly-size CNOT-NOT circuit [151], which has complexity class of  $\oplus\mathbf{L}$  (parity-L) (see more in Sec. 3.2).

## 2.7.3 Discrete Wigner functions

The discrete Wigner representation for odd dimensions enjoys a special property that all stabilizer operations can be represented non-negatively, i.e., it provides a classical probability model for the stabilizer theory [85]. Such a connection motivates one to study Wigner functions, which offer an alternative way to understand stabilizer states.

The Wigner function [159] provides a representation of the wavefunction in terms of a probability distribution in phase space. The discrete Wigner function is a direct

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<sup>7</sup>The complexity class  $\oplus\mathbf{L}$  consists of decision problems that can be solved by a nondeterministic log-space Turing machine, where the correct answer is “yes” if and only if the number of accepting computation paths is odd. A problem is  $\oplus\mathbf{L}$ -complete if it belongs to  $\oplus\mathbf{L}$  and every problem in  $\oplus\mathbf{L}$  can be reduced to it via a log-space reduction. Such problems are the hardest in  $\oplus\mathbf{L}$  under log-space reductions.



analog of it, but is only defined for quantum systems with finite, odd Hilbert space dimension [160]. Given a phase-space point  $\vec{V} = (v_1^{(1)}, v_2^{(1)}) \oplus \dots \oplus (v_1^{(N)}, v_2^{(N)})$  of  $N$ -qudits, we define the corresponding phase space operators using Pauli operators,

$$A_0 = \frac{1}{d^N} \sum_{\vec{V}} T_{\vec{V}}, \quad A_{\vec{V}} = T_{\vec{V}} A_0 T_{\vec{V}}^\dagger. \quad (2.75)$$

From here we arrive at the definition of the discrete Wigner function of a state  $\rho$  as

$$\textbf{Wigner function: } W_\rho(\vec{V}) = \frac{1}{D^N} \text{Tr}[A_{\vec{V}} \rho]. \quad (2.76)$$

The main connection between Wigner representation and stabilizer formalism is through discrete Hudson's theorem [160].

### Discrete Hudson's theorem

**Theorem 2.** *A pure state  $|S\rangle$  has a positive representation if and only if it is a stabilizer state,*

$$\forall \vec{V}, W_{|S\rangle\langle S|}(\vec{V}) \geq 0 \quad \Leftrightarrow \quad \exists U \in \mathcal{C}_N, \text{ such that } |S\rangle = U|0\rangle, \quad (2.77)$$

Moreover, Clifford unitaries act as permutations of phase space,

$$W_{U\rho U^\dagger}(\vec{V}) = W_\rho(\vec{V}'), \quad \forall \vec{V}. \quad (2.78)$$

## 2.7.4 Non-Clifford resources

Given that we know stabilizer states are easy to simulate, one may expect that a state that is not a stabilizer state will be interesting and necessary for quantum computing. Indeed, one needs some non-Clifford gates, for example, a T-gate  $\begin{pmatrix} 1 & 0 \\ 0 & e^{i\pi/4} \end{pmatrix}$  to make a universal quantum computation.

### Why are T-gates (non-Clifford) required for fault-tolerant universal quantum computing?

In fault-tolerant quantum computation, transversal gates<sup>8</sup> are commonly employed [161]. However, the set of gates that can be implemented transversally may differ between stabilizer codes. For instance, the 7-qubit Steane code [162] does not allow for transversal implementation of the T gate. Notably, no non-trivial quantum error-correcting code permits the transversal implementation of a *universal* gate set, which is a limitation formalized by the Eastin-Knill Theorem [163].

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<sup>8</sup>A transversal gate is a fault-tolerant quantum gate implemented by applying independent gates to corresponding physical qubits across different code blocks. It is called *transversal* because the operation acts across these blocks rather than within a single block, thereby preventing errors from spreading between qubits in the same code block.

### Eastin-Knill Theorem

**Theorem 3.** *For any quantum error correcting code with distance at least 2, the set of logical gates that can be implemented transversally generates a set of operations that (up to a global phase) is discrete, and is therefore not universal.*

In other words, no quantum error-correcting code can have a continuous symmetry that acts transversely on physical qubits, and we can see that the T-gate implements the smaller rotational gates.

Moreover, as stabilizer states are efficiently simulable classically, and as Clifford operations keep the stabilizer states stabilizer, one can construct a resource theory of nonstabilizerness by considering stabilizer states as free states, and Clifford operations as free operations [85]. A resource state is thus easily defined as a state that is not a stabilizer state, which we call a nonstabilizer state or a *magic state*<sup>9</sup>. One can show that T-gate is indeed optimal for generating magic resources among the class of diagonal unitary operators, both for the qubit and qutrit cases [165].

To define a resource measure of magic (let's say  $\mathcal{M}$ ), we need to fulfill the two conditions we discussed in Sec. 2.3.1:

- $|\psi\rangle \in \text{STAB} \Leftrightarrow \mathcal{M}(|\psi\rangle) = 0$ ,
- $\mathcal{M}(|\psi\rangle) \geq \mathcal{M}(U|\psi\rangle), \quad \forall |\psi\rangle \in \mathcal{H}, \quad U \in \mathcal{C}_N$ .

## 2.7.5 Stabilizer Rényi entropy

The Stabilizer Rényi entropy (SRE) captures how a state is distributed when expressed in the basis of Pauli operators  $P \in \mathcal{P}_N$ , where  $\mathcal{P}_N$  represents the Pauli group for  $n$ -qudits. The squared (normalized) expectation value  $|\langle\psi|P|\psi\rangle|^2/d^N$  forms a probability distribution<sup>10</sup> over all  $P \in \mathcal{P}_N$ , as they are positive and sum to 1:

$$\sum_{P \in \mathcal{P}_N} \frac{|\langle\psi|P|\psi\rangle|^2}{d^N} = \text{Tr}[(|\psi\rangle\langle\psi|)^2] = 1. \quad (2.79)$$

Note that the second equality comes from the fact that any state can be represented in the Pauli basis,

$$\begin{aligned} \rho &= \frac{1}{d^N} \sum_P \underbrace{c_P}_{\text{Tr}(\rho P)} P \\ \Rightarrow \rho^2 &= \frac{1}{d^{2N}} \sum_{P,Q} c_P c_Q P Q \\ \Rightarrow \text{Tr}(\rho^2) &= \frac{1}{d^N} \sum_P c_P^2 = \frac{1}{d^N} \sum_P |\langle\psi|P|\psi\rangle|^2. \end{aligned}$$

Stabilizer states, which are common eigenstates of a maximal set of mutually commuting Pauli operators ( $d^N$ -many), have  $\langle\psi|P|\psi\rangle = \pm 1$  for  $d^N$  Paulis, out of a total of  $d^{2N}$  many Paulis. That means stabilizer state is a highly concentrated

<sup>9</sup>The term *magic* probably came from magic-state distillation, a purification protocol which allows a qubit to increase the polarization only along certain ‘magic’ directions [164].

<sup>10</sup>probability of finding  $P$  in the representation of the state  $|\psi\rangle$  in Pauli basis.

probability distribution, and thus is easily simulable [148, 149, 151, 158, 166, 167]. In contrast, a Haar-random state has approximately equal weight on all  $P$ 's [168]. Thus, a natural way to quantify the nonstabilizerness of a state  $|\psi\rangle$  is by assessing the spread of this distribution.

### Stabilizer Rényi entropy

**Definition 4.** The  $q$ -th order stabilizer Rényi entropy of a state  $|\psi\rangle$  is defined as the Rényi entropy on the probability distribution formed by the Pauli expectation value [169]:

$$M_q(\psi) = \frac{1}{1-q} \log_2 \sum_{P \in \mathcal{P}_N} \frac{|\langle \psi | P | \psi \rangle|^{2q}}{d^N}. \quad (2.80)$$

The SRE is zero for the stabilizer state, as exactly  $d^N$  expectation values are  $\pm 1$ . As the Clifford group stabilizes the Pauli group, any action of unitary  $U \in \mathcal{C}_N$  keeps the Pauli group intact,  $U^\dagger P U = \pm Q$ , where  $Q \in \mathcal{P}_N$ . Thus, SRE is conserved under Clifford operations. Moreover, it is additive,

$$M_q(|\psi\rangle \otimes |\phi\rangle) = \frac{1}{1-q} \log_2 \sum_{P_1 \in \mathcal{P}_N^1, P_2 \in \mathcal{P}_N^2} \frac{|\langle \psi | P_1 | \psi \rangle|^{2q} |\langle \phi | P_2 | \phi \rangle|^{2q}}{d_1^N d_2^N} = M_q(|\psi\rangle) + M_q(|\phi\rangle).$$

Monotonicity holds only for  $q \geq 2$  [170], while it can be violated when  $q < 2$  [171]. Importantly, values of  $q > 1$  and  $q < 1$  highlight different properties of stabilizer Rényi entropies (SREs) and correspond to distinct quantum computational applications. Specifically,  $q < 1$  SREs are connected to the number of stabilizer state superpositions required to represent a given quantum state, which determines the resource cost for Clifford-based simulation algorithms and fault-tolerant state preparation. In contrast,  $q > 1$  SREs quantify the distance to the nearest stabilizer state, serving as a measure of the cost in Pauli-based fidelity certification [172].

SRE is related to many quantum information protocols, for example, it relates to phases of error-corrected circuits [173], quantifies the entanglement spectrum [174], bounds fidelity estimation [175–177], characterizes the robustness of shadow tomography [178], and characterizes pseudorandom states [179, 180]. It also provides a lower bound on T-gates [169], and experimentally computable [173].

## 2.7.6 Sum negativity and mana

The discrete Wigner function offers a robust resource measure. Early work on bound magic states [181] used this framework to show that negativity in a state's Wigner representation is a necessary condition for magic state distillation. However, this negativity was not initially explored as a quantitative measure of resource.

Intuitively, one expects that a state exhibiting only slight negativity should possess less magic than a state with large negative values in its Wigner function. This intuition is formalized in later work [85], which demonstrates that the sum of the negative entries in the Wigner function defines a valid magic monotone, capturing the degree of non-classicality relevant for quantum computation.

### Sum negativity

**Definition 5.** The sum negativity of a state  $\rho$  is the sum of the negative elements of its Wigner function:

$$\text{sn}(\rho) = \sum_{\vec{V} | W_\rho(\vec{V}) < 0} |W_\rho(\vec{V})| = \frac{1}{2} \left( \sum_{\vec{V}} |W_\rho(\vec{V})| - 1 \right). \quad (2.81)$$

Due to the discrete Hudson's theorem, free states have a magic zero, and, as Clifford unitaries keep the Wigner function unchanged (Eq. (2.78)), monotonicity is also satisfied. However, the composition law of sum-negativity follows,

$$\text{sn}(\rho^{\otimes n}) = \frac{1}{2} [(2 \cdot \text{sn}(\rho) + 1)^n + 1], \quad (2.82)$$

i.e., a linear increase in the number of resource states implies an exponential increase in the amount of resource according to the measure. One can make such dependence linear, essentially putting a log in front of similar quantities, which defines another measure called mana:

### Mana

**Definition 6.** The mana of a quantum state  $\rho$  is

$$\text{Mana}(\rho) = \log \left( \sum_{\vec{V}} |W_\rho(\vec{V})| \right) = \log(2 \cdot \text{sn}(\rho) + 1). \quad (2.83)$$

As log is a monotonic function, Mana also follows the necessary properties of a magic monotone. It is also additive, i.e.,  $\text{Mana}(\rho \otimes \sigma) = \text{Mana}(\rho) + \text{Mana}(\sigma)$ .

## 2.8 Non-Markovianity

In the classical setting, a Markovian process is a type of stochastic (random) process that satisfies the Markov property, i.e., the future state of the process depends only on the present state, and not on the sequence of past states. More mathematically, three random variables  $XYZ$  with joint probability distribution  $p_{XYZ}$  form a Markov chain ( $X \rightarrow Y \rightarrow Z$ ), if the mutual information does not remember information about  $Z$ ,

$$\text{Markov chain: } I(X : Z|Y) = I(X : Y|Z) - I(X : Y) = 0. \quad (2.84)$$

Here,  $I(A : B) = H(A) - H(A|B)$ , where  $H(A)$  is the Shannon entropy. Equivalently, the conditional probability distribution  $p_{Z|X=x}(z) := p_{ZX}(z, x)/p_X(x)$  satisfy

$$p_{Z|X=x}(z) = \sum_y p_{Z|Y=y}(z) p_{Y|X=x}(y). \quad (2.85)$$

When such past-future independence fails, i.e., some history is retained, the process is referred to as non-Markovian. The relevance of quantum non-Markovianity, or the characterization of memory effects, becomes particularly important in open quantum

systems. A quantum system is called *open* when it has interaction with its environment [182]. As time evolves, it builds up correlations, such as entanglement, and destroys them through decoherence and dissipation, which are detrimental to quantum devices. Thus, it is essential to determine whether any past information is lost over time or not, as quantified by witnesses of non-Markovianity. However, a universal definition of non-Markovianity has remained elusive; hence, we will discuss some different witnesses, which, together, can provide a hint about non-Markovianity.

### 2.8.1 Quantum channels

Open system dynamics is represented by quantum dynamical maps, also known as quantum channels, which we discuss in this section before exploring non-Markovianity witnesses. Linear maps, e.g.,  $\Phi, \Lambda$ , represent a physical evolution map from density matrices to density matrices. A linear map  $\Phi: \mathcal{H}(A) \rightarrow \mathcal{H}(B)$  is called:

$$\text{positive : } \Phi(\rho) \geq 0, \quad \forall \rho; \quad (2.86)$$

$$k - \text{positive : } \Phi \otimes \text{id}^C \geq 0, \quad \dim \mathcal{H}_C = k; \quad (2.87)$$

$$\text{completely - positive : } k - \text{positive } \forall k. \quad (2.88)$$

Additionally, a physical evolution must preserve the trace, and such completely-positive trace-preserving (CPTP) maps are called *quantum channels*, denoted as  $\mathcal{Q}$ .

There are three well-known representations of quantum channels, which we use to express quantum processes throughout the thesis on several occasions..

**1. System-Environment representation.** The evolution of an open quantum system  $A$  with environment  $E$ , is modeled by the unitary evolution of the joint system  $U^{AE}$ . Given an uncorrelated initial state of environment  $|0\rangle\langle 0|^E$ , the dynamics of the system are represented by the reduced density matrix:

$$\rho^A \mapsto \Phi(\rho^A) = \text{Tr}_{E'} \left[ U^{AE} \left( \rho^A \otimes |0\rangle\langle 0|^E \right) U^{\dagger AE} \right], \quad (2.89)$$

where  $E'$  does not need to be the same system as  $E$ . Every completely positive trace-preserving (CPTP) map admits a unitary representation; that is, any physical evolution can be viewed as a unitary process acting on a larger system that includes the environment. CPTP maps effectively describe this evolution by tracing out the environment, whose degrees of freedom are not directly accessible.

**2. Kraus operator.**  $\Phi$  can be represented by set of matrices  $\{K_j\}$  satisfying  $\sum_j K_j^\dagger K_j = \mathbb{I}^A$  in the following form-

$$\rho^A \mapsto \Phi(\rho^A) = \sum_j K_j \rho^A K_j^\dagger. \quad (2.90)$$

**3. Choi matrix.** For any CP map  $\Phi$  maps half of a maximally entangled vector  $|\phi^+\rangle^{AA'} = \sum_{j=1}^{d_A} |j\rangle^A |j\rangle^{A'}$  of  $\mathcal{H}^A \otimes \mathcal{H}^{A'}$  to another system  $B$ , it can produce the bipartite operator known as Choi matrix of  $\Phi$ ,

$$J_\Phi^{AB} := \text{id}^A \otimes \Phi(\phi^+). \quad (2.91)$$

Conversely, any bipartite positive semi-definite operator  $J^{AB}$  corresponds to a CP map  $\Phi$  given by

$$\Phi_J(\rho) = \text{Tr}_A [J^{AB}(\rho^T \otimes I^B)], \quad (2.92)$$

where  $\rho^T$  denotes the matrix transpose with respect to some basis of  $A$ . If this basis can be chosen as same as used for  $|\phi^+\rangle^{AA^*}$ , then  $\Phi_{J_\Phi} = \Phi$ , and thus defines Choi-Jamiołkowski Isomorphism between the CP maps  $(\Phi)$  ( $A \rightarrow B$ ) and Choi matrices  $J_\Phi^{AB}$ .

**Unital map.** A special class of CP maps that act invariantly on the identity, i.e.,  $\Phi(\mathbb{I}) = \mathbb{I}$ , are called *unital* maps, which will be helpful to understand as it has a close connection to the CPTP map. For every map  $\Phi$ , its dual is the adjoint map fixed by the Hilbert-Schmidt inner product,

$$\text{Tr}(\Phi^\dagger(Y)X) = \text{Tr}(\Phi(X)Y), \quad \forall X \in \mathcal{B}(A), Y \in \mathcal{B}(B). \quad (2.93)$$

Thus, one can verify that

$$\Phi \text{ is trace preserving} \Leftrightarrow \Phi^\dagger \text{ is unital}. \quad (2.94)$$

*Proof.* It is worth understanding how two-way implication works.

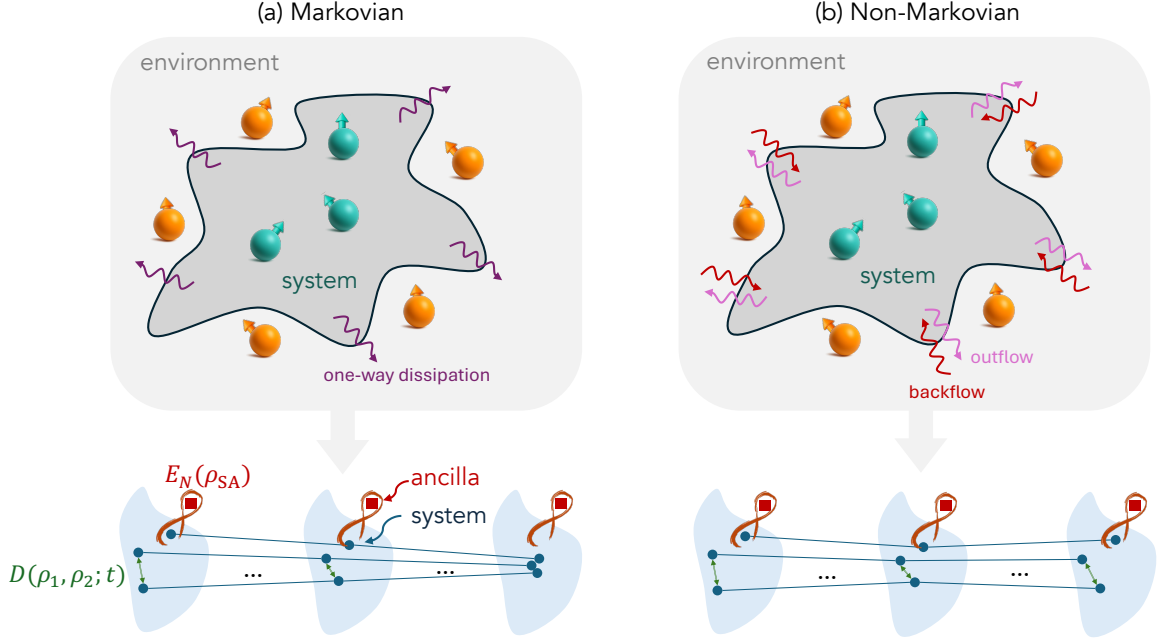
- $\Phi$  is trace preserving  $\Rightarrow \text{Tr}(\Phi(\rho)) = \text{Tr}(\rho)$ . Now, substituting  $X = \rho, Y = \mathbb{I}$  at Eq. (2.93),  $\text{Tr}(\rho\Phi^\dagger(\mathbb{I})) = \text{Tr}(\Phi(\rho)\mathbb{I}) = \text{Tr}(\rho) \Rightarrow \Phi^\dagger(\mathbb{I}) = \mathbb{I}$ .
- $\Phi^\dagger$  is unital  $\Rightarrow \text{Tr}(X \underbrace{\Phi^\dagger(\mathbb{I})}_{\mathbb{I}}) = \text{Tr}(\Phi(X))$ : Trace-preserving.

Note that we do not use any particular representation of the  $\Phi$ , yet we can prove the above statement.

## 2.8.2 Witnesses of non-Markovianity

The memory effect of a non-Markovian process is sometimes characterized by the backflow of information from the environment to the system, leading to revivals of information [183, 184][see Fig. 2.8(b)], although the precise relation between such revivals and backflow is a matter of ongoing investigations [185]. Alternatively, non-Markovianity can be described in terms of the divisibility property, positive or completely-positive, of the associated dynamical maps, where non-divisibility is linked to non-Markovian behavior [186–190]. Non-Markovian behavior is also sometimes related to nonexponential decays and dissipationless oscillations [191]. Non-Markovianity can also serve as a tool to probe quantum complexity [192]. To characterize the non-Markovianity, we use the following witnesses.

**Trace distance (BLP criterion).** As an open system evolves, it typically develops correlations with its environment and undergoes an irreversible loss of information, particularly when the system is weakly coupled to the environment. Under the strong coupling limit, the information might periodically be revived from the environment, which is known as the backflow of information. As the trace-norm is CP-contractive under a CPTP map, we can consider the trace distance as a measure to define the



**Figure 2.8.** Temporal flow of information in open Quantum Systems: (a) In the Markovian regime, the system experiences one-way dissipation, i.e., information flows irreversibly from the system to the environment. (b) In contrast, the non-Markovian regime exhibits backflow, where the environment can return information to the system, impacting its evolution. The trace distance  $D(\rho_1, \rho_2; t)$  (BLP criterion), which captures changes in distinguishability over time, reduces monotonically in Markovian dynamics, whereas it can increase again in non-Markovian dynamics. Similarly, the entanglement between system and ancilla  $E_N(\rho_{SA})$  (RHP criterion) only decreases in Markovian, but it can be revived in non-Markovian dynamics, showing the signature of memory effects arising from the backflow of information.

distance between the density of matrices, which can define the distinguishability. The trace distance between two different initial states  $\rho_1(0)$  and  $\rho_2(0)$ , defined as

$$D(\rho_1, \rho_2; t) = \frac{1}{2} \|\rho_1 - \rho_2\|_1. \quad (2.95)$$

Here,  $\|M\|_1$  is the trace norm  $\text{Tr}[\sqrt{M^\dagger M}]$ . A map  $\Phi$  is said to be Markovian if it follows the data-processing inequality [Eq. (2.22)] for trace distance, i.e., tends to reduce the distinguishability between any two states continuously.

$$D(\rho_1, \rho_2) \geq D(\Phi(\rho_1), \Phi(\rho_2)), \quad \forall t. \quad (2.96)$$

Thus, according to the Breuer–Laine–Pilo (BLP) criterion, breakdown of the monotonicity of trace distance between any two orthogonal initial states implying a temporal growth of distinguishability can be considered a hallmark of non-Markovianity [183]. Note that this criterion is only sufficient, but not necessary, since it might fail to witness non-Markovianity for some non-unital channels [193, 194] and unital channels [190].

**Logarithmic negativity (RHP criterion).** Entanglement is considered one of the pivotal resources in quantum computing [195]. One of the entanglement measures, known as logarithmic negativity [196, 197], can also be used to identify non-Markovian



evolutions following the Rivas–Huelga–Plenio (RHP) criterion [188]. Consider an initial maximally entangled state between the system and an ancilla. For the example of a system given by a qubit, this state can be written as

$$|\Phi\rangle = \frac{1}{\sqrt{2}}(|\uparrow\uparrow\rangle + |\downarrow\downarrow\rangle) \Rightarrow \rho_{\text{SA}}(0) = |\Phi\rangle\langle\Phi|. \quad (2.97)$$

Evolving the system in time with a quantum evolution operator,  $\rho_{\text{SA}}(t) = (\mathcal{E}_t \otimes \mathbb{I})\rho_{\text{SA}}(0)$ , and partially transposing the joint density matrix, one can calculate the logarithmic negativity as

$$E_N(\rho_{\text{SA}}) = \log_2 \|\rho_{\text{SA}}^{T_S}\|_1. \quad (2.98)$$

Here,  $(\ )^{T_S}$  denotes the partial transposition with respect to the system S. As Markovian dynamics will always destroy quantum correlations between ancilla and system, a temporal increase in logarithmic negativity witnesses non-Markovianity [188, 197]. Note that nonmarkovianity witnessed by the BLP criterion (which checks P-indivisibility) also implies it will be detected in the RHP criterion (related to CP-indivisibility), but not the other way around.

**Purity.** The purity of a state is defined as  $\text{Tr}[\rho^2(t)]$ . For finite-dimensional Hilbert spaces, the purity is monotonically decreasing during dynamics generated by a Lindbladian if and only if the Lindbladian is unital, i.e.,  $\mathcal{L}_t[\mathbb{I}] = 0$  [198]. Note that a Lindbladian form (where decay rates are absorbed in jump operators, thus requiring positive decay rates) guarantees a completely-positive-trace-preserving map (CPTP) [183, 190]. If the dynamics induced by any unital Lindblad-like generator<sup>11</sup> leads to an increasing purity, then the associated map is non-CPTP in the time interval. Thus, the increase in purity can witness non-Markovianity for unital maps [199].

We want to highlight that although purity-increasing maps can be unital, they cannot be represented in the exact Lindbladian form, as the CPTP conditions need to be conserved there. Thus, such non-Markovian features do not contradict the ‘if and only if’ conditions for unital Lindbladians mentioned above. For example, such a scenario can appear in the presence of a negative decay rate in a Lindblad-like master equation.

To summarize, if the map is unital, an increasing purity is a good non-Markovianity witness, but if the map is non-unital, one has to employ different measures [194] to reveal the non-Markovianity originating from the non-unital aspect of the dynamics. Regardless of the purity’s ability to reveal non-Markovianity, it remains an important figure of merit, e.g., in optimal control: a control pulse can be identified as robust against imperfections when the purity of a disorder-averaged state revives near the completion of the pulse [200, 201].

Non-Markovian states have been identified as valuable resources for various quantum information tasks such as quantum state redistribution [202], secure communication [203], and state deconstruction [204]. These applications motivate the treatment of non-Markovianity as a quantum resource [205, 206].

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<sup>11</sup>By Lindblad-like, we mean the form where the decay rate is not absorbed in the jump operators.



## 2.9 Summary

This chapter introduces the concept of quantum resources, which serves as the central theme of this thesis. We begin by outlining the desirable properties of resource measures, including axiomatic approaches, as well as practical formulations such as entropic and geometric measures. We then explore the resource theory of entanglement, with a particular emphasis on multipartite entanglement and its quantification using tools like quantum Fisher information and the geometric measure of entanglement—topics that play a key role in Chapters 4, 5, and 6. The chapter also covers the notion of nonstabilizerness and its quantifiers, including Stabilizer Rényi entropy and mana, which are relevant for Chapters 7 and 9. Finally, we discuss non-Markovianity as a crucial resource in open quantum systems and review non-Markovianity witnesses, providing supporting background for Chapter 8.



## Chapter 3

# Quantum optimization

### *quantum solutions for optimization problems*

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Quantum computing is increasingly regarded as a transformative technology for both scientific research and industrial applications. This view is supported by sustained efforts within the scientific community and by substantial investments from governments and industry stakeholders. Among the most promising applications of quantum computing is quantum optimization, a new computational paradigm that has drawn significant attention due to its potential impact across a wide range of fields.

Optimization problems arise naturally in many areas, including logistics, finance, energy systems, drug discovery, etc. These problems often become extremely difficult to solve as their size and complexity grow. While classical algorithms have advanced considerably, they continue to face limitations when dealing with large-scale or highly constrained problems. Therefore, any improvement, whether from classical heuristics, machine learning techniques, or quantum approaches, can be highly valuable.

The key motivation behind quantum optimization lies in the fundamentally different way quantum computers process information. Quantum algorithms can make use of quantum phenomena such as superposition and entanglement, allowing them to explore solution spaces in ways that are not possible for classical systems. This has led to the belief that quantum devices could offer new routes to solving problems that are currently intractable, or at least deliver improvements in speed or accuracy.

Although large-scale, fault-tolerant quantum computers are still under development, significant attention is being directed toward identifying near-term advantages. Both academic researchers and industry practitioners are actively evaluating whether quantum optimization methods can show benefits across several important metrics: the quality of the solutions, the time required to reach them, the overall cost of computation, and the general applicability of the approach to different problem types. These questions are central to understanding when and where quantum computing can provide a meaningful advantage over classical methods.

In this chapter, we begin by introducing the computational complexity associated with optimization problems to understand where the challenging problems lie (Sec. 3.2). We then present quantum optimization algorithms that are employed later in this thesis, including Quantum Annealing (QA) and the Quantum Approximate Optimization Algorithm (QAOA) in Sec. 3.3. In Sec. 3.6, we provide an overview of how quantum algorithms can be implemented on actual quantum computing hardware. The review by Abbas *et al.* [22] is our primary reference for this chapter.

## 3.1 Classical optimization

Optimization problems involve identifying the optimal solution from a potentially vast set of candidates, often under complex constraints. Mathematically, these problems are framed as the minimization or maximization of an objective function  $f(\vec{x})$ ,

$$\min_{\{\vec{x}\}} f(\vec{x}) \quad \text{or,} \quad \max_{\{\vec{x}\}} f(\vec{x}), \quad (3.1)$$

where the search domain  $\{\vec{x}\}$  can be discrete or continuous, constrained or unconstrained.

Here are some examples to understand how it is often structured.

- In portfolio optimization, the goal is to determine the optimal allocation of capital across a set of assets to maximize expected return while minimizing risk.
- Job scheduling problem in Operations Research aims to minimize total completion time given multiple jobs are scheduled across machines.
- In drug discovery, determining the lowest-energy 3D structure of a molecule is the goal.
- In power-grid optimization, the goal is to balance supply and demand to minimize loss and maintain stability.
- In many-body physics, finding the ground state can also be considered an optimization problem, for example, when using the variational principle to obtain the best solution.

The main challenge of such problems is to find a solution or a set of solutions efficiently. However, for some combinatorial optimization problems, the required effort (one can think of it as resources, but in a classical sense) using a known classical algorithm may scale exponentially with problem size, in a worst-case scenario. One may expect that a quantum computer can provide some quadratic speedup, though it may remain an exponential runtime. However, in classical optimization, one is not always worried about the worst-case scenario, and when considering a concrete problem instance, the story can be different. Many classical algorithms and heuristics can obtain almost optimal solutions in a reasonable time, even for large problems, for example, the travelling salesperson problem (TSP) [207–209]. The problem does not necessarily lie in the number of variables to optimize, as one may sometimes claim, because there are hard problems with fewer than 100 variables as well [210, 211]. The key mantra one should keep in mind is that although quantum optimization algorithms will not necessarily improve performance for all instances of a problem, they can exploit some *quantumness* that can potentially improve performance for some problem instances, if used wisely, and thus improve overall capabilities in optimization.

## 3.2 Computational complexity classes

The difficulty of quantum optimization largely depends on the nature of the optimization problems it addresses. Therefore, it is helpful to categorize problems into different complexity classes based on the computational resources they require, such as

time and memory. In theoretical computer science, these complexity classes serve as a framework for analyzing problem difficulty and provide a systematic way to approach and compare different problem types. While complexity theory primarily evaluates classical computational difficulty, it also offers insights into areas where quantum algorithms might outperform classical ones in terms of resource efficiency.

However, demonstrating a clear advantage of quantum computing over classical methods is not straightforward. Complexity theory focuses on worst-case scenarios, which do not always reflect the practical performance of algorithms. Real-world instances are often much easier than the hardest theoretical cases [212, 213]. Consequently, even if quantum algorithms do not outperform classical ones in worst-case scenarios, they may still offer significant benefits on average-case instances.

This distinction between worst-case and average-case performance suggests that the notion of quantum advantage should extend beyond traditional complexity-theoretic comparisons. In this context, we will explore various complexity classes, examine the types of problems they encompass, and highlight corresponding quantum algorithms. This overview will help identify problem domains where quantum approaches hold the potential for meaningful advantages.

### 3.2.1 Exact solutions

A decision problem involves deciding whether there exists a particular solution to the problem or not.

#### Decision problem

**Definition 7.** A decision problem is a binary function  $f_n : \{0, 1\}^n \rightarrow \{0, 1\}$  where  $f_n(x) = 1$  if and only if  $x \in L$ , which is a set of “yes” instances.

Note that a decision problem only cares about the existence of a particular solution, and not the actual solution itself. In a practical scenario, finding an actual solution is often necessary, which leads to relational problems.

#### Relational problem

**Definition 8.** Given a relation  $R \subseteq \{0, 1\}^n \times \{0, 1\}^n$  between input  $x$  and output  $y$ , the relational problem is about finding a  $y$ , that satisfies the relation, i.e.,  $(x, y) \in R$ .

For example, consider the MaxCut problem, where the goal is to bipartition the nodes of the graph in such a way that the maximum number of edges is traversed between the two sets (see Fig. 4.1). In such problems, the *decision problem* asks whether there exists a cut of size greater than a given threshold  $d$ , while the *relational problem* seeks the exact configuration, that is, which nodes belong to each partition, assuming the cut size is  $d$ . In this context, we can represent the input as  $x = d$ , and the output as  $y$ , denoting the corresponding partition configuration. It is important to note that the decision problem already encapsulates the computational complexity of the relational problem. This is because one can, in principle, solve the relational version by iteratively applying the decision problem; for instance, by successively removing edges and checking whether the maximum cut size remains unchanged.

Given such problem classes of decision and relational problems, one can define several complexity classes in terms of solvability.

### *Easy problems when asked for **exact** solution*

**Definition 9.** Concerns about whether it needs *polynomial* resources to obtain the exact solution.

#### Under deterministic computation

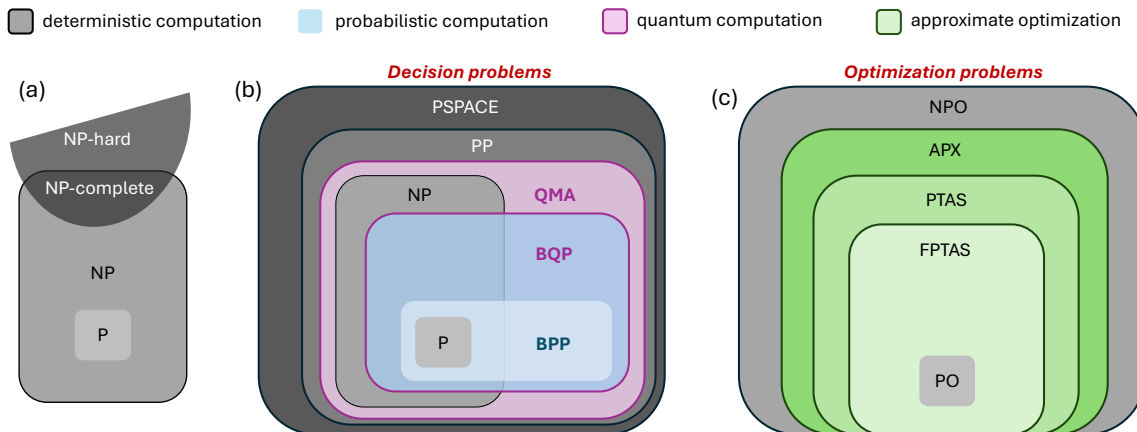
- Efficiently solvable by a deterministic machine in polynomial *time* –
  1. **P**: for decision problem,
  2. **FP**: for relational/functional problem,
  3. **PO**: for the set of optimization problems,  $\mathbf{PO} \subset \mathbf{FP}$ .
- Efficiently solvable by a deterministic machine in polynomial *space* – **PSPACE**, but no restriction on time. Thus, **PSPACE** is larger than **P**-time.

#### Under probabilistic computation

Efficiently solvable in polynomial time, with an error probability  $\leq 1/3$

- **BPP**: when solved in a classical probabilistic machine (**b**ounded-**e**rror **p**robabilistic **p**olynomial time), thus  $\mathbf{P} \subset \mathbf{BPP}$ .
- **BQP**: when solved in a quantum computer.  $\mathbf{BPP} \subset \mathbf{BQP}$ , i.e., there exists problem in **BQP**, which are not in **BPP**, example: Recursive Fourier Sampling [214].

**BQP** class already provides a set of problems where quantum advantage can be achieved, as it contains problems outside **BPP**. However, there are more useful problems for optimization, which are outside **P**, and hence hard to solve due to the



**Figure 3.1.** The inclusions of various complexity classes concerned with decision problems (b) and optimization problems (c). The well-known complexity classes concerning only deterministic computation are shown in (a).

unavailability of polynomial-time algorithms. These problems are also a test bed for quantum algorithms to be successful, such as Shor's algorithm [66].

### Hard problems when asked for *exact* solution

#### Definition 10. In classical machines.

- Although finding a solution is difficult, *checking* if it is a solution can be done in polynomial time in classical machines
  1. **NP**: for decision problem (*nondeterministic polynomial* time),
  2. **FNP**: for relational/ functional problem,
  3. **NPO**: for optimizational problem,  $\text{NPO} \subset \text{FNP}$ .
- **NP-complete**: If all other problems in **NP** can be efficiently reduced to it with a polynomial overhead in time. E.g.,- Travelling salesperson problem, graph coloring [215].
- **NP-hard**: If it is at least as hard as the hardest problems in **NP**, but not necessarily inside **NP** (see Fig. 3.1a).

Thus,  $\text{NP-complete} \subset \text{NP}$  and  $\text{NP-complete} \subset \text{NP-hard}$ .

#### In quantum machines.

- **QMA**: (*quantum Merlin-Arthur*) where the 'yes' instance can be verified efficiently and with high probability in a quantum computer (see Fig. 3.1b for its inclusion).

**Average-case scenarios.** Although one does not expect the quantum computer to solve **NP-hard** problems in the worst case, in the average-case settings, it can gain some advantage over a classical solver. In this context, two important concepts should be mentioned.

(a) **Overlap Gap Property (OGP)**: Every two solutions that are *close* (within an additive error) to optimality are either close to each other or far from each other, thus exhibiting a fundamental topological discontinuity of the set of distances of near-optimal solutions [216]. Such behavior introduces average-case hardness in classical algorithms, though it is still possible to gain superpolynomial speedups in quantum computers.

(b) **Computational statistical gaps**: The underlying statistical problem is information-theoretically possible, although no efficient algorithm exists, rendering the problem essentially unsolvable for large instances [217].

Another interesting complexity class is  $\#\text{P}^1$ , which emerges in the context of Gaussian Boson Sampling, and is interestingly related to stabilizer theory. It is con-

<sup>1</sup>The complexity class  $\#\text{P}$  consists of functions  $f : \{0,1\}^* \rightarrow \mathbb{N}$  for which there exists a non-deterministic polynomial-time Turing machine  $M$  such that for all inputs  $x \in \{0,1\}^*$ , the value  $f(x)$  equals the number of accepting computation paths of  $M$  on input  $x$ . Formally,  $f(x) = \#\text{accepting paths of } M(x)$ . Equivalently,  $\#\text{P}$  captures the complexity of counting the number of solutions to decision problems in **NP**.

cerned with counting the number of acceptable computation paths of an NP machine on input  $x$ . Its decision analogue is **PP**, and it is known that **QMA**  $\subseteq$  **PP** [218].

### 3.2.2 Approximate solutions

If finding exact solutions efficiently were considered the marker of success, the field of computing would not have been motivating enough. In reality, finding an approximate solution close to the optimal one is often sufficient. Such consideration leads to the study of approximability, i.e., how hard it is to approximate solutions to various problems. Although it may seem that allowing approximate solutions would reduce the complexity of a problem, it is not that straightforward. Hence, it is essential to understand the hardness of approximation.

A longstanding result called the **PCP-theorem** informally says that, for many problems, computing an approximate solution with arbitrary precision is as hard as computing the exact solution. The approximate ratio, a ratio between the approximate solution and the optimal solution, quantifies the success of such an approximate solution. For many **NP-hard** problems, there exist some known inapproximability bounds. A famous example is the MaxCut problem by the Goemans and Williamson algorithm, which achieves  $\approx 0.87856$  [219], and there cannot exist a polynomial-time algorithm that can achieve better than this [220] under the assumption of the Unique Games Conjecture<sup>2</sup>. In such cases, if quantum computers can go beyond it, that would imply that quantum computers can solve **NP-hard** problems or that the Unique Games Conjecture is false.

#### when asked for *approximate* solution

**Definition 11.** Depending on the approximability in terms of the approximation ratio,

- **APX:** *approximable* if there are polynomial-time approximation algorithms with approximation ratios bounded by some constant  $c$ . Example.- MaxCut, MAX-3-SAT.
- **PTAS:** *polynomial-time approximation scheme* if it is guaranteed to find a solution in polynomial-time within an arbitrary  $1 \pm \varepsilon$  factor of the optimum,  $\varepsilon > 0$ , thus the runtime is in polynomial time in system size. Example.- Euclidean Travelling Salesperson problem.
- **FPTAS:** *fully-polynomial time approximation scheme* if runtime is polynomial is both the problem size and  $1/\varepsilon$ . Example.- Knapsack problem.

Thus, **PO**  $\subseteq$  **FPTAS**  $\subseteq$  **PTAS**  $\subseteq$  **APX**  $\subseteq$  **NPO** [see Fig. 3.1(c)].

Understanding the complexity of approximability has consequences in many-body physics as well. For example, quantum **PCP-conjecture** [222–224] states that determining the ground state of a  $k$ -local hamiltonian is **QMA-complete** [225]. Even if one allows approximability, it can be **QMA-hard**.

<sup>2</sup>The conjecture postulates that problem of determining the approximate value of a certain type of game, known as a unique game, has **NP-hard** computational complexity [221].



The hardness of approximation suggests that quantum algorithms may outperform classical methods when there is a gap between known inapproximability bounds on classical algorithms and provable approximation factors.

### 3.3 Quantum optimization algorithms

Optimization problems are of different kinds, depending on the types of variables, objective functions, constraints, and allowed approximations. The choice of problem formulations also defines the solution strategy. Hence, some algorithms may only work on specific problem formulation classes. Quantum optimization algorithms are designed to solve optimization problems using a quantum hardware device, e.g.,- an analog or digital quantum computer. Thus, in such algorithms, it is crucial to encode the problem in quantum bits (or qubits), such that it is possible to exploit quantum mechanical phenomena to obtain (possibly) better solution than its classical counterparts. Grover search [226] and the Quantum Adiabatic Algorithm (QAA) [227] are the prominent examples of exact quantum optimization algorithms, which are expected to require Fault-Tolerant Quantum Computing (FTQC) [150] due to the resulting circuit sizes. In the era of noisy quantum computers [87], one may expect to exploit quantum subroutines to accelerate known classical approximation algorithms. For instance, Semi-Definite Programming (SDP) relaxations [228], or more general variants of binary optimization problems, may profit from Quantum SDP solvers [229, 230]. Similarly, the so-called Variational Quantum Algorithms (VQAs) [231] work well in the noisy hardware, though there is no provable advantage.

In this thesis, we consider a particular type of problem known as unconstrained discrete optimization problems. There also exists a large class of other optimization problems, for example, constrained discrete optimization, continuous optimization (includes convex and non-convex optimization), mixed-integer programming, and dynamic programming (see [22] for a review).

#### 3.3.1 Quadratic Unconstrained Binary Optimization (QUBO)

Quadratic unconstrained binary optimization (QUBO) is a foundational example of unconstrained discrete optimization [232–234], where the cost function is expressed as

$$\min_{x \in \{0,1\}^n} x^T Q x, \quad (3.2)$$

where  $x$  denotes the discrete decision variables,  $Q \in \mathbb{R}^{n \times n}$  is the cost matrix, and  $n \in \mathbb{N}$  is the number of decision variables. One can also consider higher-order polynomial objective functions known as PUBO (polynomial unconstrained binary optimization). Some ubiquitous problems, such as the maximum-cut (MaxCut) problem [see Fig. 4.1], are naturally formulated in this QUBO.

#### Complexity of QUBO problem

QUBO problems are **NP-hard**. Also **APX-hard**, i.e., there cannot exist a **PTAS** for QUBO unless **P=NP** [235].

A QUBO problem can be translated to an Ising-type Hamiltonian [232], where

finding an optimal solution is the same as finding the ground state of the Hamiltonian. Such a connection is fundamental to many quantum optimization algorithms. The variables  $x \in \{0, 1\}$  is substituted with spin variables  $z = 1 - 2x$ , where  $z \in \{-1, +1\}$ . By replacing the spin-variables  $z$  by the operator, Pauli-Z matrices, resulting in a diagonal Hamiltonian  $H \in \mathbb{R}^{2^n \times 2^n}$  and the problem looks similar to the Ising model [232]. Now, we will discuss two algorithms in detail: quantum annealing [227, 236] and quantum approximate optimization algorithm [81, 237].

### 3.3.2 Quantum annealing

Probably the first quantum computing approach introduced for solving QUBO-type problems was quantum annealing [227, 238, 239]. It is a quantum adiabatic algorithm method that specializes in finding the ground state of an Ising model by continuously evolving (*annealing*) from the known ground state of an *easy* Hamiltonian to the problem Hamiltonian. It is an exact optimization algorithm, reminiscent of homotopy methods of numerical mathematics<sup>3</sup>.

The known solution is chosen as the eigenstate of the so-called mixing Hamiltonian,  $\hat{H}_M$ , which should have an easy-to-prepare ground state  $|\psi_0\rangle$ , and also has a non-zero overlap with the ground state of the problem Hamiltonian  $\hat{H}_C$ . Moreover, it should be non-commuting with the problem Hamiltonian, such that they do not have common eigenstates, and allow exploration of several eigenstates (hence, it is called *mixing* Hamiltonian). A common choice is

$$\hat{H}_M = \sum_{i=1}^n X_i, \text{ where } X_i = \cdots \otimes \underbrace{\begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}}_{i^{\text{th}}} \otimes \cdots, \quad (3.4)$$

having ground state  $|+\rangle^{\otimes n} = \frac{1}{\sqrt{2^n}} \sum_{x \in \{0,1\}^n} |x\rangle$ , i.e.,  $n$ -qubit equal superposition state that has non-zero overlap with every computational basis state. Starting from  $|\psi_0\rangle = |+\rangle^{\otimes n}$ , quantum annealing slowly evolves the state according to the instantaneous Hamiltonian:

$$\hat{H}(t) = A(t)\hat{H}_C + B(t)\hat{H}_M. \quad (3.5)$$

This Hamiltonian  $\hat{H}(t)$  is slowly evolved from  $\hat{H}_M$  at  $t = 0$  to the cost Hamiltonian  $H_C$  at  $t = T$ , whose ground state encodes the solution to the target problem (see Fig. 3.2), following the time-dependent Schrödinger equation:

$$i \frac{d|\psi_t\rangle}{dt} = H(t) |\psi_t\rangle. \quad (3.6)$$

I.e.,  $B(t)$  is slowly reduced from  $B(0) = 1$  to  $B(T) = 0$ , and  $A(t)$  is slowly increased from  $A(0) = 0$  to  $A(T) = 1$  (for simplicity, we choose  $A(t) = t/T$  and  $B(t) =$

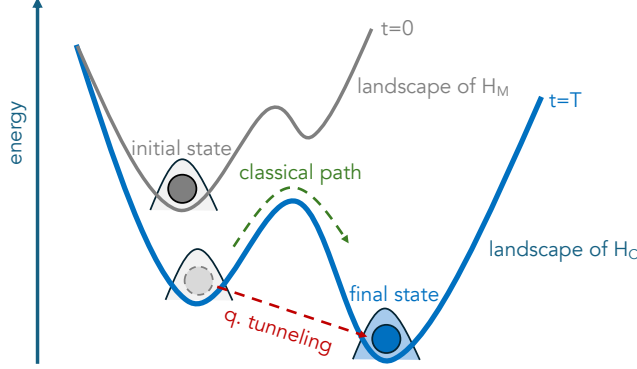
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<sup>3</sup>If one wishes to solve  $f(\mathbf{x}) = \mathbf{0}$ , then construct a system  $g(\mathbf{x}) = \mathbf{0}$ , whose solutions are known. For example, consider the *homotopy*:

$$H(\mathbf{x}, t) := (1 - t)g(\mathbf{x}) + tf(\mathbf{x}) = \mathbf{0}. \quad (3.3)$$

By continuation, trace the paths starting from the known solutions of  $g(\mathbf{x}) = 0$  to the desired solutions of  $f(\mathbf{x})$ . Such homotopy continuation methods are symbolic-numeric, where homotopy methods treat polynomials as algebraic objects and continuation methods use polynomials as functions [240].

$(1-t/T)$  throughout). According to the adiabatic theorem, if the time evolution is slow enough, the quantum state always remains in the ground state of the instantaneous Hamiltonian and reaches the solution state at the end of the sweep [236, 241].



**Figure 3.2.** Scheme of quantum annealing: starting from the ground state of the mixer Hamiltonian at  $t = 0$ , and with evolution, the cost landscape approaches  $H_C$ , and then through quantum tunneling it can cross the energy barrier to reach the final state in the absolute minima, which is otherwise harder to obtain through a classical path.

In practice, quantum annealing requires a very long sweep to successfully reach the solution state, i.e., the corresponding circuits are deep and error-prone. To execute quantum annealing on noisy digital devices with limited coherence time, one can discretize the continuous evolution as a sequence of gates, resulting in a Trotterized quantum annealing (TQA). A first-order Trotter–Suzuki decomposition transforms the continuous sweep, up to  $\mathcal{O}(\Delta t^2)$ , into

$$e^{-i\Delta t \hat{H}(t)} \simeq e^{-i\beta \hat{H}_M} e^{-i\gamma \hat{H}_C}, \quad (3.7)$$

where  $\beta = (1 - t/T)\Delta t$ , and  $\gamma = (t/T)\Delta t$ . Discretizing the entire quantum annealing schedule into  $p$  time steps requires  $\Delta t = T/p$ . The resulting discretized annealing schedule is

$$\gamma_s = \frac{s}{p}\Delta t, \quad \beta_s = (1 - \frac{s}{p})\Delta t, \quad (3.8)$$

where the time-factor of  $t/T$  is replaced by the layer-number  $s/p$ . Crucially, unlike in quantum annealing, here  $\Delta t$  and the total time  $T$  can be modified independently by changing  $p$ , which hands us an additional control knob for TQA.

The performance of annealing depends on the total sweep time  $T$ , and if  $T$  is large enough, with a suitable annealing schedule, the Adiabatic Theorem guarantees that the state  $|\psi_t\rangle$  always remains in the ground state of  $H(t)$ . To understand this in more detail, we estimate how close the time-evolved state is to the instantaneous ground state of  $H(t)$ . Let's denote the  $k$ th eigenstate of  $H(t)$  as  $|k(t)\rangle$  with eigenvalue  $\varepsilon_k(t)$ ,

$$\hat{H}(t)|k(t)\rangle = \varepsilon_k(t)|k(t)\rangle. \quad (3.9)$$

Here, denote  $|0(t)\rangle$  as the ground state, and other eigenstates are orthogonal to it. The above equation already gives us,

$$\langle j(t)| \frac{d}{dt} |k(t)\rangle = \frac{-1}{\varepsilon_j(t) - \varepsilon_k(t)} \langle j(t)| \frac{d\hat{H}(t)}{dt} |k(t)\rangle, \quad j \neq k. \quad (3.10)$$

Now, when the instantaneous state is written as a superposition of energy eigenstates,

$$|\psi(t)\rangle = \sum_j c_j(t) e^{i\Delta t \phi_j(t)} |j(t)\rangle, \quad \Delta t = T/p. \quad (3.11)$$

Then one can show that [242], the probability amplitude of being in the ground state is

$$c_0(t) \simeq 1 + \mathcal{O}(1/T^2), \quad (3.12)$$

and the error (i.e., contamination from higher excited states) is

$$c_{j \neq 0}(t) \simeq \frac{i}{T} [A_j(0) - e^{iT[\phi_j(t) - \phi_0(t)]} A_j(t)] + \mathcal{O}(1/T^2), \quad (3.13)$$

where  $\phi_j(t) \equiv \int_0^t ds \varepsilon_j(s)$ , and

$$A_j(t) \equiv \frac{1}{\Delta_j(t)^2} \langle j(t) | \frac{d\hat{H}}{dt} | 0(t) \rangle, \quad (3.14)$$

where  $\Delta_j(t) \equiv \varepsilon_j(t) - \varepsilon_0(t)$ . Thus, the adiabaticity condition is written as

$$\frac{1}{\Delta_j(t)^2} \left| \langle j(t) | \frac{d\hat{H}}{dt} | 0(t) \rangle \right| = \delta \ll 1 \Rightarrow T \sim \mathcal{O}(1/\Delta^2). \quad (3.15)$$

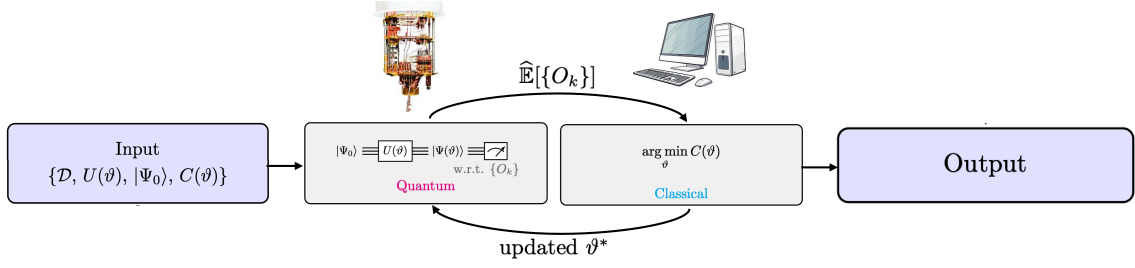
During annealing, the system can undergo phase transitions. In the spin-glass phases, the energy gap closes exponentially with system size, demanding an exponentially long runtime [243–245], which is often not possible, and hence we are limited to at most polynomial speedups.

In practical scenarios, quantum annealing is repeatedly run for a finite time [246], or potentially, diabatic annealing [247]. Thus, in its practical implementation, it often becomes a heuristic algorithm. In gate-based quantum computing, typical long annealing times imply notably long quantum circuits, which become challenging for NISQ devices.

### 3.3.3 Variational quantum algorithms

Due to the intrinsic noise and errors present in Noisy Intermediate-Scale Quantum (NISQ) devices, the behavior of the system often deviates from the idealized theoretical model. As a result, the parameter tuning that we aim to perform in an ideal setting does not translate directly to what is actually achieved on the hardware. In such cases, the so-called Variational Quantum Algorithms (VQAs) [231] have attracted a lot of attention. Such algorithms exploit the quantum hardware for the resource-intensive job of evaluating expectation values. Specifically, as Fig. 3.3 depicts, provided some problem data  $\mathcal{D}$ , a parametrized circuit  $U(\vartheta)$  is empirically evolved in the quantum computer, and then from measurement in some basis  $\{\mathcal{O}_k\}$  corresponding function  $C(\vartheta)$  is evaluated. Thereafter, classical optimization of the variational parameters is executed over the experimentally-measured cost function to obtain updated parameters, which are used as feedback for the circuit.

As finding the optimal parameters in VQA is considered **NP**-hard [249], the classical optimization becomes the main bottleneck in VQA. Yet, it can still be useful in the path of finding good solutions, even potentially offering speedups [250].



**Figure 3.3.** Variational algorithms iterate between the evaluation of experimental estimates of the expectations and classical optimization with updated parameters. (adapted from Ref. [248])

**Barren plateaus [251].** The training of a parametrized circuit, i.e., finding the optimal parameters, is a major challenge, especially if the cost landscape becomes flat, the gradients concerning the parameters are exponentially small, and the optimization process cannot proceed. Formally, a cost function  $C(\vartheta)$  exhibits a *probabilistic* barren plateau if, for some variational parameters  $\theta_i \in \vartheta$ , the variance of the partial derivative of the cost function vanishes exponentially in the number of qubits:

$$\text{Var}_{\vartheta}[\partial_i C(\vartheta)] \in \mathcal{O}(b^{-n}), \quad b > 1. \quad (3.16)$$

This inequality immediately implies (due to Chebyshev's inequality) that the probability that it will find a gradient different than a given constant  $c$  is upper bounded,

$$\Pr[|\partial_i C(\vartheta)| \geq c] \leq \frac{1}{c^2} \text{Var}_{\vartheta}[\partial_i C(\vartheta)]. \quad (3.17)$$

A type of VQA is the Variational Quantum Eigensolver (VQE) [252], which approximates the ground states of Hamiltonians and was initially proposed for quantum chemistry problems. Here, the job of a quantum computer is to evaluate  $\langle \Psi(\vartheta) | \hat{H} | \Psi(\vartheta) \rangle$  for a given parameter value of  $\vartheta$ . The variational principle guarantees that this expectation value is lower bounded by the ground state energy of  $H$ , and the equality is reached if and only if the ground state is reached:

$$\begin{aligned} \langle \Psi | \hat{H} | \Psi \rangle &= \sum_{\varepsilon_1, \varepsilon_2 \in \text{spec}(\hat{H})} \langle \Psi | \psi_{\varepsilon_1} \rangle \langle \psi_{\varepsilon_1} | \hat{H} | \psi_{\varepsilon_2} \rangle \langle \psi_{\varepsilon_2} | \Psi \rangle \\ &= \sum_{\varepsilon \in \text{spec}(\hat{H})} \varepsilon |\langle \psi_{\varepsilon} | \Psi \rangle|^2 \geq \sum_{\varepsilon \in \text{spec}(\hat{H})} E_0 |\langle \psi_{\varepsilon} | \Psi \rangle|^2 = E_0. \end{aligned} \quad (3.18)$$

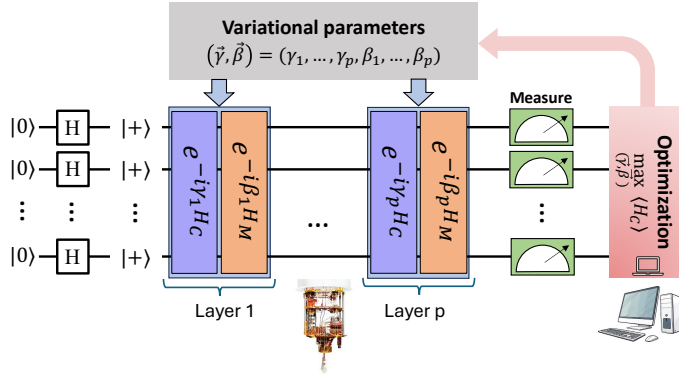
### 3.3.4 Quantum Approximate Optimization Algorithm (QAOA)

A special class of VQE is the Quantum Approximate Optimization Algorithm (QAOA) [81], which has been successfully applied to several types of optimization problems in the last decade. QAOA is motivated by both quantum annealing and VQE. The standard QAOA applies  $p$  layers of unitaries

$$\hat{U}_k(\boldsymbol{\beta}, \boldsymbol{\gamma}) = \exp(-i\beta_k \hat{H}_M) \exp(-i\gamma_k \hat{H}_C), \text{ where } k = 1, \dots, p, \quad (3.19)$$

to the ground state of a mixer Hamiltonian  $\hat{H}_M$ , e.g.,  $-\sum_i \hat{X}_i$  with  $\hat{X}_i$  being a Pauli operator) to create the trial state  $|\psi(\boldsymbol{\beta}, \boldsymbol{\gamma})\rangle$  (see Fig. 3.4). To obtain the optimal values of  $\boldsymbol{\beta} = (\beta_1, \dots, \beta_p)$  and  $\boldsymbol{\gamma} = (\gamma_1, \dots, \gamma_p)$ , which minimize the energy  $\langle \psi(\boldsymbol{\beta}, \boldsymbol{\gamma}) | \hat{H}_C | \psi(\boldsymbol{\beta}, \boldsymbol{\gamma}) \rangle$ , a classical optimizer is employed. The energy is measured on the quantum processor to find the best solution.

**Performance guarantee.** QAOA appears as a promising candidate despite being a heuristic algorithm, because it comes with worst-case performance bounds for certain problems and algorithmic settings. For example, the MaxCut problem on a 3-regular graph, there is a lower-bound on the approximation ratio of 0.692 for  $p = 1$  [81], 0.7559 for  $p = 2$ , and 0.7924 for  $p = 3$  [253]. Moreover, for  $d$ -regular graphs, numerically QAOA outperforms many known Semi-Definite Programming (SDP)-based relaxations when  $p \geq 11$  [254, 255].



**Figure 3.4.** Structure of QAOA ansatz. Every layer consists of the cost Hamiltonian ( $\hat{H}_C$ ) and the mixer Hamiltonian ( $\hat{H}_M$ ) with corresponding variational parameters  $\{\beta_k, \gamma_k\}$  for  $k = 1, \dots, p$ .

Similar to any hybrid optimization algorithm, QAOA is also bottlenecked by the training of variational parameters. Hence, considerable research has been done on different training strategies for QAOA, e.g.,-

1. initialize with random parameters and optimize them by evaluating the cost function multiple times (conventional approach).
2. bound-search the parameter space of  $(\vec{\beta}, \vec{\gamma})$  to certain regimes, depending on the symmetry of  $\hat{H}_C$  and  $\hat{H}_M$  (e.g., time-reversal symmetry,  $\mathbb{Z}_2$  symmetry) [237].
3. using Physics-inspired initialization, e.g.,-
  - (a) Trotterized quantum annealing inspired parameters [256].
  - (b) linearly interpolate the curve formed by optimized parameters at depth  $p$  to extract a set of initial parameters for level  $p + 1$  [237, 257].
  - (c) machine learning based approaches [258, 259].
4. evaluating the cost-function classically using tensor networks or Clifford perturbation theory from empirically sampled solutions obtained in hardware, which is computationally easier than sampling the expectation value from the complete quantum state [260, 261].
5. using the concentration of parameters, where the optimal parameters become problem-instance independent, and reuse the parameters for multiple different

problem-instances by training on one of them [262–264]. Such a transfer of parameters can work for the problem of the same size and sometimes also for larger sizes. It is often helpful because one can train a small system classically and then apply it to a larger system where classical optimization is not possible. It also enables reasonable solutions to a problem quickly.

Recently, the Qiskit community has released a repository called *qaoa training pipeline* [265], which is a set of tools designed to generate optimal parameters for QAOA Ansatz circuits using classical means. In Chapter 6, we use the package to obtain optimal parameters for QAOA up to 76 qubits efficiently.

**Variants of QAOA.** Several variants of QAOA, depending on different mixers and initial states, have been proposed to achieve certain goals, see [266] for a review.

1. The Quantum Alternating Operator Ansatz encodes constraints into the mixer to preserve the constraints and to restrict the algorithm to feasible states [267].
2. Using a classical algorithm to obtain an approximate solution, and then using it to warm-start QAOA to further improve upon the obtained solution [268, 269].
3. Recursive QAOA uses the quantum computer to produce a sequence of reduced problems by fixing the strongest one, and at each step, QAOA estimates the correlation between variables. The process is iterated until the reduced problem becomes small enough for classical solvers [270].
4. ADAPT-QAOA [271] starts with a pool of candidate mixers. At each iteration, it selects the mixers from the pool that give the greatest gradient improvement to the cost function, instead of fixing the mixers beforehand.

Motivated by QAOA, the Mean Field Approximate Optimization Algorithm [272] is developed, where the QAOA circuit is approximated through mean-field approximations without any quantum effects.

## 3.4 Basics of quantum computation

In the upcoming sections, we will explore how to work with real quantum computers. To prepare for that, it is important to first introduce some fundamental concepts, terminology, and standard gate sets. Quantum computers aim to exploit the uniquely quantum mechanical properties of matter to process information in ways that classical computers cannot. Superposition enables quantum bits (qubits) to represent multiple states simultaneously, while entanglement allows for non-classical correlations that can be harnessed for computational advantages. A variety of physical platforms, including trapped ions [273], superconducting circuits [274], Rydberg atoms [275], and photonic systems [276], have been developed to realize quantum processors.



### 3.4.1 Quantum bits and gates

In classical computing, the fundamental bits of information are 0 and 1. Similarly, in quantum computation, the unit of information is encoded in two objects<sup>4</sup> They reside in the Hilbert space, where superposition is allowed. The two logical states, also known as qubits, defined in the two-dimensional Hilbert space  $\mathcal{H}$  can be written as basis matrices of a 2-dimensional system. One popular choice is considering the eigenstates of the Pauli-Z matrix as a qubit basis, which are also known as the computational basis:

$$|0\rangle = \begin{pmatrix} 1 \\ 0 \end{pmatrix}, \quad |1\rangle = \begin{pmatrix} 0 \\ 1 \end{pmatrix}. \quad (3.20)$$

This allows us to write a generic state of a single qubit in the continuous space of its superposition,

$$|\psi\rangle = \alpha |0\rangle + \beta |1\rangle, \quad \alpha, \beta \in \mathbb{C}, \quad |\alpha|^2 + |\beta|^2 = 1. \quad (3.21)$$

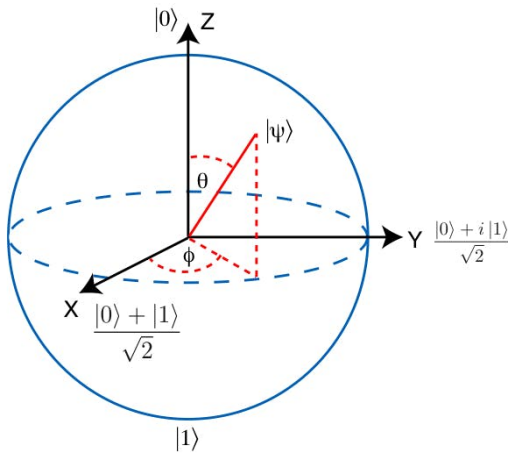
Such superposition principle implies that the probability amplitude defines the state, not the probability, which means  $(|0\rangle + |1\rangle)/\sqrt{2}$  and  $(|0\rangle - |1\rangle)/\sqrt{2}$  are different state, despite them having same probability of occurring  $|0\rangle$  and  $|1\rangle$ . In matrix notation, it is easy to see that they are orthogonal to each other, which is quite different from the classical world. Thus, the superposition of states allows us to define three sets of operators, whose eigenstates form one of the possible bases in the Hilbert space of a single qubit, with eigenvalues  $\pm 1$ . Pauli matrices define the operators-

$$Z = \begin{pmatrix} 1 & 0 \\ 0 & -1 \end{pmatrix}, \quad X = \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}, \quad Y = \begin{pmatrix} 0 & -i \\ i & 0 \end{pmatrix}. \quad (3.22)$$

We can represent a single-qubit state on the sphere, known as the Bloch sphere, where the coordinates denote the expectation value of  $\sigma = X, Y, Z$ , (see Fig. 3.5)-

$$|\psi\rangle = \cos \frac{\theta}{2} |0\rangle + e^{i\phi} \sin \frac{\theta}{2} |1\rangle, \quad \theta \in [0, \pi], \quad \phi \in [0, 2\pi). \quad (3.23)$$

$$(x, y, z) = (\sin \theta \cos \phi, \sin \theta \sin \phi, \cos \theta). \quad (3.24)$$



**Figure 3.5.** Bloch sphere: representation of the qubit state on sphere. At the opposite poles of the sphere along the axes, we find the eigenstates of the corresponding Pauli operator. Note that the orthogonal states  $|0\rangle$  and  $|1\rangle$  are  $\theta = \pi$  away from other, contrary to the geometrical understanding of  $\pi/2$ . This is an artifact of  $SU(2)$  symmetry of spins instead of  $SO(2)$ , and hence in Eq. (3.23), there is  $\theta/2$  instead of  $\theta$ .

Points lying on the surface of the sphere represent pure states, while mixed states

<sup>4</sup>It is not limited to two units only. A qudit generalizes the concept of a qubit ( $d = 2$ ) to a quantum unit of information that exists in a  $d$ -dimensional Hilbert space.



lie inside the sphere. Computational basis states (corresponding to the eigenvectors of Pauli-Z) lie at the poles:  $|0\rangle \leftrightarrow (0, 0, 1)$  and  $|1\rangle \leftrightarrow (0, 0, -1)$ . The eigenstates of  $X$  are denoted by

$$|+\rangle = \frac{1}{\sqrt{2}}(|0\rangle + |1\rangle), \quad |-\rangle = \frac{1}{\sqrt{2}}(|0\rangle - |1\rangle). \quad (3.25)$$

Similar to the logic-gates (OR, AND, etc) in classical computations, one can construct quantum gates in quantum computations that manipulate the state of qubits through mathematically defined, reversible transformations. These gates are represented by unitary matrices, which preserve the total probability (or norm) of the quantum system, a key feature of quantum mechanics. Quantum gates act on one or more qubits and are the building blocks of quantum circuits. The Pauli matrices, specifically Pauli- $X$ ,  $Y$ , and  $Z$ , are examples of single-qubit quantum gates.

Beyond the Pauli matrices, there are also other unitaries that are used as gate-sets for digital quantum computing, which allow the manipulation of qubits to modify their state. Any quantum algorithm can be reduced to a sequence of gates, called a quantum circuit, applied to a register of qubits.

Together with the identity operator, Pauli matrices form the basis of two-dimensional Hermitian matrices. Moreover, the Pauli matrices can also be used to generate a rotation of an arbitrary angle around the axes of the Bloch sphere:

$$\text{Rotation gates: } R_\sigma(\vartheta) = e^{-i\frac{\vartheta}{2}\sigma}, \quad \sigma \in \{X, Y, Z\} \quad (3.26)$$

Other single-qubit gates often used in quantum algorithms are :

$$\text{Hadamard } \hat{H} = \frac{1}{\sqrt{2}} \begin{pmatrix} 1 & 1 \\ 1 & -1 \end{pmatrix}, \quad \text{Phase } \hat{S} = \begin{pmatrix} 1 & 0 \\ 0 & i \end{pmatrix}, \quad \hat{T} = \begin{pmatrix} 1 & 0 \\ 0 & e^{i\pi/4} \end{pmatrix}. \quad (3.27)$$

They correspond to different quantum operations: the Hadamard-gate ( $\hat{H}$ ) creates a superposition state, the phase-gate ( $\hat{S}$ ) creates a phase in one of the superpositions, and the  $\hat{T}(\pi/8)$ -gates are used as non-Clifford resources.

If we expand this framework to a system composed of  $n$ -qubits, the Hilbert space of the system can be defined by the tensor product of the individual Hilbert spaces of each qubit:

$$\mathcal{H}_n = \bigotimes_{i=1}^n \mathcal{H}_i.$$

In such scenarios, there could be gates that act on more than one qubit simultaneously. In fact, two-qubit gates are necessary to create quantum correlation, for example, a CNOT gate. This gate conditionally inverts the state of the second (target) qubit depending on the state of the first (control). For example, if the control qubit is set to 0, the state of the target qubit is left unchanged; if the control qubit is set to 1, then the target qubit is flipped. Depending on the physical system used for their construction, it can be achieved by various types of gates, such as the controlled- $Z$  (CZ) or the iSWAP gate.

$$\text{CNOT} = \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 \\ 0 & 0 & 1 & 0 \end{pmatrix}, \text{CZ} = \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & -1 \end{pmatrix}, \text{iSWAP} = \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 0 & i & 0 \\ 0 & i & 0 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix}. \quad (3.28)$$

Notably, the CNOT gate is considered a fully entangling gate as it can transform a separable state into a maximally entangled one, resulting in the generation of one of the Bell states. For example, by acting on the Hadamard gate on the left qubit and then using the CNOT gate, we can obtain one of the Bell states,

$$|00\rangle \xrightarrow{H_0} \frac{1}{\sqrt{2}}(|00\rangle + |10\rangle) \xrightarrow{\text{CNOT}_{01}} \frac{1}{\sqrt{2}}(|00\rangle + |11\rangle). \quad (3.29)$$

A circuit is defined through a set of gates applied in a given sequence:

$$|\psi\rangle = \left( \prod_k \hat{U}_{i_k j_k}(\vec{\theta}_k) \right) |\psi_0\rangle, \quad (3.30)$$

where  $|\psi_0\rangle$  is an initial states (often it is  $|0\rangle^{\otimes n}$ ), and  $\hat{U}_{i_k j_k}(\vec{\theta}_k)$  denotes the unitary gate acting on qubits  $i_k$  and  $j_k$  with parameters  $\vec{\theta}_k$ , where  $k$  is the index of the gate.

### 3.4.2 Quantum measurements

Quantum measurement differs fundamentally from classical measurement in that it collapses a superposition of states into a single outcome, with probabilities governed by the system's wavefunction. Despite the probabilistic nature of quantum measurements, deterministic algorithms can still be implemented on a quantum computer. By executing the same algorithm multiple times and analyzing the resulting outcome distribution, one can extract reliable, deterministic results. Consequently, quantum computing often requires repeated execution of identical circuits to capture the full range of possible measurement outcomes. The accuracy and confidence in the results improve with the number of measurements performed.

Quantum measurement plays a central role in numerous quantum algorithms, such as quantum teleportation, error correction, and measurement-based quantum computation. In these protocols, measurement outcomes guide the choice of subsequent operations, allowing for the manipulation and transfer of quantum information. However, since measurements can introduce decoherence, reducing measurement-related errors is essential in designing reliable quantum algorithms. Additionally, post-processing methods are often required to interpret measurement results and correct for noise or inaccuracies.

To show the measurement process in quantum computing, we define a set of measurement operators  $\{\hat{M}_m\}$ , which act on the states being measured. The index  $m$  refers to the possible measurement outcomes. If  $|\psi\rangle$  is the state of the quantum system immediately before the measurement, the probability of obtaining the result  $m$  is:

$$p(m) = \langle \psi | \hat{M}_m^\dagger \hat{M}_m | \psi \rangle, \quad (3.31)$$

and since the operator  $M_m$  is not unitary, the state after the measurement needs to be normalized:

$$|\psi'\rangle = \frac{\hat{M}_m|\psi\rangle}{\sqrt{\langle\psi|\hat{M}_m^\dagger\hat{M}_m|\psi\rangle}} \quad (3.32)$$

The measurement operators satisfy the completeness relation:

$$\sum_m \hat{M}_m^\dagger \hat{M}_m = \mathbb{I} \quad \Rightarrow \quad \sum_m p(m) = 1. \quad (3.33)$$

Let us now examine a straightforward example: the measurement of a qubit in the computational basis. This scenario involves two possible outcomes, represented by the measurement operators:

$$\hat{M}_0 = |0\rangle\langle 0|, \quad \hat{M}_1 = |1\rangle\langle 1|. \quad (3.34)$$

Assuming the qubit is in a general state  $|\psi\rangle = \alpha|0\rangle + \beta|1\rangle$ , the probability of obtaining outcome 0 is:

$$p(0) = \langle\psi|\hat{M}_0^\dagger\hat{M}_0|\psi\rangle = \langle\psi|\hat{M}_0|\psi\rangle = |\alpha|^2. \quad (3.35)$$

Similarly, the probability of measuring outcome 1 is given by:

$$p(1) = |\beta|^2. \quad (3.36)$$

### Measuring observables

In quantum computing, information about a circuit is extracted by measuring observables, i.e., quantities associated with Hermitian operators, often constructed from Pauli matrices. Theoretically, the expectation value of an observable  $\hat{O}$  is obtained as:

$$\langle\hat{O}\rangle = \langle\psi|\hat{O}|\psi\rangle. \quad (3.37)$$

In an experiment, the first measurement is executed by evaluating the probability of 0 or 1 along the  $z$ -basis (projection to the  $Z$ ), and then it can be used to obtain the expectation value. For instance, the expectation value of the Pauli- $Z$  operator for a single-qubit state is:

$$\langle\hat{Z}\rangle = \langle\psi|\hat{Z}|\psi\rangle = p(0) - p(1). \quad (3.38)$$

Similarly, one can determine the expectation values of  $\hat{X}$  and  $\hat{Y}$  by first rotating the state into the respective bases and then measuring the computational basis. Hence, to measure the observable  $\mathcal{O}$ , one needs to find a unitary  $U$  such that  $\mathcal{O} = U^\dagger Z U$ . Then, projective measurement of the state is the same as measuring the expectation value  $\langle Z \rangle$  as the states are represented in the computational basis.

$$\langle\psi|\mathcal{O}|\psi\rangle = \langle\psi|U^\dagger Z U|\psi\rangle = \langle\psi'|Z|\psi'\rangle \simeq p(0) - p(1). \quad (3.39)$$

For example, in Tab. 3.1, we show how to measure different  $\mathcal{O}$ , using corresponding unitary rotations  $U$ .

This methodology naturally extends to multi-qubit systems. For example, to compute the expectation value of a product operator such as  $\hat{Z}_i\hat{Z}_j$  on a two-qubit state, one multiplies the outcomes of the individual measurements on qubits  $i$  and  $j$ . For

Measurement operator ( $\mathcal{O}$ )	Unitary transformation ( $U$ )
$Z$	$I$
$X$	$H$
$Y$	$HS^\dagger$
$Z \otimes I$	$I \otimes I$
$I \otimes Z$	SWAP

**Table 3.1.** Unitary transformation ( $U$ ) for measuring observables ( $\mathcal{O}$ ).

example,

$$\langle \psi | Z_1 Z_2 | \psi \rangle = p_1(0)p_2(0) - p_1(1)p_2(0) - p_1(0)p_2(1) + p_1(1)p_2(1), \quad (3.40)$$

where  $p_i$  denotes the probability corresponding to site  $i$ .

### Tomography

As previously discussed, one can measure specific operators to infer about the features of the quantum state. One can similarly reconstruct the full state. Another hurdles come in noisy or mixed settings that are described by a density matrix (or density operator), which generalizes the concept of a wavefunction. Given a statistical ensemble of  $n$  pure states  $|\psi_i\rangle \in \mathcal{H}$  with associated probabilities  $p_i$ , the density matrix is defined as:

$$\rho = \sum_{i=1}^n p_i |\psi_i\rangle \langle \psi_i| \quad (3.41)$$

The density operator is a Hermitian, positive semi-definite matrix with unit trace. When  $n = 1$ , the system is in a pure state. Otherwise, the state is mixed. One can distinguish between the two by evaluating:

$$\text{Tr}(\rho^2) = \begin{cases} 1 & \text{for pure states,} \\ < 1 & \text{for mixed states.} \end{cases} \quad (3.42)$$

A general mixed state of a single qubit can be decomposed in terms of the Pauli matrices:

$$\rho = \frac{1}{2} \left( \mathbb{I} + \sum_{i=1}^3 r_i \hat{\sigma}_i \right), \quad \hat{\sigma} = (\hat{X}, \hat{Y}, \hat{Z}). \quad (3.43)$$

Here, the vector  $r = (\langle \hat{X} \rangle, \langle \hat{Y} \rangle, \langle \hat{Z} \rangle)$  represent the Bloch vector. If  $|r| = 1$ , the state lies on the surface of the Bloch sphere and is pure. If  $|r| < 1$ , it corresponds to a mixed state residing inside the sphere. By estimating the expectation values of the Pauli operators, one can reconstruct the full density matrix of a qubit. This process is known as *quantum state tomography* [277].

## 3.5 Superconducting quantum computer

The fundamental ingredients for quantum computers are qubits, two-level quantum systems, robust control of gates, and the ability to measure the final output. However, to retain the quantum properties, it is also important to isolate the system while

allowing accessibility to manipulate qubits in terms of gates as well as measurements. Regardless of the physical system used for the physical realization of a qubit, a qubit Hamiltonian is generally given by the following form:

$$H = c_z(t) Z + c_x(t) X + c_y(t) Y, \quad (3.44)$$

where  $X, Y, Z$  are Pauli matrices, and  $c_x, c_y, c_z$  are their respective tunable coefficients. For example, when considered a Harmonic oscillator kind of Hamiltonian, the first two levels can form the computational basis states, i.e.,  $|0\rangle$  and  $|1\rangle$ , and are separated by an energy difference  $\hbar\omega_q$ , where  $\omega_q$  is the oscillator frequency. In this case,  $c_z = \hbar\omega_q$ . This term produces rotations around the  $z$ -axis of the Bloch sphere, representing phase gates. On the other hand, by manipulating the coefficients  $c_x(t), c_y(t)$ , a qubit can oscillate between the two states, performing arbitrary single-qubit rotations. Different physical systems can be used to implement a qubit, such as photons, electrons, ions, or neutral atoms. Rather than exploring all quantum platforms in detail, we focus on superconducting qubits, utilized in IBM's quantum devices, where our experiments are conducted. In this section, we provide a brief introduction to the topic, primarily following Krantz et al. [274] as the primary reference.

The superconducting qubits are implemented using electrical circuits, typically arranged in two-dimensional layouts, while the third dimension is reserved for interfacing with control and readout hardware. To understand how a superconducting qubit circuit works, it is useful to first understand how a linear LC resonant circuit works [Fig. 3.6(a)]. Due to the presence of both the capacitor (C) and inductor (L), the energy oscillates between the electric energy of C (think of it as kinetic energy of electrons) and the magnetic energy of L (potential energy of the oscillator). From here, one can see that the instantaneous energy is defined from the current and voltage of the capacitor or inductor,

$$E(t) = \int_{-\infty}^t V(t') I(t') dt'. \quad (3.45)$$

Similarly, we can define the flux as the time integral of the voltage

$$\Phi(t) = \int_{-\infty}^t V(t') dt'. \quad (3.46)$$

Now using the relations,  $V = L \frac{dI}{dt}$  and  $I = C \frac{dV}{dt}$ , we can write the energy terms for the capacitor and inductor in terms of the node flux:

$$T_C = \frac{1}{2} C \dot{\Phi}^2, \quad U_L = \frac{1}{2L} \Phi^2. \quad (3.47)$$

From here, we can define the Lagrangian as  $T_C - U_L$ , and can derive the Hamiltonian using Legendre transformation as [274],

$$H = \frac{1}{2} C V^2 + \frac{1}{2} L I^2, \quad \text{with } \omega = \frac{1}{\sqrt{LC}}. \quad (3.48)$$

We can further introduce the reduced charge  $n = Q/2e$  and reduced flux  $\phi = 2\pi\Phi/\Phi_0$  to obtain the Hamiltonian in terms of  $E_C = e^2/(2C)$  and  $E_L = (\Phi_0/2\pi)^2/L$ , in the

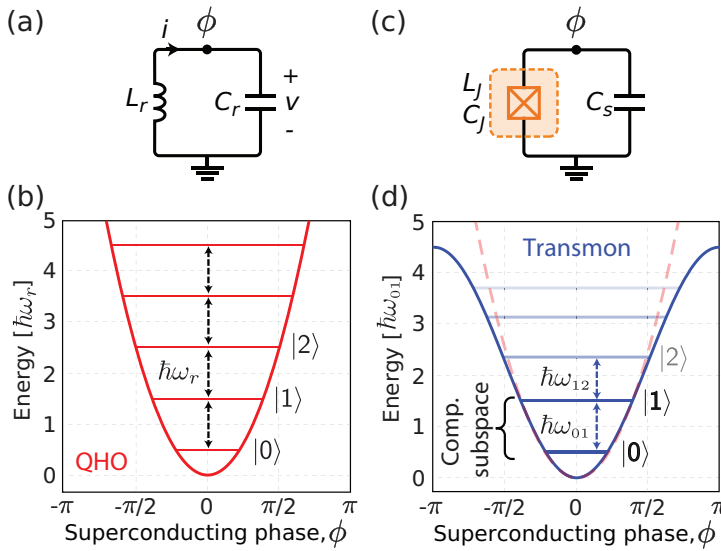
quantized form as

$$\hat{H} = 4E_C \hat{n}^2 + \frac{1}{2} E_L \hat{\phi}^2. \quad (3.49)$$

Here,  $\Phi_0 = h/(2e)$  is the superconducting magnetic flux quantum, and the number operator  $\hat{n}$  is the excess number of Cooper pairs. In this quantum scenario,  $[\hat{\phi}, \hat{n}] = i$ , and one can get Harmonic oscillator [Fig. 3.6(b)],

$$\hat{H} = \hbar \omega_r (\hat{a}^\dagger \hat{a} + \frac{1}{2}), \quad \omega_r = \sqrt{8E_L E_C} / \hbar, \quad (3.50)$$

where  $\hat{a}^\dagger(\hat{a})$  are the usual creation (annihilation) operator.



**Figure 3.6.** (a) Circuit for a parallel LC-oscillator. The superconducting phase on the island is denoted as  $\phi$ , referencing the ground as zero. (b) Energy potential for the quantum harmonic oscillator, where energy levels are equidistantly spaced  $\hbar\omega_r$  apart. (c) Josephson qubit circuit, where the nonlinear inductance  $L_J$  is shunted by a capacitance,  $C_s$ . (d) The Josephson inductance makes the potential anharmonic, forming a computational subspace of  $|0\rangle$  and  $|1\rangle$  (Taken from Ref. [274]).

However, in this case, the level spacing of all the energy levels is equal, and hence there could be transitions to other higher levels by using the same frequency of  $\omega_r$ . To mitigate this problem of unwanted dynamics involving non-computational states, we need to introduce non-linearity (anharmonicity) into the system, which can be achieved by using a Josephson junction that serves as a nonlinear inductor. The energy corresponding to it can be derived using two relations,

$$I = I_c \sin \phi, \quad V = \frac{\hbar}{2e} \frac{d\phi}{dt}, \quad (3.51)$$

resulting in a modified Hamiltonian, making the energy spectrum non-degenerate [Fig. 3.6(d)]–

$$\hat{H} = 4E_C \hat{n}^2 - E_J \cos \hat{\phi}. \quad (3.52)$$

Here,  $E_C = e^2/(2(C_s + C_J))$  is the total capacitance, including both shunt capacitance  $C_s$  and the self-capacitance of the junction  $C_J$ , and  $E_J = I_c \Phi_0 / 2\pi$  is the Josephson energy,  $I_c$  being the critical current of the junction. The system dynamics is governed by the  $E_J/E_C$  ratio. As the community has realized that charge noise is more challenging to mitigate than flux noise, it is often chosen as  $E_J \gg E_C$ . Additionally, current experimental technology enables greater flexibility in engineering the  $E_J$  component. To achieve this limit, one approach is to make  $E_C$  small by shunting the junction with

a large capacitor  $C_s \gg C_J$  [see Fig. 3.6(c)], which is known as the *transmon qubit*<sup>5</sup>. In this limit, one can express the state in terms of the superconducting phase  $\phi$ , and the  $\phi$  does not fluctuate much. In such scenario, we can expand the  $\cos \phi$  in power series,

$$E_J \cos \phi = \frac{1}{2} E_J \phi^2 - \frac{1}{24} E_J \phi^4 + \mathcal{O}(\phi^6). \quad (3.53)$$

Now, the qubit frequency becomes  $\omega_q = (\sqrt{8E_J E_C} - E_C)/\hbar$ . The excitation to higher states can be suppressed either by a large enough anharmonicity or by using robust control techniques, and thus, one can achieve a two-level system

$$\hat{H} = \frac{\omega_q}{2} \hat{Z}. \quad (3.54)$$

In practice, several architectures for superconducting quantum computers exist, such as frequency-tunable qubits [279], fixed-frequency qubits with static couplings [280], and fixed-frequency qubits with tunable coupling elements [281, 282].

## 3.6 Quantum stack for running on a quantum computer

Executing quantum optimization on noisy devices requires specialized tools and best practices to achieve good results from the hardware. We focus on gate-based cloud computing platforms, where quantum algorithms theorists can directly test their algorithms on hardware. In abstract design, Fig. 3.7 describes the common stages of the quantum stack, which also helps us to understand where scalability and performance bottlenecks arise.

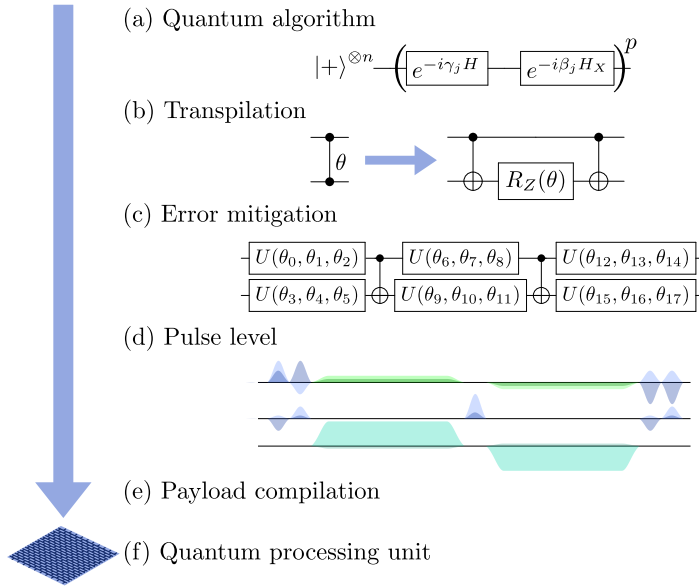
At the highest level of the quantum computing stack lies a conceptual representation of the algorithm, such as abstract, hardware-agnostic, and built from mathematical formulations and logical structures tailored to solving a specific problem. As this algorithm progresses through the stack, a transpilation process transforms it into a hardware-compatible circuit by translating abstract operations into executable instructions for the chosen quantum platform. During this phase, additional circuit elements may be added or adjusted to support error mitigation. Next, a pulse-level compiler further refines these instructions into precise electromagnetic pulses designed to manipulate quantum states and carry out the intended computation. This transformation from abstract logic to physical execution highlights how the algorithm evolves across layers of the stack, with each stage providing opportunities to optimize performance on noisy quantum hardware.

### 3.6.1 Adjusting quantum problems and algorithms

From a pen-and-paper abstract circuit to hardware-aware quantum problems or problem-dependent circuit structures—these are crucial in mitigating the effects of noise at the lower level of the quantum stack. For example, a problem with more qubits and low connectivity could have a lower noise effect compared to a model with densely

---

<sup>5</sup>The name transmon comes from ‘transmission-line shunted plasma oscillation qubit’ [278]. The suffix *-mon* refers to its similarity to the Cooper pair box (often called a quantronium) and its oscillatory behavior, reminiscent of other artificial atoms like the fluxmon or phasemon. The *trans-* prefix highlights the addition of a large shunting capacitance that suppresses charge noise, transforming the charge qubit into a more stable, weakly anharmonic oscillator.



**Figure 3.7.** Abstract levels of quantum stack from quantum algorithm to quantum processing unit (QPU). (a) Quantum algorithm in high-level quantum circuit, (b) Circuit transpiled to native-gate sets of the QPU, (c) Error mitigation methods, (d) QPU is instructed with physical pulses corresponding to the native-gate instructions, (e) finally, the circuit is compiled into QPU-executable waveforms, (f) which are run on the QPU (taken from Fig.4 of [248]).

connected variables. Alternatively, one can project a fully-connected QUBO problem onto the sparse connectivity of a quantum computer [283].

To efficiently utilize the quantum resources in the hardware, it is often helpful to select the algorithms wisely. For example,

- Warm-start can reduce the number of QAOA layers [268, 269].
- Including the constraints of the optimization problem in the mixer instead of the cost function [267].
- Counter-diabatic terms can improve the quality of the solution [284].
- Using circuit-cutting to reduce the depth and fight against coherence time [285].

Nevertheless, it is crucial from a hardware perspective to choose the methods that limit the final circuit depth and gate counts, without reducing the solution quality.

### 3.6.2 Transpilation

A transpiler plays a crucial role in quantum computing by converting a quantum circuit into another version that adheres to the constraints and capabilities of the target quantum hardware. This transformation typically results in a lower-level circuit expressed in domain-specific languages for continuous-variable platforms. Because every quantum device has unique physical and architectural characteristics, transpilers must be adapted to each hardware platform to ensure optimal performance.

To generate circuits that can be executed effectively on a given device, the transpilation process must consider several constraints, including the hardware's native gate set and the qubit connectivity. For the NISQ devices, additional factors, such as gate error rates, qubit decoherence times (e.g.,  $T_1$  and  $T_2$ )<sup>6</sup>, and cross-talk play a significant role. Thus, the effectiveness of the transpiler has a substantial impact

<sup>6</sup> $T_1$ , or relaxation time, is the average time it takes for a qubit to decay from the excited state  $|1\rangle$  to the ground state  $|0\rangle$ .  $T_2$ , or dephasing time, measures how long a qubit maintains phase coherence between the states  $|0\rangle$  and  $|1\rangle$ .  $T_2$  is always less than or equal to  $T_1$ , and both are critical for determining how long quantum information can be stored and manipulated reliably.



on both the total gate count and the overall execution fidelity of the algorithm on hardware.

```

1  # Tips: Check your gate counts and circuit depth after transpilation
2  print("Gate Counts:", qc.count_ops())
3  print("Circuit Depth:", qc.depth())

```

Generally, the transpilation process involves three main steps:

## 1. Qubit selection

This step is influenced by both the hardware's characteristics and the specific requirements of the algorithm. When not all qubits on the chip are needed, it is advantageous to select the most reliable ones based on a suitable performance metric. Metrics might include gate fidelity [286], cross-talk levels [287], or layer fidelity [288], depending on the circuit's structure. For circuits dominated by two-qubit operations, choosing qubits with high two-qubit gate fidelity is appropriate [286]. Conversely, if the algorithm requires many mid-circuit measurements, measurement fidelity should influence the selection [289]. In some quantum optimization problems, individual decision variables are mapped to specific qubits, and the mapping itself can affect the circuit depth [290].

```

1  ## Qubit selection
2  import networkx as nx
3  import numpy as np
4  from typing import Tuple, Dict, List, Any
5  from networkx.algorithms.isomorphism import GraphMatcher
6
7  def make_subgraph(num_qubits: int) -> nx.Graph:
8      """Example: Constructs linear structure."""
9      edges = [(i, i + 1) for i in range(2 * num_qubits - 1)]
10     return nx.from_edgelist(edges)
11
12  def evaluate_mapping(graph: nx.Graph, mapping: Dict[int, int], backend:
13     ↪ Any) -> Tuple[float, List[int]]:
14     """
15     Computes circuit fidelity based on gate errors and returns the list of
16     ↪ measured physical qubits.
17     """
18     reverse_map = {v: k for k, v in mapping.items()}
19     fidelity = 1.0
20     for edge in graph.edges():
21         physical_edge = (reverse_map[edge[0]], reverse_map[edge[1]])
22         try:
23             gate_error = backend.properties().gate_error("ecr",
24                 ↪ physical_edge)
25         except:
26             gate_error = backend.properties().gate_error("ecr",
27                 ↪ physical_edge[:-1])
28         fidelity *= (1 - gate_error)
29     # All logical qubits are measured now

```

```
26     measured_qubits = list(mapping.values())
27     return fidelity, measured_qubits
28
29 def find_best_qubits(
30     num_qubits: int,
31     backend: Any,
32     minimum_fidelity: float,
33     readout_threshold: float = 0.99
34 ) -> Tuple[int]:
35     """
36     Returns the best mapping tuple that satisfies fidelity and readout
37     ↪ threshold.
38     If no mapping meets criteria, returns an empty tuple and prints a
39     ↪ message.
40     """
41     logical_graph = make_subgraph_for_two_copies(num_qubits)
42     hardware_graph = nx.from_edgelist(backend.coupling_map.get_edges())
43     matcher = GraphMatcher(hardware_graph, logical_graph)
44
45     used_qubits = set()
46
47     for mapping in sorted(matcher.subgraph_isomorphisms_iter(), key=lambda
48     ↪ m: hash(frozenset(m.items()))):
49         ordered_mapping = tuple(mapping[i] for i in range(2 * num_qubits))
50         if set(ordered_mapping) & used_qubits:
51             continue
52
53         fidelity, measured_qubits = evaluate_mapping(logical_graph,
54         ↪ mapping, backend)
55
56         readout_fidelities = [
57             1 - backend.properties().readout_error(q) for q in
58             ↪ measured_qubits
59         ]
60         avg_readout = np.mean(readout_fidelities)
61
62         if fidelity >= minimum_fidelity and avg_readout >=
63         ↪ readout_threshold:
64             print("map_used:", ordered_mapping)
65             return ordered_mapping
66
67     used_qubits.update(ordered_mapping)
68
69     print("No valid mapping found above fidelity and readout thresholds.")
70     return ()
```

## 2. Gate decomposition

In this phase, high-level operations are broken down into sequences of native gates supported by the hardware [291–293]. Achieving this requires a universal gate set [294], and the choice of decomposition strategy depends on the desired trade-off between circuit accuracy and depth. Some techniques are exact, while others allow for approximations that reduce gate count and depth at the cost of some fidelity. For instance, exponentials of Pauli operators can be implemented using Pauli-Z rotations along with local two-qubit gates and single-qubit Clifford gates [295]. In noisy environments, approximate compilation methods may be preferred to keep the circuit shallower [296, 297].

```

1  # transpile from the qc to backend using basis_gates
2  transpile(qc,
3            basis_gates= backend.basis_gates,
4            initial_layout=[idx for idx in range(0,n_qc,1)],
5            [idx for idx in range(0,n_qc,1)],
6            inplace=True)

```

## 3. Qubit routing

Limited connectivity between qubits requires routing strategies to enable interactions between non-adjacent qubits. This typically involves inserting SWAP gates to bring qubits into proximity. General-purpose routing algorithms exist for arbitrary circuits [298–300]. However, some quantum algorithms, such as QAOA, produce structured circuits with blocks of commuting two-qubit gates. Leveraging this structure allows for more efficient routing and shallower circuits [290, 301–303]. Still, when the problem graph is denser than the hardware’s coupling map, a larger number of SWAP gates may be needed, increasing depth [301]. Consequently, quantum algorithms that map to sparse graphs tend to perform better on devices with limited connectivity [286, 304], highlighting the importance of identifying classically complex problems that remain sparse.

Some modern hardware platforms support dynamic circuits, which include mid-circuit measurements and conditional logic executed within the coherence time of the qubits [305]. These capabilities can be leveraged to address connectivity issues more efficiently [306]. For example, a QAOA circuit that would traditionally require SWAP operations to respect the layout of a grid-like coupling map can be replaced with a constant-depth implementation that uses auxiliary qubits at a quadratic overhead [307]. This area is still under active research, with the intention of reducing both circuit depth and width.

It is also important to note that hardware limitations vary across quantum technologies. For example, trapped-ion systems often feature all-to-all qubit connectivity, which allows them to execute dense circuits without increasing depth [308]. However, these systems currently scale to fewer qubits than superconducting platforms, limiting the size of problems they can handle.

### 3.6.3 Error suppression and mitigation

Once a quantum circuit has been transpiled, it may be further modified to incorporate instructions aimed at suppressing or mitigating errors.

**Error suppression** reduces the effects of noise in quantum circuits without requiring additional measurement shots. A well-known method in this category is Dynamical Decoupling (DD), which mitigates non-Markovian noise by inserting additional pulses during idle times in a circuit [309–311]. This can improve circuit performance with minimal resource overhead. However, when the transpiled circuit is dense, such as in QAOA circuits employing SWAP networks [286], the limited idle time may restrict the use of DD. In contrast, circuits with longer idle periods, such as those native to hardware with heavy-hex connectivity, can benefit substantially from DD [312].

Dynamical decoupling (DD) is a family of open-loop quantum control techniques used to suppress dephasing in quantum systems by averaging out environmental noise through time-dependent control pulses. It originates from refocusing techniques in NMR (like spin-echoes) and involves applying a carefully timed sequence of unitary control pulses to reverse or cancel out the effect of unwanted system-environment interactions.

```

1  # using dynamical decoupling
2  from qiskit.circuit.library import XGate
3  from qiskit.transpiler import PassManager, InstructionDurations
4  from qiskit_ibm_runtime.transpiler.passes.scheduling import
   ↪  ALAPScheduleAnalysis, PadDynamicalDecoupling
5
6  durations = InstructionDurations.from_backend(backend)
7  dd_sequence = [XGate(), XGate()]
8  pm = PassManager([ALAPScheduleAnalysis(durations),
9                    PadDynamicalDecoupling(durations, dd_sequence)])
10 circ_dd = pm.run(tqc)

```

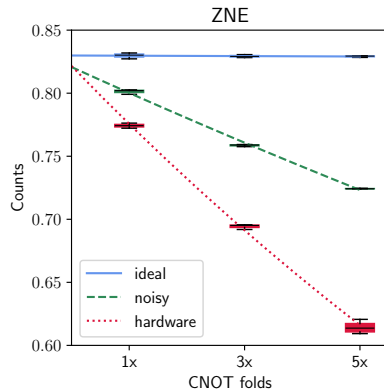
**Error mitigation** methods aim to reduce the impact of noise by applying classical pre- and post-processing around the quantum circuit execution. These approaches do not require full-scale quantum error correction, which involves encoding logical qubits into large arrays of physical ones [313]. While error mitigation does not eliminate errors, it can enhance computational results at a lower classical cost [314–316]. Common techniques in this category include Probabilistic Error Cancellation (PEC) [317] and Zero-Noise Extrapolation (ZNE) [44, 315, 318, 319], both of which are designed to yield more accurate expectation values.

- PEC involves characterizing the noise channel  $\Lambda$  associated with layers of Pauli-twirled gates [317]. Although the inverse channel  $\Lambda^{-1}$  is non-physical, it can be approximated using quasi-probability decompositions to yield an unbiased estimate of a noiseless observable  $O$ . The trade-off is a variance increase by a factor of  $\gamma^2$ , where  $\gamma \geq 1$  reflects the strength of the noise in the learned model and serves as a proxy for the circuit’s execution fidelity. The value of  $\gamma$  is closely

related to metrics like Layer Fidelity (LF) and Error Per Layered Gate (EPLG), which characterize hardware noise in large-scale systems [288]. If the average noise factor per qubit is  $\bar{\gamma}$ , then a circuit with  $n$  qubits and  $d$  layers will incur a mitigation cost scaling as  $\bar{\gamma}^{2 \times n \times d}$ . PEC becomes impractical for large-scale circuits due to its exponential variance overhead; for instance, applying PEC to 60 CNOT layers on 127 qubits yielded a  $\gamma^2$  value of  $10^{128}$  in Kim *et al.* [44]. To address this, Probabilistic Error Amplification (PEA) was introduced [44], which instead amplifies the learned noise to enable extrapolation, potentially reducing bias in the final estimate. PEA and PEC are particularly efficient when circuits consist of a small number of repeating gate layers, as is the case with SWAP-network-based designs [301].

- ZNE, on the other hand, estimates the zero-noise expectation value by executing equivalent circuits at different noise levels and extrapolating the results. Early implementations involved pulse stretching [315], which introduced significant calibration overhead, or gate folding, which increased circuit depth. Newer strategies, such as partial gate folding [320] and pulse-level techniques like cross-resonance stretching [321], help reduce these drawbacks. Unlike PEC, ZNE's resource requirements do not scale with the problem size, although deep circuits limit the stretch factors that can be used effectively (see sec. 6.3.1 for an example).

**Zero Noise Extrapolation (ZNE)** works by deliberately increasing the noise in a quantum circuit, typically through methods like gate folding, then executing the circuit at multiple noise levels and mathematically extrapolating the results back to the zero-noise limit using polynomial or exponential fits. This allows researchers to approximate the expected value of observables as if they were measured on a perfect quantum computer. ZNE was first formally introduced by Temme *et al.* [314], and has since become a widely adopted approach in the NISQ era. To show an example, we show Fig.



**Figure 3.8.** Example of ZNE. 1x CNOT folds represent the actual data, and then 3x and 5x imply that the number of CNOTs is increased without changing the unitary to incorporate the effect of noise coming from the CNOT gates. Then by extrapolation, we can look at 0x noise, or the *noiseless* case, where it approaches close to the ideal (noiseless) data.

- Conditional Value at Risk (CVaR). Due to noisy hardware, the estimation of energy can be biased by the bad samples, and then minimizing it would not provide a good solution. Instead, if one cares about sampling good solutions, then one can use alternative cost functions to increase the robustness against noise or relax the requirements on an ansatz. In CVaR, good samples are sorted by taking the best  $\alpha$ -fraction of samples, where  $\alpha = 1$  corresponds to the full expectation value and  $\alpha = 0$  corresponds to the single best observed sample. By

choosing  $\alpha$  based on the noise of the device, CVaR can provably bound noise-free expectation values [322].

Expectation-value-based mitigation methods like PEC and ZNE are especially valuable during variational parameter optimization [286, 315]. However, for quantum optimization tasks that require generating high-quality samples, e.g., finding a bit-string  $x$  that minimizes a cost function  $f(x)$ . It is essential to mitigate noise not just in expectation values but in actual circuit outputs. Therefore, advancing techniques that can produce noise-resilient samples will be critical to the success of QAOA.

## 3.7 Hardware benchmarks

Quantum hardware performance is commonly characterized by three primary aspects: scale, quality, and speed [323].

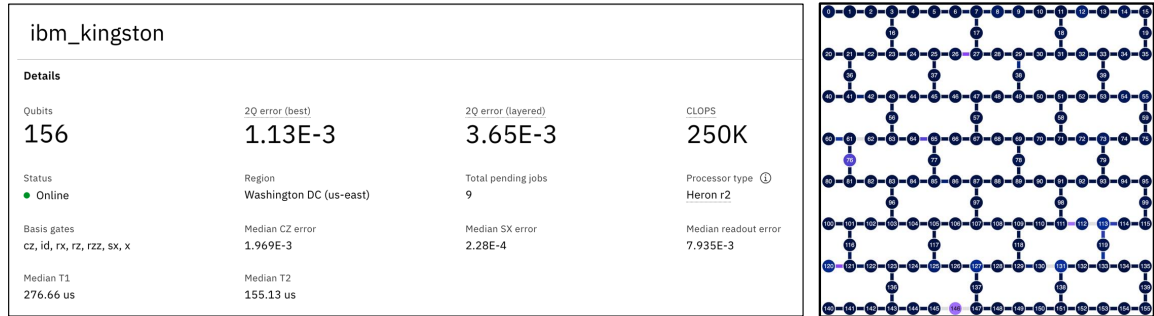
*Scale* refers to the total number of available qubits.

*Quality* encompasses both low-level device metrics, such as qubit coherence times and gate fidelities, and more comprehensive indicators like Layer Fidelity (LF), Error per Layered Gate (EPLG) [288], Quantum Volume (QV) [324], and the  $\gamma$  and average  $\bar{\gamma}$  parameters from Probabilistic Error Cancellation (PEC) [317]. For instance, a QPU with a QV of  $2^n$  can reliably execute an  $n$ -qubit circuit with  $n$  layers of random  $SU(4)$  gates [324] [see Fig. 4.6], effectively capturing performance contributions from gate fidelities, qubit connectivity, and the effectiveness of noise suppression and mitigation techniques.

However, these quality-focused metrics do not reflect the *speed* at which quantum computations are performed. Speed is a critical factor for optimization tasks, especially when a quantum device is expected to outperform classical systems by delivering high-quality solutions more quickly. Moreover, faster execution allows for more accurate observable estimates within time constraints and helps pay off the cost of error mitigation techniques, which typically require additional sampling. To address this, the Circuit Layer Operations Per Second (CLOPS) metric was introduced as a quantum analog to classical FLOPS. CLOPS quantifies a device's throughput by measuring how many layers of gates can be executed per unit time, including the classical overhead involved in updating circuit parameters. This makes CLOPS especially relevant for variational workloads where runtime is dominated by repeated circuit executions with parameter updates.

Metrics like QV, EPLG, and CLOPS are designed to provide general-purpose benchmarks that allow comparisons across different quantum hardware platforms and to monitor progress over time. For example, on *ibm\_kingston*, EPLG and CLOPS are reported as 0.113% and 250000, respectively (see Fig. 3.9). While these figures offer useful insight, they are ultimately only proxies and may fall short in predicting real-world performance on specific tasks.

The most informative benchmark for a quantum processor is its performance on application-relevant tasks, evaluated using the appropriate performance criterion, be it solution quality, execution time, or sample efficiency. In this context, our research in Chapter 4 shows that the ability to generate spin-squeezed states is directly related to solving fully connected MaxCut instances with QAOA [325]. As many quantum algorithms are heuristic in nature, we anticipate the growing importance of problem-specific benchmarking frameworks that more accurately reflect their practical utility.



**Figure 3.9.** Benchmarking properties of *ibm\_kingston*: 2Q error (EPLG), CLOPS, and readout error are among the most important metrics. Right: One can also decide which qubits to choose by looking at the quality of the connectivity (CZ error) and readout error.

## 3.8 Summary

This chapter provides an overview of quantum optimization, a major focus of this thesis. To identify where quantum advantage may emerge, we begin by reviewing quantum computational complexity classes. We then introduce key quantum optimization algorithms, including quantum annealing, relevant to Chapter 5, and the quantum approximate optimization algorithm (QAOA), which plays a central role in Chapters 4, 6, and 7. In the final section, we discuss tools and techniques for implementing quantum algorithms on noisy quantum devices, which are employed in Chapter 6.





# **Part II**

## **Resources in Quantum Optimization**



## Chapter 4

# Squeezing and quantum approximate optimization

*where quantum metrology meets quantum optimization*

---

## 4.1 Introduction

The Quantum Approximate Optimization Algorithm (QAOA) [326] is a promising approach for solving combinatorial optimization problems using digital quantum computers [327, 328] as discussed in Chapter 3. It uses constructive interference to find solution states [329], and it has better performance than finite-time adiabatic evolution [330]. However, such a hybrid algorithm heavily uses classical optimization techniques to obtain suitable parameters, and one may wonder if quantum effects, such as entanglement, play a role in QAOA.

Here, we show how concepts from quantum metrology shed light on this question through squeezing and how it connects to the performance of QAOA. Illustratively, the connection is established through the insight that (a) the aim of QAOA is to obtain the ground state as precisely as possible, while (b) quantum metrology leverages entanglement between particles to generate states that permit precision beyond the capacities of any comparable classical sensor [331–333]. For example, squeezed states can increase sensitivity for detecting phases [334], magnetic fields [335], and gravitational waves [336]. The most sensitive states for phase estimation are Dicke states [332, 337], where all qubits are equally entangled. We substantiate this connection through numerically exact calculations and data gathered on IBM Quantum hardware with up to eight qubits. Our analysis shows how the search for an optimal solution to the MaxCut problem on a complete graph through QAOA generates Dicke states, with squeezing [338, 339] and multipartite entanglement [139, 140]. Based on this, we propose the amount of squeezing generated as an application-tailored performance benchmark of QAOA, which is able to capture the depth dependency of QAOA better than quantum volume [324, 340, 341], and can be used to detect errors. Our work thus further strengthens the intimate links between quantum metrology and quantum information processing [132, 342–344]. Moreover, we show that warm-starting the optimization of a weighted MaxCut problem with a squeezed state can provide enhanced performance. We also discuss the parameter regimes, compared for different platforms, where a metrological task using a multi-party-entangled squeezed state generated through QAOA could become advantageous over fast repetitions with simple unsqueezed states.

In the rest of this chapter, we first provide some background on the MaxCut problem and spin-squeezing. Then, QAOA is formally connected to the generation of en-

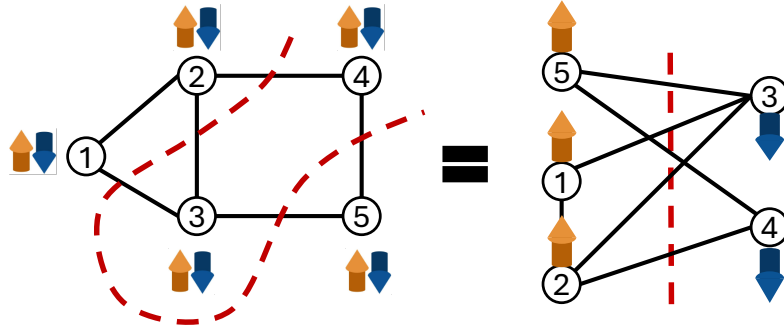
tangled squeezed states (Sec. 4.4), which was subsequently demonstrated numerically. Based on squeezing, then, we develop a benchmark tailored for QAOA (Sec. 4.5), and assess the ability of superconducting qubits to run QAOA on fully connected problems while simultaneously creating Dicke states and estimating the number of entangled particles (Sec. 4.6). Finally, we extend this connection for application in metrology and quantum optimization (Sec. 4.7).

## 4.2 MaxCut problem in QAOA

Recall from sec. 3.3.1 that universal quantum computers can address hard classical problems such as quadratic unconstrained binary optimization (QUBO), which is defined through

$$\min_{x \in \{0,1\}^n} x^T \Sigma x \quad \text{with } \Sigma \in \mathbb{R}^{n \times n}. \quad (4.1)$$

Then, the binary variables  $x_i$  are mapped to qubits through  $x_i = (1 - z_i)/2 \rightarrow (1 - \hat{Z}_i)/2$ , where  $\hat{Z}_i$  is a Pauli spin operator with eigenstates  $|0\rangle$  and  $|1\rangle$ . The result is an Ising Hamiltonian  $\hat{H}_C$  whose ground state is the solution to Eq. (4.1) [345]. The standard QAOA then applies  $p$  layers of the unitaries  $\exp(-i\beta_k \hat{H}_M) \exp(-i\gamma_k \hat{H}_C)$ , with  $k = 1, \dots, p$ , to the ground state of a mixer Hamiltonian  $\hat{H}_M$ , such as  $-\sum_i \hat{X}_i$  where  $\hat{X}_i$  is a Pauli operator, to create the trial state  $|\psi(\beta, \gamma)\rangle$ . A classical optimizer seeks the  $\beta = (\beta_1, \dots, \beta_p)$  and  $\gamma = (\gamma_1, \dots, \gamma_p)$  that minimize the energy  $\langle \psi(\beta, \gamma) | \hat{H}_C | \psi(\beta, \gamma) \rangle$ , which is measured in the quantum processor (see Fig. 3.4).



**Figure 4.1.** Example of a MaxCut Problem. Left: A five-node graph with the maximum cut indicated by the red dashed line, which partitions the graph into two disjoint subsets. Right: The corresponding bipartition, where nodes in each subset represent opposite spin orientations, up or down, in the qubit encoding. In the QUBO formulation of MaxCut [Eq. (4.2)], this corresponds to finding an antiferromagnetic ground state of the associated Ising Hamiltonian, where graph edges represent spin-spin interactions.

MaxCut aims at bipartitioning the set of nodes  $V$  in a graph  $G(V, E)$  such that the sum of the weights  $\omega_{i,j}$  of the edges  $(i, j) \in E$  traversed by the cut is maximum, i.e.

$$\max_{z \in \{-1,1\}^n} \frac{1}{2} \sum_{(i,j) \in E} \omega_{i,j} (1 - z_i z_j). \quad (4.2)$$

Note that  $z_i = \pm 1$  and that means the above equation is maximum only when  $z_i z_j = -1$ , which occurs when edges become anti-ferromagnetic, see Fig. 4.1 for an example.

Another extreme problem instance is when  $\omega_{i,j} = 1, \forall(i,j)$ , i.e., an unweighted fully connected graph labelled  $\mathcal{G}_n$ , see Fig. 4.4(a). Dividing  $V$  into two sets of size as equal as possible creates a maximum cut.

**Optimization method.** In this chapter, the Qiskit QAOA Runtime program [301] is used to simulate QAOA. To optimize the  $\{\gamma, \beta\}$ , we use the simultaneous perturbation stochastic approximation (SPSA) algorithm [346], which simultaneously optimizes multiple parameters and can handle noisy environments. We do not initialize the optimizer with values for the learning rate and a perturbation. Instead, we let SPSA calibrate itself in the first 25 iterations. To obtain good solutions, we allow SPSA a maximum of 500 iterations and gather a total of  $2^{15}$  shots per iteration.

## 4.3 Squeezing

Squeezed states are entangled states with a reduced variance of one observable at the cost of an increased variance in non-commuting observables. A large body of experimental work exists addressing the generation of squeezing in various platforms [343, 347–351]. Squeezing can also detect multipartite entanglement among qubits [352–356].

### 4.3.1 Squeezing in qubit ensemble

In our setting, we are interested in squeezing within an ensemble of  $n$  qubits (whose symmetric subspace can be seen as a qudit with length  $\ell = \frac{n}{2}$ ). Consider a coherent state, such as the collective superposition  $|+\rangle^{\otimes n}$ , where  $|+\rangle = (|0\rangle + |1\rangle)/\sqrt{2}$ . This state has a variance of  $\sigma_{\text{css}}^2 = \frac{n}{4}$ , commonly called the shot-noise, in the collective observable  $\hat{L}_z = \frac{1}{2} \sum_{i=1}^n \hat{Z}_i$ . By evolving  $|+\rangle^{\otimes n}$ , e.g., under the non-linear one-axis-twisting (OAT) operator [338]

$$\hat{L}_z^2 = \frac{1}{4} \left( n + \sum_{i \neq j} \hat{Z}_i \hat{Z}_j \right), \quad (4.3)$$

the state is stretched over the collective Bloch sphere<sup>1</sup>. The direction with reduced variance can be transferred to the  $z$  coordinate by rotating the state around the  $x$ -axis with [338, 349, 357] (see Fig. 4.2)

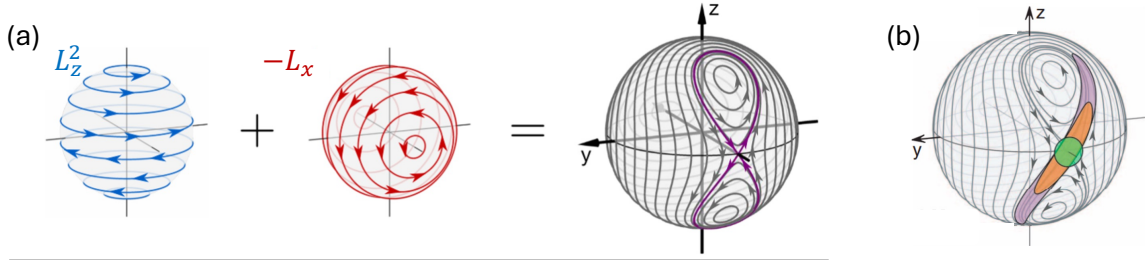
$$\hat{L}_x = \frac{1}{2} \sum_{i=1}^n \hat{X}_i. \quad (4.4)$$

<sup>1</sup>The collective Bloch sphere is a generalization of the standard Bloch sphere used to represent the quantum states of multiple qubits, typically in a system with  $n$  two-level systems. It provides a way to visualize the evolution and properties of these multi-qubit states. One can also think of it as a qudit Bloch sphere, where instead of 2 poles, one would also have multiple rings on the sphere to denote the  $d$  levels. Given the collective spin observable,  $L_z = \frac{1}{2} \sum_{i=1}^n Z_i$ , its eigenvalues  $\{-\frac{n}{2}, \dots, 0, \dots, \frac{n}{2}\}$  would define  $n+1$  level qudits. For example, the north (south) pole would be  $|0\rangle^{\otimes n}$  ( $|1\rangle^{\otimes n}$ ), and the equator would be superposition of equal number of 0 and 1:  $D_{n/2}^n$  for even  $n$  (see Eq. (4.9)).

The resulting  $n$  particle state is called number squeezed along  $z$  when the observed variance is below  $\sigma_{\text{css}}^2$ , i.e., if the squeezing parameter

$$\mathcal{S} [\text{dB}] = 10 \log_{10} \left( \frac{\text{Var}(\hat{L}_z)}{\sigma_{\text{css}}^2} \right) \quad (4.5)$$

is negative [338, 339].



**Figure 4.2.** Spin-squeezing via one-axis twisting Hamiltonian. (a) Pictorial mean-field representation of the two contributions on the collective Bloch sphere, coming from the nonlinear term  $\hat{L}_z^2$  and rotation term  $\hat{L}_x$ , which leads in combination to the bifurcated phase space, where the arrow indicates the existence of an unstable point. (b) If a state is initiated in that unstable fixed point, under the dynamics (will follow the arrow), its quasi-probability distribution will stretch along the lines, and in the orthogonal direction, it will be squeezed. Adapted from Ref. [358].

### 4.3.2 Entanglement from squeezing

Measurements of collective spin observables can reveal entanglement. In particular, separable states satisfy [356]

$$(E1) : \quad \langle \hat{L}_x^2 \rangle + \langle \hat{L}_y^2 \rangle \leq \frac{n}{2} \left( \frac{n}{2} + \frac{1}{2} \right) \iff \frac{n}{4} \leq \langle \hat{L}_z^2 \rangle = \text{Var}(\hat{L}_z). \quad (4.6)$$

The second implication is reached using the identity  $\langle \hat{L}^2 \rangle = \langle \hat{L}_x^2 \rangle + \langle \hat{L}_y^2 \rangle + \langle \hat{L}_z^2 \rangle = \frac{n}{2}(\frac{n}{2} + 1)$ , and  $\langle \hat{L}_z \rangle = 0$  for our target states. Any squeezed state defined through Eq. (4.5) violates the relation above and is thus entangled. Moreover,  $\langle \hat{L}_x^2 \rangle + \langle \hat{L}_y^2 \rangle$  reaches the maximum  $\frac{n}{2}(\frac{n}{2} + 1)$  in the Dicke state [356], which is the same as having  $\text{Var}(\hat{L}_z) = 0$ .

### 4.3.3 Multipartite entanglement: quantum Fisher information and squeezing

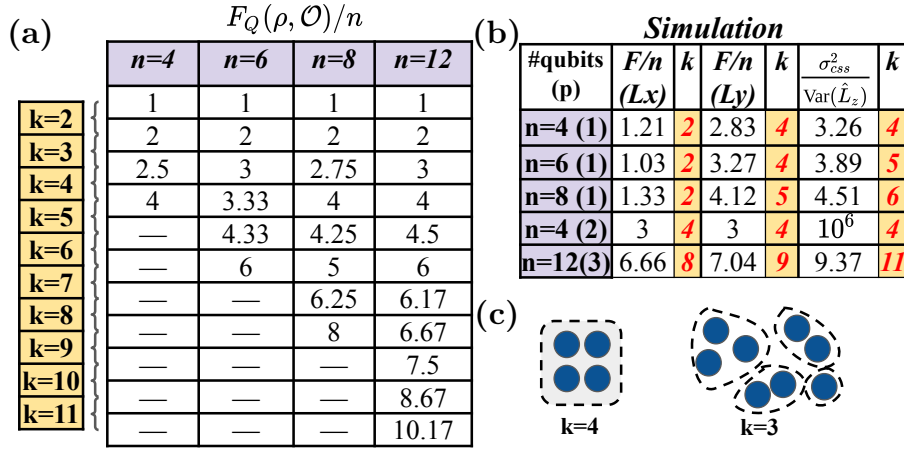
Here, we recall some concepts for multipartite entanglement from Sec. 2.6. A pure state of  $n$ -qubits, written as a product  $|\psi\rangle = \otimes_{j=1}^M |\psi_j\rangle$ , is  $k$ -partite entangled when at least one state  $|\psi_j\rangle$  contains non-factorizable  $k$ -qubits ( $k$ -producible) [139, 359]. This definition is the same as the entanglement depth [360], see Fig. 4.3(c). A sufficient condition for  $(k+1)$ -partite entanglement stems from the quantum Fisher information  $F_Q$  (see Sec. 2.6.3): a state reaching

$$(E2) : \quad F_Q[\rho_n, \mathcal{O}] > (sk^2 + r^2), \quad (4.7)$$

where  $s = \lfloor n/k \rfloor$  denotes the integer division of  $n$  by  $k$ , and  $r$  is the remainder—is at least  $(k+1)$ -partite entangled [139, 140] (see Fig. 4.3(a) for a concrete example).

While  $F_Q$  is becoming a useful witness for entanglement in quantum many-body systems [361–366], its origin is as a key figure of merit in quantum metrology, where  $F_Q$  quantifies the distinguishability of a state  $\rho$  from  $\rho' = e^{-i\theta\mathcal{O}}\rho e^{i\theta\mathcal{O}}$ , generated by the Hermitian operator  $\mathcal{O}$  with infinitesimal  $\theta$ . Thus, a large  $F_Q$  implies a high measurement precision for estimating the value of  $\theta$  [137, 367]. For pure states  $\psi$ , the quantum Fisher information becomes simply  $F_Q[\psi, \mathcal{O}] = 4\text{Var}(\mathcal{O})_\psi$  [368], whereas for mixed states it provides a lower bound on the variance.

The target state of the QAOA for MaxCut on  $\mathcal{G}_n$ , the Dicke state, is invariant under a unitary evolution generated by  $\hat{L}_z$  but is highly sensitive to rotations around the  $x$ - or  $y$ -axes of the collective Bloch sphere [349]. Thus, to obtain a large  $F_Q$ , it is advantageous to choose  $\mathcal{O} = \hat{L}_x$  or—even more so— $\mathcal{O} = \hat{L}_y$ , as can be observed in Fig. 4.3(b).



**Figure 4.3.** Multipartite entanglement from quantum Fisher information and number squeezing. (a)  $F_Q$  witnessing  $k$ -partite entanglement for different  $n$ , (b) In the simulations,  $F_Q$  obtained with  $\hat{L}_y$  is larger than  $\hat{L}_x$ . The numbers of entangled particles ( $k$ ) estimated from squeezing ( $\sigma_{\text{css}}^2/\text{Var}(\hat{L}_z) = 10^{-S/10}$ ) are close to the numbers obtained from  $F_Q[\hat{L}_y]$  for most cases. In a proper Dicke state as obtained with  $n = 4(p = 2)$ , the  $\text{Var}(\hat{L}_z)$  becomes extremely small, leading to the very large value of  $\sigma_{\text{css}}^2/\text{Var}(\hat{L}_z)$  seen in the fourth row. (c) Illustrative examples of  $k$ -partite entanglement as entanglement depth.

In the hardware, where the system is no longer in a pure state, it is considerably more challenging to directly access  $F_Q$  [361, 369]. However, for Gaussian states, one can nevertheless use the empirical relation [349]

$$(E3) : \quad F_Q/n[\hat{L}_y] \simeq \sigma_{\text{css}}^2/\text{Var}(\hat{L}_z) = 10^{-S/10} \quad (4.8)$$

between  $F_Q$  and squeezing. For the simulation, the estimated  $k$  using this relation is close to the exact estimation from  $F_Q$  in most of cases, see Fig. 4.3(b), except for the depth-three QAOA, where the states are no longer Gaussian [370]. Assuming that the above relation holds for depth-one QAOA, where the states are expected to be Gaussian, we obtain the estimates for  $k$ -partite entanglement in the hardware implementation reported in Fig. 4.12(b).

## 4.4 Connecting QAOA to Squeezing

In a quantum circuit representation, the steps described above for the generation of squeezed states coincide with a single-layer QAOA sequence, see Fig. 4.4(c): (i) The application of Hadamard gates to  $|0\rangle^{\otimes n}$  initialize the system in  $|+\rangle^{\otimes n}$ , the ground state of the mixer Hamiltonian  $\hat{H}_M$ . (ii) The evolution under the OAT operator corresponds to applying the unitary  $\exp(-i\gamma_1 \hat{H}_C)$  with the cost function  $\hat{H}_C \propto \hat{L}_z^2$ . On the qubit level, this corresponds to controlled-Z gates generated by  $\hat{Z}_i \hat{Z}_j$  between all qubits  $i$  and  $j$ . (iii) The rotation around the  $x$ -axis to reveal the squeezing corresponds to the unitary  $\exp(-i\beta_1 \hat{H}_M)$ , i.e., an application of the mixer.

Interestingly, the cost function  $\hat{H}_C \propto \hat{L}_z^2$  is a special instance of the MaxCut problem (Eq. 4.2). For even  $n$ , the set of all maximum cuts corresponds to the symmetric Dicke state [337] (see Section. 2.6.1 for more details about Dicke state):

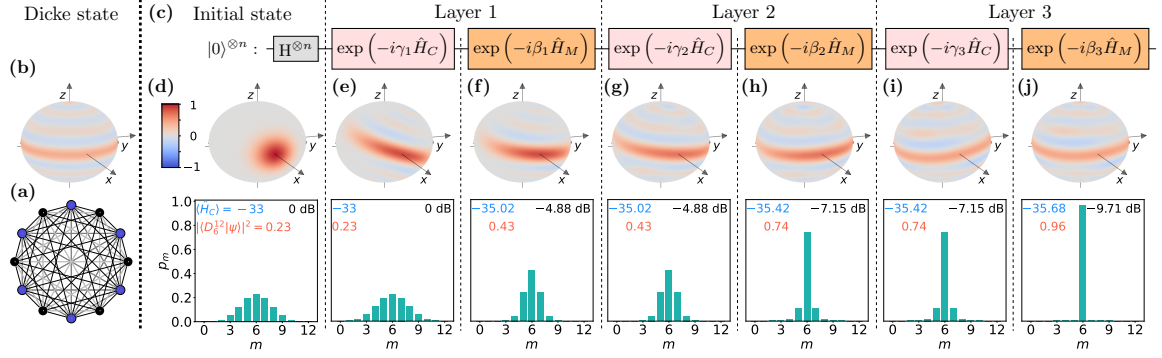
$$D_k^n = \binom{n}{k}^{-1/2} \sum_i P_i \left( |1\rangle^{\otimes k} \otimes |0\rangle^{\otimes (n-k)} \right), \quad (4.9)$$

with  $k = \frac{n}{2}$ . Here,  $P_i(\cdot)$  denotes a permutation of all states with  $k$  particles in  $|1\rangle$  and  $n - k$  particles in  $|0\rangle$ . For odd  $n$ , the set of all maximum cuts corresponds to  $(D_{\lfloor n/2 \rfloor}^n + D_{\lfloor n/2 \rfloor + 1}^n)/\sqrt{2}$ . These states are maximally squeezed along  $z$  and are useful for metrological applications [332]. The QAOA cost function Hamiltonian to minimize in this problem is  $\hat{H}_C = \frac{1}{2} \sum_{i < j} (\hat{Z}_i \hat{Z}_j - 1) = \hat{L}_z^2 - \frac{n^2}{4} \mathbb{I}$ . Therefore, QAOA is tasked to find the maximum cut of a fully connected unweighted graph that maximizes the squeezing. This relation thus translates analog metrological protocols [349] to a digital quantum processor. In addition, by formulating the constraints that an arbitrary Dicke state  $D_k^n$  imposes on the spins as a QUBO, we can generate  $D_k^n$  for arbitrary  $k$  (see Sec. 4.7.1).

### 4.4.1 QAOA as generator of squeezing

To illustrate the connection between QAOA and squeezing, we numerically simulate a system with  $n = 12$  qubits and follow the usual QAOA protocol, using  $\hat{H}_C = \hat{L}_z^2 - \frac{n^2}{4} \mathbb{I}$ ,  $\hat{H}_M = -2\hat{L}_x$ , and  $p = 3$ . We depict the generated collective-spin/ qudit state using the Wigner quasi-probability distribution as well as the probability distribution over the spin eigenvalues  $\{m = \langle \hat{L}_z \rangle + \frac{n}{2}\}$  at each step, see Fig. 4.4(d)-(j). Each application of  $\hat{H}_C$  stretches the Wigner distribution, making it resemble an ellipse with the major axis tilted with respect to the equatorial plane of the qudit Bloch sphere. As  $[\hat{H}_C, \hat{L}_z] = 0$ , this operation does not alter the distribution of  $\langle \hat{L}_z \rangle$ . Next, the mixer operator rotates the Wigner distribution back towards the equator, thereby transferring the squeezing to the operator  $\hat{L}_z$ . After three layers, the final state has an overlap with the symmetric Dicke state of 96% and the squeezing number reaches  $\mathcal{S} = -9.71$  dB. Crucially, noiseless QAOA with fewer layers produces less squeezing, e.g., see depth-one QAOA in Sec. 4.4.2.





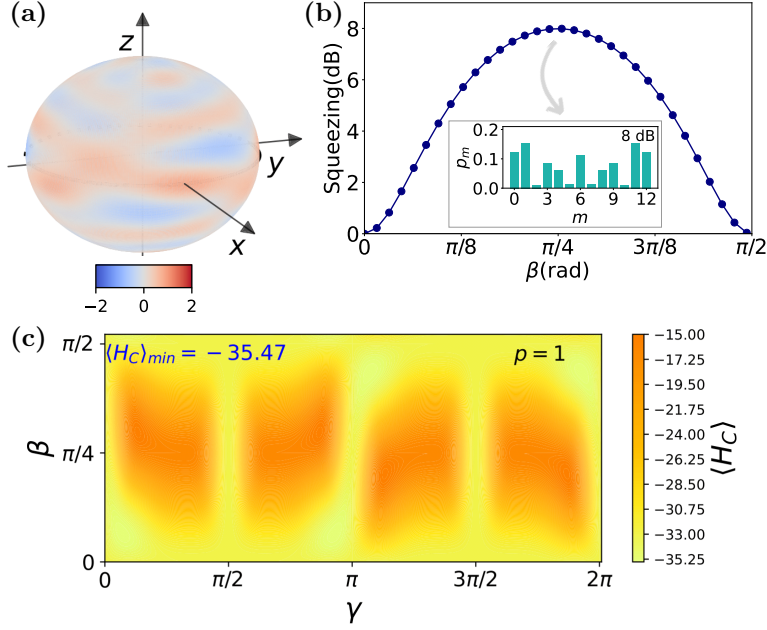
**Figure 4.4.** Metrologically useful squeezing generated by a depth-three QAOA for the MaxCut problem in a complete graph of 12 nodes. (a) Fully connected unweighted graph with the nodes and edges colored according to one of the 924 possible maximum cut configurations. (b) Wigner quasi-probability distribution of the symmetric Dicke state  $D_6^{12}$ , an idealized example of a squeezed state, and the target for our QAOA. (c) Circuit representation of QAOA with alternating application of cost-function and mixer Hamiltonian. The Bloch spheres and histograms from (d) to (j) show the state after the corresponding gate in the optimized QAOA circuit with  $(\gamma_i, \beta_i)$  given by  $(0.199, 0.127)$ ,  $(0.306, 0.087)$ , and  $(4.592, 1.518)$  for  $i = 1, 2$ , and  $3$ , respectively. Negativity in the Wigner distribution indicates that the states are non-Gaussian [370]. The squeezing (in black), energy expectation  $\langle \hat{H}_C \rangle$  (in blue), and overlap probability density  $|\langle D_6^{12} | \psi \rangle|^2$  with the target Dicke state (in orange) are shown inside each histogram.

#### 4.4.2 Advantage of using QAOA to generate squeezing

**Merely increasing the duration of  $\hat{H}_C$  does not increase squeezing.** Squeezing is generated by  $\hat{L}_z^2$  [349], which suggests that simply applying  $\hat{H}_C \propto \hat{L}_z^2$  for a longer duration, corresponding to a larger coefficient  $\gamma$  in the QAOA, may transform the coherent state to a squeezed state, after which we can use the mixer  $\hat{H}_M$  to reveal the squeezing along  $\hat{L}_z$ . In this way, one layer of QAOA would suffice to create any squeezing, which would also require fewer CNOT gates than when  $p > 1$ . To test this hypothesis, we run depth-one QAOA using  $\gamma = \gamma_1 + \gamma_2 + \gamma_3$  where the  $\gamma_i$  are taken from Fig. 4.4, as they contain the source of “total” squeezing. The result is a fragmented Wigner distribution on the Bloch sphere without observable squeezing in any direction, see Fig. 4.5(a). Furthermore, no squeezing is detected along  $z$  for any value of the tomography angle  $\beta$ , see Fig. 4.5(b). This finding is in agreement with the known observation that over-squeezing can be detrimental for precision [358], however, the states here do not wrap around points near poles because  $\hat{L}_z^2$  and  $\hat{L}_x$  are not applied simultaneously as in Ref. [358].

**Advantages of multi-layer QAOA.** One may object to the arguments in the preceding paragraph that the  $\gamma$  we chose is sub-optimal. To address that, in Fig. 4.5(c), we numerically map the energy landscape of depth-one QAOA in the  $\{\gamma, \beta\}$  plane. The results reveal a minimum energy of  $\langle \hat{H}_C \rangle_{\min} = -35.47$  which corresponds to  $|\langle D_6^{12} | \psi \rangle|^2 = 98.53\%$ . These results are inferior to those we obtain from the depth-three QAOA, i.e.,  $\langle \hat{H}_C \rangle_{\min} = -35.68$  and  $|\langle D_6^{12} | \psi \rangle|^2 = 99.08\%$ . Alternating multiple layers of  $\hat{H}_C$  and  $\hat{H}_M$  is therefore advantageous over a single application of the one-axis-twisting operator.

To quantify the obtainable improvement as a function of the number of layers used, we can define a new performance metric  $\Delta_p^n(\%)$ , which compares the energy reduction over the initial ansatz obtained by  $p$ -layers of QAOA with the one achieved



**Figure 4.5.** Illustration of the importance of alternating the cost-function and the mixer operator. (a) A fragmented Wigner distribution on the Bloch sphere is obtained when  $\hat{H}_C$  is applied with  $\gamma = \gamma_1 + \gamma_2 + \gamma_3$  from Fig. 4.4. (b) For the state in (a) no squeezing is observed at any  $\beta$ . The inset shows the probability distribution at  $\beta = \pi/4$ , corresponding to  $\mathcal{S} = 8$  dB, i.e. over-squeezing. (c) The energy landscape of the depth-one QAOA reveals that the lowest energy it can reach is  $-35.47$ , which is inferior to  $-35.68$  obtained in depth-three QAOA.

by the ideal target state. In the  $n = 12$  case, the initial coherent state and the target Dicke state have  $\langle \hat{H}_C \rangle = -33$ , and  $-36$ , respectively. Thus, a depth-one QAOA (corresponding to the usual squeezing protocol) can reach a maximum  $\Delta_1^{12} = 2.47/3$ . In contrast, the depth-three QAOA can reach  $\Delta_3^{12} = 2.68/3$ , as shown in Fig. 4.4. Thus, according to this metric, a depth-three QAOA is  $0.21/3 = 7\%$  better than a depth-one QAOA.

### 4.4.3 Multipartite entanglement in QAOA

The squeezing in collective spin observables can further be related to entanglement, using the notions introduced in Sec. 4.3.3. We employ three different criteria:

- (E1) If  $\frac{n}{4} > \langle \hat{L}_z^2 \rangle$ , the state violates a bound on separable states [356]. Any squeezed state ( $\text{Var}(\hat{L}_z) < \frac{n}{4}$ ) at the equator ( $\langle \hat{L}_z \rangle = 0$ ) is witnessed as entangled by this criterion (Sec. 4.3.2). Here,  $\langle \hat{L}_z^2 \rangle = 0.32 < \frac{n}{4} = 3$ , which is close to the minimal value of 0 achieved by the Dicke state.
- (E2) If the quantum Fisher information for a pure state  $\psi$ ,  $F_Q[\psi, \mathcal{O}] = 4\text{Var}(\mathcal{O})$ , is larger than  $(sk^2 + r^2)$ , where  $s = \lfloor n/k \rfloor$  denotes the integer division of  $n$  by  $k$ , and  $r$  is the remainder, at least  $(k+1)$  particles are entangled [139, 140]. Here,  $F_Q[\psi, \hat{L}_y] = 84.48$  and the final state has multipartite entanglement between at least 9 out of 12 particles (see Fig. 4.12(b) to compare with Fig. 4.3(b)). In Fig. 4.3(b), we also report  $F_Q$  for both  $\mathcal{O} = \hat{L}_x, \hat{L}_y$  and the resulting  $k$ -partite entanglement witnessed by it for the ideal simulations. In this ideal scenario of noiseless numerical simulations, the large values of  $F_Q[\psi, \hat{L}_{x,y}]$  are directly

related to the anti-squeezing of the final state along the equator of the Bloch sphere.

- (E3) Specifically for Gaussian states, one can approximately estimate the number of entangled particles  $k$  assuming the identity  $F_Q/n \simeq \sigma_{\text{css}}^2/\text{Var}(\hat{L}_z)$  [349] (see Sec. 4.3.3 for more discussion), which here yields  $k = 11$ . We have to use this estimate for the hardware results where direct access to  $F_Q$  is not possible (Sec. 4.6).

#### 4.4.4 Comparative metrological gain via QAOA in quantum platforms

The law of large numbers suggests that increasing the number of measurements leads to more accurate phase estimation by reducing statistical variance. It is thus important to analyze when QAOA-prepared squeezed states can be convenient for a metrological task, as compared to simply repeating various shots with an unsqueezed coherent time. With that motivation, we study the time taken to measure a phase  $\theta$  with the coherent spin state (CSS) versus QAOA-generated Dicke states within different hardware architectures (but without an attempt at comparing the different architectures to each other, which often have different aims and boundary conditions that are difficult to compare on an even footing). The coherent state is easily prepared by a single rotation around the  $y$ -axis. It has no entanglement, and  $m$  measurements of the phase  $\theta$  have a variance bound by  $\Delta^2\theta \geq 1/(mN)$ . By contrast, a QAOA-prepared state with  $F_Q$  above the  $k + 1$ -partite entangled limit takes more time to prepare than the CSS, but it requires a smaller number of measurements to reach the same variance since  $\Delta^2\theta \geq 1/(mkN)$  is lowered by a factor of  $1/k$  [140].

One may then wonder whether, in a practical application, the improved precision can offset the larger preparation time. Crucially, the optimization cost of QAOA can be ignored in these considerations since the optimal  $\gamma$  and  $\beta$  parameters are reusable across different measurements and experiments. We therefore compute the duration of a single measurement repetition  $t_{\text{repet.}}$ , which is the sum of the duration of the gates in the circuit to prepare the state  $t_{\text{gates}}$  and the readout time including the reset of the measurement apparatus  $t_{\text{rr}}$ , i.e.,  $t_{\text{repet.}} = t_{\text{gates}} + t_{\text{rr}}$ . The gate duration for the coherent spin state  $t_{\text{gate}}^{\text{CSS}}$  is the duration of a single-qubit gate, while the QAOA protocol requires two-qubit gates, whose number can depend on the available universal gate set and the hardware connectivity.

The QAOA-generated states are advantageous when the time  $t^{\text{QAOA}} = t_{\text{repet.}}^{\text{QAOA}} \times m_{\text{QAOA}}$  to achieve a certain precision is smaller than the time  $t^{\text{CSS}} = t_{\text{repet.}}^{\text{CSS}} \times m_{\text{CSS}}$  to achieve the same precision with coherent states. If the QAOA-prepared state achieves  $F_Q = kN$ , we have for equal precision  $m_{\text{QAOA}} \times k = m_{\text{CSS}}$ , i.e., the QAOA-prepared states are advantageous if

$$t_{\text{gates}}^{\text{QAOA}} + t_{\text{rr}} < k (t_{\text{gates}}^{\text{CSS}} + t_{\text{rr}}). \quad (4.10)$$

**Superconducting qubits:** The duration of a QAOA layer is impacted by the qubit connectivity. Each QAOA layer on  $N$  linearly connected superconducting qubits requires  $3(N - 2)$  layers of simultaneously executable CNOT gates, which include SWAP gates [301]. Under the assumption that QAOA can create  $k + 1$ -partite entanglement in  $p = \log_2(k)$  layers [371], the duration  $t_{\text{gates}}^{\text{QAOA}} = 3(N - 2) \log_2(k) t_{\text{cx}}$  with  $t_{\text{cx}}$  the duration of a CNOT gate. Here, we neglected the duration of single-qubit gates. With

$k = N$ , Eq. (4.10) yields

$$3(N - 2) \log_2(N) t_{\text{cx}} + t_{\text{rr}} \lesssim N t_{\text{rr}}, \quad (4.11)$$

which, for large  $N$  and the durations in Tab. 4.1, amounts to

$$N \lesssim 2^{t_{\text{rr}}/(3t_{\text{cx}})} = 2^{1000/3}. \quad (4.12)$$

Although the linear layout of the hardware poses a limit on when QAOA-generated states remain useful, this limit is extremely high. Assuming noisy hardware achieves only a finite  $k$ , one has

$$(N - 2) \log_2 N < (k - 1) t_{\text{rr}} / (3 t_{\text{cx}}). \quad (4.13)$$

For  $k = 2$ , e.g., the QAOA-generated states would remain advantageous up to about 60 qubits arranged in a linear chain, which lies at the size limit of the current hardware. These numbers are conservative estimates that can be significantly increased by improved noise resilience and higher hardware connectivity.

**Trapped-ion qubits:** Large multipartite entangled states of trapped-ions can be generated by a single application of the Mølmer–Sørensen gate (MS) [372, 373] where interaction strength among all qubit pairs is equal [374]. Therefore, if we neglect the duration of single-qubit gates,  $t_{\text{gates}}^{\text{QAOA}}$  only depends on the number of QAOA layers  $p = \log_2(k)$ , and  $t_{\text{gates}}^{\text{QAOA}} = \log_2(k) t_{\text{ms}}$  with  $t_{\text{ms}}$  being the duration of a MS gate. Following Tab. 4.1, QAOA-generated  $k + 1$ -partite entangled states are therefore advantageous when

$$\log_2(k) t_{\text{ms}} \lesssim (k - 1) t_{\text{rr}} \Rightarrow k^{\frac{1}{k-1}} \lesssim 2^{t_{\text{rr}}/t_{\text{ms}}} = 2^{25}. \quad (4.14)$$

Since  $k^{1/(k-1)}$  is a decreasing function, QAOA-generated states are always advantageous in trapped-ion setups.

**Cold-atoms:** We now consider cold-atoms in Bose-Einstein condensates, which can, e.g., manipulate states with a number of the order of 400 atoms [349]. Following QAOA, we assume that  $\log_2(k)$  layers of the one-axis-twisting Hamiltonian interleaved with  $x$ -rotations can generate  $k + 1$ -partite entanglement. The squeezed state is thus created in a time  $t_{\text{gates}}^{\text{QAOA}} = \log_2(k) t_{\text{OAT}}$ . We neglect the duration of  $x$  and  $y$ -rotations. Equation (4.10) implies  $\log_2(k) t_{\text{OAT}} < (k - 1) t_{\text{rr}}$ , showing that given a Bose-condensed atom cloud of fixed size it is always favorable to create spin squeezed states for metrology since  $t_{\text{OAT}} \ll t_{\text{rr}}$ , see Tab. 4.1.

Platform	Single-qubit	Entanglement	Readout & Reset $t_{\text{rr}}$
Transmons	10 ns [375]	100 ns [340]	100 $\mu\text{s}$ [376]
Trapped ions [377]	15 $\mu\text{s}$	200 $\mu\text{s}$	300 $\mu\text{s}$ + 5 ms [378]
Cold atoms (BEC)		10 ms [349]	1 s

**Table 4.1.** Duration of key operations presented as orders of magnitude only. The entangling operation for the transmons, trapped-ions, and cold atoms is the two-qubit CNOT gate, the Mølmer–Sørensen gate, and the one-axis twisting Hamiltonian, respectively.

## 4.5 QAOA-tailored hardware benchmarks

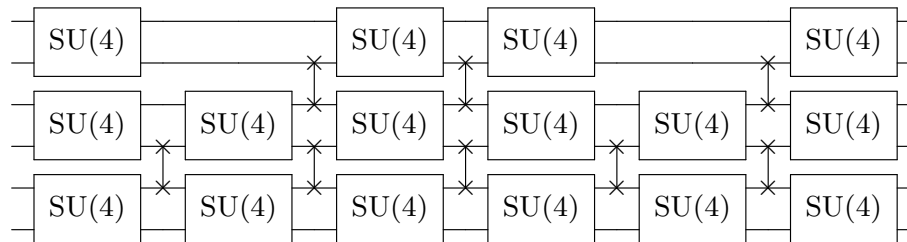
The performance of quantum computing hardware is often measured by metrics such as randomized benchmarking [379–381] and quantum process tomograph [382, 383], which focus on gates acting on typically one to two qubits, while Quantum Volume (QV) (Sec. 4.5.1) is designed to measure the performance of a quantum computer as a whole [324, 340, 341]. For certain applications, these are complemented by specifically designed benchmarks, e.g., for quantum chemistry [384], generative modelling [385, 386], variational quantum factoring [387], Fermi-Hubbard models [388], and spin Hamiltonians [389]. It is of particular interest to identify such application-tailored benchmarks also for variational algorithms, as these employ highly structured circuits. This necessity is well illustrated by considering the QV: the circuit complexity of a  $2^n$  QV system is equivalent to a  $p = 2$  QAOA running on  $n$  linearly connected qubits.

### 4.5.1 Quantum Volume

A processor with a Quantum Volume of  $QV = 2^n$  can reliably, as defined by the generation of heavy output bit-strings, execute circuits that apply  $n$  layers of  $SU(4)$  gates on random permutations of  $n$  qubits [324]. When transpiled to a line of  $n$  qubits, QV circuits result in  $n$  layers of  $SU(4)$  gates that have at most  $\lfloor \frac{n}{2} \rfloor$  individual  $SU(4)$  gates simultaneously executed on the qubits [340]. In between these  $SU(4)$  layers, there are at most  $\lfloor \frac{n}{2} \rfloor$  SWAP gates, see Fig. 4.6. Furthermore, each  $SU(4)$  and SWAP gate require at most and exactly three CNOT gates, respectively [390]. Under these conditions, the total number of CNOT gates is at most

$$3n \left\lfloor \frac{n}{2} \right\rfloor + 3(n-1) \left\lfloor \frac{n}{2} \right\rfloor, \quad (4.15)$$

which approaches  $3n^2$  as  $n$  becomes large. By comparison, the cost operator of QAOA circuits of complete graphs transpiled to a line requires exactly  $\frac{3}{2}n(n-1) - n + 1$  CNOT gates, approaching  $3n^2/2$  for large  $n$ . This suggests that a  $2^n$  Quantum Volume is a good performance indicator for a depth  $p = 2$  QAOA on  $n$  qubits. Importantly, this comparison is only possible as long as the QAOA circuit is executed using the same error mitigation and transpilation methods as those employed to measure QV [341]. However, QV fails to capture the depth dependency  $p$  of QAOA. The benchmark that we develop overcomes this limitation as the QAOA depth should be chosen such that the measured squeezing is maximum. This also provides the maximum  $p$  for which it makes sense to run QAOA on the benchmarked noisy hardware.



**Figure 4.6.** Example of a six-qubit quantum volume circuit as presented in Ref. [340], which shows the layers of  $SU(4)$  and SWAP gates.



### 4.5.2 Proposed benchmark

As the above section shows, QV fails to properly capture the dependency on  $p$  as QAOA circuits on complete graphs are deeper than their width. Ref. [391] shows using entropic inequalities, if the circuit is too deep, a classical computer can sample in polynomial time from a Gibbs state while achieving the same energy as the noisy quantum computer. That bound is based on the fidelity of layers of gates, which is, however, often overestimated when built from fidelities of gates benchmarked in isolation, e.g., due to cross-talk [301].

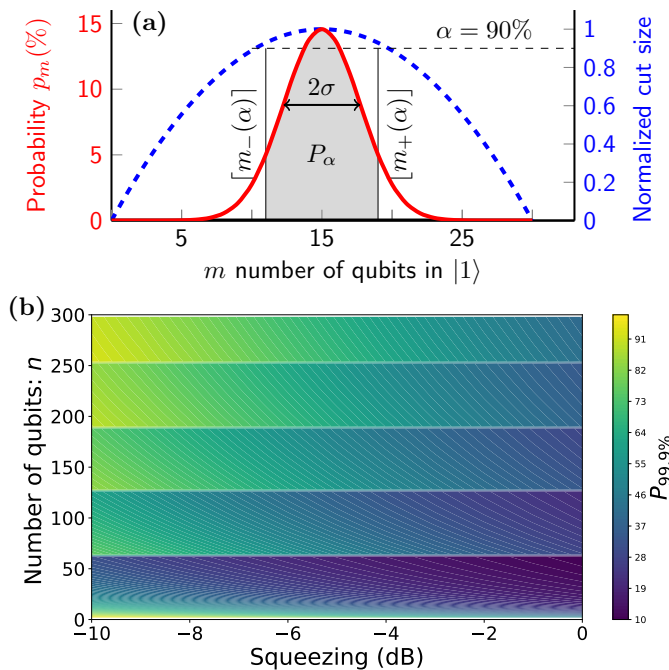
Since the solution to the MaxCut problem on the fully connected unweighted graph  $\mathcal{G}_n$  is known, we propose squeezing as a good hardware benchmark for QAOA to complement other performance metrics. For our proposed benchmark, we first label the quantum numbers of  $\hat{L}_z + \frac{n}{2}$  by  $m \in \{0, 1, \dots, 2\ell\}$ , which correspond to cuts of size  $c(m) = m(n-m)$  on  $\mathcal{G}_n$ . We relate squeezing to a QAOA performance metric through the question:

*Given the squeezing  $\mathcal{S}$  in the trial state, what is the probability  $P_\alpha(n, \mathcal{S})$  of sampling a cut with size  $c(m)$  greater than a given  $\alpha$ -fraction of the maximum cut size  $c_{\max} = n^2/4$ ?*

Here,  $\alpha$  can be seen as an approximation ratio. By definition, cuts with  $c(m) > \alpha c_{\max}$  must satisfy  $m_-(\alpha) < m < m_+(\alpha)$  for even  $n$ , where  $m_\pm(\alpha) = \frac{n}{2}(1 \pm \sqrt{1 - \alpha})$ . Under a QAOA trial state  $|\psi(\beta, \gamma)\rangle$  with a distribution  $p_m$  over  $m$ , see Fig. 4.7(a), the probability to sample cuts larger than  $\alpha c_{\max}$  is thus

$$P_\alpha(n) = \sum_{m=\lceil m_-(\alpha) \rceil}^{\lfloor m_+(\alpha) \rfloor} p_m. \quad (4.16)$$

We now make the simplifying assumption that the distribution  $p_m$  is a Gaussian



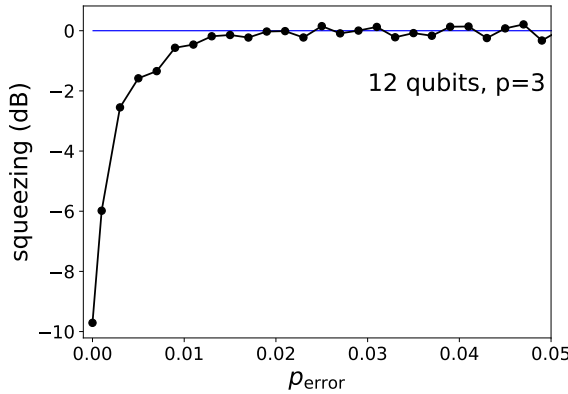
**Figure 4.7.** Benchmarking QAOA with squeezing. (a) Probability distribution ( $p_m$ , red solid line) and normalized cut size ( $c_m/c_{\max}$ , blue dashed line) simultaneously plotted against  $m = \langle \hat{L}_z \rangle + \frac{n}{2}$  for  $n = 30$ . States with normalized cut-size more than  $\alpha$  lie in  $m \in [\lceil m_-(\alpha) \rceil, \lfloor m_+(\alpha) \rfloor]$ . These yield the shaded area under the probability  $p_m$ , which is the figure of merit  $P_\alpha$  defined in Eq. (4.16). (b)  $P_\alpha$  shows how the probability of sampling high-value cuts changes with squeezing  $\mathcal{S}$  and qubit count  $n$ , based on trial Gaussian distributions.

$\mathcal{N}(\frac{n}{2}, \sigma)$ , where the standard deviation  $\sigma$ —the only free variable for fixed  $n$ —is, by definition, in one-to-one correspondence to the squeezing  $\mathcal{S} = 10 \log_{10}(4\sigma^2/n)$ . In summary, the benchmark (i) relates squeezing to the probability of sampling good solutions, a QAOA performance metric, (ii) captures the ability of QAOA to create entangled states (Sec. 4.4.3), and (iii) is as susceptible to hardware noise as other fully connected QAOA circuits (Sec. 4.5.3).

We illustrate the benchmark by numerically computing  $P_\alpha(n, \mathcal{S})$  as a function of  $n$  and the squeezing  $\mathcal{S}$  in the Gaussian distribution  $p_m(\mathcal{S})$ . Since the ground state of  $\mathcal{G}_n$  is highly degenerate, we select a high value of  $\alpha$ , e.g., 99.9%. At fixed  $n$ , an increased squeezing (more negative) increases  $P_\alpha$ , see Fig. 4.7(b), as cuts with a larger size receive more weight. In addition,  $P_\alpha$  has discontinuous jumps at  $n_{\text{dis.}}$ , where  $z = \lfloor \frac{n_{\text{dis.}}}{2} \sqrt{1 - \alpha} \rfloor \in \mathbb{Z}^+$ . In between discontinuities,  $P_\alpha$  diminishes with increasing  $n$  because  $\sigma$  increases  $\propto \sqrt{n}$  for fixed  $\mathcal{S}$ , which reduces the weight attributed to high value cuts.

**Explaining the discontinuity.** According to Eq. (4.16), the states in the domain  $(\lceil m_- \rceil, \lfloor m_+ \rfloor)$  are included in  $P_\alpha$ , where  $m_\pm(n, \alpha) = \frac{n}{2} \pm \frac{n}{2} \sqrt{1 - \alpha}$ . Since  $\lfloor m_+ \rfloor$  and  $\lceil m_- \rceil$  must both be integers, the span of the domain  $\lfloor m_+ \rfloor - \lceil m_- \rceil$  remains constant over a large  $n$  range and changes abruptly when  $\lfloor \frac{n}{2} \sqrt{1 - \alpha} \rfloor \in \mathbb{Z}$  changes value. We denote the values of  $n$  at which such changes occur as  $n_{\text{dis.}}$ , which correspond to the discrete jumps along the  $n$ -axis in Fig. 4.7(b). For  $\alpha = 99.9\%$  and  $n$  even, we obtain discontinuities in  $P_{99.9\%}$  at  $n_{\text{dis.}} = 64, 128, 190, 254$ .

### 4.5.3 Error detection by spin-squeezing

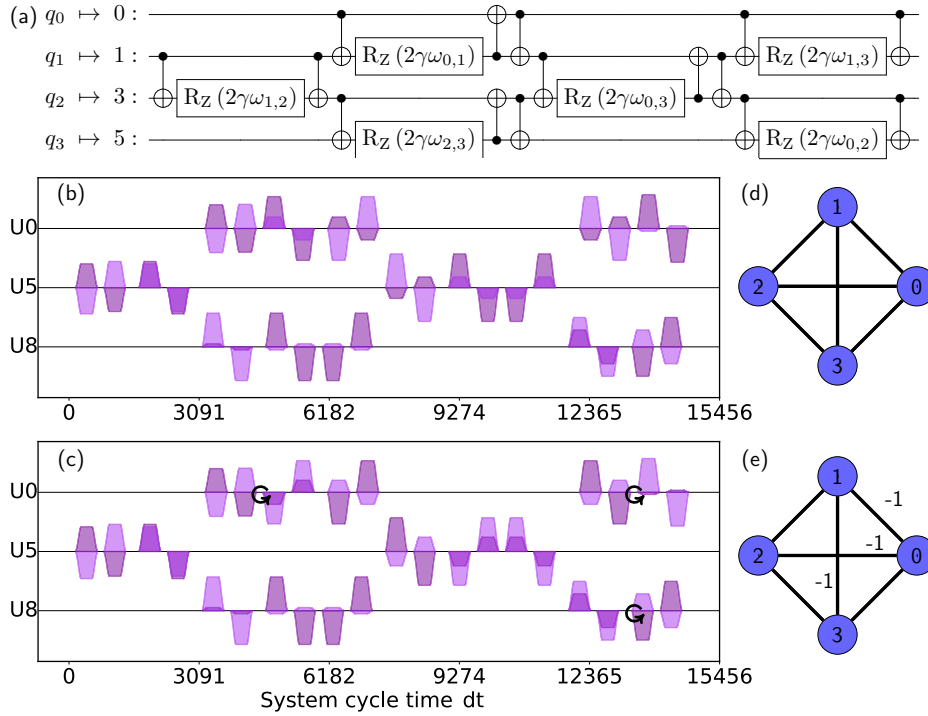


**Figure 4.8.** Squeezing as a function of the strength of gate noise modeled as bit-flip errors. A bit-flip error is added to every CNOT gate by 2-qubit Pauli channels,  $\mathcal{P}_2 = \mathcal{P}_1 \otimes \mathcal{P}_1$ , where  $\mathcal{P}_1 = \sqrt{p_{\text{error}}}X + \sqrt{1 - p_{\text{error}}}I$ . The squeezing approaches zero as gate errors increase, showing the validity of squeezing as an application-tailored hardware benchmark.

As undesired processes destroy fragile quantum superpositions, the degree of squeezing is sensitive to the noise levels of the quantum device. To illustrate this, we run our depth-three QAOA example of Fig. 4.4 on a noisy simulator. After each CNOT gate, we include a Pauli bit-flip error with strength  $p_{\text{error}}$ . We observe that the squeezing rapidly decays towards zero as  $p_{\text{error}}$  increases, resulting in a non-squeezed state, see Fig. 4.8. This simple simulation demonstrates that gate errors destroy the fragile correlations needed to create a squeezed state. Conversely, the ability (or inability) to realize squeezing through QAOA can be used as a tool to estimate the errors in a circuit experiment.

#### 4.5.4 Squeezing as a good benchmark for arbitrary QUBO problem

From a hardware perspective, although  $\mathcal{G}_n$  is a specific problem, its QAOA circuit is representative of the noise of an arbitrary fully-connected QUBO problem since the gates constituting a generic cost function  $\exp(-i\gamma_k \hat{H}_C)$  can be implemented with virtual Z-rotations and CNOT gates [392]. Indeed, the difference between the pulse schedules only amounts to phase changes, indicated by circular arrows in Fig. 4.9(c). The duration of the QAOA pulse schedule and the absolute amplitude of the pulses are thus independent of the variables  $\Sigma$  in the QUBO, see Eq. (4.1), and the variational parameters  $\gamma$  and  $\beta$  (compare Fig. 4.9(b) and (c)). Therefore, much like Quantum Volume, the hardware benchmark based on squeezing captures effects such as limited qubit connectivity, unitary gate errors, decoherence, and cross-talk. Furthermore, from a hardware perspective, the squeezing circuit is also the hardest to implement since QUBOs that are not fully connected, i.e.,  $\exists (i, j) \mid \Sigma_{i,j} = 0$ , require fewer pulses.



**Figure 4.9.** Cross-resonance pulse schedules of the squeezing circuit and an arbitrary QUBO. (a) Quantum circuit of a general four-qubit fully connected cost operator  $e^{-i\gamma \hat{H}_C}$  transpiled to qubits 0, 1, 3, and 5 of *ibm\_lagos*. (b) Pulse schedule of the cost operator used to generate the symmetric Dicke state, i.e.,  $\omega_{i,j} = 1 \forall i, j$ . (c) Pulse schedule of a MaxCut instance with edge weights  $\omega_{0,1} = \omega_{0,2} = \omega_{1,3} = -1$  and  $\omega_{0,3} = \omega_{1,2} = \omega_{2,3} = 1$ . The circular arrows show where the phase shifts differ from the pulse schedule in (b). (d) and (e) MaxCut graph corresponding to the pulse schedule in (b) and (c), respectively. The duration of a single sample of the arbitrary waveform generators is  $dt = 0.222$  ns. The light and dark pulses show the in-phase and quadrature of each complex amplitude pulse applied to control channels U0, U5, and U8 of *ibm\_lagos*.

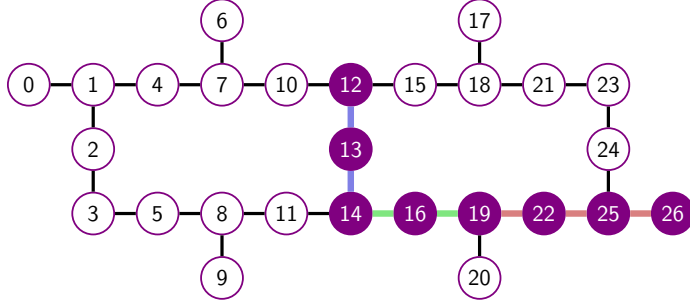
## 4.6 Squeezing in superconducting qubits

We now evaluate the benchmark on gate-based superconducting transmon qubits [393] with the following characteristics.



### 4.6.1 Hardware details

The superconducting qubit data is gathered on the *ibmq\_mumbai* system, which has 27 fixed-frequency qubits connected through resonators; its coupling map is shown in Fig. 4.10. We chose a set of qubits that form a line with the smallest possible CNOT gate error. Each circuit is measured with 4000 shots. The properties of the device, such as  $T_1$  times and CNOT gate error, are shown in Tab. 4.2.



**Figure 4.10.** Coupling map of *ibmq\_mumbai* with the qubits used shown in violet. The four, six, and eight-qubit data were measured on the linearly connected qubits  $\{19, 22, 25, 26\}$ ,  $\{14, 16, 19, 22, 25, 26\}$ , and  $\{12, 13, 14, 16, 19, 22, 25, 26\}$ , respectively, chosen based on the CNOT gate fidelity.

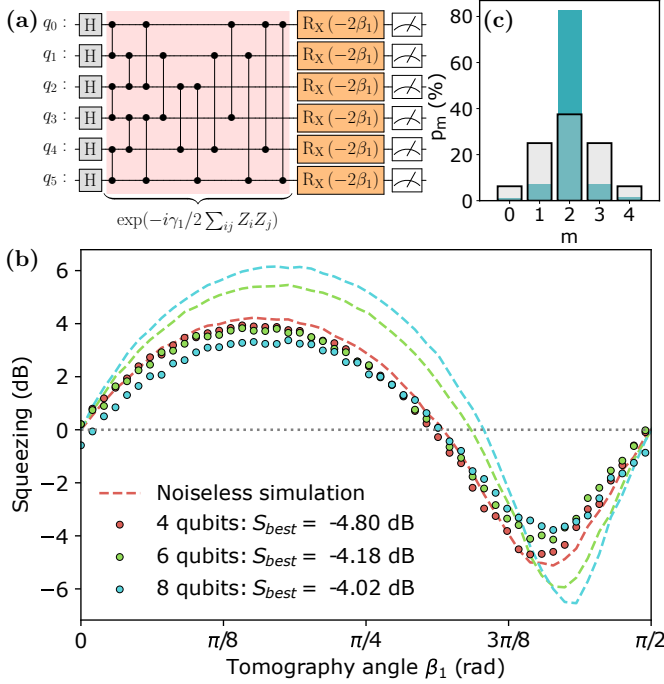
Qubit pair	CNOT gate		Qubit	$T_1$ ( $\mu$ s)
	error (%)	duration (ns)		
(12, 13)	0.77	548	12	166
(13, 14)	1.26	320	13	137
(14, 16)	1.04	348	14	174
(16, 19)	0.77	747	16	118
(19, 22)	0.66	363	19	227
(22, 25)	0.58	484	22	122
(25, 26)	0.50	348	25	194
	$0.80 \pm 0.27$	$451 \pm 155$	26	103

**Table 4.2.** Properties of the relevant CNOT gates as reported by *ibmq\_mumbai* on the date of the circuit execution. The average  $T_1$  of the selected qubits is  $155 \pm 43$   $\mu$ s.

### 4.6.2 Measurement of squeezing on IBMq hardware

We measure the squeezing on the IBM Quantum system *ibmq\_mumbai* using Qiskit [394] for four, six, and eight qubits. Since the chosen qubits have a linear connectivity, we use a line swap strategy [301, 395] to create the all-to-all qubit connectivity required by the squeezing circuit, shown in Fig. 4.11(a) for  $p = 1$ . This circuit is then transpiled to the cross-resonance-based hardware [396, 397] employing a pulse-efficient strategy instead of a CNOT decomposition [398] using Qiskit Pulse [399]. The optimal value of the variational parameter  $\gamma$  is found with a noiseless simulation for each  $n$ . We use readout error mitigation [400, 401], which on average improves the best measured squeezing by  $-0.7 \pm 0.1$  dB averaged over all three  $n \in \{4, 6, 8\}$ . At depth one, a sweep of the tomography angle  $\beta_1$  reveals a squeezing of  $-4.80$ ,  $-4.18$ , and  $-4.02$  dB whereas noiseless simulations reach  $-5.14$ ,  $-5.90$ , and  $-6.56$  dB for  $n = 4, 6$ , and  $8$ , respectively. These metrological gains are comparable to prior works in trapped ions [373, 402–406] (see at the end of this section). Given the measured squeezing, we compute a  $P_{99.9\%}(n, \mathcal{S})$  of 61.5%, 49.1%, and 42.6%, respectively. Furthermore, we

run a depth-two QAOA on the fully connected four-qubit graph to create a state with a  $-5.96$  dB squeezing, see Fig. 4.11(c), which results in  $P_{99.9\%}(4, -5.96) = 68.2\%$ . These results indicate that the potential to generate squeezing in a four-qubit system is limited by the variational form at depth one. By contrast, in systems with six and eight qubits, the squeezing generated in practice is limited by the large number of CNOT gates at depth one (40 and 77, respectively).



**Figure 4.11. Squeezed states generated on superconducting qubits.** (a) A quantum circuit implementing a single QAOA-layer for the MaxCut problem on a six-qubit system, producing a state with a reduced variance along the  $z$ -axis. (b) Squeezing measured with the quantum circuit in (a). The dashed lines show a noiseless QASM simulation. (c) The  $p_m$  distribution of the four-qubit state with  $-5.96$  dB squeezing generated by a depth-two QAOA with optimal parameters  $\gamma_1 = 0.918$ ,  $\gamma_2 = -0.257$ ,  $\beta_1 = -0.711$ , and  $\beta_2 = -2.175$ . The gray histogram shows the state  $(H|0\rangle)^{\otimes n}$ .

### 4.6.3 Entanglement in hardware

Here we use the entanglement criteria from Sec. 4.3.3. The criterion (E1) witnesses the generated states in both simulation and hardware as entangled, see Fig. 4.12(a). In a noiseless simulation of a depth-one QAOA of system sizes  $n = 4, 6, 8$ , criterion (E2) witnesses at least 4, 4, 5 qubit entanglement, respectively. In the noisy hardware implementation, estimate (E3) suggests these numbers to still reach 4, 3, 3, respectively, see Fig. 4.12(b).

(a)		$\langle \hat{L}_z^2 \rangle$		$n/4$	(b)		Simulation (E2)		Hardware (E3)	
#qubits (p)	Simulation	Hardware	Upper bound		qubits (p)	$F/n(L_y)$	$k$	$S(\text{dB})$	$k$	
n=4 (1)	0.31	0.33	1		n=4 (1)	2.83	4	-4.80	4	
n=6 (1)	0.38	0.57	1.5		n=6 (1)	3.27	4	-4.18	3	
n=8 (1)	0.44	0.79	2		n=8 (1)	4.12	5	-4.02	3	
n=4 (2)	0	0.25	1		n=4 (2)	3	4	-5.96	4	
n=12(3)	0.32	—	3		n=12 (3)	7.04	9	—	—	

**Figure 4.12. Entanglement from squeezing and quantum Fisher information.** (a) The values of  $\langle \hat{L}_z^2 \rangle$  (E1) obtained in simulation and hardware are close to 0, indicating that the states are in the vicinity of entangled Dicke states. (b) Number of entangled particles  $k$  calculated from  $F_Q[\hat{L}_y]$  (E2) in simulation, and estimated for hardware using (E3).

#### 4.6.4 Comparison of metrological gain

The squeezing generated by QAOA in the superconducting hardware (see Fig. 4.12) is comparable to prior trapped ion works whose aim was to generate highly entangled states (see Fig.2 of Reference [332]). Even more, our depth-three QAOA simulations with 12 qubits indicate that with lower CNOT error rates, superconducting qubit hardware may reach  $-9.7$  dB of squeezing. The same method could also be applied in trapped ions, where the QAOA circuit—thanks to the trapped-ion all-to-all connectivity—is particularly compact (Sec. 4.4.4). In addition, our method has a number of advantages over variational ansätze developed explicitly for a metrological phase-estimation scenario [343, 407–409]. For example, Marciniak et al. [343], (i) use a cost function particularly tailored for sensing tasks that optimize the phase sensitivity of their quantum sensor, (ii) use a very general variational ansatz consisting of entangling and decoding unitaries containing rotation once and one-axis twisting operations twice in different directions, following the established knowledge in quantum metrology, and (iii) use a linear phase estimator with an unknown parameter  $a$  to estimate the phase from spin-measurement, where  $a$  also has to be optimized.

By contrast to (i), our cost function is the energy that also creates an entangled state. By contrast to (ii), our variational ansatz is grounded in QAOA, i.e., a Trotterized version of adiabatic computing with a classical optimization that inherits the performance guarantee in the limit  $p \rightarrow \infty$ . By contrast to (iii), our results can be applied to enhance phase-sensitivity but are not limited to it. Indeed, the optimization does not include any phase estimator, potentially making the approach useful for other problems while also reducing the number of parameters to optimize. These three points result in a variational form with fewer parameters to optimize than a general variational form and thus are, in principle, easier to optimize. Furthermore, the approach chosen here allows us to benefit from the vast literature on optimizing QAOA parameters, of which the TQA initialization is just one example.

## 4.7 Extension

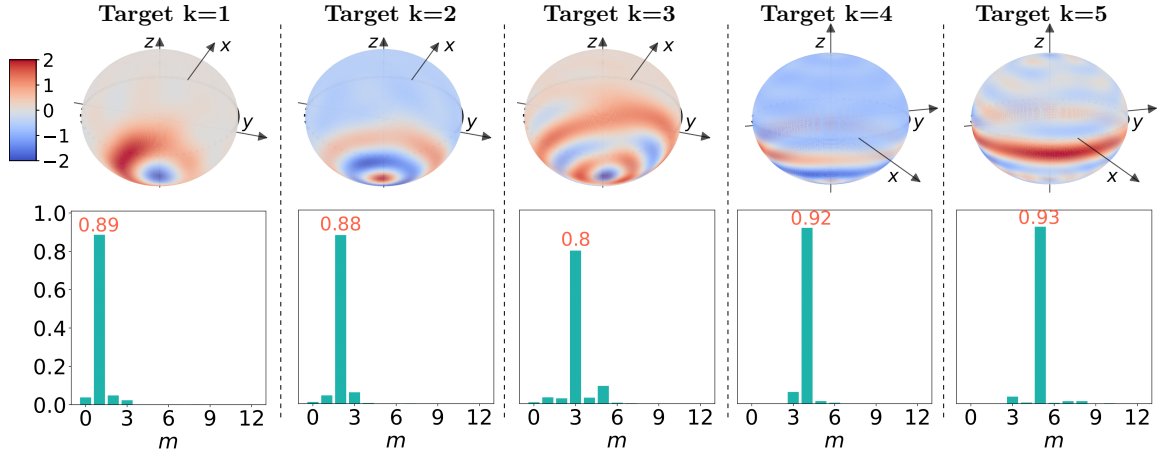
In this section, we extend the connection between QAOA and metrology to create arbitrary Dicke states and warm-starting QAOA with partially squeezed states for random MaxCut problems.

### 4.7.1 Creating arbitrary Dicke states

We can create arbitrary Dicke states (Eq. 4.9) by minimizing a QUBO cost function with QAOA. Let  $|x_{n-1} \dots x_0\rangle$  be a basis state in which qubit  $i$  is in state  $x_i \in \{0, 1\}$ . Each basis state in  $D_k^n$  satisfies the equation  $\sum_{i=0}^{n-1} x_i = k$ , which is a constraint on the binary variables  $x_i$ . We express this constraint as the QUBO problem

$$\min_{x \in \{0,1\}^n} \left( k - \sum_{i=0}^{n-1} x_i \right)^2. \quad (4.17)$$

The solution to this optimization problem is a superposition of all basis states with  $k$  qubits in the excited state, i.e.,  $D_k^n$ . We apply the change of variables  $x_i = (z_i + 1)/2$



**Figure 4.13.** Metrologically useful arbitrary Dicke states generated by a depth-three QAOA by minimizing the cost Hamiltonian Eq. (4.19). The top panels show the Wigner quasi-probability distribution on the Bloch spheres. The bottom panels show the corresponding histograms of the total spin operator  $\langle \hat{Z} \rangle = m$ . The orange numbers in each histogram show the overlap probability density  $|\langle D_k^{12} | \psi \rangle|^2$  with the target Dicke states.

to rewrite  $(k - \sum_i x_i)^2$  as

$$k^2 - kn + \frac{n}{4}(n+1) + \left(\frac{n}{2} - k\right) \sum_{i=0}^{n-1} z_i + \frac{1}{2} \sum_{i<j} z_i z_j. \quad (4.18)$$

After promoting each  $z_i$  variable to a Pauli spin operator  $\hat{Z}_i$ , Eq. (4.18) yields a cost Hamiltonian to minimize

$$\hat{H}_C = \left(\frac{n}{2} - k\right) \sum_{i=0}^{n-1} \hat{Z}_i + \frac{1}{2} \sum_{i<j} \hat{Z}_i \hat{Z}_j. \quad (4.19)$$

When  $k = n/2$ , we recover the MaxCut problem on the symmetric graph. For  $k \neq n/2$ , we have an extra term  $(n/2 - k) \sum \hat{Z}_i$  that biases the total spin towards  $\langle \hat{Z} \rangle = k$ . The Hamiltonian in Eq. (4.19) can therefore be used to generate the Dicke state  $D_k^n$  with QAOA.

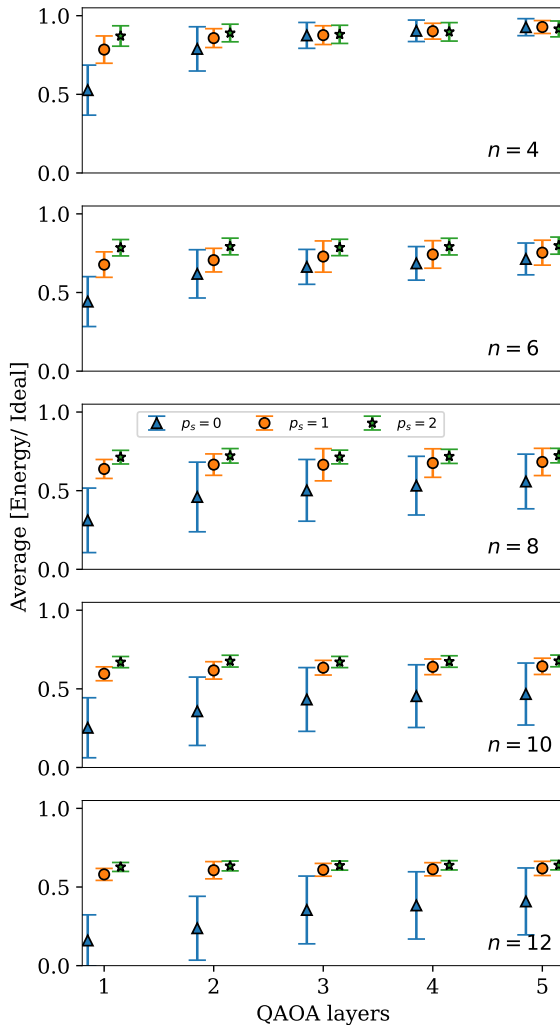
For  $n=12$  qubits, we use the cost Hamiltonian in Eq. (4.19) to simulate the generation of Dicke states with  $k = 1, 2, 3, 4, 5$ . With three QAOA layers, we obtain fidelities  $> 80\%$ , see Fig. 4.13. The corresponding QAOA parameters  $\gamma$  and  $\beta$  are shown in Tab. 4.3.

Num. spin up $k$	$\gamma_1$	$\beta_1$	$\gamma_2$	$\beta_2$	$\gamma_3$	$\beta_3$
1	0.101	0.903	0.317	1.324	1.506	-0.155
2	0.093	1.106	0.427	1.409	1.457	-0.068
3	0.149	1.205	1.645	1.576	0.472	-0.076
4	0.111	1.220	0.441	1.690	1.028	0.062
5	0.231	1.340	1.643	1.500	1.774	0.004

**Table 4.3.** The parameters  $(\gamma_i, \beta_i)$  of an optimized depth-three QAOA circuit to create  $k = 1, 2, 3, 4, 5$  Dicke states that are shown in Fig. 4.13.

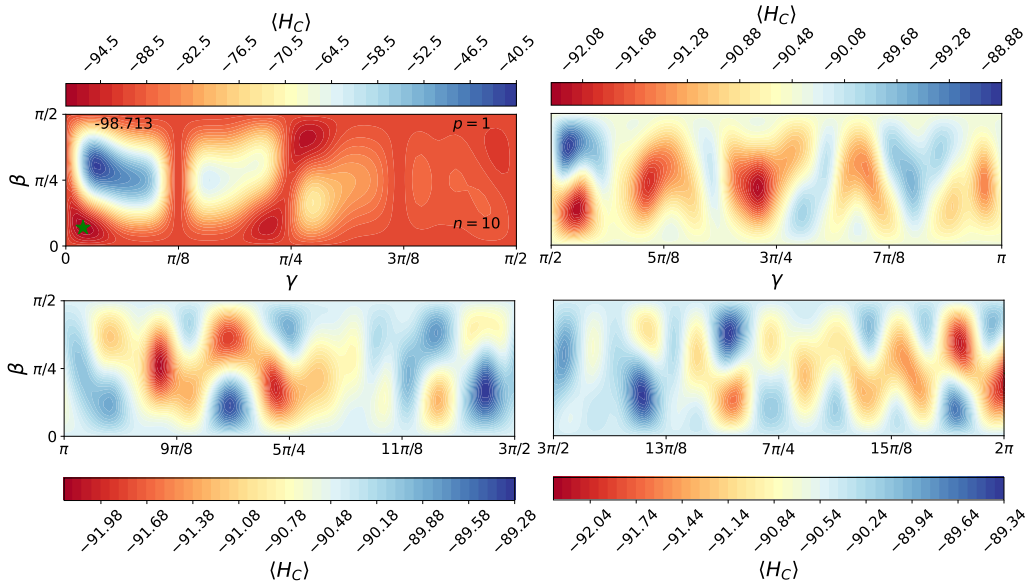
### 4.7.2 Warm-starting QAOA with squeezed states

In this section, we explore how far the symmetric MaxCut problem that has a Dicke state as a ground state can help solve non-trivial MaxCut problems with asymmetric edge-weights. We show how such squeezed states increase the likelihood of sampling good cuts on graphs with random edge weights. Each edge  $\omega_{i,j}$  of a graph is sampled from a Gaussian distribution  $\mathcal{N}(\mu, \epsilon)$  and then rounded to one decimal place to increase the separation in the cut-values of the graph. We compare standard QAOA with  $p$  layers to a QAOA with  $p_s + p$  layers in which the first  $p_s$  layers have fixed parameters to produce a squeezed state. Both methods, therefore, have  $2p$  parameters that require optimization for each graph instance. For the second approach, in addition,  $2p_s$  parameters are optimized once with the symmetric MaxCut problem as a target and are reused for different problem instances. For each  $n \in \{4, 6, 8, 10, 12\}$ , we sample 100 graph instances from  $\mathcal{N}(\mu, \epsilon)$  for which we chose  $\mu = 4$  and  $\epsilon = 0.5$  and optimize the cut-value for varying  $p$ . The resulting energy, normalized to the minimum energy and averaged over the 100 graph realizations, is used to compare both methods. To ensure that  $p$  layers always produce a result that is at least as good as the one for  $p - 1$  layers, we bootstrap the optimization parameters. The initial guess of the parameters for layer  $p$  are based on the optimized parameters of layer  $p - 1$ , i.e.,  $(\beta_1, \beta_2, \dots, \beta_p, \gamma_1, \gamma_2, \dots, \gamma_p)_{\text{initial}} = (\beta_1^{\text{opt}}, \beta_2^{\text{opt}}, \dots, \beta_{p-1}^{\text{opt}}, 0, \gamma_1^{\text{opt}}, \gamma_2^{\text{opt}}, \dots, \gamma_{p-1}^{\text{opt}}, 0)$ .



**Figure 4.14.** Advantages of QAOA initialized with squeezed states. The blue triangles show standard QAOA initialized from a coherent spin state. The orange circles and green stars show QAOA initialized with a spin-squeezed state created with  $p_s = 1$  and 2 QAOA layers, respectively. The  $x$ -axis is the QAOA depth after the initial state, and the  $y$ -axis is the energy normalized to the ideal value. The markers and error bars indicate the average and variance of 100 graph instances drawn from  $\mathcal{N}(4, 0.5)$  with different sizes  $n = 4$  (top) to  $n = 12$  (bottom). Squeezed initial states boost the average energy of the QAOA optimized state, as shown by the orange and green markers having an energy that is closer to the ideal energy than the blue markers. The energy increases only modestly as  $p$  increases due to the complexity of the optimization landscape.

QAOA initialized with squeezed states, shown as orange circles and green stars in Fig. 4.14, significantly improves the average energy when compared to QAOA initialized from an equal superposition, shown as blue triangles in Fig. 4.14. We observe a slight improvement in solution quality with increasing  $p$ . We attribute this to the complexity of the optimization landscape, which has many local minima, even at depth-one, due to the interference of the frequencies generated by the different edge weights, see Fig. 4.15. In the four-qubit case, the energy for  $p \geq 3$  layers of both methods is comparable. As the system size is increased, we observe a greater advantage for QAOA initialized with a squeezed state. These results indicate that, when solving a family of problems, it may be advantageous to initialize QAOA with a state that corresponds to the average problem even when such a problem is trivial to solve.



**Figure 4.15.** Complexity of the optimization landscape for a  $n = 10$  vertices graph with edge weights  $[3.1, 3.5, 3.9, 3.6, 4.6, 4.2, 4.8, 3.8, 4.1, 4.8, 5.0, 4.8, 4.0, 3.8, 3.2, 4.1, 4.2, 4.6, 4.3, 3.5, 3.9, 3.8, 3.2, 3.2, 4.7, 3.7, 4.1, 3.5, 4.1, 4.0, 4.2, 3.6, 4.4, 4.1, 3.5, 4.2, 3.7, 3.4, 4.4, 4.4, 3.6, 4.0, 4.3, 4.9, 4.1]$  and QAOA depth  $p = 1$ . Out of the full landscape, we show the first  $0$  to  $2\pi$  portion of the  $\gamma$  landscape in four subplots  $\gamma \in [0, \pi/2], [\pi/2, \pi], [\pi, 3\pi/2], [3\pi/2, 2\pi]$  with different color scales to increase the contrast between the local minima and maxima. This reveals a large number of local minima. The small improvement in solution quality with increasing  $p$  can therefore be attributed to the many local minima, created from the interference of the frequencies generated by the different edge weights.

## 4.8 Summary

In summary, the generation of squeezed states that are useful for metrology can be cast as a MaxCut problem, which in turn can be addressed with variational algorithms. Such squeezing also unveils the multipartite entanglement structure, as we discussed in detail. The procedure that we illustrated in the creation of a 12-qubit Dicke state can be implemented on universal quantum computing platforms, such as superconducting qubits or trapped ions, as well as on special-purpose machines such as BECs trapped in optical tweezers [349]. Interestingly, an enhancement of squeez-

ing within the multilayer QAOA protocol is not equivalent to simply applying the  $\hat{L}_z^2$  operator for a longer period, as the mixer Hamiltonian periodically intervenes. Our results show how variational algorithms may generalize existing protocols and provide systematic guidance for the creation of highly squeezed states for metrology. By contrast to, e.g., Ref [343, 407–409], which uses variational quantum algorithms with a hardware native ansatz to enhance phase sensitivity, the QAOA approach to create squeezing encapsulates the structure of the target state in the variational form, which may reduce the number of parameters to optimize. In a similar vein, we show that custom states beyond Dicke states can be generated by QAOA by casting them as solutions of a combinatorial optimization problem. In addition, we suggested squeezing as a QAOA-specific hardware benchmark. This benchmark is both portable across hardware platforms and captures hardware-specific properties such as limited qubit connectivity and cross-talk. On the other hand, warm-started QAOA using squeezed states seems to have an advantage as well.





## Chapter 5

# Genuine multipartite entanglement in quantum optimization

*entanglement barrier and relation to success probability*

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## 5.1 Introduction

Quantum effects are expected to play a central role in quantum algorithms designed to address specific computational tasks [195]. However, understanding their precise contribution to the algorithm’s performance is a subtle question. For instance, a key question is whether entanglement is necessary or sufficient in pure-state quantum computing [88, 410, 411]. Among the various applications of quantum computers, quantum combinatorial optimization stands out as particularly promising [22]. Here, the algorithm is tasked to find an optimal solution  $x^*$ , or a near-optimal one, of a classical optimization problem  $\min f(x)$  [412]. Interestingly, in quantum optimization approaches, such as quantum annealing (QA) [236, 413], the quantum approximate optimization algorithm (QAOA) [326], and variational quantum algorithms (VQA) [231], the initial state is typically a product state, while the solution state should encode a classical problem and thus (barring superpositions of degenerate solutions) may not contain any entanglement. Having both the initial and solution state with little to no entanglement raises the question of how much entanglement is genuinely required to make a quantum-optimization algorithm successful [414].

In previous years, most analyses focused exclusively on bipartite entanglement [414–418], where the entanglement between two non-overlapping partitions of the quantum state is computed. However, the behavior of bipartite entanglement can vary significantly depending on the chosen partitions [419, 420]. Additionally, a state may be entangled across all possible bipartitions or may exhibit global or multipartite entanglement, making it impossible to describe the total entanglement through contributions from individual bipartitions [421]. Since quantum optimization is, in essence, concerned with navigating a many-body system through a quantum spin-glass transition [422, 423], one may expect entanglement shared between multiple parties to play an important role. In the context of pure-state quantum computing, it is known that merely increasing bipartite entanglement with system size is insufficient for an exponential speed-up [88]. In contrast, multipartite entanglement shared among many parties can be a valuable resource for quantum information processing, such as in quantum secret sharing [424], multi-party quantum teleportation [425], quantum key distribution [426], quantum metrology [427], and measurement-based quantum computation [428]. However, its precise role in ensuring successful performance remains unclear for quantum optimization algorithms [325, 414, 429].

Here, we analyze the role of genuine multipartite entanglement in quantum optimization using the generalized geometric measure (GGM) of entanglement [115, 116, 141–143]. The GGM is an attractive measure for multipartite entanglement as it is experimentally accessible [430]. From detailed numerical benchmarks, we find—similar to the bipartite entanglement barrier [414, 418]—a multipartite entanglement barrier in quantum annealing. Initially, the circuit builds up multipartite entanglement as it approaches the minimal gap, where the system enters approximately a GHZ state between the two lowest-energy states. Subsequently, in successful optimization schedules, this multipartite entanglement is removed as the system approaches the solution state. We further analytically derive how the GGM in the optimization schedule is upper-bounded by the distances between the instantaneous eigenstate and both the initial and final product states, which can also be used as a proxy for bounding multipartite entanglement in experiments. Through our work, we shed light on the role and existence of multipartite entanglement in quantum optimization, and its relation to the success probability of the algorithm.

Our studies are complementary to a vigorous ongoing effort to understand the role of quantum resources in quantum optimization. E.g., there are works that study whether quantum optimization still performs well with a cap on entanglement [417, 431–436] or if the success of the algorithm requires other quantum resources [47].

The rest of this chapter is structured as follows. First, we provide a general overview of how entanglement manifests in quantum optimization, along with a definition of the GGM. From Sec. 5.2.3 onwards, we introduce quantum annealing and the MaxCut problem used in our simulation. Section 5.3 details the results from Trotterized quantum annealing, focusing on success probability, bipartite entanglement, and the GGM. In Sec. 5.4, we explore the emergence of the entanglement barrier and its connection to success probability. Finally, we conclude in Sec. 5.5.

## 5.2 Background

Here, we briefly summarize earlier works that relate bipartite entanglement and the performance of certain classes of quantum algorithms. Thereafter, we motivate the necessity of analyzing multipartite entanglement properties in quantum algorithms and define a computable measure for it, the GGM of multipartite entanglement. Finally, we describe the quantum optimization algorithm and the problem we investigate.

### 5.2.1 Entanglement in quantum optimization

Various efforts have already demonstrated how bipartite entanglement manifests in quantum optimization algorithms. The presence of von Neumann entanglement entropy has been observed numerically during quantum annealing sweeps in the exact cover problem [415]. Furthermore, entanglement witnesses in experiments have detected non-zero entanglement during quantum annealing [437]. Reference [414] explores a potential connection between bipartite entanglement and the success probability of adiabatic quantum optimization. It shows that high entanglement during the optimization does not necessarily mean a high success probability. On the contrary, in clean systems, significant final entanglement after slow sweeps suggests a superposition state rather than a separable ground state and thus implies a reduced probability of successfully finding the classical ground state. Therefore, measuring bi-

partite entanglement alone is insufficient to gauge the efficiency of adiabatic quantum optimization.

The subtle role of bipartite entanglement is also observed in the variational quantum eigensolver (VQE), where distributing the entangling gates according to the problem's topology increases success rates and reduces runtime [438]. Moreover, the ADAPT variant of QAOA uses a non-standard mixer, which allows a larger amount of entanglement in the earlier part of the circuit and accelerates convergence in the later stages [439]. To further understand the role of bipartite entanglement, one can simulate a quantum algorithm with matrix-product states with finite bond dimension ( $\chi$ ), since  $\chi$  bounds the maximum bipartite entanglement. Using this approach, reference [417] shows that the state fidelity in QAOA can be limited by  $\chi$ . In particular, QAOA needs a bond dimension that scales exponentially with the system size  $N$  to create high-fidelity states [418]. However, increasing the bond dimension beyond a certain threshold does not appear to improve the fidelity [435]. Also, reference [325] demonstrates numerically, as well as on IBM Quantum hardware, how QAOA can generate high entangled states with multipartite entanglement witnessed through squeezing inequalities and the quantum Fisher information.

### 5.2.2 Generalized geometric measure of entanglement

Recall from sec. 2.6.4 that the GGM of entanglement generalizes the notion of geometric measure of entanglement and quantifies genuine multipartite entanglement. Here we recapitulate the definition for the sake of convenience. For pure states, the GGM takes the simple form

$$G_2(|\psi\rangle) = 1 - \max_{|\pi\rangle_{\mathcal{A}:\mathcal{B}} \in \mathcal{S}_2} |\langle \pi | \psi \rangle|^2 = 1 - \max_{\mathcal{A}:\mathcal{B}} (\lambda_{\mathcal{A}:\mathcal{B}}^{\max})^2. \quad (5.1)$$

Here,  $|\psi\rangle$  belongs to the Hilbert space  $\mathcal{H}_{A_1} \otimes \mathcal{H}_{A_2} \otimes \cdots \otimes \mathcal{H}_{A_N}$ . The bipartition  $\mathcal{A} : \mathcal{B}$  satisfies  $\mathcal{A} \cup \mathcal{B} = \{1, 2, \dots, N\}$ , and  $\mathcal{A} \cap \mathcal{B} = \emptyset$  [142, 143]. In other words, maximizing  $|\langle \pi | \psi \rangle|$  across the bipartition  $\mathcal{A} : \mathcal{B}$  is equivalent to finding the maximum Schmidt coefficient,  $\lambda_{\mathcal{A}:\mathcal{B}}^{\max}$ , for that bipartition. For qubits, the theoretical maximum value of GGM is  $1/2$ .

Reference [440] investigates the GGM in Grover's algorithm. Here, the GGM initially increases and reaches a maximum in the first half of the optimal number of iterations and then decreases towards the end of the algorithm. In the Bernstein–Vazirani algorithm, the presence of geometric entanglement in the initial state prevents it from reaching an optimal performance (the key resource for the Bernstein–Vazirani algorithm is coherence rather than entanglement, and maximally entangled states cannot be maximally coherent) [436]. One can thus expect that the presence of an exaggerated amount of multipartite entanglement can have a detrimental effect, and understanding the nature of such entanglement may help us design more efficient quantum algorithms.

### 5.2.3 Trotterized quantum annealing

We have already discussed quantum annealing in detail in Sec. 3.3.2. Here, we use first-order Trotter–Suzuki decomposition of the continuous sweep, up to  $\mathcal{O}(\Delta t^2)$ , into

$$e^{-i\Delta t H(t)} \simeq e^{-i\beta H_M} e^{-i\gamma H_C}, \quad (5.2)$$

where  $\beta = (1 - t/T)\Delta t$ , and  $\gamma = (t/T)\Delta t$ . Discretizing the entire quantum annealing schedule into  $p$  time steps requires  $\Delta t = T/p$ . The resulting discretized annealing schedule is

$$\gamma_s = \frac{s}{p}\Delta t, \quad \beta_s = (1 - \frac{s}{p})\Delta t, \quad (5.3)$$

where the time-factor of  $t/T$  is replaced by the layer-number  $s/p$ . Crucially, unlike in quantum annealing, here  $\Delta t$  and the total time  $T$  can be modified independently by changing  $p$ , which hands us an additional control knob for Trotterized annealing.

### 5.2.4 Maximum-Cut problem

As a paradigmatic example, we focus on the Maximum Cut problem (MaxCut). MaxCut aims at partitioning the set of  $N$  nodes  $V$  in a graph  $\mathcal{G}(V, E)$ , such that the sum of the weights  $\omega_{i,j}$  of the edges  $(i, j) \in E$  traversed by the cut is maximum. By introducing Ising variables  $z_i \in \{-1, 1\}$  to describe which side of the cut node  $i$  falls on, this task is mathematically equivalent to maximizing the cost function (see Fig. 4.1 as an example)

$$\max_{z \in \{-1, 1\}^N} \frac{1}{2} \sum_{(i,j) \in E} \omega_{i,j} (1 - z_i z_j). \quad (5.4)$$

Since  $z_i$  can also be seen as the eigenvalues of the Pauli  $Z_i$  operator acting on a qubit  $i$ , the maximization is equivalent to finding the ground state of the Ising model [345, 441]

$$\tilde{H}_C = \frac{1}{\sqrt{N}} \sum_{i < j} J_{ij} Z_i Z_j. \quad (5.5)$$

Here,  $J_{ij}$  encode the edge weights of  $\mathcal{G}$  with  $J_{ij} = 0$  if  $(i, j) \notin E$ .

Due to the  $\mathbb{Z}_2$  symmetry of  $\tilde{H}_C$ , there are two equivalent degenerate solutions. A ground-state search may thus result in varying superpositions of the two solutions, whose entanglement content can depend significantly on the details of the annealing sweep. To ensure that the entanglement we observe is solely a signature of the performance of the quantum algorithm, we remove this degeneracy by adding a small perturbation  $fZ_0$  to  $\tilde{H}_C$  to break the symmetry. We choose  $f = 0.05$  for all problems. The modified Hamiltonian is thus

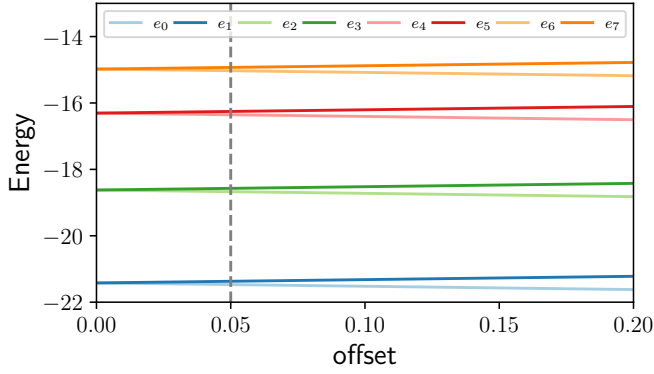
$$H_C = fZ_0 + \frac{1}{\sqrt{N}} \sum_{i < j} J_{ij} Z_i Z_j, \quad (5.6)$$

which will be the problem on which we study genuine multipartite entanglement.

### 5.2.5 Degeneracy breaking and pairing of states

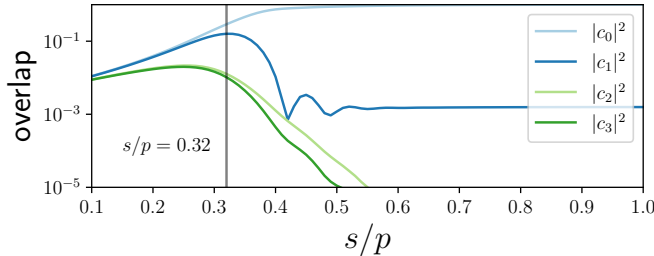
To break the  $\mathbb{Z}_2$  symmetry in MaxCut, we add an offset term as described in Eq. (5.6). Here, we show how the degeneracy in the spectrum of  $H_C$  is lifted with increasing offset. We choose 0.05 as an offset to lift the degeneracy while preserving the eigenlevel structure.

At this level of bias, although the  $\mathbb{Z}_2$ -symmetric states are no longer degenerate, they are still energetically close and thus have a pairing effect during the annealing



**Figure 5.1.** Effect of offset  $f$  on degeneracy of  $H_C$ . A non-zero magnitude  $f$  of the offset  $fZ_0$  lifts the degeneracy in the eigenvalue of  $H_C$ , as illustrated for the instance described in Fig. 5.3. As  $f$  increases, the paired states become increasingly distinguishable. For the data shown, we choose an offset of  $f = 0.05$  (grey vertical line) at which the states are non-degenerate but still form pairs close to each other in energy.

sweep. For example, let us consider the graph instance (Fig. 5.3) which we study in detail later. We study the overlap probability of the instantaneous state with the four lowest energy states  $|e_i\rangle$ ,  $i \in \{0, 1, 2, 3\}$  of  $H_C$  during the sweep. Until the sweep reaches the GGM peak, as we will see in Fig. 5.5(a), the population of the ground and the first excited states is almost equal, see Fig. 5.2. The same holds for the third and fourth excited states. This hints at a pairing of  $|e_0\rangle$  with  $|e_1\rangle$  and of  $|e_2\rangle$  with  $|e_3\rangle$ . Moreover, again due to the  $\mathbb{Z}_2$  symmetry,  $|e_0\rangle$  and  $|e_1\rangle$  are bit-wise orthogonal basis states (and the same for  $|e_2\rangle$  and  $|e_3\rangle$ ). An equal superposition of such states creates a GHZ-like state with a high GGM. For example, with the graph considered here  $|e_0\rangle = |10101001\rangle$  and  $|e_1\rangle = |01010110\rangle$  form the state  $|\text{GHZ}_8\rangle = \frac{1}{\sqrt{2}}(|e_0\rangle + |e_1\rangle)$  state.

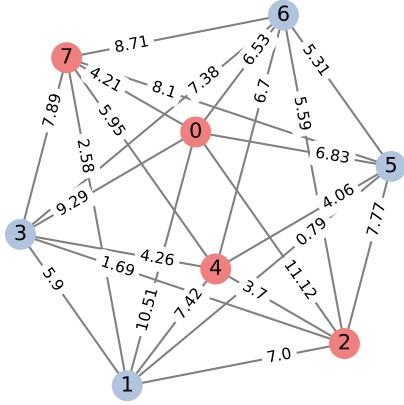


**Figure 5.2.** Population of the four lowest-energy instantaneous eigenstates, for the MaxCut problem from Fig. 5.3 and with  $p = 5000$ . Pairing is observed until the barrier, after which the pairs break, and  $|c_0(s)|^2$  approaches 1.

## 5.3 Genuine multipartite entanglement for a fixed instance of MaxCut

In this section, we illustrate how, for an instance of the MaxCut problem, the success probability, the von Neumann entanglement entropy, and the generalized geometric measure of multipartite entanglement behave during an optimization schedule. Unless stated otherwise, we choose the MaxCut problem on six-regular graphs on eight vertices, with edge weights  $J_{ij}$  randomly chosen from a Gaussian distribution with mean  $\mu = 6$  and standard deviation  $\sigma = 3$ . Figure 5.3 shows the resulting six-regular eight-node graph instance used. The solution of this MaxCut instance is defined by the separation into vertex sets  $(1, 3, 5, 6)$  and  $(0, 2, 4, 7)$ .

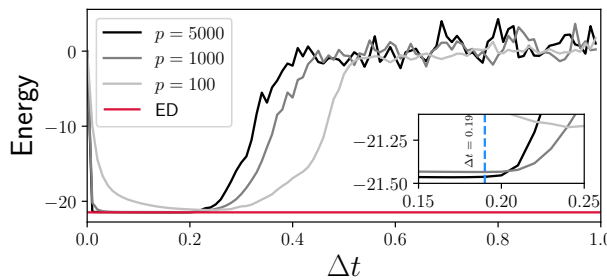
For illustration purposes, in this section, we choose the above graph instance, and in the next section, we choose from many instances to study the average behavior. We solve the problem with Trotterized quantum annealing and an increasing number



**Figure 5.3.** Details of the MaxCut problem for the fixed graph instance used in Sec. 5.3. Weights  $J_{ij}$  are indicated on the edges. The MaxCut solution is denoted by red and blue node colors.

of Trotter steps, or circuit layers, ranging from  $p = 100$  to  $p = 10000$ . We select  $\Delta t = 0.19$ , which allows us to achieve a good minimum energy across all layer counts (see Fig. 5.4). The total time is  $p\Delta t$ , which amounts to, e.g.,  $T = 950$  for  $p = 5000$ .

**$\Delta t$  for TQA.** In Trotterized quantum annealing (TQA), the quality of the annealing, which is related to the success probability, depends on the total number of Trotter layers  $p$  and the chosen time step  $\Delta t$ , see Eq. (5.3). In principle,  $\Delta t$  should be chosen to maximize the success probability. However, to study the impact of the sweep time (i.e., the number of layers multiplied by the step size  $\Delta t$ ) on the behavior of the GGM, we choose a value of  $\Delta t$  that keeps the energy of the final state small for all considered numbers of layers ( $p = 100, 1000, 5000$ ). Our simulations show that there is no  $\Delta t$  value that simultaneously minimizes the energy of all layers  $p$ , see Fig. 5.4. As a compromise, we choose  $\Delta t = 0.19$  for Fig. 5.5, for which the final energy is close to the absolute ground state for all  $p$  considered. In Fig. 5.9, when we average over 100 instances, we will use  $\Delta t = 0.2$  for all instances instead of optimizing  $\Delta t$  for each instance.



**Figure 5.4.** Energy of the final state of Trotterized quantum annealing as a function of  $\Delta t$  for different Trotter layers  $p$  for the MaxCut problem of Fig. 5.3, compared to the ground-state energy obtained from exact diagonalization (ED).

### 5.3.1 Success probability for different sweep speeds

As the number of Trotter steps increases, the evolution becomes more adiabatic, and thus the success probability is expected to improve with larger  $p$ . We define the success probability as the degree of overlap between the final state  $|\psi(p)\rangle_{\text{TQA}}$  and the true ground state  $|e_0\rangle$  of  $H_C$ . To understand the evolution of the state at each Trotter step, we evaluate the instantaneous probability of finding the ground state  $|e_0\rangle$  and first excited state  $|e_1\rangle$  of the final cost Hamiltonian at each layer  $s$ . Expressing the



instantaneous state in the eigenbasis of  $H_C$ ,

$$|\psi(s)\rangle_{\text{TQA}} = \sum_i c_i(s) |e_i\rangle, \quad (5.7)$$

the instantaneous probability of being in the ground and first excited states are  $|c_0(s)|^2$  and  $|c_1(s)|^2$ , respectively. Here, the basis states  $\{|e_i\rangle\}$  are ordered by increasing energy. The probability of reaching the final ground state,  $|c_0(s=p)|^2$ , defines the success probability of the algorithm.

For all  $p \in [1000, 10000]$ , the instantaneous ground-state probability  $|c_0|^2$  increases with layer index  $s$  and reaches the highest value at the end of the sweep  $s = p$ , see solid lines in Fig. 5.5(b). As the number of layers  $p$  increases, so does the success probability, i.e., slower sweeps are more successful. The first excited state probability  $|c_1|^2$  (dashed lines) increases until  $s/p \approx 0.32$ , after which it decreases if  $p$  is sufficiently large. This results in a peak, whose location is around the minimal energy gap between the first excited and the ground state of the instantaneous Hamiltonian [see Fig. 5.5(d)]. The peak becomes sharper as the sweep speed is reduced, i.e., as  $p$  increases.

For the smallest considered number of layers ( $p = 100$ , i.e.,  $T = 19$ ), an oscillation appears after crossing the minimum energy gap between the instantaneous ground and the first excited states. In such fast sweeps, the adiabatic conditions break down, causing the instantaneous state to become a superposition of multiple eigenstates. The dynamical phases between these lead to oscillations. One can also see these as the formation of defects, as described by the Kibble-Zurek mechanism (KZM) [442–444]. As the adiabatic condition becomes valid for slower sweep speeds, the defects and oscillations disappear.

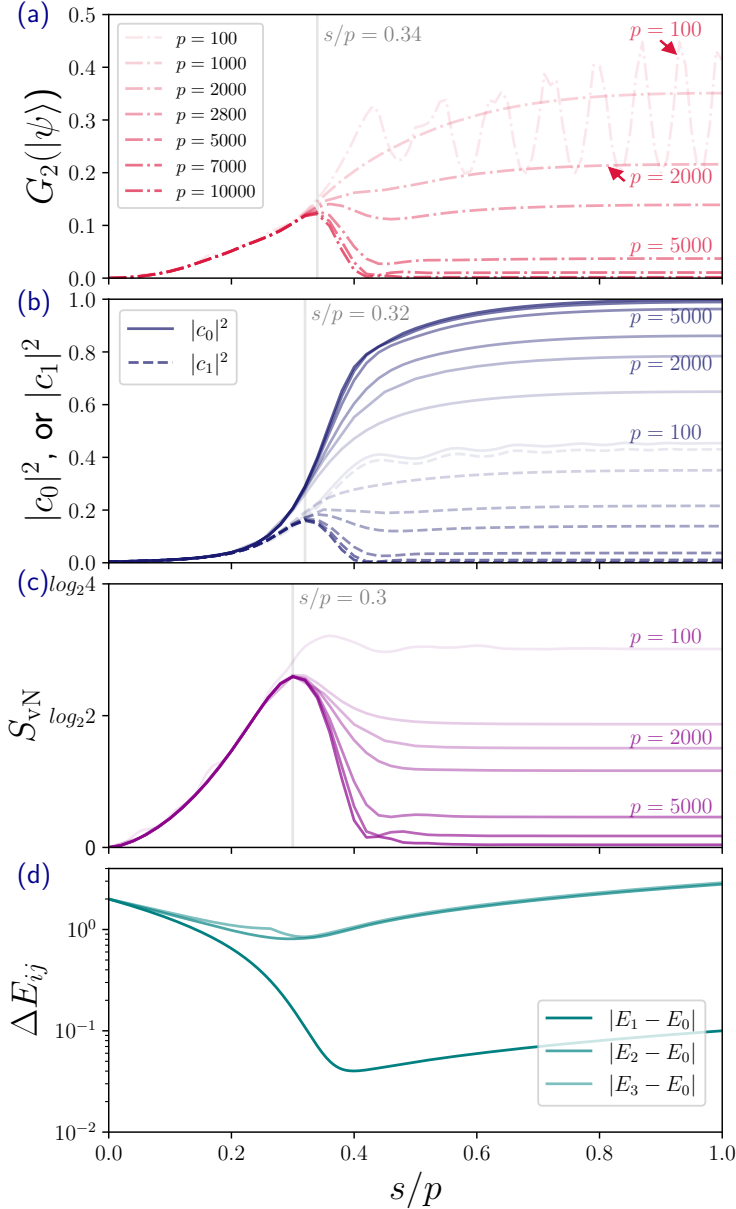
### 5.3.2 von Neumann entanglement entropy for bipartite entanglement

Our modified MaxCut problem, given by Eq. (5.6), has by design a non-degenerate solution state. Therefore, we expect the ideal quantum anneal to reach this desired product state solution with vanishing entanglement. However, any realistic sweep within finite time has some degree of non-adiabaticity, thereby populating higher energy states, which generates undesired final-state entanglement [414]. For comparison purposes to the GGM, we quantify the bipartite entanglement across the half–half bipartition of the system, i.e.,  $(0, 1, 2, 3) : (4, 5, 6, 7)$ , using the von Neumann entanglement entropy  $S_{vN} = -\text{Tr}[\rho_A \log_2(\rho_A)]$ . Note that  $S_{vN}$  depends on the chosen bipartition, whereas the GGM is optimized over all bipartitions.

As shown in Fig. 5.5(c), we observe a maximum in  $S_{vN}$  around the gap closing, reaching almost half of its theoretical maximum of  $4 \log_2(2)$  for the given subsystem size at  $s/p \simeq 0.3$ . Similar to reference [414], slow sweeps (large  $p$ ) reduce the entropy after this point as the quantum fluctuations produced by  $H_M$  become weaker, while fast sweeps (small  $p$ ) are unable to remove the entropy, resulting in a loss of success probability.

### 5.3.3 Generalized geometric measure for multipartite entanglement

For the fastest sweep considered,  $p = 100$ , the GGM oscillates in accord with the oscillations of the overlaps with the ground and excited states, see Fig. 5.5(a,b). The



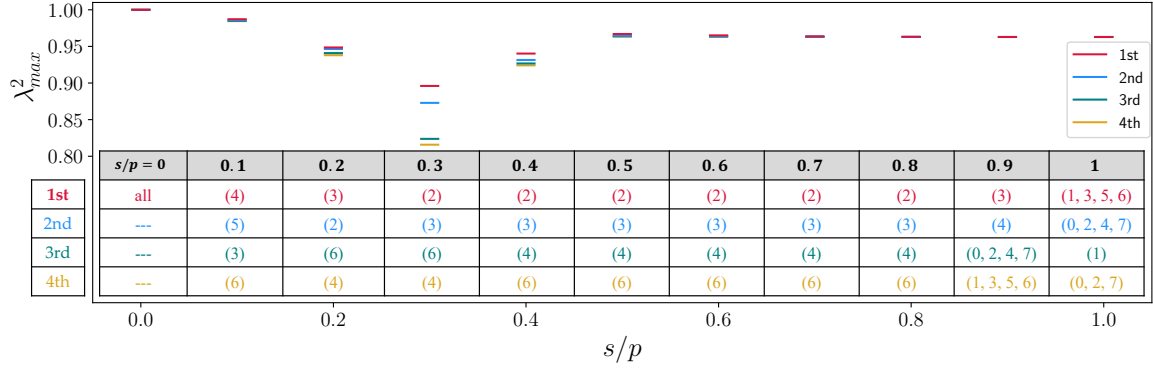
**Figure 5.5.** Results shown as a function of normalized layer number  $s/p$  for a single MaxCut instance, for various total TQA layers  $p$ . (a) For low  $p$ , the GGM saturates at high  $s/p$ , while for high  $p$ , a peak appears near  $s/p = 0.34$  followed by a dip. Fast sweeps ( $p = 100$ ) show oscillations due to non-adiabatic transitions. (b) Overlap of  $|\psi(s)\rangle$  with the ground and first excited state of  $H_C$ , with  $|c_1|^2$  peaking near max GGM. (c) The mid-graph von Neumann entropy peaks for all  $p$  but saturates lower as  $p$  increases. (d) Instantaneous energy gaps; key features in (a)–(c) align with low-lying gap closings.

GGM peak near  $s/p = 0.34$  becomes more visible with slower sweeps, which resembles the observed peak in  $|c_1|^2$ . As the GGM follows qualitatively the same structure as the von Neumann entropy on the fixed bipartition, one may wonder if the optimal bipartition for the GGM is static or if the half–half bipartition captures the GGM on average. For the particular problem instance we considered, we found that the optimal bipartition remains constant for  $0.3 \leq s/p \leq 0.8$  and is  $\mathcal{A} = (2), \mathcal{B} = (0, 1, 3, 4, 5, 6, 7)$ , which differs from the half–half bipartition (see Fig. 5.6).

### Optimal bipartitions corresponding to the GGM

To find the value of the GGM, we need to look for the bipartition that yields the maximum Schmidt coefficient. A question that may naturally arise is whether such a bipartition is unique or changes with the annealing sweep. At the start of the anneal, at  $s/p = 0$ , we have  $\lambda_{max}^2 = 1$  in all bipartitions. This implies a vanishing GGM, see Fig. 5.6. As the annealing schedule proceeds,  $\lambda_{max}^2$  of different partitions starts





**Figure 5.6.** Largest Schmidt coefficients along the annealing sweep, at selected times for the partitions  $(\mathcal{A}) : (\mathcal{B})$  yielding the four largest values, for the MaxCut problem of Fig. 5.3. The table gives the corresponding sets of qubits in  $\mathcal{A}$ . The maximum Schmidt values decrease from the initial states until  $s/p = 0.3$ , implying an increasing GGM. Later, the Schmidt values increase while the optimal bipartitions remain unchanged until  $s/p = 0.8$ . Subsequently, the GGM remains constant while the optimal bipartition changes to finally reach  $\mathcal{A} = (1, 3, 5, 6)$ .

to differ; for example, at  $s/p = 0.2$ , partition  $\mathcal{A} = (3), \mathcal{B} = (0, 1, 2, 4, 5, 6, 7)$  and partition  $\mathcal{A} = (2), \mathcal{B} = (0, 1, 3, 4, 5, 6, 7)$  have different values of  $\lambda_{max}^2$ . The individual Schmidt values decrease during the sweep until they reach the lowest  $\lambda_{max}^2$  corresponding to the GGM barrier at around  $s/p = 0.3$ . After this point, the GGM decreases, and the partition producing the  $\lambda_{max}^2$  —  $\mathcal{A} = (2), \mathcal{B} = (0, 1, 3, 4, 5, 6, 7)$  — does not change until  $s/p = 0.8$ . Towards the end of the anneal, in the range  $s/p > 0.8$ , the maximum Schmidt values and the GGM do not change considerably. However, the optimal bipartition varies until it finally reaches  $\mathcal{A} : \mathcal{B} = (1, 3, 5, 6) : (0, 2, 4, 7)$ . This bipartition has the maximum Schmidt values (maximal bipartite entanglement). In summary, it is interesting to see that from  $s/p = 0.1$  to  $0.3$ , the GGM changes, but the corresponding partition does not. Thereafter, and until the end of the anneal, the GGM remains constant, but the underlying partition changes.

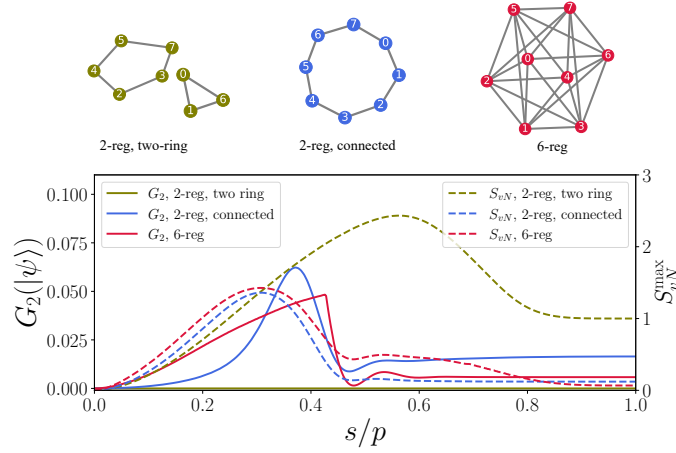
### Effect of graph's connectivity

One might naturally question whether a graph's connectivity affects the value of the GGM, for instance, if a more connected graph would display greater entanglement. Perhaps surprisingly, even a two-regular graph, with just two edges per node, can exhibit nearly the same amount of GGM as a six-regular graph, see Fig. 5.7. The same is true for the von Neumann entanglement entropy. Thus, it is crucial to understand in what sense bipartite and multipartite entanglement differ.

In general, bipartite entanglement for fixed bipartitions does not need to follow the behavior of the GGM, as we now show through some counterexamples. As discussed in Sec. 5.2.2, multipartite entanglement exists only when there is a non-zero bipartite entanglement for *all* possible bipartitions. This implies that a state can have a large bipartite entanglement for a particular bipartition and yet can have a significantly smaller GGM, even zero. To exemplify the strong dependence of the von Neumann entropy on the bipartition, we consider  $S_{vN}^{\max}$  as  $S_{vN}$  maximized over  $\binom{N}{\lfloor N/2 \rfloor}$  equal-sized (half-half) bipartitions,

$$S_{vN}^{\max} = \max_{|\mathcal{A}|=N/2:|\mathcal{B}|=N/2} S_{vN}[\mathcal{A}]. \quad (5.8)$$

For the 6-regular graph studied above, we observe that the GGM peaks later than  $S_{vN}^{\max}$ , and the same happens in the example of a connected two-regular graph, see Fig. 5.7. This implies that in both cases, the bipartition corresponding to the GGM is not the same as the maximal bipartition of  $S_{vN}^{\max}$ . An even more substantial difference between half-half von Neumann entanglement entropy and GGM can be found in the example of a disconnected two-regular graph, see the “2-reg, two ring” in Fig. 5.7. In this extreme example, while the maximum bipartite entanglement entropy is large throughout the entire annealing schedule, the optimal bipartition of the GGM corresponds to the disconnected subgraphs, resulting in an identically vanishing GGM.



**Figure 5.7.** The von Neumann entanglement entropy of a half-half bipartition and the GGM can peak at different times during an annealing sweep. A two-regular graph may exhibit similar entanglement strength as a six-regular graph. In disconnected graphs, a fixed bipartition may show large bipartite entanglement while the multipartite entanglement (GGM) can vanish.

### 5.3.4 Relation between state occupation probability and GGM

In Fig. 5.5(a,b), the GGM and the occupation probability of the first excited state  $|c_1|^2$  are quite comparable throughout the sweep, especially for larger  $p$ . This similarity becomes even more apparent by plotting  $G_2$  and  $|c_1|^2$  together in Fig. 5.8. This quantitative comparison shows that the GGM and  $|c_1|^2$  do neither strictly coincide with each other nor do one strictly bound the other. Instead,  $G_2$  is strictly bounded from above by the overlap probability with the initial state and the exact solution state, as we now prove.

Calculating the GGM of a given state  $|\psi\rangle$  is equivalent to finding the two-party product state  $|\pi\rangle_{\mathcal{A}:\mathcal{B}}$  with minimal distance from  $|\psi\rangle$ , see Eq. (5.1). As the annealing sweep terminates with the cost Hamiltonian, whose eigenstates  $|e_i\rangle$ , defined in Eq. (5.7) are product states, it seems reasonable to estimate the GGM by using these as a basis. The true GGM has to be smaller than the distance to any of these, i.e.,

$$G_2(|\psi(s)\rangle) \leq 1 - \underbrace{|\langle e_i | \psi(s) \rangle|^2}_{|c_i(s)|^2}, \quad \forall i. \quad (5.9)$$

Towards the end of a slow sweep, the expansion of Eq. (5.7) will have the largest weight in the ground-state of the cost Hamiltonian  $|e_0\rangle$ . We thus estimate the GGM by the upper bound

$$G_2(|\psi(s)\rangle) \leq 1 - |c_0(s)|^2. \quad (5.10)$$

Similarly, during the early stages of the sweep, the wave function will have a considerable weight in the initial product state  $|+\rangle^{\otimes N}$ . It is thus reasonable to expect

the distance to this state to provide a good upper bound in the initial part of the anneal,

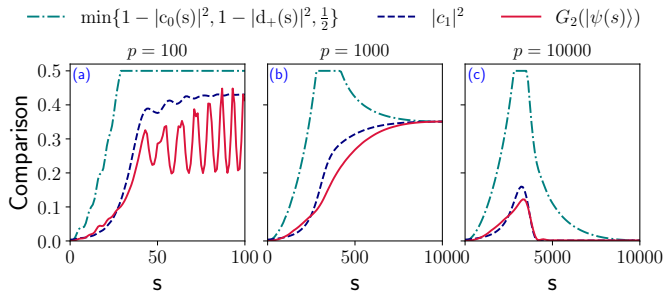
$$G_2(|\psi(s)\rangle) \leq 1 - |(\langle +|^{\otimes N})|\psi(s)\rangle|^2 \equiv 1 - |d_+(s)|^2. \quad (5.11)$$

Moreover, the GGM for qubits has a theoretical maximum of  $1/2$ . Putting all three bounds together, the GGM is upper bounded by

$$G_2(|\psi(s)\rangle) \leq \min \left\{ 1 - |c_0(s)|^2, 1 - |d_+(s)|^2, \frac{1}{2} \right\}. \quad (5.12)$$

Numerical experiments exemplify this bound, see Fig. 5.8. Considering the simplicity of the bound, the qualitative agreement is very satisfactory. In particular, the bound is saturated by  $1 - |c_0|^2$  towards the end of a successful anneal, as  $|c_0|^2$  becomes the most significant probability among all  $|c_i|^2$ s. Later, in Sec. 5.4.3, we further use Eq. (5.10), to relate the success probability, i.e., the final  $|c_0|^2$ , to the final GGM. Importantly, the values entering Eq. (5.12) are easily obtained experimentally by projecting the state onto the computational basis or a locally rotated basis. It can thus be used as a physically-informed proxy for bounding multipartite entanglement.

As remarked above, in addition, there seems to be a qualitative similarity between the first excited state overlap probability  $|c_1(s)|^2$  and the GGM  $G_2(\psi)$ , see Fig. 5.8. Towards the very end of the sweep for  $p = 1000, 10000$ , this similarity can be explained as only the states  $|e_0\rangle$  and  $|e_1\rangle$  remain prominent, i.e.,  $|c_0|^2 + |c_1|^2 \simeq 1$ , and the upper bound of the GGM saturates, thus giving  $G_2 = 1 - |c_0|^2 \simeq |c_1|^2$ . However, their similarity in the middle of the sweep cannot be explained solely by this reasoning. Nonetheless, we speculate that such a non-zero  $|c_1|^2$  has a role in keeping the instantaneous state entangled, as—assuming  $|c_1|^2 < |c_0|^2$ —it heralds the presence of an entangled superposition state.



**Figure 5.8.** The GGM  $G_2$  (solid red) is upper bounded by  $\min \{1 - |c_0(s)|^2, 1 - |d_+(s)|^2, \frac{1}{2}\}$  (dash-dotted teal). For slow sweeps ( $p = 1000, 10000$ ), the final state becomes a superposition of a few eigenstates, leading to  $G_2 \approx |c_1|^2$  (blue dashed), as seen in panels (b) and (c).

## 5.4 Genuine multipartite entanglement barrier in Random Max-Cut ensemble

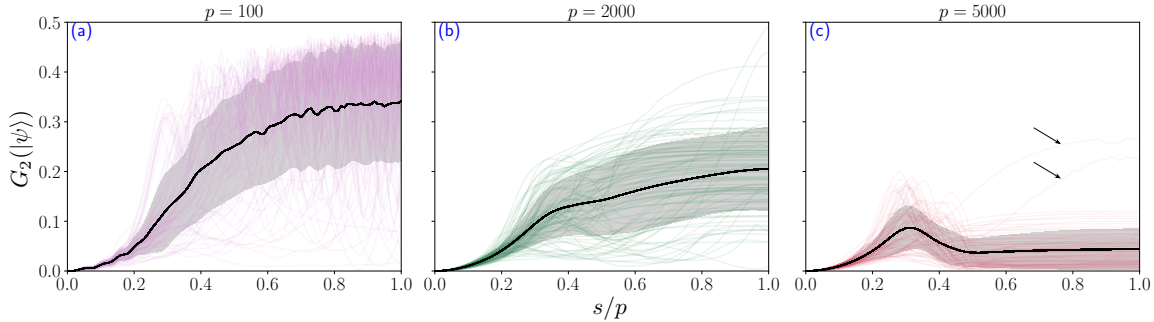
In Sec. 5.3, we have observed that the GGM assumes a peak during a slow quantum-annealing sweep, see Fig. 5.5(d). We call this a multipartite entanglement barrier since the inability to build up and then reduce the GGM prevents the algorithm from preparing the non-entangled solution state. We now investigate whether the observed barrier is (i) a universal feature not specific to the MaxCut problem instance used above and (ii) whether it is related to the instantaneous energy gap.

### 5.4.1 For random instances

We analyze 100 random MaxCut instances for eight vertices of 6-regular graphs with edge weights sampled from a Gaussian distribution  $\mathcal{N}(\mu = 6, \sigma = 3)$ . We run TQA with annealing sweep speeds  $p = 100, 2000, 5000$ . For  $p = 100$ , many instances display a peak or an oscillation in the GGM. However, there is no peak on average since the peaks occur at different times for different instances, see Fig. 5.9(a). Notably, the spread in the GGM at the beginning of the protocol is small. This trend is also observed for  $p = 2000$  and  $p = 5000$ .

For  $p = 2000$ , we observe a small hump also in the instance-averaged GGM around  $s/p = 1/3$ , see Fig. 5.9(b). The final state's GGM is lower as compared to  $p = 100$ . The entanglement peak, or barrier, becomes more strongly apparent at  $p = 5000$ , see Fig. 5.9(c). In this case, the multipartite entanglement remains relatively low throughout the sweep, and the final GGM is minimal, indicating that the protocol has reached the solution state with high probability. Nevertheless, there are problem instances for which the GGM increases until  $s/p = 1$ , e.g., see the two red lines in Fig. 5.9(c), which do not exhibit a multipartite entanglement barrier.

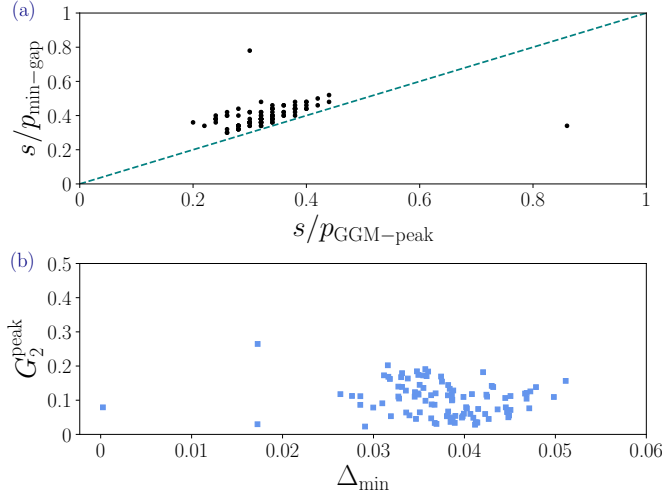
For comparison, we also investigate the GGM in a QAOA protocol with few layers but classically optimized angles  $\beta$  and  $\gamma$  in Eq. (5.2), see Sec. 5.4.4 for details. Here, we do not observe a peak in the GGM but rather a build-up of the GGM with each additional QAOA layer. This behavior is similar to a short-depth quantum anneal, as in Fig. 5.9(a), which is reasonable considering the few layers used in QAOA.



**Figure 5.9.** GGM from numerical benchmarks of 100 MaxCut instances. Panels (a) to (c), correspond to TQA with  $p = 100, 2000, 5000$  layers, respectively, i.e., increasing annealing time. For  $p = 100, 2000$ , although some individual instances have peaks in the GGM (refer to Fig. 5.5 for an example), the average GGM does not show any peak (panel a) or only a small hump (b). In contrast, for  $p = 5000$  (c), a peak in the GGM is also visible on average. With increasing  $p$ , the fluctuations in GGM between different instances decrease. Problem instances with no barrier are indicated by arrows. Again, strong oscillations typical for the defects generated by fast sweeps appear in  $p = 100$ .

### 5.4.2 Relation between GGM barrier and energy gap

In equilibrium, high entanglement can appear at quantum critical points with small energy gaps [361, 415]. However, this connection between low-lying spectrum and entanglement may not necessarily be expected in the out-of-equilibrium context of quantum optimization. Here, we investigate (a) whether the point where the energy gap closes correlates with the position of the GGM peak and (b) whether the size of the energy gap relates to the amount of GGM generated.



**Figure 5.10.** Relation between the multipartite entanglement barrier and the energy gap for 100 instances solved with TQA with  $p = 5000$  layers and  $dt = 0.2$ . (a) The time of minimum energy gap and the time of peak GGM are strongly correlated (99% confidence level). (b) The peak magnitude of GGM and the minimum energy gap show only weak anti-correlation (80% confidence level).

In Fig. 5.10(a), we plot the time  $s/p$  at which the energy gap and the GGM are minimum and maximum, respectively, for the same 100 problem instances as in Fig. 5.9. We report the first peak in the GGM to avoid ambiguities in case the GGM oscillates. The position of the minimum gap has a linear correlation with the position of the first GGM peak. Quantitatively, Pearson’s correlation coefficient is  $r = 0.28$  for the whole dataset, including the two extreme points. The corresponding  $t$ -score is  $t = |r|\sqrt{(n-2)/(1-r^2)} = 2.89$ , with  $n = 100$  problem instances. This implies a statistically significant correlation at the 99% confidence level. Excluding the two outliers increases Pearson’s correlation coefficient to  $r = 0.525$ . By contrast, we do not observe such a strong correlation between the magnitude of the energy gap and the size of the GGM peak, see Fig. 5.10(b). Indeed, the associated correlation coefficient of  $r = -0.16$  is only significant at the 80% confidence level. As in this out-of-equilibrium situation, the near-degeneracy between ground and first excited state is not the sole reason for entanglement, the absence of a strong correlation between the amount of entanglement and the energy gap is not unexpected.

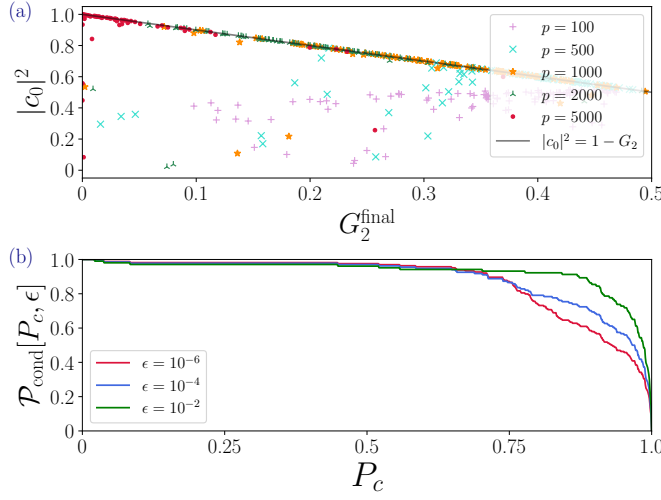
### 5.4.3 Relation to success probability

Generating and removing entanglement in quantum circuits are complex operations. For our MaxCut problems with a non-degenerate solution, quantum annealing must eliminate any existing entanglement to achieve the final product state. This raises an interesting question: Does the non-zero entanglement generated during quantum annealing hinder the optimization algorithm’s ability to find the optimal solution? Specifically, does a higher GGM in the final state limit the probability of success? If so, given the presence of an entanglement barrier, could a drop in the GGM from its peak enhance the chances of achieving a better success probability?

#### Final GGM and success probability

As a corollary to the Sec. 5.3.4, the success probability, i.e., the probability corresponding to the ground-state of the cost Hamiltonian,  $|c_0|^2$ , is bounded above by  $1 - G_2^{\text{final}}$ . In Fig. 5.11(a), we see that with an increasing number of layers (slower sweep), the success probability saturates the bound. In contrast, with fewer layers, the success probability does not saturate the bound, as the final state has support on multiple

eigenstates, making the bound of Eq. (5.10) using a single amplitude too loose. These results are similar to the observed relation between *bipartite* entanglement for a fixed bipartition and the success probability in Ref. [414].



**Figure 5.11.** Relation of entanglement and success probability, over 100 instances of 6-regular graphs with 8 vertices (a) Final success probability  $|c_0|^2$  is upper bounded by  $1 - G_2^{\text{final}}$ , and large  $p$  values approach this bound. (b) The conditional probability of achieving a success probability  $> P_c$  increases with greater disentanglement  $\Delta G$ , shown for  $\epsilon > \{10^{-6}, 10^{-4}, 10^{-2}\}$  across 100 instances with  $p = 2000$  and  $p = 5000$ .

### Disentanglement and success probability

As a complementary analysis, we also study how the amount of disentanglement, i.e., the reduction of GGM after its peak,  $\Delta G = G_2^{\text{max}} - G_2^{\text{final}}$ , is related to the success probability. To this end, we calculate the conditional probability ( $\mathcal{P}_{\text{cond}}[P_c, \epsilon]$ ) of obtaining a success probability greater than some  $P_c$ , given that the sweep has shown some minimum amount of disentanglement,  $\Delta G > \epsilon$

$$\mathcal{P}_{\text{cond}}[|c_0|^2 > P_c | \Delta G > \epsilon] = \frac{\mathcal{P}[(|c_0|^2 > P_c) \cap (\Delta G > \epsilon)]}{\mathcal{P}[\Delta G > \epsilon]}. \quad (5.13)$$

We study only the cases where a GGM barrier is observed, i.e.,  $G_2^{\text{max}} > G_2^{\text{final}}$ . We choose  $p = 2000, 5000$ , computing the TQA of 100 instances for both numbers of layers. To show reasonable comparison, we need to choose small values of  $\epsilon = (10^{-6}, 10^{-4}, 10^{-2})$ , as the final GGM is itself small in terms of absolute values. We find a larger  $\Delta G$  (increasing  $\epsilon$ ) to result in a larger success probability, see Fig. 5.11(b). In particular, large values of  $P_c$  are capped by the given fixed minimum disentanglement, leading to a downward bend in the curves [Fig. 5.11(b)] and suggesting the necessity of a large amount of disentanglement for a good success probability.

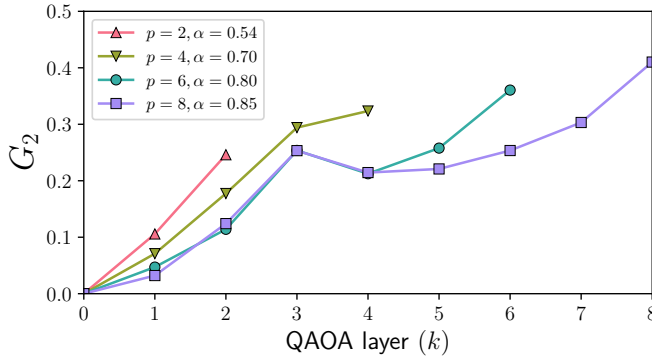
#### 5.4.4 GGM in Quantum Approximate Optimization Algorithm

In this chapter, we are primarily interested in Trotterized quantum annealing. Nonetheless, it is interesting to compare the results with those obtained with a QAOA protocol. The standard QAOA applies  $p$  layers of the unitaries  $\exp(-i\beta_k H_M) \exp(-i\gamma_k H_C)$ , with  $k = 1, \dots, p$  on the initial state  $|+\rangle^{\otimes N}$  to create a trial state  $|\psi(\gamma, \beta)\rangle$ . Then, a classical optimizer seeks the optimal parameters  $\beta = (\beta_1, \dots, \beta_p)$  and  $\gamma = (\gamma_1, \dots, \gamma_p)$  that minimize the energy  $\langle \psi(\beta, \gamma) | \hat{H}_C | \psi(\beta, \gamma) \rangle$ , which is measured by the quantum



processor. As optimizing over  $2p$  parameters in the classical optimizer is a bottleneck, we limit our simulation to small depths  $p$ .

In its spirit, QAOA is an approximate algorithm. It is designed to provide samples of a good set of solutions using only short-depth circuits (in contrast to quantum annealing, where one is typically more interested in slow but near-optimal runs). Therefore, as a figure of merit to quantify the performance of QAOA, we use the approximate ratio  $\alpha = \langle E \rangle / E_{\text{exact}}$  instead of success probability. We run QAOA with different depths  $p$  for the same MaxCut instance we consider in Sec. 5.3.



**Figure 5.12.** The GGM ( $G_2$ ) of the state obtained after layer  $k = 0, \dots, p$  of a depth- $p$  QAOA. The approximation ratio (see legend) and final GGM both increase with  $p$ .

For fixed depth  $p$ , the GGM increases with each QAOA layer  $k$ ; however, for larger  $p$ , there is a kink at  $k = 3$ . Nonetheless, the approximation ratio  $\alpha$  and the final GGM increase with QAOA depth  $p$  without a clear presence of an entanglement barrier. This is akin to the observed feature in the GGM when the number of TQA layers is small, see  $p = 100$  in Fig. 5.9(a). Recall that the multipartite entanglement barrier appears when the instantaneous overlap probability of the ground state  $|c_0|^2$  and the first excited state  $|c_1|^2$  starts to become distinguishable, see Fig. 5.5(b) and Fig. 5.2. Thus, a large GGM may not necessarily hinder its performance as long as it stems from the superposition of the actual ground state  $|e_0\rangle$  with a few excited states  $|e_{i>0}\rangle$ . In that case, there is a high probability that a manageable number of repetitions of the algorithm will then permit us to measure the optimal state on the computational basis at least once.

## 5.5 Summary and outlook

To summarize, multipartite entanglement arises in Trotterized quantum annealing even when the solution of the classical optimization problem is non-degenerate. While fewer layers—or a faster sweep—generate more entanglement, increasing the number of layers causes the GGM to grow, reach a peak, and then decline to zero during the sweep—a phenomenon one can call a multipartite entanglement barrier. We prove that the success probability is upper-bounded by  $1 - G_2^{\text{final}}$ , which is saturated when the final state is close to a product state (such as the desired solution state), as illustrated by numerical results from MaxCut instances. Thus, achieving a higher success probability typically requires lower final-state entanglement. Such a relation can be leveraged to benchmark quantum hardware [325, 443, 445, 446]. From the phenomena of the entanglement barrier, we quantify the relation between disentanglement and success probability through a conditional probability and find that a stronger drop in multipartite entanglement after the GGM peak improves the success probability.

Moreover, as the GGM corresponds to an optimal bipartition with minimal bipartite entanglement [447], insights about GGM can be used in circuit-cutting protocols to find an optimal cut such that the multipartite entanglement loss is minimal [448].

Our work has consequences for short-depth protocols such as QAOA and its variants. The classical parameter optimization loop is time-consuming [301], and navigating the optimization landscape in variational quantum algorithms is NP-hard [449]. Therefore, recent research investigates methods to obtain QAOA parameters by classical means. In particular, tensor networks may help to obtain good parameters [260]. Here, it is sufficient to approximate the loss landscape to produce good QAOA parameters qualitatively. Our work shows that multipartite entanglement can build up in QAOA. An interesting research question is thus whether tensor-network methods can provide a qualitative approximation of the loss landscape that is good enough to produce good QAOA parameters at a classically tractable bond dimension.

Further research could explore how the multipartite entanglement barrier manifests if the Hamiltonian is engineered using the tools from shortcuts to adiabaticity [450]. By exploring possible links to the graph's spectral properties, one may hope to understand better why GGM barriers cluster in specific regions [451]. Moreover, understanding the source of such entanglement in terms of physical phenomena, e.g., the Landau–Zener transition [444] or multiqubit quantum tunneling [452], one may hope to better predict the possibility or impossibility of reaching the solution state [243]. Our work focuses on pure states. However, some quantum algorithms may leverage mixed states [453–456], and it will be interesting to use the GGM to study the role of multipartite entanglement in such cases [141, 146, 147].



## Chapter 6

# Validating quantum effects in quantum algorithm

*Do quantum processing units generate entanglement in QAOA?*

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## 6.1 Introduction

In recent years, quantum computation has seen sustained and accelerating progress in both algorithmic development [457–459] and experimental platforms [288, 460–462]. Among the many potential applications, quantum optimization stands out as a particularly promising area, where even modest improvements over the best-known classical methods could lead to significant advancements across scientific and industrial domains. Hybrid quantum-classical algorithms, such as the Quantum Approximate Optimization Algorithm (QAOA) [81, 237], have emerged as viable approaches for harnessing the capabilities of noisy intermediate-scale quantum (NISQ) devices. These algorithms combine quantum circuits with classical optimization routines, leveraging the strengths of both computational paradigms. Despite notable improvements in quantum hardware, including increased qubit counts and extended coherence times, the performance of QAOA is often limited by the efficiency and effectiveness of the classical optimization component. This raises a critical question: to what extent does the quantum processing unit (QPU) contribute to the overall performance of the algorithm? In other words, if classical routines primarily perform the heavy lifting, are there genuinely quantum effects at play that are essential to the algorithm’s success? Empirically investigating such “quantumness” within QAOA is crucial, as any claims of quantum advantage hinge upon it. Without meaningful quantum contributions, the potential for QAOA to surpass classical methods would be fundamentally undermined.

We address this question by measuring entanglement on IBMQ hardware for classically optimized QAOA. We find that the optimized QAOA on hardware exhibits entanglement, which is qualitatively comparable to that in a noiseless simulation up to a certain circuit depth. To detect entanglement, we employ an entanglement measure inspired by the geometric measure of entanglement of Chapter. 5.

In this chapter, first, in Section 6.2, we describe our measurement protocol in detail, including how we estimate entanglement from a fixed bipartition using a hardware-efficient SWAP test, and how Schmidt coefficients can be extracted from Rényi entropy. Section 6.3 presents benchmarks of our method against exact simulations on small systems, analyzing its robustness under realistic noise models. We then move to implementation in Section 6.4, where we discuss the practical challenges of executing the protocol on IBM quantum devices. Finally, in Section 6.5, we apply our protocol to QAOA circuits across various system sizes up to 76 qubits and analyze

how entanglement evolves in QAOA. This progression allows us to assess the presence of entanglement in QAOA circuits experimentally.

## 6.2 Methods: Obtaining GGM in experiment

Measuring entanglement can offer profound insights into quantum correlations, extending beyond the conventional benchmarking of individual qubit performance in noisy intermediate-scale quantum (NISQ) devices [463–466]. Yet, its role in quantum optimization remains largely unexplored. As hybrid quantum algorithms gain attention, understanding entanglement becomes essential for two reasons: (1) to quantify how much genuine quantum resource the algorithm is using, and (2) to assess whether any observed advantage truly stems from quantum effects beyond classical optimization capabilities.

One established method for probing entanglement is the SWAP test [467–470], which compares copies of a quantum state to measure the purity of the subsystem. From this, one can compute quantities such as the Rényi entropy or, in ideal scenarios, the von Neumann entropy, provided that multiple Schmidt coefficients can be resolved. However, creating multiple identical copies is challenging on current small-scale hardware.

### 6.2.1 SWAP test to Rényi entropy

Consider a system composed of two subsystem  $\mathcal{A}$  and  $\mathcal{B}$ , which can be written as

$$|\psi\rangle = \sum_{ij} c_{ij} |a_i\rangle \otimes |b_j\rangle, \quad (6.1)$$

where the states  $|a_i\rangle$  and  $|b_j\rangle$  form orthonormal bases of  $\mathcal{A}$  and  $\mathcal{B}$ . The reduced density matrix for subsystem  $\mathcal{A}$  is defined by tracing over  $\mathcal{B}$ ,

$$\rho_A = \text{Tr}_B(|\psi\rangle \langle\psi|), \quad (6.2)$$

which contains the information about entanglement between  $\mathcal{A}$  and  $\mathcal{B}$ . For example, one can define the  $n$ th Rényi entropy,

$$S_n = \frac{1}{1-n} \log(R_n), \quad R_n = \text{Tr}(\rho_A^n). \quad (6.3)$$

Similarly, one can also calculate von Neumann entanglement entropy,

$$S_{\text{vN}} = -\text{Tr}[\rho_A \log \rho_A]. \quad (6.4)$$

Now, the main challenge in the experiment is to obtain  $R_n$  without estimating the state  $\rho_A$ , which is costly in terms of tomography. On the other hand, it is clear that to find the trace of the power of a matrix, knowing the entire matrix is overkill. In this scenario, the SWAP test provides a way.

**SWAP test.** In quantum computation, the SWAP test is a well-known method to check how much two states differ [471,472]. To demonstrate the method, let's consider two states  $|\phi\rangle$  and  $|\psi\rangle$ . Now, if we follow the circuit in Fig. 6.1, we can see that after Hadamard and C-SWAP, the state becomes

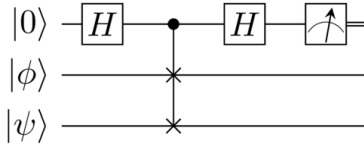
$$\frac{1}{\sqrt{2}}(|0, \phi, \psi\rangle + |1, \psi, \phi\rangle), \quad (6.5)$$

and then the second Hadamard results in

$$\frac{1}{2} |0\rangle (|\phi, \psi\rangle + |\psi, \phi\rangle) + \frac{1}{2} |1\rangle (|\phi, \psi\rangle - |\psi, \phi\rangle). \quad (6.6)$$

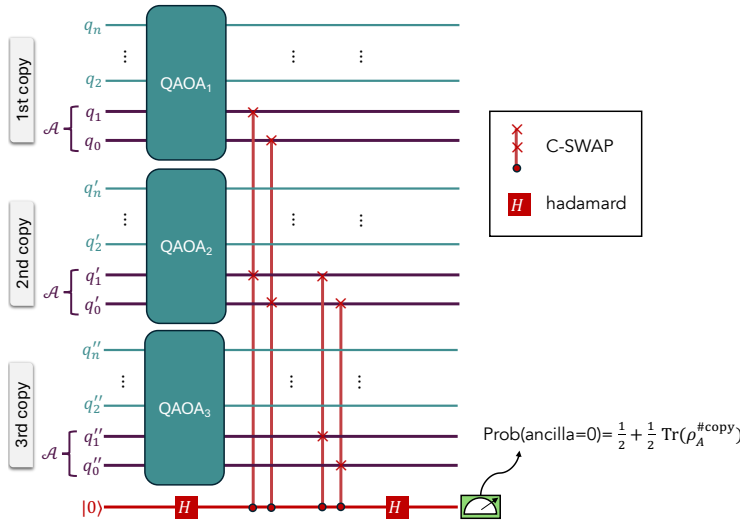
Thus, the probability of ancilla being 0 captures their overlap,

$$\text{Prob}(\text{ancilla} = 0) = \frac{1}{2} + \frac{1}{2} |\langle \psi | \phi \rangle|^2. \quad (6.7)$$



**Figure 6.1.** Circuit for the quantum SWAP test: used to estimate the overlap between two quantum states. An ancilla qubit controls a SWAP operation between the two input states, and the probability of measuring  $|0\rangle$  on the ancilla is related to the fidelity between the states (from Wikipedia).

The SWAP test, originally used to assess similarity between states, can also estimate subsystem purity—e.g., applying it to subsystem  $\mathcal{A}$  yields  $\text{Tr}(\rho_A^2)$ , and with  $n$  copies, one can access  $\text{Tr}(\rho_A^n)$ . We elaborate on this below.



**Figure 6.2.** Multi-copy SWAP test: By preparing multiple copies of a quantum state and applying the SWAP test on a subset of qubits ( $\mathcal{A}$ ), one can estimate the Rényi entropy of order  $n$ , which is proportional to  $\text{Tr}(\rho_A^n)$ . This procedure requires  $n$  copies of the state. The estimation is obtained from the probability of measuring the ancilla qubit in the  $|0\rangle$  state. The SWAP test involves a controlled-swap (or controlled-permutation) operation between corresponding qubits in the copies, with the ancilla (control) qubit encoding the interference information.

Let's consider two copies of the state:  $|\psi\rangle = \sum_{ij} c_{ij} |a_i\rangle |b_j\rangle$  and  $|\psi'\rangle = \sum_{i'j'} c_{i'j'} |a_{i'}\rangle |b_{j'}\rangle$ . The full circuit for measuring  $R_n$  involves applying a Hadamard gate to an ancilla qubit, followed by a controlled-SWAP (C-SWAP), and then another Hadamard gate,

as shown in Fig. 6.2. Let's consider the simplest case with just two copies:

$$\begin{aligned}
 |\psi\rangle |\psi'\rangle |0\rangle &\xrightarrow{\mathbb{I} \otimes \mathbb{I} \otimes H} |\psi\rangle |\psi'\rangle |+\rangle \\
 &\xrightarrow{C\text{-SWAP}_A} \frac{1}{\sqrt{2}} (|\psi\rangle |\psi'\rangle |0\rangle + \sum_{ij} \sum_{i'j'} c_{ij} c_{i'j'} |a_{i'}\rangle |b_j\rangle |a_i\rangle |b_{j'}\rangle |1\rangle) \\
 &\xrightarrow{\mathbb{I} \otimes \mathbb{I} \otimes H} \frac{1}{2} \left[ \left( |\psi\rangle |\psi'\rangle + \sum_{ij} \sum_{i'j'} c_{ij} c_{i'j'} |a_{i'}\rangle |b_j\rangle |a_i\rangle |b_{j'}\rangle \right) |0\rangle \right. \\
 &\quad \left. + \left( |\psi\rangle |\psi'\rangle - \sum_{ij} \sum_{i'j'} c_{ij} c_{i'j'} |a_{i'}\rangle |b_j\rangle |a_i\rangle |b_{j'}\rangle \right) |1\rangle \right]. \tag{6.8}
 \end{aligned}$$

From here we can see the probability of ancilla being 0 is given by

$$\begin{aligned}
 \text{Prob}(\text{ancilla} = |0\rangle) &= \frac{1}{4} \left\| \left( |\psi\rangle |\psi'\rangle + \sum_{ij} \sum_{i'j'} c_{ij} c_{i'j'} |a_{i'}\rangle |b_j\rangle |a_i\rangle |b_{j'}\rangle \right) \right\|^2 \\
 &= \frac{1}{2} + \frac{1}{2} \sum_{ij} \sum_{i'j'} c_{ij} c_{i'j'} \langle \psi | \langle \psi' | a_{i'} \rangle |b_j\rangle |a_i\rangle |b_{j'}\rangle \\
 &= \frac{1}{2} + \frac{1}{2} \sum_{kl} \sum_{k'l'} \sum_{ij} \sum_{i'j'} c_{kl}^* c_{k'l'}^* c_{ij} c_{i'j'} \underbrace{\langle a_k | a_{i'} \rangle}_{\delta_{ki'}} \underbrace{\langle b_l | b_j \rangle}_{\delta_{lj}} \underbrace{\langle a_{k'} | a_i \rangle}_{\delta_{k'i}} \underbrace{\langle b_{l'} | b_{j'} \rangle}_{\delta_{l'j'}} \\
 &= \frac{1}{2} + \frac{1}{2} \sum_{ii'jj'} c_{ij}^* c_{i'j'}^* c_{ij} c_{i'j'}. \tag{6.9}
 \end{aligned}$$

We can identify the last term as  $\text{Tr}(\rho_A^2)$ ,

$$\begin{aligned}
 \rho_A &= \sum_{ii'jj} c_{ij}^* c_{i'j} |a_{i'}\rangle \langle a_i| \Rightarrow \rho_A^2 = \sum_{kk' ll} \sum_{ii'jj} c_{kl}^* c_{k'l} c_{ij}^* c_{i'j} |a_{k'}\rangle \langle a_k| a_{i'}\rangle \langle a_i| \\
 &\Rightarrow \text{Tr}(\rho_A^2) = \sum_{ii'jl} c_{il}^* c_{i'l} c_{ij}^* c_{i'j} = \sum_{ii'jj'} c_{ij}^* c_{i'j'}^* c_{ij} c_{i'j'}. \tag{6.10}
 \end{aligned}$$

Thus, by measuring the probability of the ancilla being in  $|0\rangle$ , one can obtain the information about  $\text{Tr}(\rho_A^2)$  in the experiment.

## 6.2.2 $R_n$ to Schmidt values

To obtain von Neumann entropy, one needs to have the eigenvalues of  $\rho_A$  instead of only the trace of it. Or, one can extract the Schmidt eigenvalues from there. We know that for a state  $|\psi\rangle \in \mathbb{C}^{ab} = \mathbb{C}^a \otimes \mathbb{C}^b$ , we can do Schmidt decomposition to obtain

$$|\psi\rangle = \sum_{k=1}^D \lambda_k |u_k\rangle \otimes |v_k\rangle, \quad \{|u_k\rangle\} \in \mathbb{C}^a, \{|v_k\rangle\} \in \mathbb{C}^b. \tag{6.11}$$

Now, we need to obtain the Schmidt values, i.e., eigenvalues of  $\rho_A$  from the information of  $R_n = \text{Tr}(\rho_A^n)$ . We can use the Newton–Girard formula to obtain the roots of a polynomial made of  $R_n$  [469, 473].

**Newton-Girard formula.** It connects power sums  $\sum_i x_i^k$  to the elementary symmetric polynomials made of those, for example,

$$e_0 = 1, \quad e_1 = \sum_i x_i, \quad e_2 = \sum_{1 \leq i < j \leq n} x_i x_j. \quad (6.12)$$

The Newton-Girard formula shows how the roots form the symmetric polynomials,

$$\prod_{i=1}^n (x - x_i) = \sum_{k=0}^n (-1)^k e_k x^{n-k}. \quad (6.13)$$

To obtain the eigenvalues of  $\rho_A$ , we can replace the LHS of Eq. (6.13), by  $\det\{(xI - \rho_A)\}$ , where  $x$  will be the eigenvalues or Schmidt probabilities. Then it is evident that  $e_1 = \text{Tr}(\rho_A)$ , and we can understand what is  $e_2$ , by investigating the structure of

$$\begin{aligned} R_2 &= \text{Tr}(\rho_A^2) = \sum_i x_i^2, \\ e_2 &= \sum_{1 \leq i < j \leq n} x_i x_j = \left( \underbrace{\left( \sum_i x_i \right)^2}_{=1} - \sum_i x_i^2 \right) / 2 = \frac{1}{2}(1 - R_2). \end{aligned} \quad (6.14)$$

Similarly, we can see that for  $n = 3$ ,

$$R_3 = \sum_i x_i^3, \quad e_3 = \sum_{1 \leq i < j < k \leq n} x_i x_j x_k, \quad (6.15)$$

and using basic algebra, we can see that,

$$\begin{aligned} \left( \sum_i x_i \right)^3 &= \sum_{i=1}^n x_i^3 + 3 \underbrace{\sum_{i,j=1, i \neq j}^n x_i^2 x_j}_{= (\sum_{i=1}^n x_i) (\sum_{i=1}^n x_i^2) - \sum_{i=1}^n x_i^3} + 6 \sum_{1 \leq i < j < k \leq n} x_i x_j x_k \\ &\Rightarrow 1 = R_3 + 3(R_2 - R_3) + 6e_3 \\ &\Rightarrow e_3 = \frac{1 - 3R_2 + 2R_3}{6}. \end{aligned} \quad (6.16)$$

One can generalize this relation between  $e_k$  and  $R_k$  by defining a matrix,

$$E_k = \begin{pmatrix} 1 & 1 & 0 & \dots \\ R_2 & 1 & 2 & \dots \\ R_3 & R_2 & 1 & \dots \\ \dots & \dots & \dots & \dots \end{pmatrix} \Rightarrow e_k = \frac{\det E_k}{k!}, \quad (6.17)$$

where  $E_k$  is defined as the  $(k \times k)$  matrix of the top-left corner, and we can easily recover the above relations between  $e_k$  and  $R_k$ . As a final equation,

$$\det(xI - \rho_A) = \sum_{k=0}^n \frac{(-1)^k}{k!} \det(E_k) x^{n-k}, \quad (6.18)$$

and the roots of the above polynomials (RHS) are nothing but the Schmidt coefficients. Now, we can see that to obtain all the Schmidt coefficients (in order to either calculate von-Neumann entropy or entanglement spectrum), one needs to have multiple copies of the same state, which becomes complicated in the experiment. However, measurement of  $R_n$  is dominated by  $\lambda_1$  as  $\lambda_1^{2n}[1 + (\lambda_2^2/\lambda_1^2)^n + \dots]$ .

For  $p$  largest eigenvalues ( $\lambda_1^2 > \lambda_2^2 > \dots > \lambda_p^2$ ) need upto  $p$ -copies, and hence  $\mathcal{O}(pN)$  qubits with a parallel circuit depth of  $\mathcal{O}\left[p\left(\frac{\lambda_1^2}{\lambda_p^2}\right)^p\right]$  to resolve the Schmidt values [469].

### 6.2.3 Schmidt values to GGM

From the complication of obtaining all the Schmidt coefficients, it is obvious that von-Neumann entropy needs more resources. Instead, either one can estimate the Rényi entropy, or, due to the ordering of the Schmidt values (coming from roots of the polynomial), the maximum of it can be used to obtain  $1 - \lambda_{\max}^2$ , which becomes the GGM (sec. 2.6.4) on the optimal bipartition. As it requires only two copies of the state, it is more hardware-friendly.

Recall the definition of GGM from Eq. (2.66),

$$G_2(|\psi\rangle) = 1 - \max_{\mathcal{A}:\mathcal{B}}(\lambda_{\mathcal{A}:\mathcal{B}}^{\max})^2. \quad (6.19)$$

Now, the challenge is to maximize over all bipartitions  $\mathcal{A}:\mathcal{B}$ , which is experimentally expensive because:

1. the number of possible bipartitions increases with system size,
2. with increasing size of partition  $\mathcal{A}$ , a larger number of C-SWAPs need to be implemented, which increases depth
3. given the limited connectivity in the devices, the distance between qubits in partitions to the ancilla can increase, and qubit routing becomes necessary, which also affects the quality of the result.

All such problems can be set aside if we consider a fixed partition, where the qubits in the partition are at a minimal distance from the ancilla on the device. Although this does not confirm genuinely multipartite entanglement, as we have not optimized over all possible bipartitions, it still reveals meaningful entanglement characteristics of the system. Moreover, as observed in Fig. 5.6, in quantum optimization algorithms such as QAOA and quantum annealing, the Schmidt values do not differ significantly across different bipartitions; hence, empirically, we may not be far away from the actual GGM<sup>1</sup>.

To be more concrete, instead of optimizing over all partitions, we define on partition  $\mathcal{A}$  as

$$G_{\mathcal{A}} = 1 - (\lambda_{\mathcal{A}:\mathcal{A}}^{\max})^2. \quad (6.20)$$

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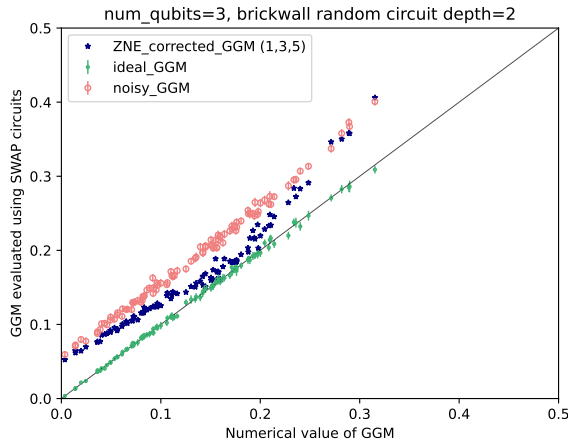
<sup>1</sup>Note, this will be an upper bound on the actual GGM

## 6.3 Benchmarking the method

To obtain the entanglement from the circuit in Fig. 6.2, one has to run the circuit multiple times to get enough count statistics, such that one can trust the probability to obtain a good result, with less error bar. To test if this way of circuit evaluation works, we first compare the  $G_{\mathcal{A}}$  obtained from the circuit with the numerical value obtained by doing singular-value decomposition.

### 6.3.1 Fixed partition

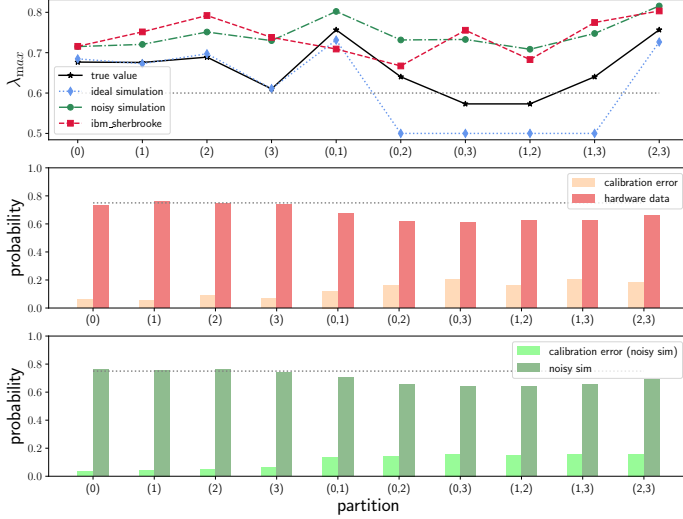
We consider a random brick-wall circuit and generate states with variable depth, obtaining 100 different circuits with distinct GGM. Note that we choose only those circuits where the optimal partition is  $\mathcal{A} = [0]$ , so  $G_{\mathcal{A}} = G_2$ . In Fig. 6.3, we can observe that in an ideal noiseless simulation using 10000 shots, we can obtain the same GGM as numerically obtained, with a small errorbar, which benchmarks our method to be accurate for a wide range of GGM values. However, if we perform a noisy simulation using the noise model of the IBM hardware, we can observe a constant upward shift, which we believe is reminiscent of the noise introduced by the measurement circuit involving the C-SWAP part. We will see later that if one can account for this bias error, one can get closer to the ideal values. We also check if one can obtain a better result using zero-noise extrapolation (see Sec. 3.6.3) on a noisy simulation; however, as we can see, even after 3X and 5X CNOT insertion, the results do not improve sufficiently, compared to the cost of doing it in hardware.



**Figure 6.3.** Benchmarking the circuit method of estimating GGM: From a random circuit of 3 qubits, we generated states with numerical values of GGM spanning from 0 to 0.4. We then estimate the GGM using the circuit by measuring the ancilla 10,000 times to obtain the y-axis values. The noiseless data matches well with the numerical GGM, while the noisy simulation data shows an upward shift. Zero-noise extrapolation does not consistently correct the deviation across all GGM values.

### 6.3.2 All partitions

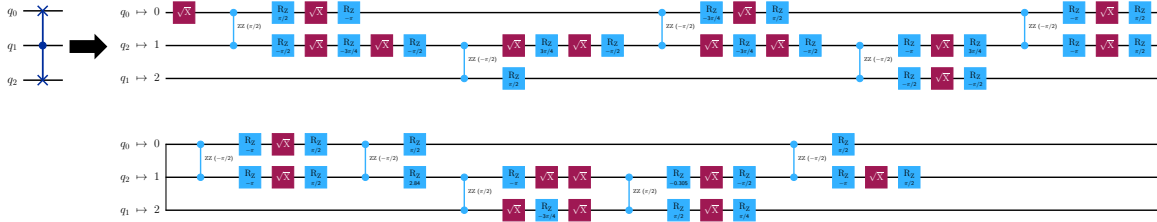
In Section 6.2.3, we explained why working with all partitions is complicated and more errors can be introduced due to their distance from the ancilla. In Fig. 6.4, we show that through experimental data. Comparing with (0), (1), (2), we can see that as the distance increases from the ancilla, which is near (0), the calibration error increases. With a larger number of qubits in the subsystem, the number of C-SWAPs has to increase, and the error also increases, as can be seen by comparing one-qubit partition with two-qubit partitions.



**Figure 6.4.** Performance over all bipartitions on quantum hardware. The protocol was implemented across several partitions (for 4 qubits, this corresponds to 10 partitions), and the results reflect the noise accumulation due to C-SWAPs and limited hardware connectivity, as we can see in the calibration error increases as the partition is far away from (0), and also when more qubits are involved in partition.

### 6.3.3 Noise accumulation from C-SWAP

As indicated by both the noisy simulations and hardware results in Fig. 6.4, the C-SWAP gate can be a significant source of error. This is because, when executed on hardware using its native gate set, it involves several error-prone two-qubit gates, which significantly contribute to the overall noise.



**Figure 6.5.** Transpiled C-SWAP circuit: A simple C-SWAP acting on only one qubit in the partition for each copy becomes a long circuit with multiple  $R_{ZZ}$  gates on the hardware.

### 6.3.4 Can circuit cutting help in measuring entanglement?

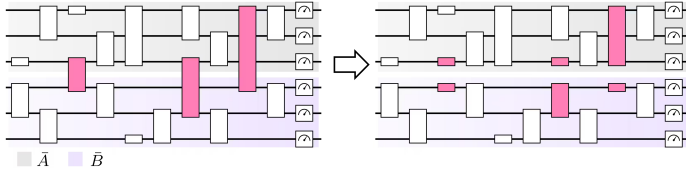
The main challenge in using the SWAP circuit is its increased circuit depth after transpilation. In this scenario, one can consider advanced techniques, such as circuit cutting [285, 474]. It is a technique that was devised to allow longer circuits to run on quantum hardware by cutting into sub-circuits that are suitable for execution on hardware, at the cost of additional sampling overhead [475, 476]. Then, one has to measure part of the circuit multiple times with a sampling overhead and post-process to obtain the target quantum channel [477, 478],

$$U = \sum_i a_i F_i, \quad (6.21)$$

where  $U$  is the target quantum channel, and each  $F_i$  is executable on hardware. The coefficients  $a_i$  come from quasi-probability decomposition (qpd).

Circuit cutting can perform two types of cuts:

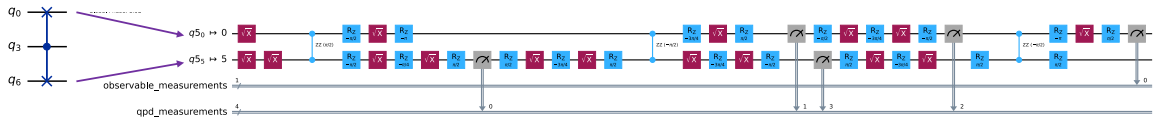




**Figure 6.6.** Example of circuit cutting via a gate cut. The full circuit is divided into two smaller sub-circuits,  $\bar{A}$  and  $\bar{B}$ , each with fewer qubits. The overall output is reconstructed by mid-circuit measurements on each subpart (Adapted from Qiskit Circuit Cutting [479]).

- Gate (space-like) cut: cut through a multi-qubit gate, to make smaller subcircuits with fewer qubits
- Wire (time-like) cut: cut directly through a qubit wire (essentially a single-qubit identity gate that has been cut into two pieces) to fight against coherence time

Cutting comes with sampling overhead, which is exponential in the number of cuts. For example, a CNOT gate cut incurs a sampling overhead of  $9X$ , compared to an original CNOT.

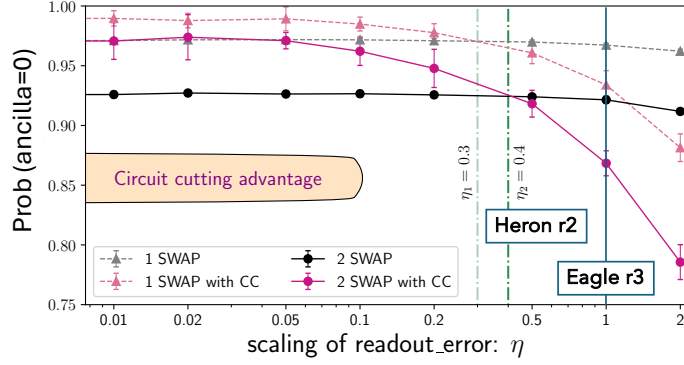


**Figure 6.7.** Example of a cut-circuit for C-SWAP: We can see the number of  $R_{ZZ}$  gates significantly reduces, and mid-circuit measurements of quasi-probability decompositions are introduced.

We use circuit cutting on the C-SWAP gate to obtain the cut circuits in a way that eliminates the need for ancilla, and the circuit depth is significantly reduced (see Fig. 6.7). However, we need to run 8 more such circuits, with their sampling overheads, which makes the method costly. Despite the overhead, we implemented the circuit-cutting approach but found that it offers no clear advantage over standard C-SWAP transpilation. This is primarily because circuit cutting requires repeated measurements of both qubit subsets coming from mid-circuit measurements (see Fig. 6.7), which are highly sensitive to readout errors. To examine this effect, we used a product state in a test setup and scaled the readout error channel by a factor of  $\eta$ . Using noisy simulations, we then extracted the ancilla measurement probabilities (see Fig. 6.8). The results show that while the standard (uncut) method is relatively insensitive to readout errors, the circuit cutting (CC) approach improves only as  $\eta$  decreases, highlighting its strong dependence on readout fidelity. Since current IBM devices typically exhibit readout errors that are 3–4 times higher than the  $R_{ZZ}$  gate error, introducing additional  $R_{ZZ}$  gates proves to be a more practical alternative than relying on readout-intensive techniques like circuit cutting. Therefore, for the remainder of the chapter, we do not employ circuit-cutting techniques and instead rely on the standard transpilation of the C-SWAP gate.

## 6.4 Obtaining entanglement in QAOA from hardware

With these methods in place, we are now equipped to measure the entanglement generated in QAOA and to explore how it is utilized during the algorithm. We consider



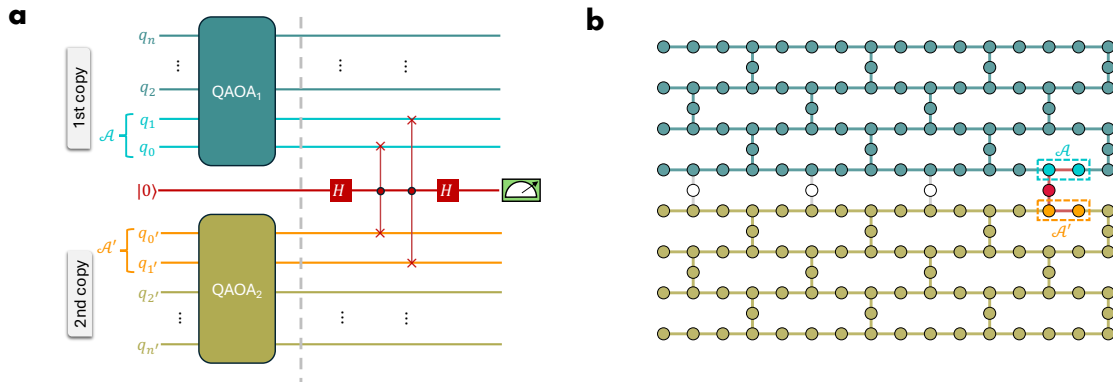
**Figure 6.8.** Readout error-regions of the advantage of circuit cutting. Circuit cutting offers advantages over without CC for 2-CSWAP at  $\eta = 0.4$ , which is just below where our current Heron r2 devices stand, and 1-CSWAP needs even better readout error.

the hardware-native MaxCut problem<sup>2</sup> with randomly chosen coefficients on the edges  $\{-1, +1\}$ , with a small bias field  $Z$  to break the degeneracy on the qubit considered in the bipartition  $\mathcal{A}$  -

$$H_C = fZ_{[\mathcal{A}]} + \sum_{i,j \in \text{coupling map}} J_{ij}Z_iZ_j; \quad J_{ij} \in \{-1, +1\}, \quad f = 0.01. \quad (6.22)$$

### 6.4.1 Choosing qubit arrangement

As the device has limited connectivity, in this case heavy-hex [Fig. 6.9(b)], keeping the ancilla close to the qubits in bipartitions  $\mathcal{A}$  would avoid needing to add multiple SWAPs for connectivity. We choose qubits such that ancilla (red) can be close to and in between  $\mathcal{A}$  (blue) and  $\mathcal{A}'$  (orange) [see Fig. 6.9(b)], which implies in the circuit the 1st copy of the QAOA is flipped [see Fig. 6.9(a)]. Out of multiple possibilities of such subgraphs or substructures in the full heavy-hex map, we choose the best one comparing total gate-fidelity and readout-error of ancilla qubit, as detailed in Sec. 3.6.2.

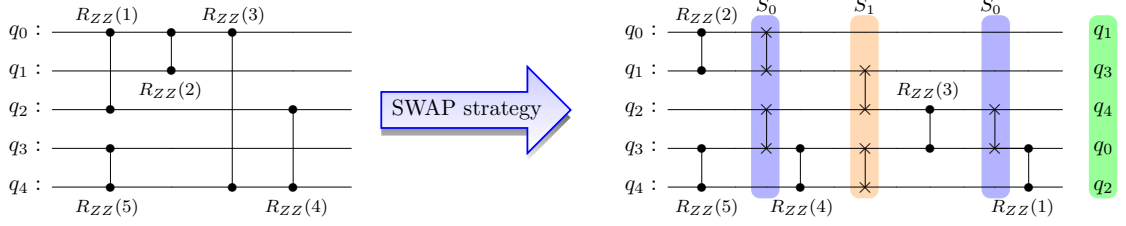


**Figure 6.9.** Qubit positions are chosen based on Sec. 3.6.2 from the possible set of subgraph structures where the ancilla qubit is in between the qubits of  $\mathcal{A}$  and  $\mathcal{A}'$  to keep them closer. (a) In the circuit, the order of the qubits in the 1st copy is flipped, so that  $\mathcal{A}$  becomes closer to the ancilla. (b) One example of chosen qubit positions, with the ancilla being in between  $\mathcal{A}$  and  $\mathcal{A}'$ .

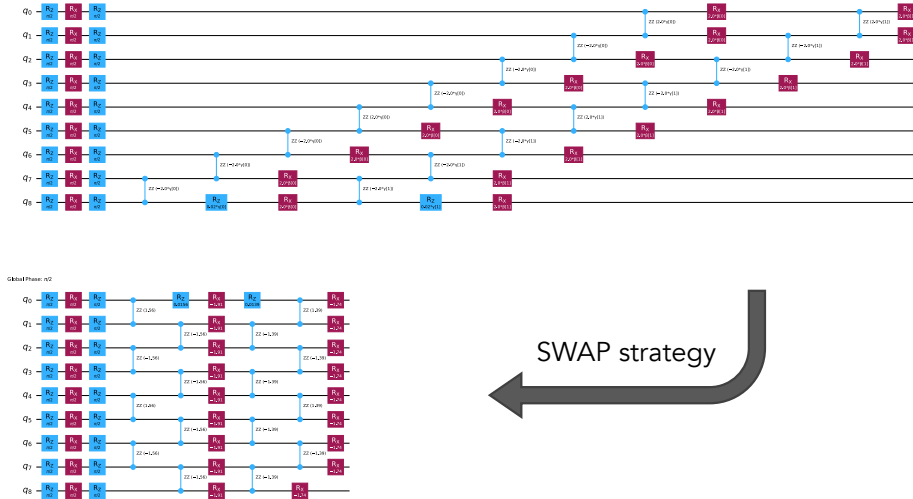
<sup>2</sup>A hardware-native MaxCut problem is a MaxCut instance specifically designed or mapped to align with a quantum device's native qubit connectivity and gate set, minimizing overhead and enabling more efficient quantum execution.

### 6.4.2 Reducing circuit depth using SWAP-strategies

Superconducting qubit devices have limited connectivity and cross-talk, usually resulting in a planar coupling map where two-qubit gates can only be applied between neighboring qubits. To overcome this limitation, additional SWAP gates are inserted for qubit routing, that is, to bring distant qubits next to each other so they can effectively interact. If not transpiled efficiently, the circuit depth and gate count can increase significantly; however, this can be mitigated by using optimized SWAP strategies [290,301]. The basic idea of this method is shown in Fig. 6.10. Fortunately, as we are already considering hardware-native problems, we do not need any extra SWAPs for qubit routing; however, we use a strategy to reorder the gates and reduce the circuit depth, as shown in Fig. 6.11.



**Figure 6.10.** A five-qubit  $\exp(-i\gamma H_C)$  circuit (left) is transpiled to a linear coupling map using the swap sets  $\mathcal{S} = \{S_0, S_1\}$ , where  $S_0 = \{\text{SWAP}_{0,1}, \text{SWAP}_{2,3}\}$  and  $S_1 = \{\text{SWAP}_{1,2}, \text{SWAP}_{3,4}\}$  alternate. In the final transpiled circuit (right), a redundant  $\text{SWAP}_{0,1}$  from the last layer is removed. The final qubit mapping is shown in green. (Taken from Ref. [301])



**Figure 6.11.** A 9-qubit QAOA with depth 2, if transpiled without SWAP strategies has a longer sparse circuit compared to using SWAP strategies, which provides a shorter dense circuit, advantageous for experimental run.

For more such tools for running quantum optimization algorithms on superconducting qubits using Qiskit, please refer to the *q-optimization-best-practices* guide in the Qiskit community [480].

### 6.4.3 Optimization strategies for QAOA

We optimize the QAOA parameters using *qaoa-training-pipelines* [265]. The idea is to grid-search over the full parameter space of  $(\beta_1, \gamma_1)$  for depth-one QAOA, and interpolate the parameter to higher depth as an initial guess. Then, we optimize the parameters using COBYLA [481]. For larger system sizes beyond the exact diagonalization capability, we use Matrix Product State (MPS) [74] with low bond-dimension to evaluate the expectation value of energy.

---

**Algorithm 1** QAOA Training Loop (Pseudocode)

---

```

1: Input: Cost operator: sp_op, max depth:  $p_{\max}$ 
2:
3: Initialize logs  $\leftarrow []$ , opt_params  $\leftarrow []$ , old_energy  $\leftarrow \text{None}$ 
4:
5: for  $p = 1$  to  $p_{\max}$  do
6:   Create run_log with  $p$  and cost_operator
7:
8:   if  $p = 1$  then
9:     Initialize evaluator  $\leftarrow \text{EfficientDepthOneEvaluator}()$ 
10:    scan_trainer  $\leftarrow \text{DepthOneScanTrainer}(\text{evaluator})$ 
11:    scan_result  $\leftarrow \text{scan\_trainer.train}(\text{sp\_op}, \text{num\_points}=100)$ 
12:    opt_params  $\leftarrow \text{scan\_result}[\text{optimized\_params}]$ 
13:    Store scan_result in run_log
14:    scipy_trainer  $\leftarrow \text{ScipyTrainer}(\text{evaluator}, \text{method}=\text{COBYLA})$ 
15:    result  $\leftarrow \text{scipy\_trainer.train}(\text{sp\_op}, \text{opt\_params})$ 
16:   else
17:     evaluator  $\leftarrow \text{MPSEvaluator}(\text{bond\_dim}, \text{use\_vidal\_form}, \text{threshold})$ 
18:     scipy_trainer  $\leftarrow \text{ScipyTrainer}(\text{evaluator}, \text{method}=\text{COBYLA})$ 
19:     recursion_trainer  $\leftarrow \text{RecursionTrainer}(\text{scipy\_trainer}, \text{interpolate})$ 
20:     result  $\leftarrow \text{recursion\_trainer.train}(\text{sp\_op}, \text{opt\_params}, \text{reps}=p)$ 
21:
22:   end if
23:
24:   opt_params  $\leftarrow \text{result}[\text{optimized\_params}]$ 
25:   energy  $\leftarrow \text{result}[\text{energy}]$ 
26:   Store opt_params, energy, and result in run_log
27:
28:   Compute approx_ratio using solve_max_cut(sp_op, energy)
29:   Store approx_ratio in run_log
30:
31:   Append run_log to logs
32:   old_energy  $\leftarrow \text{energy}$ 
33:
34: end for
35:
36: Save logs to disk as qaoa_training_logs_{num_qubits}_p_{p_max}.pkl

```

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## 6.5 Entanglement from IBM device

### 6.5.1 Results for 9 qubits QAOA on a line

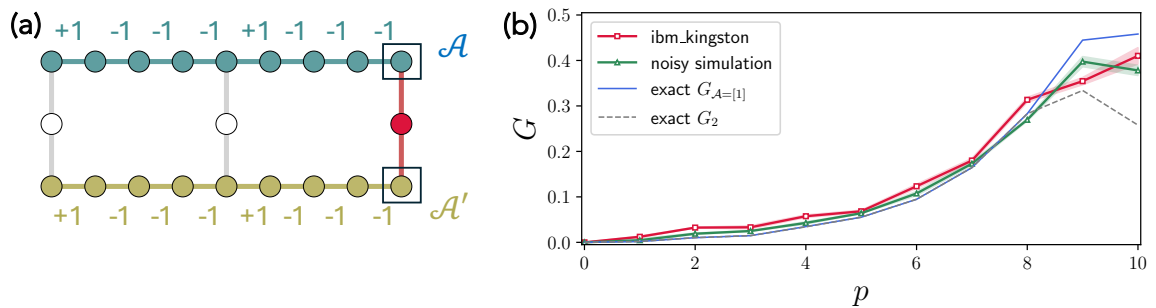
For all the experimental data, we obtain three sets of data, each with 10000 shots, to obtain the mean and error bar on the result. Due to the long C-SWAP measurements, it introduces error, as we have observed in Fig. 6.3. To tackle this issue, we first measure the initial state  $|+\rangle^N$  ( $p = 0$  in figures), which is a product state, and renormalize the probability counts (the probability of the ancilla being  $|0\rangle$ ) for all  $p$  with respect to the  $p = 0$  counts. For example,

$$P_{\text{raw}} = [P_0, P_1, \dots, P_d] \rightarrow P_{\text{renorm}} = [1, \frac{P_1}{P_0}, \dots, \frac{P_d}{P_0}]. \quad (6.23)$$

The motivation behind this renormalization is to ensure that the initial product states are unentangled, as expected at  $P = 1$ , and so any deviation can be attributed to the C-SWAP measurement circuit. More fundamentally, the renormalized probability can be interpreted as a conditional probability to the condition that the circuit produces the correct product state with no entanglement for  $p = 0$  (denoted as  $P_0$ ).

#### For a fixed partition and fixed depth

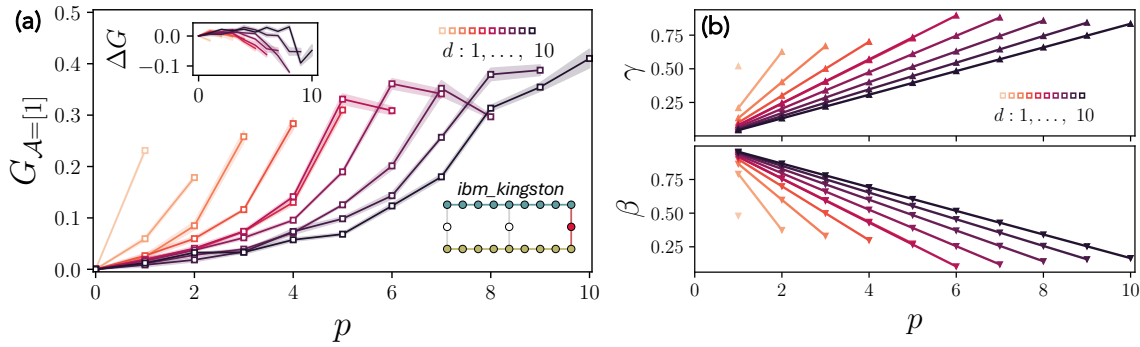
We begin by analyzing smaller systems, such as 9-qubit chains, using a bipartition of the form  $[1] : [2, 3, 4, 5, 6, 7, 8, 9]$ , see Fig. 6.12(a). As shown in Fig. 6.12(b), the entanglement grows steadily with QAOA-layer up to  $p = 10$ . We compute the quantity  $G_{\mathcal{A}=[1]}$  numerically (denoted as exact) and compare it with experimental data from the *ibm kingston* quantum device, as well as with noisy simulations based on its hardware noise model. Up to  $p = 8$ , the experimental results closely follow the exact value, indicating that the device generates entanglement as expected. Beyond that point, deviations appear likely due to limitations in the device's coherence time. Additionally, we observe that the numerically obtained GGM,  $G_2$ , optimized over all bipartitions, remains close in value to  $G_{\mathcal{A}=[1]}$  in our results, suggesting that the choice of bipartition does not significantly affect the outcome in this case. The fact that the difference only emerges toward the end of the circuit suggests that using a fixed bipartition, such as  $[1] : [\text{rest}]$ , already provides sufficient information to benchmark the quantumness of the hardware.



**Figure 6.12.** GGM on 9 qubits QAOA: (a) The chosen qubits are in filled circles, qaoa copy-1 in teal ( $\mathcal{A}$ ) and qaoa copy-2 in olive color ( $\mathcal{A}'$ ), where only one qubit is in the partition. (b) The experimental data and noisy simulation follow the exact value up to the 8th layer, after which entanglement starts decreasing, as expected from the finite coherence time of noisy hardware.

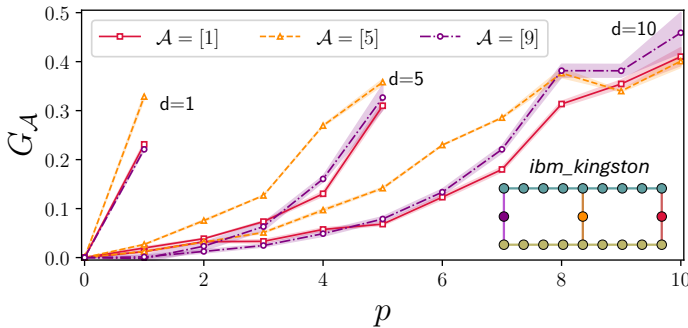
### Dependence on depth

For QAOA, increasing depths typically improves the solution quality [81], captured by the approximation ratio<sup>3</sup>. To understand whether the entanglement also changes depending on the different depths considered, we also measured entanglement at these depths. In Fig. 6.13(a), we can see that with more allowed depth, the final  $G$  can increase to higher entanglement, although the slope of entanglement growth in layers is slower for larger depth. This behavior of the slope can be understood by examining how  $\{\gamma_p\}$ s change with different depths, as it controls the strength of the cost function, i.e., the entangling part of the evolution. We can see in Fig. 6.13(b) that the behavior of  $\gamma$  is similar to the growth of entanglement. However, the entanglement does not follow the noiseless value up to arbitrary layers, and it deviates to a lower entanglement close to larger layers, as expected, see inset of Fig. 6.13(a).



**Figure 6.13.** Entanglement for different depths. (a) With increasing depth  $d$ , the final entanglement reaches higher values, although with slower growth slopes, similar to the optimal  $\gamma_p$  parameters of QAOA, shown in (b). The measured entanglement deviates from the ideal noiseless one at larger depth, as shown in the inset of (a), where  $\Delta G = G[\text{experiment}] - G[\text{ideal}]$ . The numerically obtained approximation ratios for different depths are: 0.729, 0.784, 0.836, 0.867, 0.883, 0.898, 0.908, 0.918, 0.928, 0.934, respectively.

### Dependence on partitions



**Figure 6.14.** Entanglement from  $\mathcal{A} = [1]$  is similar or lower than other chosen partitions for all the layers for depth  $d = 1, 5, 10$ .

<sup>3</sup>The approximation ratio is defined as

$$\text{Approximation Ratio} = \frac{C_{\text{QAOA}} - C_{\min}}{C_{\max} - C_{\min}},$$

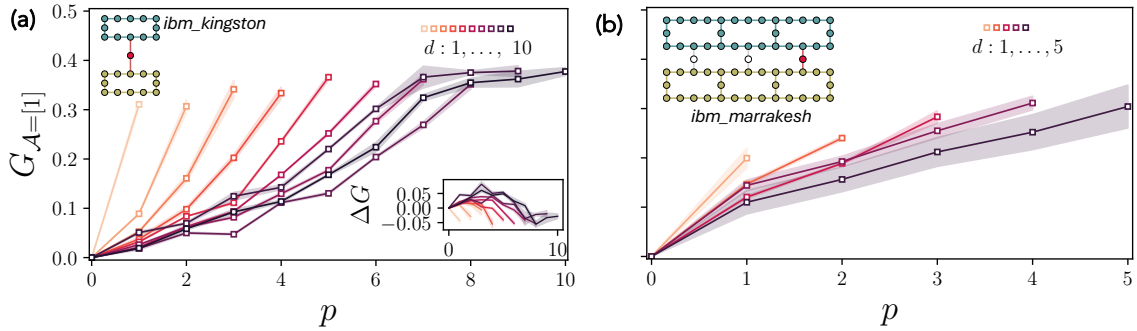
where  $C_{\text{QAOA}} = E + \frac{1}{2} \sum_{(i,j)} w_{ij}$ , with  $E$  representing the QAOA energy and  $w_{ij}$  the edge weights. This ratio measures how close the QAOA solution is to the optimal maximum cut.

We motivated earlier that instead of calculating  $G_{\mathcal{A}}$  for all partitions, choosing one is probably enough. In experiments, we utilize the ancilla between the two copies (see purple, orange, red in Fig. 6.14) without compromising the quality of the result due to their close distance. Though the results for  $\mathcal{A} = [1], [5], [9]$  qualitatively follow similar patterns, there are some quantitative differences, with  $\mathcal{A} = [1]$  exhibiting the lowest or comparable  $G_{\mathcal{A}}$  compared to the other partitions, for all depths considered.

### 6.5.2 Results for 12 and 30 qubit QAOA

As the method works well, we extended it to 12 and 30-qubit QAOA. For the 12-qubit case, even the small depth can reach comparatively stronger entanglement, and when we compare with exact values, the deviation is higher than before at larger depths. We speculate if it can be related to the closed structure (heavy-hex) of the subgraph [Fig. 6.15(a)].

On the other hand, for 30 qubits (3 heavy-hex), the entanglement for smaller depth almost overlaps with the entanglement for the larger depth, i.e., they are of similar slopes [Fig. 6.15(b)]. Note that calculating entanglement in a 30-qubit system is already beyond the scope of a laptop simulation.



**Figure 6.15.** (a) For 12 qubits QAOA, entanglement grows with layer. A smaller depth has a faster slope of growth. Final  $G$  is similar for all depths. The approximation ratios corresponding to depths  $d = 1, 2, 3, 4, 5, 6, 7, 8, 9, 10$  are 0.747, 0.824, 0.879, 0.912, 0.926, 0.927, 0.931, 0.939, 0.958, 0.970, respectively. (b) For 30-qubit QAOA, entanglement growth in smaller depth overlaps with the entanglement growth in larger depth. The approximation ratios corresponding to depths  $d = 1, 2, 3, 4, 5$  are 0.752, 0.832, 0.876, 0.907, 0.925, respectively.

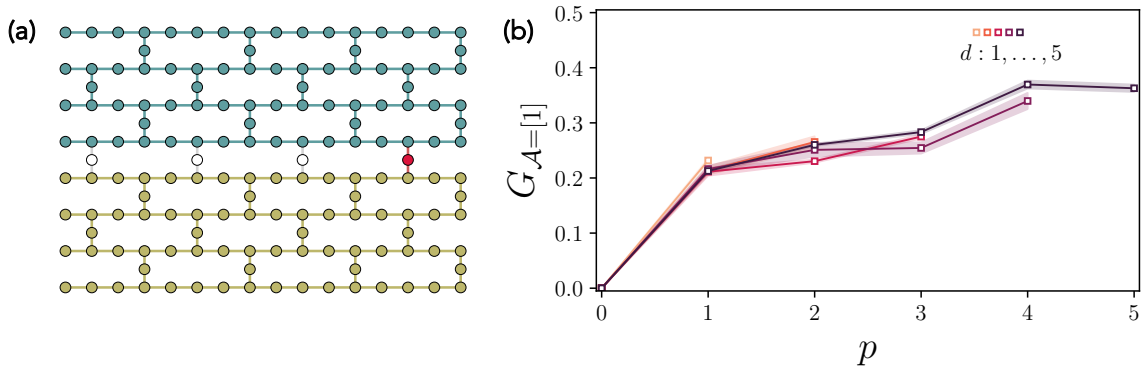
### 6.5.3 Results for 76 qubit QAOA

For the 156-qubit IBM Heron device, we push the limit of measuring the entanglement of a 76-qubit system, which can be nicely fit with its second copy on the chip [Fig. 6.16(a)]. In this case, entanglement grows rapidly during the first layer across all depths  $d$ , followed by a gradual increase up to the final layer [Fig. 6.16(b)]. The similarity in entanglement across different depths resembles the 30-qubit case, though it is even more pronounced here.

## 6.6 Summary and outlook

In this chapter, we addressed a fundamental question in the study of hybrid quantum-classical algorithms: to what extent does the Quantum Approximate Optimization





**Figure 6.16.** Measuring entanglement in 76-qubit QAOA. (a) Chosen coupling map for QAOA-1, ancilla, and QAOA-2 fits to the full device. (b) Entanglement for each depth almost overlaps with the others as it grows up to the final depths. The approximation ratios here are 0.752, 0.836, 0.886, 0.924, 0.947 for  $d = 1, 2, 3, 4, 5$ , respectively.

Algorithm (QAOA) exploit genuinely quantum effects from the quantum processing unit (QPU)? Using a SWAP-circuit-based method, we estimated a geometric measure of entanglement (GGM)-like measure, all within the constraints of current noisy intermediate-scale quantum (NISQ) devices.

We built our approach around a SWAP-test-based method, carefully designed to extract a GGM-like entanglement measure that could be implemented within the tight constraints of NISQ devices. We first tested this method on random quantum circuits, validating our results against exact numerical simulations. We then shifted focus to QAOA, applying the same technique to circuits with up to 76 qubit QAOA (and the whole circuit consists of 153 qubits) on IBM’s Heron device. Across different depths and system sizes, the entanglement we measured closely mirrored what noiseless simulations predicted, for the systems where we could simulate until hardware noise began to dominate at larger depths. This provided direct experimental evidence that QAOA is not simply following classical heuristics, but generating genuine quantum entanglement.

Along the way, we discovered practical insights. A fixed bipartition was often enough to capture the essential structure of entanglement, reducing the overhead of optimization over multiple partitions. We also saw how entanglement growth correlated with the evolution of QAOA parameters and how hardware-aware strategies like optimized qubit mapping, careful transpilation of C-SWAP gates, and calibration of measurement outcomes were vital for success. Techniques like circuit cutting and zero-noise extrapolation offered limited benefit due to dominant readout noise. But perhaps the most compelling outcome was not just that we could measure entanglement, but that we could do so at scales where classical simulation becomes intractable. This opens the door to using entanglement not only for evidence for quantumness but also as a benchmarking tool to guide the design and improvement of future quantum algorithms and hardware.

Looking forward, the methods we have developed can be extended to other variational quantum algorithms [231] or quantum machine learning models [482], helping us understand how and when quantum correlations play a role in performance. The entanglement may become a cornerstone in certifying quantum advantage as a physically meaningful and experimentally accessible way to verify that a quantum processor is doing something no classical computer can.



## Chapter 7

# Role of nonstabilizerness in quantum optimization

*Can quantum optimization outperform stabilizer computations?*

---

## 7.1 Introduction

Combinatorial optimization problems are ubiquitous, spanning from science to industry [232, 483–487]. Hence, achieving any improvement over state-of-the-art classical algorithms through quantum computing could yield a substantial impact, making quantum optimization a leading candidate for practical uses of quantum technologies [248]. In particular, the quantum approximate optimization algorithm (QAOA) [81] offers a promising strategy in the era of noisy quantum devices [87, 301, 488–490]. Despite many successful applications [237, 491–499], it remains unclear to what extent such algorithms truly leverage non-classical quantum resources [47]. This is essential to understand as it defines whether quantum optimization can outperform classical computations or not. In this context, entanglement—a defining feature and a key resource in quantum information processing [89, 500, 501]—has been extensively studied, both in the form of bipartite [414, 417, 435, 437–439] and multipartite entanglement [325, 361, 502, 503]. However, entanglement alone does not ensure a quantum advantage. Stabilizer states can exhibit large entanglement yet remain efficiently simulable classically via Clifford circuits, as guaranteed by the Gottesman–Knill theorem [148–150]. Nonstabilizerness [504]—the degree to which a nonstabilizer state, also called a magic state, deviates from the set of stabilizer states—is thus a crucial property to unlock a potential quantum advantage [160, 504–508]. Based on the discrete Wigner function [509], many measures for this quantum resource have been introduced [181, 510–515]; among which Mana [85, 181] and the Stabilizer Rényi Entropy (SRE) [169, 171, 516] are notable for being efficiently computable and experimentally accessible. In recent years, the study of nonstabilizerness has seen many applications from many-body physics [517–523] over random circuits [172, 524, 525] to conformal field theory [526–528]. Yet, its role in variational quantum optimization remains largely unexplored.

In this chapter, we address this gap by analyzing nonstabilizerness in QAOA applied to combinatorial problems, focusing on both SRE and Mana. We study the performance of QAOA on the paradigmatic Sherrington–Kirkpatrick (SK) model [529], which combines computational hardness with statistical complexity. Our simulations reveal the presence of a “magic barrier”—a transient build-up of nonstabilizerness that occurs during the QAOA run, akin to previously observed entanglement barriers [414, 418, 503]. We find that while QAOA begins and ends with low-magic (sta-

bilizer) states, it must pass through a regime of increased magic in order to achieve high fidelity with the target solution. This behavior appears in both qubit and qutrit versions of the algorithm. We find that magic curves corresponding to different depths collapse under a simple rescaling, and we identify a similar magic barrier in quantum annealing. Furthermore, we analyze how the success probability of QAOA correlates with the final nonstabilizerness, uncovering characteristic trends and structure in the fidelity-magic plane. Using analytical calculations on *ansatz* states, made of few-component superpositions, we explain the observed features and clarify under what conditions low-magic solutions may or may not correspond to high-fidelity outcomes.

Through this chapter, we illuminate the role and significance of nonstabilizerness in quantum optimization, providing deeper insight into its influence on algorithmic performance and complementing existing studies on entanglement and coherence [418, 439, 503]. The results also have practical relevance, as measurements of magic can provide a figure of merit for estimating the performance of variational or optimization algorithms performed on concrete quantum hardware, such as superconducting qubits [530], trapped ions [173], and Rydberg atoms [531].

## 7.2 Quantifying nonstabilizerness for qudits

Here we recall two measures of nonstabilizerness from Sec. 2.7.4: the stabilizer Rényi entropy (SRE) [169, 171] and Mana [85, 181].

### 7.2.1 Stabilizer Rényi Entropy

The SRE [169] characterizes how a pure state  $|\psi\rangle$  of  $N$  qudits spreads over the basis of Pauli strings  $\mathcal{P}_N$ :

$$M_n(|\psi\rangle) = \frac{1}{1-n} \log_2 \left[ \sum_{P \in \mathcal{P}_N} \frac{|\langle \psi | P | \psi \rangle|^{2n}}{D^N} \right]. \quad (7.1)$$

$M_n$  is non-negative and equals zero if and only if  $|\psi\rangle$  is a stabilizer state [171, 532]. An important feature of Eq. (7.1) is that it is experimentally computable [173]. The SRE defined in Eq. (7.1) satisfies all properties (i-iii) necessary to be a valid magic measure [169]. For  $d > 2$ , the Pauli operators are no longer Hermitian, and thus their expectation values can be complex. For this reason, the absolute values are taken in Eq. (7.1). Throughout this chapter, we use  $M_2$  as our measure.

### 7.2.2 Mana

Defining Mana requires knowledge of two other concepts: Heisenberg–Weyl operators, and discrete Wigner functions, which we have discussed in Sec. 2.7.3. Then, the negativity of the discrete Wigner function provides another quantifier of nonstabilizerness [160, 533]. Specifically, the log-negativity of the Wigner function  $W_\rho$  associated with a quantum state  $\rho$  defines a quantity known as Mana [85]:

$$\text{Mana} = \log \sum_{\vec{v}} |W_\rho(\vec{v})|. \quad (7.2)$$

For pure states  $\rho = |\psi\rangle\langle\psi|$ , as considered here, the set of states with non-negative Wigner representation coincides precisely with the set of pure stabilizer states [160], in which case the Mana vanishes.

## 7.3 QAOA on SK model

As illustrated in Fig. 7.2(a), QAOA prepares a trial state  $|\psi(\beta, \gamma)\rangle$  by applying  $d$  layers of alternating unitaries,  $\exp(-i\beta_p \hat{H}_M) \exp(-i\gamma_p \hat{H}_C)$ , starting from the ground state of the mixer Hamiltonian  $\hat{H}_M$  [534], see Sec. 3.3.4 for more details. Originally developed for qubits, the Quantum Approximate Optimization Algorithm (QAOA) can be generalized to qudit systems with any local dimension  $D$  [497], thereby expanding its range of potential applications [497, 499, 535, 536]. As the initial state is non-magical<sup>1</sup> and the solution state of a classical problem with a unique optimum is also non-magical, QAOA offers a natural setting to test the buildup and role of magic during the quantum optimization procedure.

The model of our interest is the Sherrington–Kirkpatrick (SK) model, a variant of the Ising model with all-to-all interactions randomly drawn from independent Gaussian distributions. Initially introduced as a solvable mean-field model to describe the spin-glass phase [529, 537], it serves as a prototype for real-world optimization problems, which often involve a diverse set of random variables [538, 539]. We use the SK model generalized to qudits (specifically, we study qubits and qutrits),

$$\hat{H}_{\text{SK}} = \sum_{i \neq j} J_{ij} (Z_i Z_j^\dagger + Z_i^\dagger Z_j) + \hat{H}_{\text{bias}}, \quad (7.3)$$

where the coefficients  $J_{ij}$  are chosen randomly from a normal distribution with unit variance, and  $Z_i$  is the generalized phase operator. The Hermitian conjugate term is necessary to ensure the Hermiticity of the Hamiltonian in the qutrit case. Since we aim to understand the magic generated solely through QAOA, it is crucial to remove any degeneracy, as it may contribute additional magic arising from a superposition of equally valid solution states rather than the algorithm itself. For that purpose, we add bias magnetic field terms  $\hat{H}_{\text{bias}}$ .

### 7.3.1 Degeneracy breaking in SK model

The Hamiltonian term without bias  $\sum_{i \neq j} J_{ij} (Z_i Z_j^\dagger + Z_i^\dagger Z_j)$  has symmetry under reflection,  $Z_i \leftrightarrow -Z_i$ , thus creating two degenerate states. The reflection degeneracy can be lifted by adding a longitudinal field,  $\hat{H}_{\text{bias},2} = \sum_i h_i (Z_i + Z_i^\dagger)$ , which is reduced to  $\sum_i 2h_i Z_i$  for the qubit case.

However, for qutrits, this term is furthermore symmetric under  $Z_i \rightarrow cZ_i$ , where  $c \in \{\pm 1, \pm\omega, \pm\omega^2\}$ , which introduces 4 additional symmetries. For example,  $Z_i Z_j^\dagger \xrightarrow{+\omega} (\omega Z_i)(\omega Z_j)^\dagger = (\omega Z_i)(\omega^2 Z_j^\dagger) = Z_i Z_j^\dagger$ . Thus, the qutrit Hamiltonian contains 6 degenerate eigenstates, equivalent to permutation among the local basis of qutrits,

<sup>1</sup>The  $|+\rangle = \frac{1}{\sqrt{D}} \sum_{i=0}^{D-1} |i\rangle$  qudit state is a stabilizer state because it is a  $(+1)$  eigenstate of a  $d$ -element subgroup of the Pauli group; for example, it is  $\{I, X\}$  for  $d = 2$ . Using the additive property,  $\mathcal{M}(|+\rangle^{\otimes N}) = N\mathcal{M}(|+\rangle) = 0$ .

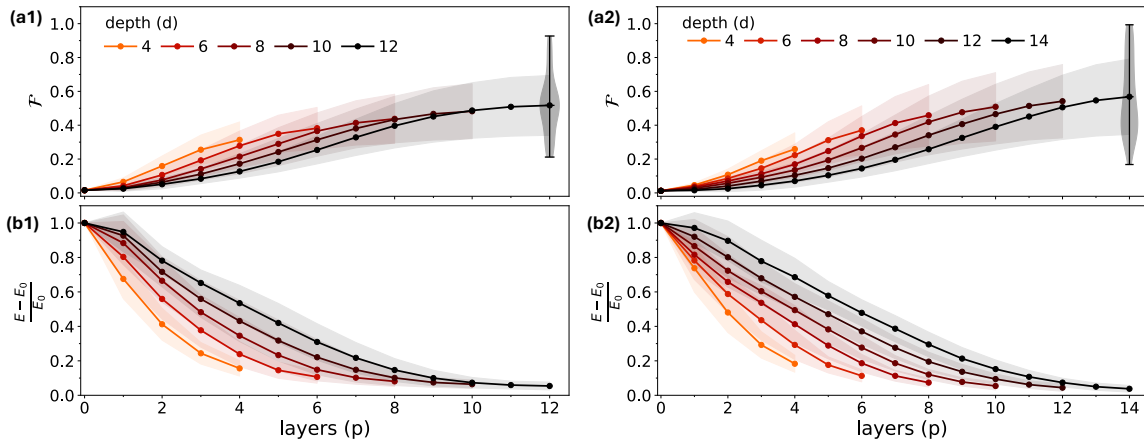
which contains  $3! = 6$  degenerate states. The longitudinal field  $\hat{H}_{\text{bias},2}$  removes the degeneracy between  $|0\rangle$  and  $\{|1\rangle, |2\rangle\}$  but retains the degeneracy between  $|1\rangle$  and  $|2\rangle$ , as  $(Z + Z^\dagger) = \text{diag}(2, -1, -1)$ . To lift this degeneracy, we add another term  $\sum_i i h'_i (Z_i - Z_i^\dagger)$ , as  $i(Z - Z^\dagger) = \text{diag}(0, -\sqrt{3}, +\sqrt{3})$ . In our numerics,  $h_i$  and  $h'_i$  are chosen randomly from a normal distribution with variance 0.3.

### 7.3.2 Optimization procedure

We execute all QAOA simulations using a custom implementation. To find the optimal parameters  $\{\beta, \gamma\}$ , we employ the Constrained Optimization BY Linear Approximation (COBYLA) algorithm [481], a gradient-free method that solves the problem by linearly approximating the cost function. The parameters are initialized using the Trotterized Quantum Annealing (TQA) strategy [256]. The classical optimizer stops when the trust-region radius decreases below the specified tolerance of  $10^{-4}$ . For each random realization of the SK model, we simulate a QAOA protocol using exact numerics, where we select the best outcome from 20 independent QAOA runs initialized using Trotterized quantum annealing initialization [256], and average the results over 50 such realizations. The SRE is computed exactly, with computational cost scaling as  $D^{2N}$  due to the exponential growth in the number of Pauli strings, where  $D$  is the qudit dimension and  $N$  is the number of qudits.

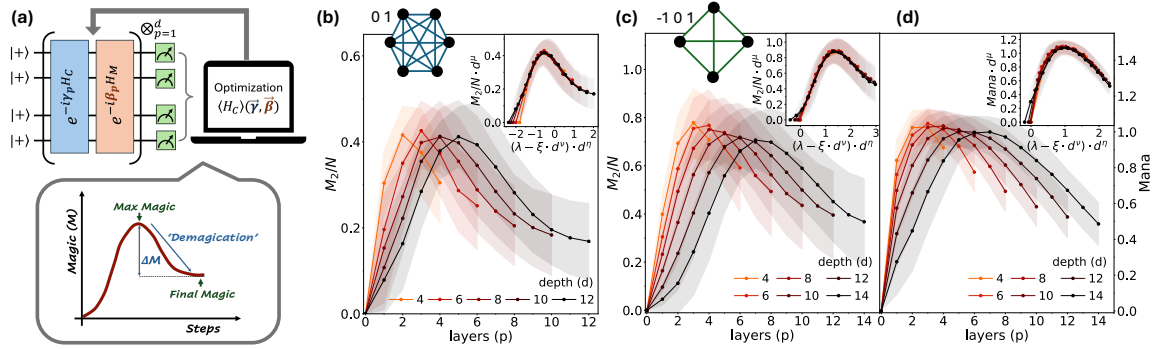
## 7.4 Magic barrier

When targeting the non-degenerate ground state of the SK model, QAOA starts—and ideally ends—in stabilizer states with zero magic. The relevant performance metrics—such as the relative energy  $(E_{\text{QAOA}}(p) - E_{\text{exact}})/E_{\text{exact}}$  (in Fig. 7.1 (b1,b2)) reflects that the QAOA energy reaches values close to the ground state, and the fidelity with the ground state  $\mathcal{F}(p) = |\langle \psi_{\text{exact}} | \psi_{\text{QAOA}}(p) \rangle|^2$  (in Fig. 7.1 (a1,a2)) improve monotonically with the number of QAOA layers  $p$ , though being far from equal to one.



**Figure 7.1.** (a1) Mean value of the fidelity over 50 different realizations, with relative standard deviation, for each layer at different depths of the 6-qubit system. The violin plot for the curve at depth 12, using the full 12-layer wave function, highlights the high fidelity achieved by some realizations. (b1) Mean value of the relative energy under the same conditions. (a2) and (b2) show the corresponding results for a 4-qutrit system.

In contrast, the evolution of magic, quantified by SRE and Mana, follows a non-monotonic behavior: it initially rises rapidly as the first QAOA layers are applied, reaches a maximum, and then decreases as the algorithm approaches the final state. We define this structure as the *magic barrier*, marking the necessity of traversing highly nonstabilizer states during the QAOA algorithm. Therefore, the quantum computational resources required during the intermediate steps of QAOA significantly exceed what would be expected from monotonic growth. This additional “cost” in non-stabilizerness emerges inherently from the QAOA ansatz structure; thus, it becomes crucial to account for this elevated resource demand in the design and benchmarking of fault-tolerant quantum devices.



**Figure 7.2. Magic barrier in QAOA.** (a) Quantum Approximate Optimization Algorithm (QAOA) scheme and a pictorial representation of the magic barrier: initially, magic rises up to a maximal value, after which it decreases, an effect we call “demagication.” (b) SRE density as a function of the layer number during a QAOA protocol, for a system of 6 qubits and different depths (i.e., total number of layers). Inset: Using a simple scaling function, the magic barrier collapses onto a universal curve. (c,d) Same plot of panel b for a system of 4 qutrits, showing SRE density (c) and Mana (d) as a function of the layer number. The universal collapses in the insets in (b-d) use the scaling function  $\mathcal{M} = d^{-\mu} f[(\lambda - \xi \cdot d^{\nu}) \cdot d^{\eta}]$ .

A pictorial representation of the magic barrier is shown in Fig. 7.2(a). Remarkably, this phenomenon is observed consistently across different systems and for both SRE and Mana. For instance, Fig. 7.2(b) depicts the evolution of the SRE density ( $M_2/N$ ) for a 6-qubit system at varying QAOA depths. A distinct peak emerges at approximately half the total depth, indicating the point of maximal nonstabilizerness. A comparable scenario arises in the 4-qutrit system, for both SRE and mana [see Fig. 7.2(c,d)].

Notably, for a fixed system size, the maximum value of magic encountered remains approximately constant across different QAOA circuit depths, suggesting that the magic peak is independent of circuit depth. Moreover, even at its peak, the generated magic remains below that of a typical state in the corresponding Hilbert space, such as given by the Haar-random value—  $M_2^{\text{Haar}} = -\log_2(\frac{4}{2^N+3})$  for qubit, and  $-\log_2(\frac{3}{3^N+2})$  for qutrit systems [168, 524]. This suggests that for a device to successfully run a QAOA, it is sufficient if it can build up a limited amount of magic.

### 7.4.1 Scaling of nonstabilizerness

The states generated by QAOA strongly depend on the circuit depth, optimization parameters, and system size, and yet we observe a qualitatively uniform behavior of

the magic barrier in Fig. 7.2. In particular, for a fixed  $N$ , while the magic curve varies with the QAOA depth, certain patterns emerge consistently: (1) the barrier height remains nearly unchanged, and (2) as the depth increases, the curve appears to stretch horizontally. These observations motivate the search for a scaling collapse of the magic barrier, which could offer valuable insights into the underlying behavior.

We begin by seeking a simple scaling function  $f[\cdot]$  for fixed  $N$ ,

$$\frac{\mathcal{M}(|\psi_{\text{QAOA}_d}(p)\rangle)}{N} = d^{-\mu} f[(\lambda - \xi \cdot d^\nu) \cdot d^\eta]; \mu, \xi, \nu, \eta \in \mathbb{R}; \quad \lambda = \frac{p}{d}. \quad (7.4)$$

where  $\mu$  and  $\eta$  control the stretching along the  $y$ - and  $x$ -axes, respectively. The curves only collapse onto each other when centered around a depth ( $d$ )-dependent parameter, which we achieve through the introduction of  $\xi$  and  $\nu$ . We use these parameters to rescale the data for different depths and observe whether the curves align. The collected scaling parameters for different *qubit* system sizes are reported in Table 7.1, which includes the values of  $\mu$ ,  $\xi$ ,  $\nu$ , and  $\eta$  obtained from numerical fits for  $N = 4, 6, 8$  qubit systems. Similarly, Table 7.2 shows scaling parameters for *qutrits*. Indeed, shown in the insets of Fig. 7.2(b–d), the data for different depths, both for average SRE density and Mana, collapse onto a single curve.

$N$	$\mu$	$\xi$	$\nu$	$\eta$
4	−0.07	0.02	0.76	0.23
6	0.01	1.28	−0.34	0.60
8	−0.07	0.02	0.95	0.86

**Table 7.1.** Scaling parameters  $\mu$ ,  $\xi$ ,  $\nu$ , and  $\eta$  used for data collapse at fixed system size  $N$  (qubits), using Eq. (7.4). Values obtained from fitting QAOA magic curves for  $N = 4, 6, 8$ .

$N = 4$	$\mu$	$\xi$	$\nu$	$\eta$
SRE	−0.08	0.00	3.49	0.46
Mana	0.03	0.00	5.50	0.34

**Table 7.2.** Scaling parameters  $\mu$ ,  $\xi$ ,  $\nu$ , and  $\eta$  used for data collapse at  $N = 4$  *qutrits* using Eq. (7.4). Values are shown for both SRE and mana.

For qubit systems, we find a scaling exponent  $\mu$  close to zero, suggesting that the peak value of SRE remains independent of both the QAOA circuit depth and the number of qubits. Interestingly, for the qubit cases, we find that the scaling formulation can be simplified by replacing the combined term  $\xi \cdot d^\nu$  with a constant critical point  $\lambda_c$ , leading to the alternative scaling form

$$\frac{\mathcal{M}(|\psi_{\text{QAOA}_d}(p)\rangle)}{N} = d^{-\mu} \tilde{f}[(\lambda - \lambda_c) \cdot d^\eta]; \mu, \lambda_c, \eta \in \mathbb{R}; \quad \lambda = \frac{p}{d}. \quad (7.5)$$

This revised form uses the scaling variable  $\lambda = p/d$  and centers the collapse around a critical point  $\lambda_c$ . In systems exhibiting universality—such as those undergoing phase transitions—the central parameter (around which the scaling occurs) becomes a constant, signaling a universal or critical point, similar to  $\lambda_c$ . The goal is then to determine whether such a  $\lambda_c$  can be found consistently across system sizes, as a signature of universal behavior. Table 7.3 reports the corresponding values of  $\mu$ ,  $\lambda_c$ , and  $\eta$  obtained from scaling collapses performed using Eq. (7.5) for  $N = 4, 6, 8$  qubit systems. Remarkably, the extracted values of  $\lambda_c$  are rather consistent around 0.2 across system sizes, suggesting the existence of a constant point in the qubit case, even at small system sizes. This behavior is in contrast to the qutrit case, where no such consistent value of  $\lambda_c$  is observed. The exponent  $\eta$ , which governs the width rescaling, lies



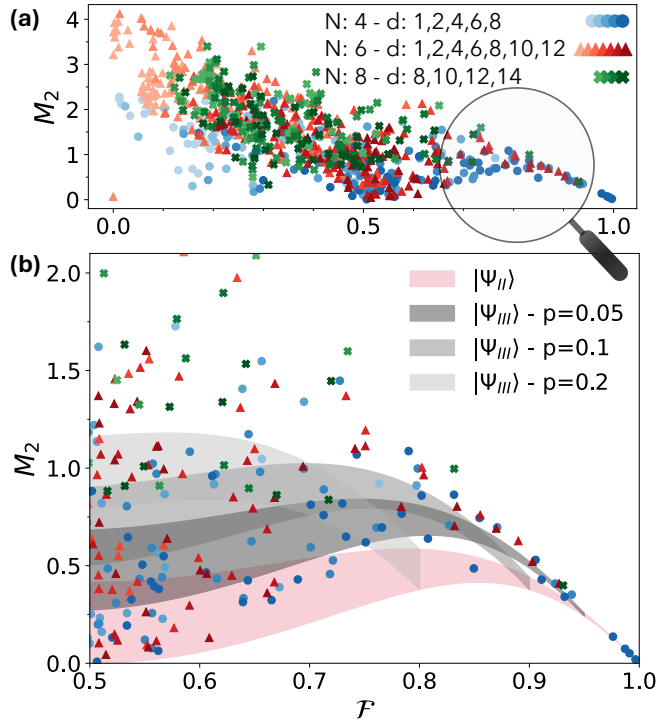
within a relatively narrow range for both the 6-qubit and 4-qutrit systems. Moreover,  $\eta$  increases with the number of qubits, suggesting a sharper magic barrier in larger systems.

$N$	$\mu$	$\lambda_c$	$\eta$
4	-0.07	0.27	0.23
6	0.01	0.27	0.59
8	-0.10	0.29	0.92

**Table 7.3.** Fitted scaling parameters  $\mu$ ,  $\lambda_c$ , and  $\eta$  using the critical-point form in Eq. (7.5). Results shown for  $N = 4, 6, 8$  qubits.

## 7.5 Final magic and fidelity

The ultimate determinant of the success of QAOA lies in the fidelity of the final state with the target solution. As the QAOA approaches the final layer, successful instances reach a high overlap with a classical state, implying—in the absence of degenerate solutions—one can associate high-fidelity QAOA instances with low magic. This behavior is indeed what we observe in randomly sampled instances for different system sizes and depths, shown in Fig. 7.3. The opposite, however, is not necessarily true: a QAOA sweep can end up in a stabilizer state, including any computational basis state distinct from the solution state, and thus can have small magic while reaching only low fidelity.



**Figure 7.3. Relation between final magic and fidelity (qubits).** (a) Magic of the final state as a function of final fidelity  $\mathcal{F}$  for randomly sampled instances with varying numbers of qubits and QAOA depths. (b) Zoom-in on the region  $\mathcal{F} \geq 0.5$ . The red shaded area marks the upper limit of SRE achievable by superpositions of two computational basis states. The grey regions correspond to the SRE of wavefunctions spanning three basis states with varying amplitudes. While higher regions could be reachable with more complex *ansatz* states, the empty area below is forbidden.

Focusing on instances with  $\mathcal{F} \geq 0.5$  [Fig. 7.3(b)], one observes a void region at the lower axis. The pattern can be explained as follows. At the end of a rather successful QAOA, we expect the state to consist of the final solution  $|\phi_0\rangle$  with potentially a small contamination, typically from the first excited state, which is another computational basis states  $|\phi_1\rangle$  or/and  $|\phi_2\rangle$ . In those cases, we can obtain an analytical expression relating magic and final-fidelity as shown in the following sections.

## 7.5.1 For superposition of two states (qubit)

Here, we consider an *ansatz* state

$$|\psi_{\text{II}}\rangle = \sqrt{\mathcal{F}} |\phi_0\rangle + e^{i\theta} \sqrt{1 - \mathcal{F}} |\phi_1\rangle, \quad \theta \in [0, 2\pi], \quad (7.6)$$

where the probability amplitude to reach the desired solution is  $\sqrt{\mathcal{F}}$ , and where we allowed for a relative phase  $e^{i\theta}$ . Using the additivity of  $M_2$  and permutational invariance, one can show that for any number of qubits:

$$M_2(|\psi_{\text{II}}\rangle) = -\log_2 [1 - 4\mathcal{F}(1 - \mathcal{F}) + 2\mathcal{F}^2(1 - \mathcal{F})^2(7 + \cos 4\theta)]. \quad (7.7)$$

The result is symmetric around  $\mathcal{F} = 0.5$ . The minima (maxima) at  $\theta = 0$  ( $\theta = \pi/4$ ) describe a bounded region in the fidelity–magic plane, within which all instances that reach the above *ansatz* are confined, independent of peculiarities of the protocol such as qubit number or circuit depth.

**Proof.** Consider a system composed of  $n$  qubits. Since the states  $|\phi_0\rangle$  and  $|\phi_1\rangle$  are in the computational basis, we can define Hamming weight distance, let's say  $h$ . Assuming  $h < n$ , we can write the state  $|\psi_{\text{II}}\rangle$  as  $|\psi'_{\text{II}}\rangle_h \otimes |d\rangle_{n-h}$ . Here,  $|d\rangle = |0\rangle^{\otimes(n-h)}$  or  $|1\rangle^{\otimes(n-h)}$ . By the additive property of SRE,  $M_n(|\psi_{\text{II}}\rangle) = M_n(|\psi'_{\text{II}}\rangle_h) + M_n(|d\rangle_{n-h})$ .  $M_n(|d\rangle_{n-h}) = 0$ , so only  $M_n(|\psi'_{\text{II}}\rangle_h)$  contributes to SRE. We can write  $|\psi'_{\text{II}}\rangle_h$  as  $\sqrt{\mathcal{F}} |a\rangle_n + e^{i\theta} \sqrt{1 - \mathcal{F}} |\bar{a}\rangle_n$ . This is SRE-equivalent to  $\sqrt{\mathcal{F}} |0\rangle^{\otimes n} + e^{i\theta} \sqrt{1 - \mathcal{F}} |1\rangle^{\otimes n}$  via bit-flip operations  $X_{\vec{j}}$  at the appropriate positions ( $\vec{j}$ ). This is equivalent to using  $P'_i = X_{\vec{j}} P_i X_{\vec{j}}$  on the earlier states. As the Pauli group is invariant under such an operation, we can compute the magic for the latter state without any loss of generality. The formula is  $n$ -independent because what counts is the number of states in superposition, not the dimension of the system or which two basis states constitute the state. We can write all four terms appearing in the expectation values as,

$$\begin{aligned} & \mathcal{F} \cdot \langle 0|^{\otimes n} P_i |0\rangle^{\otimes n}, \quad (1 - \mathcal{F}) \cdot \langle 1|^{\otimes n} P_i |1\rangle^{\otimes n}, \\ & \sqrt{\mathcal{F}(1 - \mathcal{F})} e^{i\theta} \cdot \langle 0|^{\otimes n} P_i |1\rangle^{\otimes n}, \quad \sqrt{\mathcal{F}(1 - \mathcal{F})} e^{-i\theta} \cdot \langle 1|^{\otimes n} P_i |0\rangle^{\otimes n}, \quad \forall i. \end{aligned} \quad (7.8)$$

The first two terms are non-zero only when the involved string  $P_i$  is built only from combinations of  $\{I, Z\}$  Pauli operators. In that case, the last two terms are zero. When  $P_i$  contains a number  $k$  of  $Z$  operators, there are  $\binom{n}{k}$  terms contributing to the sum. These will cumulatively contribute to

$$\sum_{k=0}^n \binom{n}{k} \left| \mathcal{F} + (-1)^k (1 - \mathcal{F}) \right|^4 = 2^{n-1} [1 + (2\mathcal{F} - 1)^4]. \quad (7.9)$$

Similarly, the last two terms are non-zero when the  $P_i = Y^{\otimes k} X^{\otimes n-k}$ . Thus,

$$\sum_{k=0}^n \binom{n}{k} \mathcal{F}^2 (1 - \mathcal{F})^2 |e^{i\theta} (-i)^k + e^{-i\theta} (+i)^k|^4 = 2^{n-1} \mathcal{F}^2 (1 - \mathcal{F})^2 2^4 [\cos^4 \theta + \sin^4 \theta]. \quad (7.10)$$

By summing these two contributions and performing some simple algebra, one obtains the final equation of Eq. (7.7).



### 7.5.2 For superposition of three states (qubit)

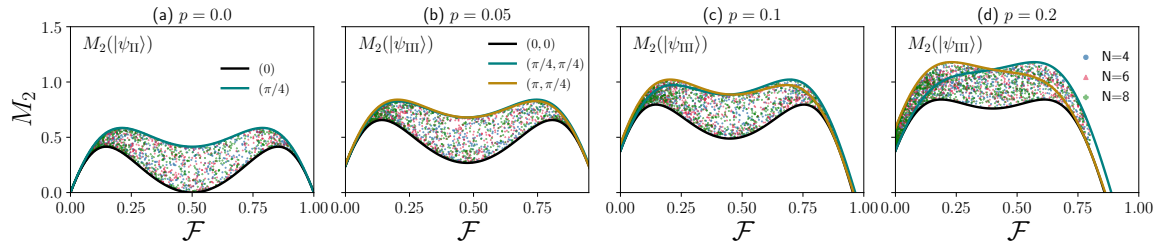
Motivated by the  $n$  independence of the previous result, we further refine this *ansatz* by considering a superposition of three computational basis states,

$$|\psi_{\text{III}}\rangle = \sqrt{\mathcal{F}} |\phi_0\rangle + e^{i\theta_1} \sqrt{p} |\phi_1\rangle + e^{i\theta_2} \sqrt{1 - \mathcal{F} - p} |\phi_2\rangle, \quad p \in [0, 1 - \mathcal{F}], \quad \theta_{1/2} \in [0, 2\pi], \quad (7.11)$$

for which we can again calculate  $M_2$  analytically. For different system sizes, we find an expression consistent for different  $N$  with the following:

$$\begin{aligned} M_2 = -\log_2 \Big[ & 1 + 14f^4 + 28f^3(p-1) + 2(p-1)p(2 + 7p(p-1)) \\ & + 6f^2(3 + p(7p-6)) + 4f(p-1)(1 + p(7p-6)) \\ & + 2f^2(f+p-1)^2 \cos(4\phi) + 4p^2 f^2 \cos(2\phi) \cos(2\phi - 4\theta) \\ & + 2p^2(p-1)(2f+p-1) \cos(4\phi - 4\theta) \Big]. \end{aligned} \quad (7.12)$$

Similar to the case of magic computed for a superposition of two computational basis states, the analytical expression derived here remains valid for states of arbitrary dimension. The generalization of the Eq. (7.7) and (7.12) to any dimension systems is shown in Fig. 7.4, where random benchmarks are used for this purpose.



**Figure 7.4.** SRE vs. fidelity for a superposition of two (a) and three (b,c,d) states at different values of  $p$ . Continuous lines represent analytic bounds given by Eq. (7.7) and (7.12), respectively. Dots represent numerically computed SRE for random states of the form  $|\psi_{\text{II}}\rangle$  [Eq. (7.7) for (a)] and  $|\psi_{\text{III}}\rangle$  [Eq. (7.12) for (b,c,d)]. The points are clearly enclosed within the bounds given by the continuous lines, certifying the validity of Eq. (7.7) and Eq. (7.12).

### 7.5.3 Relation to the QAOA fidelity

The shaded regions in Fig. 7.3(b) correspond to  $M_2$  contained in the *ansatz* states  $|\psi_{\text{II,III}}\rangle$  for varying  $\mathcal{F}, \theta, p$ , as detailed in Eq. (7.7) and Eq. (7.12). Since *ansätze* with a further increased number of basis states will only lead to higher magic, these analytic considerations also explain why no instances are found numerically in the empty region at the bottom of the figure. Moreover, they suggest a reduced likelihood of obtaining a medium-to-high-fidelity state  $\mathcal{F} \sim (0.6 - 0.9)$  with low magic, further indicating that QAOA has to go through a high-magic state to reach a good solution.

### 7.5.4 For superposition of qutrit states

Motivated by the qubit superposition, we also calculated how SRE behaves for the superposition of two qutrit states. The pauli group for qutrits is defined as  $P =$

$\{X^r Z^j | r, j \in (0, 1, 2)\}$ . So, for a qutrit state  $|\psi\rangle = \sqrt{\mathcal{F}}|a\rangle + e^{i\theta}\sqrt{1-\mathcal{F}}|b\rangle$ , we get

$$\langle\psi|P_i|\psi\rangle = \mathcal{F}\langle a|P_i|a\rangle + (1-\mathcal{F})\langle b|P_i|b\rangle \quad (7.13)$$

$$+ e^{i\theta}\sqrt{\mathcal{F}(1-\mathcal{F})}\langle a|P_i|b\rangle + e^{-i\theta}\sqrt{\mathcal{F}(1-\mathcal{F})}\langle b|P_i|a\rangle. \quad (7.14)$$

The first two terms will be different from zero when  $a = 0$ , while the second two terms will be different from zero when  $a \neq 0$ . Thus, we have

$$e^{-M_2} = \sum_{j=0,1,2} |\mathcal{F}\langle a|Z^j|a\rangle + (1-\mathcal{F})\langle b|Z^j|b\rangle|^4 \quad (7.15)$$

$$+ \sum_{r=1,2;j=0,1,2} |e^{i\theta}\sqrt{\mathcal{F}(1-\mathcal{F})}\langle a|X^r Z^j|b\rangle + e^{-i\theta}\sqrt{\mathcal{F}(1-\mathcal{F})}\langle b|X^r Z^j|a\rangle|^4$$

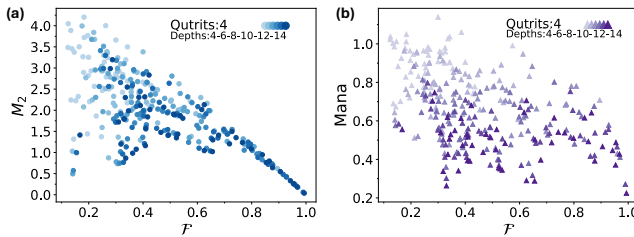
$$= 1 + |\omega^a \mathcal{F} + (1-\mathcal{F})\omega^b|^4 + |\omega^{2a} \mathcal{F} + (1-\mathcal{F})\omega^{2b}|^4 \quad (7.16)$$

$$+ \mathcal{F}^2(1-\mathcal{F})^2 \sum_{r=1,2;j=0,1,2} |e^{i\theta}\omega^{jb}\delta_{a,b\oplus r} + e^{-i\theta}\omega^{ja}\delta_{b,a\oplus r}|^4. \quad (7.17)$$

Notice that for a qutrit,  $a = b \oplus r$  and  $b = a \oplus r$  cannot be satisfied at the same time. This will remove any dependence on the relative phase  $\theta$  between the two basis states ( $\oplus$  indicated sum mod  $d$ ). Considering  $a = 0, b = 1$  we get

$$e^{-M_2} = 1 + |\mathcal{F} + (1-\mathcal{F})\omega|^4 + |\mathcal{F} + (1-\mathcal{F})\omega^2|^4 + 3\mathcal{F}^2(1-\mathcal{F})^2. \quad (7.18)$$

Figure 7.5 (a-b) presents the values of nonstabilizerness vs. fidelity of the optimized QAOA state for 4-qutrit systems. However, we do not find any similarity of our QAOA data with such an equation, suggesting that it may be due to the less successful instance of QAOA, or it is too naive to expect a superposition of two states in the qutrit basis without any knowledge of the cost function.



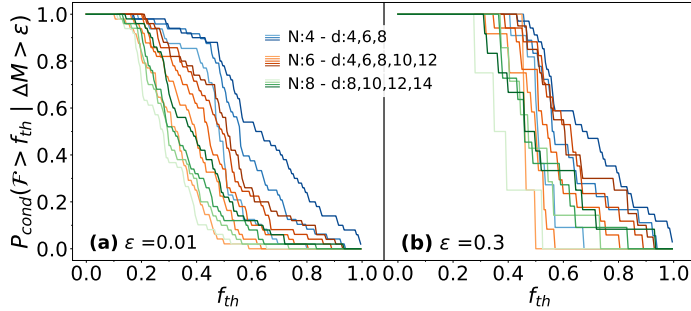
**Figure 7.5.** SRE (a) and Mana (b) as a function of fidelity for a system of four qutrits. Different shades correspond to different depths of the QAOA algorithm.

## 7.6 Demagication and success of QAOA

One may wonder whether the effect of demagication, i.e., the amount by which SRE decreases after the barrier,  $\Delta M = M_2^{\max} - M_2^{\text{final}}$ , is related to the success fidelity. To analyze this question, we calculate the conditional probability ( $\mathcal{P}_{\text{cond}}[f_{\text{th}}, \epsilon]$ ) of obtaining a fidelity greater than some  $f_{\text{th}}$ , provided that the QAOA has shown some minimum amount of demagication,  $\Delta M > \epsilon$ :

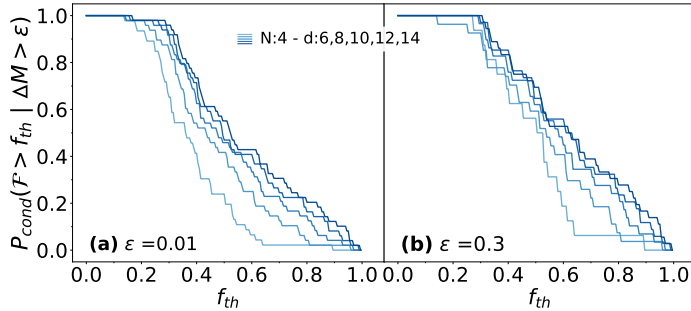
$$\mathcal{P}_{\text{cond}}[F > f_{\text{th}} | \Delta M > \epsilon] = \frac{\mathcal{P}[(F > f_{\text{th}}) \cap (\Delta M > \epsilon)]}{\mathcal{P}[\Delta M > \epsilon]}. \quad (7.19)$$

**For qubits.** As we see in Fig. 7.6, a larger demagication (larger  $\epsilon$ ), appears to result in a higher probability of solving the QAOA with high fidelity. Additionally, the probability of reaching or surpassing a given fidelity also increases (shifts to the right) as the depth increases, which is to be expected.



**Figure 7.6.** Conditional probability of achieving high final fidelity  $f \geq f_{th}$  given a demagication  $\Delta M$  (difference between peak and final magic) exceeding a threshold  $\epsilon$ , for different system sizes and QAOA depths. (a)  $\epsilon = 0.01$ . (b)  $\epsilon = 0.3$ . The rightward shift in (b) relative to (a) shows that stronger demagication correlates with a higher chance of high fidelity. Darker lines represent deeper QAOA circuits.

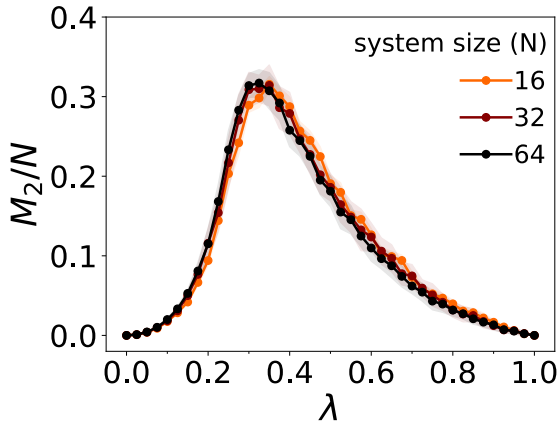
**For qutrits.** Similarly, for qutrits, Fig. 7.7 (a-b) shows the conditional probability of achieving a fidelity above a given threshold, provided the QAOA reaches a minimum amount of demagication.



**Figure 7.7.** (a) Conditional probability curve for  $\epsilon = 0.01$  and (b) for  $\epsilon = 0.3$ , obtained for four qutrits at QAOA depths 6, 8, 10, 12, and 14.

## 7.7 Magic barrier in quantum annealing

To assess the generality of the magic barrier beyond QAOA, we analyze its emergence in continuous-time quantum annealing protocols [236, 242, 540] (see Sec. 3.3.2). To simulate the stationary regime of the annealing protocol, we employ tensor network techniques, specifically matrix product state (MPS) methods [91, 541–543], which allow us to compute numerically exact ground the ground state  $|\psi(\lambda)\rangle$  of the instantaneous Hamiltonian  $H(\lambda = t/T)$ , at arbitrary points along the interpolation, with small bond dimension  $\chi$  (we fixed  $\chi = 60$  in our simulation, at which value the result is converged). This framework also enables efficient computation of SRE using the Pauli Matrix Product State (Pauli MPS) formalism [544], with bond dimension up to  $\chi_P = 1024$ . To make the computation tractable, we truncate the fully connected spin-glass interactions in the cost Hamiltonian to fifth-neighbor couplings. We verified that this truncation does not qualitatively affect the key features of the observed magic dynamics.



**Figure 7.8. Magic barrier in quantum annealing** The SRE density, computed on the ground state of the instantaneous Hamiltonian (ideal adiabatic quantum annealing,  $\lambda \in [0, 1]$ ), exhibits a magic barrier around  $\lambda \sim 0.35$ . The nearly overlapping curves for  $N = 16, 32, 64$  indicate the SRE density traces a barrier largely independent of system size.

The resulting SRE profile, shown in Fig. 7.8, reveals a clear magic barrier: magic rises during the early stages of the sweep, peaks around intermediate  $\lambda$ , and decreases as the system approaches the classical solution. This behavior closely mirrors the magic evolution observed in QAOA.

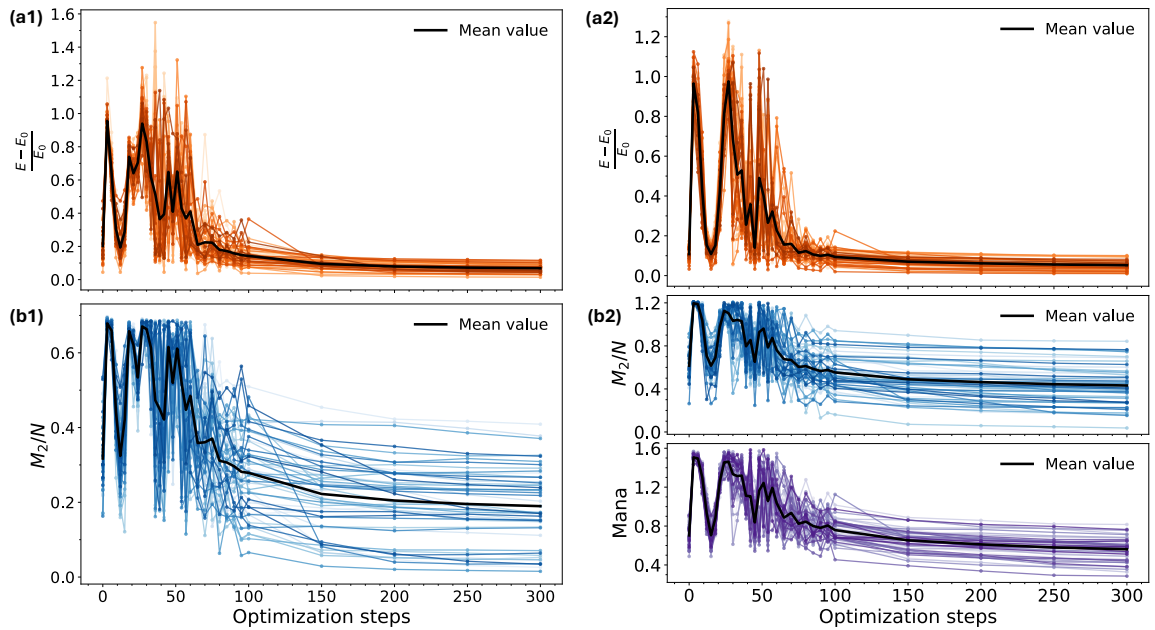
## 7.8 Evolution of magic during classical optimization loop

To reach a more complete understanding of the role of nonstabilizerness in solving optimization problems using variational quantum algorithms, we analyze how it evolves throughout the optimization process. QAOA heavily relies on classical optimization to determine optimal parameters, requiring repeated execution of quantum circuits [Fig. 7.2(a)] that include non-Clifford gates, making them resource-intensive in terms of quantum magic. By studying how magic evolves during the optimization of circuit parameters, we extract information about the resources required to achieve convergence. In particular, we consider 50 problem instances, each defined by different Hamiltonian coefficients, and run 50 QAOA optimizations per instance, initialized with annealing-inspired parameters (increasing  $\gamma$ , decreasing  $\beta$ ) [256]. For each instance, we select the run achieving the lowest energy as the best-performing optimization.

Figure 7.9 shows energy values and nonstabilizerness measures as a function of the optimization steps for the best-performing QAOA run of each of the 50 realizations. Panels (a1) and (b1) refer to results for a 6-qubit system using a QAOA with a circuit depth of 12, while panels (a2) and (b2) report results for a 4-qutrit system using a QAOA circuit with depth 14. The success of QAOA is measured here using the approximation ratio, defined as  $\frac{E-E_0}{E_0}$ , where  $E$  is the expectation value of the Hamiltonian for the optimized state, and  $E_0$  represents the minimum energy obtained through exact diagonalization. During optimization, the energy initially increases, crosses a barrier, and eventually converges close to  $E_0$  for nearly all problem instances. Despite the initial parameterization ensuring a good starting point, the optimization process allows the algorithm to escape local minima and reach a lower-energy minimum, closer to the ground state. This energy barrier is accompanied by a peak in magic, suggesting that crossing a high-nonstabilizerness region is required for convergence. A similar pattern is observed for the Mana in the 4-qutrit case, shown in Fig. 7.9(b2).

Although magic, like energy, shows a barrier during optimization, it captures a

different property: the distance from stabilizer states in the Clifford group. As such, it is not necessarily expected to follow the energy landscape directly. While the energy approximation ratio consistently approaches zero across different instances, indicating convergence toward the ground state [254], the final magic values remain widely spread. This highlights a key point: although the ground state has zero magic, optimized QAOA states often retain finite magic, showing that energy convergence does not imply convergence to the exact solution. Moreover, our Hamiltonians are non-degenerate and thus their ground state are product-states with zero magic. This suggests that even near the exact solution energy, QAOA may converge to a superposition of low-lying eigenstates. This occurs when energy level splittings—which depend on the external field strength—are small, preventing the algorithm from entirely collapsing onto the actual ground state. We explore this further in the next section through fidelity analysis with the exact solution.



**Figure 7.9.** Energy and magic barrier during classical optimization. The black curves represent the mean values calculated over 50 realizations. (a1), (a2) Energy ratio as a function of the number of optimization steps during a QAOA run of each problem realization for 6-qubit and 4-qutrit systems, respectively. (b1) Magic as a function of optimization steps of each QAOA run for the qubit system. (b2) Magic in the upper plot and Mana in the lower plot as a function of optimization steps for the qutrit system.

## 7.9 Summary and outlook

To summarize, magic arises in QAOA even when the target problem has a non-magical solution. Irrespective of the depth of the QAOA, magic rises to a peak and, in a well-performing run, falls towards the end of the protocol, incurring an extra resource cost to overcome, a phenomenon we call the magic barrier. This peak consistently appears near the middle of the QAOA, and upon rescaling, the average magic collapses onto a universal curve across different depths, for both qubits and qutrits, and for SRE and Mana. The fidelity of the final state has a characteristic relation to the final magic,

whose qualitative behavior can be explained analytically through the dependence on probability amplitude and local phase of a simple *ansatz* state.

While in current noisy devices non-Clifford gates are not necessarily more expensive than Clifford gates, the situation is different in early fault-tolerant machines, where non-Clifford operations constitute a significant bottleneck [505, 545–548]. In such cases, the emergence of a magic barrier in QAOA becomes a crucial factor. This motivates the question of whether QAOA can be parameterized to reach the solution state while remaining close to the convex hull of stabilizer states, thus minimizing magic generation.

A natural next step is to test these findings on a quantum processor such as one based on superconducting qubits, cold atoms, or trapped ions, which would also provide a figure of merit for the performance characteristics of a given device. Trapped ions, e.g., appear well suited for a direct test, as they naturally implement long-range interactions as occur in the Sherrington–Kirkpatrick model studied here, and they have recently permitted for measuring nonstabilizerness [173], and for the implementations of quantum optimization protocols [549, 550]

## **Part III**

# **Resources in Disordered Systems**





## Chapter 8

# Disorder-averaged qudit dynamics

*emergence of non-Markovian dynamics from symmetry and disorder*

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## 8.1 Introduction

The effective use of the quantum resources within an engineered quantum device requires precise control. However, even in cases where the device is almost perfectly isolated from its surroundings, uncontrolled and uncertain parameters can significantly restrict the accuracy and efficiency of the quantum device, a challenge shared across experimental platforms. The effect of the lack of certainty and control on the system's evolution is often modeled via disorder, described by random system parameters [551–557]. In condensed matter, disordered systems are ubiquitous, and their fundamental and technological importance makes them a longstanding subject of research [362, 558–564]. In quantum-information-processing devices, the disorder may arise because of imperfect sample production, finite accuracy of control electronics, imprecise gate rotations, or slow drifts of system parameters. In this context, understanding disordered dynamics may be key for the mitigation of errors, e.g., suppressing charge-noise decoherence in superconducting charge qubits [565], boosting coherence via non-ergodicity [566], determining optimal readout times, reducing gate errors [567], tracking drift errors in quantum processors [568], and mitigating sample-to-sample fluctuations in quantum dots [569]. Disorder also plays a key role in quantum annealing [236, 243, 570, 571], environment enhanced quantum transport [572, 573], and the memory effect of disorder-induced localization can even be used to protect against errors [574–576].

While looking for a solution to a disordered model, one is often not interested in an individual disorder realization, either because it is too specific or because the disorder in the physical system is not sufficiently characterized. In such situations, one can derive information about the typical behavior of the quantum system by investigating disorder-averaged ensembles. However, the dynamics of disordered systems are difficult to compute in general, and reaching a converged average often requires considerable numerical effort to acquire sufficient independent samples. In such scenarios, it can be useful to have an effective equation for describing the averaged dynamics, either in the form of a dynamical map or a differential evolution equation. Interestingly, the effective dynamics of the disorder-averaged ensemble can behave as if it were an open system [577]. For various cases, a master equation has been formulated that governs the disorder-averaged evolution, e.g., in the context of a qubit under spectral disorder [578], anisotropic decoherence of qubits [579], relaxation of a qubit under coupling [580], Dirac particles with mass perturbations [201], many-body boson dynamics [581], simulations of the spin-boson model using a disordered

ensemble [582], transport properties in imperfect waveguides [583], obtaining robust control pulses [200], quantum parameter estimation [584], equilibration dynamics of the Sachdev–Ye–Kitaev model [585], and operator spreading [586]. For the case where the dynamical map describing the time-evolved ensemble density matrix can be inverted, Kropf *et al.* [578] discuss a formalism to obtain a master equation of Lindblad form using a matrix-based approach. However, deriving such an exact master equation is not always possible [587].

Here, we discuss an analytical method to find dynamical maps to describe the exact disorder-averaged evolution dynamics. Our framework is valid for arbitrary times and for any unbiased disorder distribution (mean zero), without necessitating a master equation. The Hamiltonians of the considered systems can be Hermitian or non-Hermitian, but they have to be periodic modulo a scalar factor, i.e.,  $H^p \propto \mathbb{I}$  or, more generally,  $H^p \propto H^q$ , with  $p$  and  $q$  integers such that  $q < p$ . In particular, this property covers periodic Hamiltonian matrices with non-trivial base  $q$ , also called  $(q, p)$ -potent [588, 589]. Prominent examples are the tensor-product Hamiltonians of two-level systems, corresponding to involutory matrices with  $p = 2$  and  $q = 0$ , or multi-level clock operators [590, 591], such as  $p = 3$ ,  $q = 0$ . Periodic Hamiltonians are ubiquitous in quantum technologies, e.g., as generators of single- and multi-qubit [372, 592, 593] or qudit gates in quantum computing [594], as stabilizer operators in quantum error correction [148, 595, 596], in error modeling of noisy quantum devices [597], in noisy quantum algorithms [598], etc.

In what follows, we analytically derive the dynamical map giving the disorder-averaged density matrix for such periodic Hamiltonians at any given time and extract common structures for three example cases. The effective equations are characterized by several quantities relevant in quantum information theory, including various witnesses of non-Markovianity [183, 188, 599]. Although solving for the dynamics of each individual realization of the chosen example Hamiltonians is relatively simple, they are non-self-averaging<sup>1</sup>. Therefore, already for somewhat restricted evolution times, numerous independent disorder realizations can be needed to obtain converged results; in one of our example cases, even up to more than  $10^7$  realizations. In such scenarios, having the exact analytic results at one’s disposal provides significant benefits. In some cases, we can also derive a master equation [182] by inverting the underlying dynamical map, the result of which matches with Ref. [578]. Thus, our results further our understanding of how dynamics typically associated with dissipative systems can emerge in deterministic disorder averages, and it may be useful in extracting exact behavior in simple but ubiquitous situations.

The structure of the rest of this chapter is as follows: Section 8.2 presents the analytical approach for deriving the disorder-averaged dynamics, supplemented by examples as well as several quantities that we use to characterize the dynamics, including non-Markovianity witnesses. Next, we delve into three specific cases in detail:  $(p = 2, q = 0)$ , as described, e.g., by single-qubit operators or products of Pauli matrices (Sec. 8.3);  $(p = 3, q = 0)$ , which we exemplify through non-Hermitian qutrit operators (Sec. 8.4); and  $(p = 3, q = 1)$ , describing, e.g., spin-1 operators (Sec. 8.5). Finally, Sec. 8.7 concludes by highlighting the key features of the analytical method and discussing potential future research directions.

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<sup>1</sup>A self-averaging property in a disordered system is one that can be accurately described by averaging over a sufficiently large number of samples [600].

## 8.2 Disorder-averaged dynamics

To set the stage, in this section, we formally derive the dynamical map<sup>2</sup> describing a disorder-averaged ensemble (Sec. 8.2.1). Finding a closed analytical solution is, in general, infeasible, but from its form, one can recognize a drastic simplification if the Hamiltonian matrix fulfills a specific structure, namely if it has a period (Sec. 8.2.2). Before discussing several examples in detail, which we postpone to the next section, we also introduce several quantities estimating quantum informational content of the state, including some witnesses of non-Markovianity (Sec. 8.2.3), which we will use in the later part of the chapter to characterize the disorder-averaged dynamics.

### 8.2.1 Disorder-averaged dynamical map: formal solution

Given a Hamiltonian  $\hat{H}$ , irrespective of it being Hermitian or non-Hermitian, and an initial state  $\hat{\rho}(0)$ , the final state after evolution until a time  $t$  is given by

$$\hat{\rho}(t) = e^{-i\hat{H}t} \hat{\rho}(0) e^{i\hat{H}^\dagger t}, \quad (8.1)$$

(note the dagger at the right of  $\hat{\rho}(0)$ , necessary for the correct description of non-Hermitian dynamics [601]). The system's dynamics follow the differential equation

$$\partial_t \hat{\rho}(t) = -i\hat{H}\hat{\rho}(t) + i\hat{\rho}(t)\hat{H}^\dagger, \quad (8.2)$$

which in the Hermitian case ( $\hat{H} = \hat{H}^\dagger$ ), turns into the well-known von Neumann equation:  $\partial_t \hat{\rho}(t) = -i[\hat{H}, \hat{\rho}(t)]$ .

For a general Hamiltonian  $\hat{H}$ , solving Eq. (8.1) is not effortless. The situation becomes even more complicated if  $\hat{H} = \hat{H}[h]$  describes an ensemble of Hamiltonians with a random variable  $h$  over which we have to average, a procedure we denote by  $\mathbb{E}[\cdot]$ . In particular, one has in general  $\mathbb{E}[e^{-i\hat{H}t}] \neq e^{-i\mathbb{E}[\hat{H}]t}$ , making it challenging to apply the ensemble average to the evolved state. This chapter aims to get a handle on the disorder-averaged density matrix for a specific but ubiquitous class of Hamiltonians useful for quantum information processing.

To calculate the disorder average irrespective of the initial state, it is advantageous to use the super-operator techniques [182, 602] that map between the matrix  $\hat{\rho}$  and a vector  $\vec{\rho}$ . Any left and right multiplications to  $\hat{\rho}$  can be vectorized as column vectors by the following rules

$$\hat{A}\hat{\rho} \leftrightarrow (\mathbb{I} \otimes \hat{A})\vec{\rho} \quad \text{and} \quad \hat{\rho}\hat{B} \leftrightarrow (\hat{B}^T \otimes \mathbb{I})\vec{\rho}. \quad (8.3)$$

By series expanding on both exponentials in Eq. (8.1) and using the super-operator notation above, we derive (see below for full derivation)

$$\vec{\rho}(t) = \sum_{n=0}^{\infty} \frac{(-it)^n}{n!} \sum_{k=0}^n \binom{n}{k} (-\hat{H}^*)^{n-k} \otimes \hat{H}^k \vec{\rho}(0), \quad (8.4)$$

where  $(\cdot)^*$  denotes complex conjugation.

---

<sup>2</sup>The dynamical map of a quantum system describes how an initial density matrix evolves over time, mapping it to the system's state at a later moment.

**Derivation.** By series expanding the evolved density operator  $\rho(t)$  of Eq. (8.1), we obtain

$$\rho(t) = e^{-iHt} \rho(0) e^{+iH^\dagger t} \quad (8.5)$$

$$= \sum_{k=0}^{\infty} \frac{(-it)^k}{k!} H^k \rho(0) \sum_{j=0}^{\infty} \frac{(it)^j}{j!} (H^\dagger)^j \quad (8.6)$$

$$= \sum_{n=0}^{\infty} \frac{(-it)^n}{n!} \sum_{k=0}^n \frac{n!}{(n-k)!k!} H^k \rho(0) (-H^\dagger)^{n-k}. \quad (8.7)$$

Now, we can go to the super-operator notation by using Eq. (8.3) to take the initial state towards the right and extract an evolution term that is independent of the initial state. Note that, the term  $((-H^\dagger)^{n-k})^T = ((-H^\dagger)^T)^{n-k} = (-H^*)^{n-k}$ , and thus we obtain

$$\vec{\rho}(t) = \sum_{n=0}^{\infty} \frac{(-it)^n}{n!} \sum_{k=0}^n \binom{n}{k} (\mathbb{I} \otimes H^k) ((-H^*)^{n-k} \otimes \mathbb{I}) \vec{\rho}(0) \quad (8.8)$$

$$= \sum_{n=0}^{\infty} \frac{(-it)^n}{n!} \sum_{k=0}^{\infty} \binom{n}{k} (-H^*)^{n-k} \otimes H^k \vec{\rho}(0). \quad (8.9)$$

This equation facilitates the Hamiltonian terms to be separated from the initial state by taking  $\vec{\rho}(0)$  outside operator products. Such an expansion of the exponential in the power series of the density operator has previously been used to derive exact equations for the dynamics in multiple scenarios, including the evolution of a classical many-body system [603] as well as an exact generalized master equation for a quantum-mechanical system [604]. It is a procedure quite analogous to the diagrammatic cluster expansion used in statistical mechanics [605].

We denote the disorder-averaged density matrix as  $\mathbb{E}[\vec{\rho}(t)] \equiv \tilde{\vec{\rho}}(t)$ . Assuming no disorder in the initial state  $\rho(0)$ <sup>3</sup>, we arrive at an equation where disorder averaging acts directly on different powers of the Hamiltonian and its complex conjugate,

$$\tilde{\vec{\rho}}(t) = \sum_{n=0}^{\infty} \frac{(-it)^n}{n!} \sum_{k=0}^n \binom{n}{k} \mathbb{E} \left[ (-\hat{H}^*)^{n-k} \otimes \hat{H}^k \right] \vec{\rho}(0) \equiv \Lambda_t[\vec{\rho}(0)]. \quad (8.10)$$

Here, a family of linear quantum dynamical maps  $\{\Lambda_t\}_{t \geq 0}$  from the set of density matrices to itself formally describes the dynamics. In Sec. 8.3-8.5 below, we derive explicit expressions of the disorder-averaged evolution from this equation for the specific cases when the Hamiltonian is periodic, the property we introduce in the next section.

Before that, it is illustrative to establish a formal connection between the ensemble-averaged evolution and that of an open quantum system. In the time-local description of dynamics in an open quantum system, the family of dynamical maps satisfies a differential equation of the form,  $\partial_t \Lambda_t[\cdot] = \mathcal{L}_t \circ \Lambda_t[\cdot]$ , where  $\mathcal{L}_t$  is the Lindbladian that generates the dynamics [182]. Alternatively, using the Nakajima–Zwanzig projective techniques, one can derive an integro-differential master equation involving a memory

<sup>3</sup>Note that our derivation can be extended for the disordered initial state as well, where the disorder averaging should be done together with  $\rho(0)$  if it follows the same disorder distribution as the Hamiltonian or has a correlated disorder.

kernel ( $\mathcal{K}$ ) [182]. The general conditions on  $\mathcal{L}_t$  or  $\mathcal{K}$  that ensure the resulting dynamical map  $\Lambda_t$  is valid, i.e., completely positive and trace-preserving, is still under research, with only certain cases fully understood (see, e.g., Refs. [606–616]). As a result, a general rule for constructing master equations is not always available. However, when the dynamical map is invertible, it is always possible to construct the time-local description, i.e., find  $\mathcal{L}_t$  [190]. For example, from Eq. (8.10), we can obtain the following equations by replacing  $\tilde{\rho}(0) = \Lambda_t^{-1}[\tilde{\rho}(t)]$  in the second equality,

$$\partial_t \tilde{\rho}(t) = \partial_t \Lambda_t [\tilde{\rho}(0)] = \partial_t \Lambda_t \circ \Lambda_t^{-1} [\tilde{\rho}(t)] \equiv \mathcal{L}_t [\tilde{\rho}(t)], \quad (8.11)$$

where  $\circ$  denotes map composition, and we can identify  $\mathcal{L}_t$  as the Lindbladian. Kropf et al. [578] obtain similar master equations based on a matrix formalism [190, 587] when  $\Lambda_t^{-1}$  exists, which forms the foundation for the exact dynamics in their cases. Here, instead, we focus on obtaining the dynamical map  $\Lambda_t$ , which is enough to exactly describe the ensemble-averaged dynamics (i.e.,  $\tilde{\rho}(t)$ ), and our method is independent of the representation of the Hamiltonian in any particular basis. When  $\Lambda_t^{-1}$  exists, we also write down the corresponding master equation.

## 8.2.2 Periodic Hamiltonians

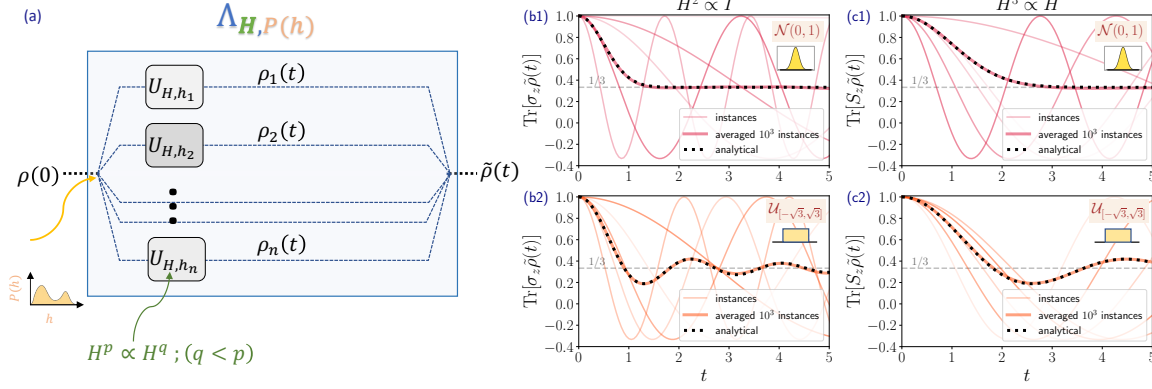
The general formal solution achieved above in Eq. (8.10) does often not help much to extract the physical properties of the system, as one has to calculate arbitrary powers of  $\hat{H}$  to obtain  $\tilde{\rho}(t)$ , at least when one wants to go beyond a perturbative short-time expansion. However, for Hamiltonian matrices with a special structure, namely, if there exists a pair of integers  $p$  and  $q$  (with  $p > q$ ) for which

$$\hat{H}^p = h^{p-q} \hat{H}^q, \quad (8.12)$$

Eq. (8.10) can be solved exactly.

In the literature, matrices with such a property (modulo the prefactor  $h^{p-q}$ ) are called periodic with period  $p - q$  and base  $q$  [588] or  $(q, p)$ -potent [589]. Special cases are periodic matrices with  $q = 1$ , called  $p$ -potent, and with  $q = 1, p = 2$  (i.e.,  $\hat{H}^2 = \hat{H}$ ) called idempotent. If the matrix is non-singular,  $\hat{H}^{p'} = h^{p'-q'} \hat{H}^{q'}$  implies the simpler relation  $\hat{H}^p = h^p \mathbb{I}$ , with  $p = p' - q'$ , which can be seen as a special case of Eq. (8.12) with  $q = 0$ . Matrices with  $q = 0$  and  $p = 2$  are called involutory. Physically, the periodicity of the Hamiltonian implies that the full dynamics are generated by a finite number of distinct operators. Here, we will use this property to reduce the infinite sums in Eq. (8.10) to a finite, tractable number of terms.

As some instructive examples, in Sec. 8.3 to Sec. 8.5, we will consider single-qubit Pauli and single-qutrit clock operators ( $q = 0$  and  $p = 2, 3$ ) as well as spin-1 operators ( $q = 1$  and  $p = 3$ ). For each case, we analytically derive the dynamics of the disorder-averaged density operator and calculate several quantities to characterize it, including non-Markovianity witnesses, as described in the next section. Such analysis can be helpful in determining noise channels [617] and error correction protocols [584]. As the spin-1 operator can be used as an alternative representation for qutrits [497], we can also compare how the symmetry ( $\mathbb{Z}_3$  for clock operators, and  $\text{SU}(2)$  for spin-1) affects its dynamics. Before we go into the details of the results, we introduce several quantities, including witnesses of non-Markovianity, that we will use to analyze the ensuing dynamics.



**Figure 8.1.** (a) An initial state  $\rho(0)$  is evolved under an ensemble of disordered Hamiltonians with a finite period ( $H^p \propto H^q$ ,  $q < p$ ). Each instance is generated from a closed-system time evolution, with disorder  $h_i$  chosen from an arbitrary disorder distribution  $P(h)$ . The disorder averaging induces the effective dynamics of  $\tilde{\rho}(t)$  defined by the dynamical map  $\Lambda_{H,P(h)}$ , which is determined by the period of the Hamiltonian and the probability distribution  $P(h)$  of the disorder. (b1-c2) Time evolution of the expectation value of an observable [(b1,b2)  $\sigma_z$ , (c1,c2)  $S_z$ ] for dynamics governed by periodic Hamiltonians [(b1,b2):  $H^2 \propto \mathbb{I}$ , (c1,c2):  $H^3 \propto H$ ] with Gaussian (top) and uniform (bottom) disorder distributions. The initial state corresponds to  $|+1\rangle$  of the corresponding observables. The evolutions numerically averaged over  $10^3$  samples match very well with the analytical expectation values derived with the disorder-averaged state  $\tilde{\rho}$ . The averaged observables for Gaussian disorder decay monotonically (b1,c1), whereas for the uniform disorder, their expectation value oscillates, with twice the period for  $H^3 \propto H$  (c2) compared to  $H^2 \propto \mathbb{I}$  (b2). Nonetheless, in all the cases, the observables reach  $1/3$  for a long time. Directly using the averaged evolution operator can be advantageous, as it obviates the need for sufficient sampling over individual realizations to reach full convergence.

### 8.2.3 Quantum information theoretic quantities and witnesses of non-Markovianity

Disorder averaging can generate effective dynamics that reproduce effects typical of dissipative systems [578, 585], see Fig. 8.1. It is interesting to question whether the effective dynamics are compatible with a Markovian description. To explore this query, we will analyze several key quantities that serve as important measures in quantum information, which can also act as witnesses of non-Markovianity [599, 618]. Different and inequivalent notions of non-Markovianity exist. We have discussed such quantities in Sec. 2.8.2. In this chapter, we will use trace distance (BLP criterion), logarithmic negativity (RHP criterion), and Purity.

Note that all the above-mentioned witnesses are non-linear functions of  $\rho(t)$ . This property implies that the disorder-averaged witness, calculated from the disorder-averaged state  $\tilde{\rho}$ , differs from the average witness of individual disorder realizations  $\rho(t)$ , i.e.  $W[\tilde{\rho}] \neq \mathbb{E}[W[\rho]]$ , where  $W$  denotes the non-linear witness function (in contrast to standard observables that are linear functions of the state). Which averaging procedure is the more relevant one can depend on the situation at hand. E.g., often the disorder varies from shot to shot, making  $\rho$  hardly accessible and rendering the ensemble  $\tilde{\rho}$  a potentially better description of the system. As each independent sample describes a fully unitary evolution,  $\mathbb{E}[W[\rho]]$  will return only trivial results. In contrast, as we will see further below,  $W[\tilde{\rho}]$  can yield striking differences between the various disorder distributions considered. The non-Markovian witnesses can thus serve as a tool to illuminate the characteristics of the emergent quantum dynamics.



### 8.3 Case I: $\hat{H}^2 = h^2\mathbb{I}$

In this section, we derive the exact quantum dynamics for one of the simplest possible classes covered by our framework,  $\hat{H}^2 = h^2\mathbb{I}$ , i.e., involutory matrices with  $p = 2, q = 0$ . Particular examples are

1. A single spin  $S = 1/2$  or qubit,

$$\hat{H}_1 = h\vec{n} \cdot \vec{\sigma}, \quad (8.13)$$

where  $\vec{n}$  is a three-dimensional unit vector and  $\vec{\sigma} = (\sigma_x, \sigma_y, \sigma_z)$  is the vector of Pauli matrices. For concreteness, in what follows, we choose without the restriction of generality  $\vec{n} = (1, 1, 1)/\sqrt{3}$ ;

2. Many-body tensor products of Pauli matrices

$$\hat{H}_N = h \bigotimes_{i=1}^N (\hat{H}_{1i}/h); \quad (8.14)$$

3. Tensor products of  $\sigma_\alpha$  at different positions,  $P_\alpha$ , where  $\sigma_\alpha = \{\mathbb{I}, \sigma_x, \sigma_y, \sigma_z\}$ . For example,  $\sigma_x \otimes \mathbb{I} \otimes \sigma_y$ .

Such Hamiltonians are ubiquitous in quantum mechanics, ranging from textbook examples such as Stern–Gerlach experiments on single spins to building blocks of advanced quantum technologies in cold-atom and quantum-computing platforms (e.g., single qubit operations are described by  $\hat{H}_1$ , while two-qubit gates such as the Mølmer–Sørensen gate are described by  $\hat{H}_2$  [619]). The parity checking operator for quantum error detection and corrections belongs to  $P_\alpha$  [595]. Here, we want to analyze analytically the effects of disorder on such systems in general.

#### 8.3.1 Effective evolution and master equations

We only consider mean-zero disorder distributions. As a consequence,  $\mathbb{E}[h^{2m+1}] = 0$  with  $m \in \mathbb{N}$ , and we are left with the terms of even  $n$ 's of Eq. (8.10). On top of that, as  $\hat{H}^2 = h^2\mathbb{I}$ , all powers of  $\hat{H}$  are either proportional to  $\mathbb{I}$  (even) or  $\hat{H}$  (odd). These constitute the dynamics of  $\tilde{\rho}(t) = \mathbb{E}[\rho(t)]$

$$\tilde{\rho}(t) = \left[ \left( \frac{1+G(t)}{2} \right) \mathbb{I} \otimes \mathbb{I} + \left( \frac{1-G(t)}{2} \right) \tilde{H}^* \otimes \tilde{H} \right] \tilde{\rho}(0) \equiv \Lambda_t \tilde{\rho}(0), \quad (8.15)$$

where  $\tilde{H} = \hat{H}/h$  is the disorder-free part of the Hamiltonian. Note that Eq. (8.15) needs only two operators out of four possible combinations possible from  $\{\mathbb{I} \text{ and } \tilde{H}\}$  because both  $\mathbb{I} \otimes \tilde{H}$  and  $\tilde{H}^* \otimes \mathbb{I}$  are associated to odd powers of  $h$ , and thus vanish after disorder averaging for a distribution with mean 0. For completeness, we present the derivation of the above equation below.

**Derivation.** To start with, we note that only terms even in  $t$  (i.e., even in  $h$ ) in Eq. (8.9) will survive disorder averaging. So, we only need to take care of the even  $n$  terms. Further splitting these terms into even and odd  $k$ , we find

$$\begin{aligned}
& \left[ \sum_{k=0}^{2n} \binom{2n}{k} (-H^*)^{2n-k} \otimes H^k \tilde{\rho}(0) \right] \\
&= \left[ \sum_{k=0}^n \binom{2n}{2k} (-H^*)^{2n-2k} \otimes H^{2k} + \sum_{k=0}^{n-1} \binom{2n}{2k+1} (-H^*)^{2n-2k-1} \otimes H^{2k+1} \right] \tilde{\rho}(0) \\
&= \left[ h^{2n} \mathbb{I} \otimes \mathbb{I} \left( \frac{2^{2n} + \delta_{n0}}{2} \right) - h^{2n-2} \tilde{H}^* \otimes \tilde{H} \left( \frac{2^{2n}}{2} \right) \right] \tilde{\rho}(0).
\end{aligned} \tag{8.16}$$

Note that the entire simplification of the terms is due to the periodicity of the Hamiltonian. Now, we can perform the average over the disorder,

$$\tilde{\rho}(t) = \left[ \sum_{n=0}^{\infty} \frac{(-it)^{2n}}{(2n)!} \mathbb{E}[h^{2n}] \left( \frac{2^{2n} + \delta_{n0}}{2} \right) \mathbb{I} \otimes \mathbb{I} - \sum_{n=1}^{\infty} \frac{(-it)^{2n}}{(2n)!} \mathbb{E}[h^{2n}] \left( \frac{2^{2n}}{2} \right) \tilde{H}^* \otimes \tilde{H} \right] \tilde{\rho}(0). \tag{8.17}$$

From here, using the moments of the disorder distributions, we will derive the exact analytical form of the equation Eq. (8.15).

The general form in Eq. (8.15) is independent of the type of disorder distribution, whose entire effect is encapsulated in the time-dependent function  $G(t)$ . For the present case of  $p = 2, q = 0$ , we have  $G(t) = \mathbb{E}[e^{-2iht}] = \int dh p(h) e^{-2iht}$ , i.e., it is given by the Fourier transform of the probability distribution with time  $t' = 2t$ , and thus corresponds to the characteristic function<sup>4</sup> containing the behavior and properties of the probability distribution. In Tab. 8.1, we give the exact form of  $G(t)$  for Gaussian and homogeneous uniform disorder distributions.

Quantities	Gaussian $\mathcal{N}(0, \sigma^2)$	Uniform $\mathcal{U}_{[-b,b]}$
Prob. dist. function: $P(h)$	$\frac{1}{\sigma\sqrt{2\pi}} e^{-\frac{1}{2}h^2/\sigma^2}$	$\frac{1}{2b}$
Characteristic function: $\varphi_x(t')$	$e^{-\sigma^2 t'^2/2}$	$\frac{\sin bt'}{bt'}$
Disorder moments: $\mathbb{E}[h^{2n}]$	$\sigma^{2n} (2n-1)!!$	$b^{2n}/(2n+1)$
Time-dependent function: $G(t)$	$e^{-2\sigma^2 t^2}$	$\frac{\sin 2bt}{2bt}$
Decay rate: $\gamma(t)$	$2\sigma^2 t$	$\frac{1}{2} \left( \frac{1}{t} - 2b \cot 2bt \right)$

**Table 8.1.** Relevant quantities describing the exact evolution depending on the disorder distributions: probability distribution function, the characteristic function of the distribution, disorder-averaged moments,  $G(t)$ , emergent decay rate governing the master equation (8.26). The last two quantities are specific to the case  $p = 2, q = 0$ , such as the single-qubit Hamiltonian  $H_1$ .

<sup>4</sup>The characteristic function is the Fourier transform of a probability distribution, encoding all its statistical moments.



**Gaussian disorder:** In the special case of a Gaussian distribution  $(0, \sigma^2)$ , we have  $\mathbb{E}[h^{2n}] = \sigma^{2n}(2n-1)!!$ , and therefore

$$\frac{(-it)^{2n}}{(2n)!} \mathbb{E}[h^{2n}] = \frac{(-i\sigma t)^{2n}}{(2n)!!} = \frac{1}{n!} \left( \frac{-\sigma^2 t^2}{2} \right)^n. \quad (8.18)$$

Inserting this to Eq. (8.17), we obtain

$$\sum_{n=0}^{\infty} \frac{1}{n!} \left( \frac{-\sigma^2 t^2}{2} \right)^n \left( \frac{2^{2n} + \delta_{n0}}{2} \right) = \frac{1}{2} \left[ 1 + e^{-2\sigma^2 t^2} \right] \quad (8.19)$$

and

$$\sum_{n=1}^{\infty} \frac{1}{n!} \left( \frac{-\sigma^2 t^2}{2} \right)^n \left( \frac{2^{2n}}{2} \right) = \frac{1}{2} \left[ e^{-2\sigma^2 t^2} - 1 \right]. \quad (8.20)$$

The final equation for the average time-evolved state in the case of Gaussian disorder thus becomes

$$\tilde{\rho}(t) = \frac{1}{2} \left[ \left( 1 + e^{-2\sigma^2 t^2} \right) \mathbb{I} \otimes \mathbb{I} + \left( 1 - e^{-2\sigma^2 t^2} \right) \tilde{H}^* \otimes \tilde{H} \right] \tilde{\rho}(0). \quad (8.21)$$

**Uniform disorder:** For a uniform box distribution in the interval  $[-b, b]$ , we have  $\mathbb{E}[h^{2n}] = b^{2n}/(2n+1)$ . Thus,

$$\frac{(-it)^{2n}}{(2n)!} \mathbb{E}[h^{2n}] = \frac{(-ibt)^{2n}}{(2n+1)!}. \quad (8.22)$$

Hence, we obtain

$$\sum_{n=0}^{\infty} \frac{(-ibt)^{2n}}{(2n+1)!} \left( \frac{2^{2n} + \delta_{n0}}{2} \right) = \frac{1}{2} \left[ 1 + \frac{\sin 2bt}{2bt} \right]. \quad (8.23)$$

In this case, the disorder-averaged time-evolved state is

$$\tilde{\rho}(t) = \frac{1}{2} \left[ \left( 1 + \frac{\sin 2bt}{2bt} \right) \mathbb{I} \otimes \mathbb{I} + \left( 1 - \frac{\sin 2bt}{2bt} \right) \tilde{H}^* \otimes \tilde{H} \right] \tilde{\rho}(0). \quad (8.24)$$

To illustrate the accuracy of our analytical method, we consider the Hamiltonian  $H_1$  given above, with  $\tilde{H} = \vec{n} \cdot \vec{\sigma}$ , and calculate the average magnetization  $\text{Tr}[\sigma_z \tilde{\rho}(t)]$  using the disorder-averaged state  $\tilde{\rho}(t)$  from the analytical solution. For the considered initial state  $|\uparrow\rangle = (1, 0)^T$ , the analytical calculation gives

$$\text{Tr}[\sigma_z \tilde{\rho}(t)] = \frac{1}{3} (1 + 2G(t)). \quad (8.25)$$

In Fig. 8.1(b1,b2), we plot the cases of Gaussian disorder (top) as well as uniform disorder (bottom). For a reasonable comparison, we set both the variances of the uniform and the Gaussian distribution to unity, i.e.,  $b^2/3 = 1$  and  $\sigma^2 = 1$ . While the ensemble-averaged magnetization dynamics for the Gaussian disorder decays monotonically compared to the value corresponding to an infinite-temperature state [Fig. 8.1(b1)], the dynamics for other distributions can be more complicated, illustrated by the oscillations visible in the case of uniform disorder [Fig. 8.1(b2)]. For both types of

distributions, the exact dynamics over the considered time range match very well with the numerically calculated disorder-averaged magnetization of  $10^3$  instances.

Furthermore, from Eq. (8.15), we can derive a dynamical master equation for the disorder-averaged density matrix following Eq. (8.11). To this end, by inverting the dynamical map in Eq. (8.15), we obtain a time-local master equation

$$\partial_t \tilde{\rho}(t) = -\frac{\partial_t G(t)}{2G(t)} \left[ \tilde{H} \tilde{\rho}(t) \tilde{H}^\dagger - \frac{1}{2} \left\{ (\tilde{H})^\dagger \tilde{H}, \tilde{\rho}(t) \right\} \right]. \quad (8.26)$$

**Deriving the master equation:** From the analytic expression of the time-evolved state, given above and in Eq. (8.15), we can also obtain a von-Neumann type evolution equation for the disorder-averaged  $\tilde{\rho}(t)$ . We rewrite Eq. (8.15) in terms of  $\hat{A}$  and  $\hat{B}$  as defined below,

$$\begin{aligned} \tilde{\rho}(t) &= \frac{1}{2} \left[ \underbrace{(\mathbb{I} \otimes \mathbb{I} + \tilde{H}^* \otimes \tilde{H})}_{\hat{A}} + G(t) \underbrace{(\mathbb{I} \otimes \mathbb{I} - \tilde{H}^* \otimes \tilde{H})}_{\hat{B}} \right] \tilde{\rho}(0) \\ \Rightarrow \Lambda_t &= \frac{1}{2} (\hat{A} + G(t) \hat{B}). \end{aligned} \quad (8.27)$$

Following the steps in Eq. (8.11), we first obtain time derivative of  $\tilde{\rho}(t)$ ,

$$\partial_t \tilde{\rho}(t) = \frac{\partial_t G(t)}{2} \hat{B} \tilde{\rho}(0). \quad (8.28)$$

To replace  $\tilde{\rho}(0) = \Lambda_t^{-1} \tilde{\rho}(t)$ , we evaluate the inverse, which has the form,

$$\Lambda_t^{-1} = \frac{1}{2} \left( \hat{A} + \frac{\hat{B}}{G(t)} \right), \quad (8.29)$$

following from the fact that,  $\tilde{H}^2 = \mathbb{I}$  and thus  $\hat{A}^2 = 2\hat{A}$ ,  $\hat{B}^2 = 2\hat{B}$ ,  $\hat{A}\hat{B} = \hat{B}\hat{A} = 0$ , as well as from  $\Lambda_0 = \mathbb{I}$  whence  $\frac{1}{2}(\hat{A} + \hat{B}) = \mathbb{I}$ . Thus, finally, we obtain,

$$\partial_t \tilde{\rho}(t) = \frac{\partial_t G(t)}{2G(t)} \hat{B} \tilde{\rho}(t). \quad (8.30)$$

By replacing  $\hat{B} = (\mathbb{I} \otimes \mathbb{I} - \tilde{H}^* \otimes \tilde{H})$  and going back from super-operator to matrix notation, we obtain Eq. (8.26).

From this equation, it becomes apparent that the  $\tilde{H}$  coupled to the disordered field assumes the role of jump operators of a corresponding master equation in Lindblad form. Moreover, the effective decay rate  $\gamma(t) \equiv -\frac{\partial_t G(t)}{2G(t)}$  is given entirely by the temporal rate of change of the time-dependent function  $G(t)$ , which in turn is determined by the disorder distribution (see Tab. 8.1 for explicit forms).

For Gaussian disorder,  $G(t)$  and  $\gamma(t)$  are always non-negative analytic functions (see Tab 8.1). In contrast, for uniform disorder,  $G(t)$  periodically switches sign, and  $\gamma(t)$  can have negative values [see Fig. 8.2(b)]. Such dynamics can no longer be  $P$ -divisible [620], and hence it is also not  $CP$ -divisible (see more in Ref. [599]), and thus a negative decay rate can also witness non-Markovianity [190], as we discuss in more detail in Sec. 8.3.3.

### 8.3.2 Relation to error channels

Quantum states are highly fragile and can easily be disrupted by environmental interactions [621, 622], hardware imperfections, or operational gate errors [623]. Adequately understanding the dominant error channels of a given quantum hardware is essential for designing fault-tolerant architectures and error-correction codes, making it one of the critical steps in building reliable quantum computing firmware. To estimate the effects of such sources on quantum devices, one typically constructs noise models using quantum channels, e.g., depolarizing, dephasing, amplitude damping, etc. The present framework can be used to derive analytic descriptions of error channels that originate from uncontrolled gate operations.

To illustrate the main idea, consider a generic dephasing channel defined by

$$\rho \rightarrow \rho' = (1 - p_d)\rho + p_d\sigma_z\rho\sigma_z, \quad (8.31)$$

where with probability  $p_d$  the state dephases, while with probability  $(1 - p_d)$  it remains unchanged. Interestingly, Eq. (8.31) coincides with Eq. (8.15) when setting  $\tilde{H} = \sigma_z$  and choosing the dephasing probability  $p_d = \frac{1-G(t)}{2}$ . As mentioned above,  $G(t)$  is the ensemble-averaged rotation generated by the disorder,  $G(t) = \mathbb{E}[e^{-2iht}]$ . Thus, if the system is subject to uncontrolled disorder, one can immediately determine the associated noise channel and its probability  $p_d$  by only characterizing the type of disorder, here  $\exp(-iht\sigma_z)$ , and the distribution of the pulse areas  $ht$ . This knowledge of disorder can further be used to determine suitable parameters for counteracting disorder-induced dephasing [581]. Similar considerations also hold for qudit systems [587, 617].

### 8.3.3 Non-Markovianity from disorder distributions

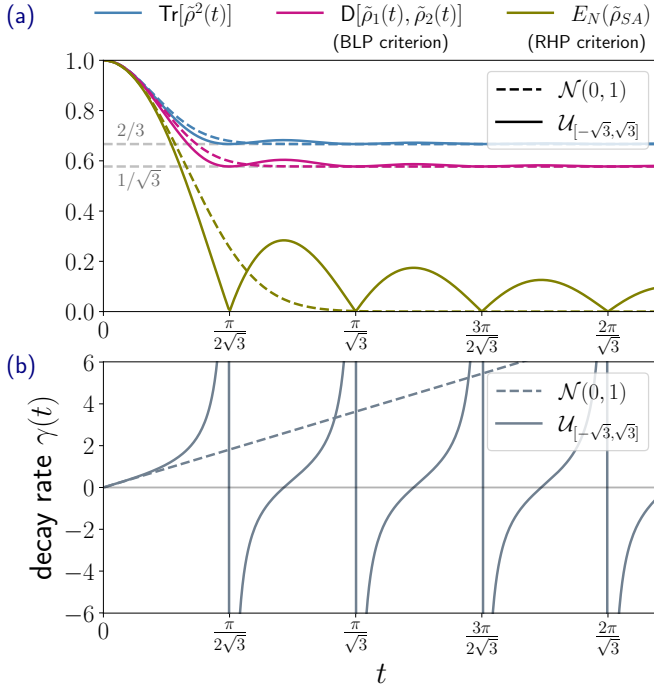
The analytical form of the disorder-averaged state [Eq. (8.15)] permits us to analytically compute the witnesses of non-Markovianity introduced in Sec. 8.2.3. They can be expressed as a function of  $G(t)$ , starting from any initial state, for example,—

$$\begin{aligned} \text{Tr}(\tilde{\rho}^2) &= \frac{1}{3}(2 + G^2(t)), \quad |\rho_0\rangle = |0\rangle\langle 0|; \\ D(\tilde{\rho}_1, \tilde{\rho}_2; t) &= \sqrt{\frac{1 + 2G^2(t)}{3}}, \quad |\rho_0\rangle = |0\rangle\langle 0|, |\rho_1\rangle = |1\rangle\langle 1|; \\ E_N(\tilde{\rho}_{\text{SA}}) &= \log_2(1 + |G(t)|) \quad \rho_{\text{SA},(t=0)} = \frac{1}{\sqrt{2}}(|00\rangle + |11\rangle). \end{aligned} \quad (8.32)$$

In order to see the difference between Gaussian and uniform distribution, we insert the corresponding form of  $G(t)$  as detailed in Eq. (8.32) into the witness functions.

With Gaussian disorder, purity, trace distance, and entanglement negativity decrease and saturate to a particular constant (see Fig. 8.2). This behavior is compatible with a complete Markovianity of the evolution. In contrast, with uniform disorder, the non-Markovian witnesses show significant oscillations and only at late times settle to the same long-time value as that obtained for Gaussian disorder. The periodicity of the oscillation is determined by the periodicity of  $|G(t)|$ , i.e.,  $\frac{\pi}{2b}$ . One can define the strength of non-Markovianity as the number of times the revival of these witnesses happens. This number is inversely proportional to the periodicity of  $|G(t)|$  and thus proportional to the standard deviation of the disorder distribution [624]. Thus, the

revival frequency can be controlled by controlling the standard deviation of the disorder, which might be exploited for information-processing tasks [625, 626]. Moreover, with non-Markovian features, the evolution dynamics can be sped up [627], or the capacity of a quantum channel can be enhanced [628]. Note that, in this case, since the dynamical map is unital (see Section 8.6), purity can be used as a witness of non-Markovianity. It functions just as effectively as other witnesses and exhibits similar behavior.



**Figure 8.2.** (a) Non-Markovian witnesses evaluated for the time-evolved disorder-averaged state for the case  $p = 2, q = 0$  (qubit Hamiltonian) for Gaussian (dashed line) and uniform (solid line) distribution: Purity (blue), trace distance (violet), and entanglement negativity (green). All quantities are consistent with Markovian behavior for Gaussian disorder, but they show distinct non-Markovian features (revivals) for uniform disorder with a period of  $\pi/2b$ . (b) The decay rate in the master Eq. (8.26) also detects non-Markovianity. For the Gaussian disorder, it is always positive, indicating Markovianity, while for the uniform disorder, it assumes negative values in regions where the non-Markovian witnesses show revivals.

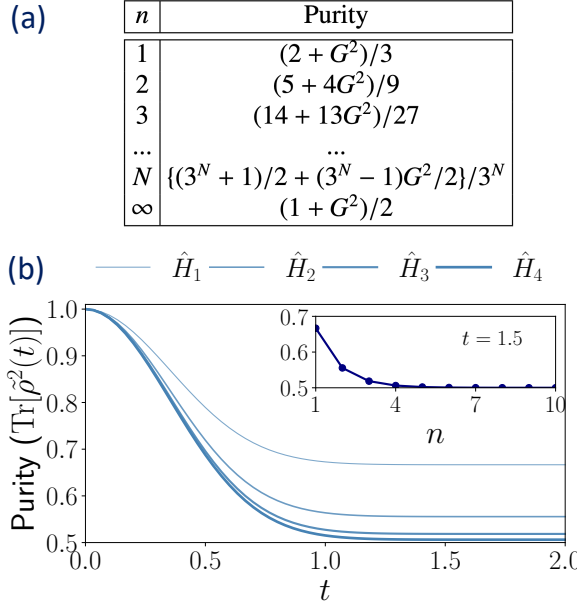
### 8.3.4 Multi-qubit operators

A multi-qubit operator such as  $\hat{H}_N$  as given in Eq. (8.14) also fulfills  $\hat{H}^2 \propto \mathbb{I}$ . Thus, the same analytical solutions as above hold. For concreteness, we consider the multi-qubit Hamiltonian

$$\hat{H}_N = h \bigotimes_{i=1}^N \left( \frac{1}{\sqrt{3}} \sum_{\alpha=x,y,z} \sigma_{\alpha} \right)_i. \quad (8.33)$$

To illustrate the effect of increasing  $N$ , we calculate the purity for Gaussian disorder.

As the table in Fig. 8.3(a) details, the purity consists of a time-independent constant and a time-dependent part given by  $G(t)$ . With increasing  $N$ , the ratio between the corresponding coefficients decreases, e.g., from 2 for  $N = 2$  to  $5/4$  for  $N = 3$ . The general ratio is  $(3^N + 1)/(3^N - 1)$ , showing that both contributions attain equal weight at  $N \rightarrow \infty$ . As  $N$  increases, the purity saturates at a lower value (see Fig. 8.3) and ultimately reaches  $1/2$  (see inset of Fig. 8.3). (Due to the special form of  $\hat{H}_N$ , the dynamics is non-ergodic, and the saturation value remains far from  $1/2^N$ , the value corresponding to a maximally mixed state.)



**Figure 8.3.** (a) Exact formulas for the purity of the disorder-averaged state under the evolution of  $\hat{H}_N$  for the initial state  $|\uparrow\rangle^{\otimes N}$ . The ratio between the terms proportional to  $\mathbb{I} \otimes \mathbb{I}$  and  $\hat{H}^T \otimes \hat{H}$  is  $(3^N + 1)/(3^N - 1)$ , which approaches 1 as  $N$  increases. (b) Exact dynamics of the purity for  $N = 1, 2, 3, 4$  with Gaussian disorder. As  $N$  increases, the final purity plateau approaches  $1/2$ .

## 8.4 Case II: $\hat{H}^3 = h^3 \mathbb{I}$

Until now, we have focused on Hamiltonians generating gates on single or multiple qubits. However, there is currently a significant drive towards developing the use of quantum-information carriers with more than  $d > 2$  levels, so-called qudits [629–636]. The basic operations for qudit computations [594] are typically defined via the generalized Pauli operators, also known as Potts or clock operators in a statistical physics context. These operators present examples of  $(q, p)$ -potent Hamiltonians with  $p = d, q = 0$ .

Qudit operations can be defined by two generators, the shift operator  $\tau$ , which moves the qudit state from  $|i\rangle$  to  $|i - 1\rangle$ , and the phase operator  $\sigma$ , which feeds  $d$ -different eigenvalues that are defined by the  $d$ -th root of unity,  $\omega \equiv e^{2\pi i/d}$ . In matrix form, they read

$$\sigma = \begin{pmatrix} 1 & 0 & 0 & \dots & 0 \\ 0 & \omega & 0 & \dots & 0 \\ 0 & 0 & \omega^2 & \dots & 0 \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & 0 & \dots & \omega^{d-1} \end{pmatrix}, \quad \tau = \begin{pmatrix} 0 & 0 & \dots & 0 & 1 \\ 1 & 0 & \dots & 0 & 0 \\ 0 & 1 & \dots & 0 & 0 \\ \vdots & \vdots & \ddots & 0 & \vdots \\ 0 & 0 & \dots & 1 & 0 \end{pmatrix}, \quad (8.34)$$

and they fulfill the relations

$$\sigma^d = \tau^d = 1, \quad \sigma^\dagger = \sigma^{d-1}, \quad \tau^\dagger = \tau^{d-1}, \quad \sigma\tau = \omega\tau\sigma. \quad (8.35)$$

Notice that the operators  $\sigma, \tau$  are both non-Hermitian. Such non-Hermitian operators have been considered, e.g., in constructing a model of Feynman’s clock [637] and in the context of topological insulators [638]. Another example of such operators also appears in the context of  $\mathbb{Z}_d$  para-fermions, which have instead non-local commutation relations and can also be used for quantum computing [639]. Recently, effective non-Hermitian qudit Hamiltonians have been realized in trapped ions [640, 641].

In this chapter, for the analytical calculation’s tractability, we stick to the case of

qutrits, i.e.,  $d = 3$ . We consider a generic single qutrit Hamiltonian as a sum of both shift and phase operators,

$$\hat{H} = \frac{h}{2^{1/3}}(\sigma + \tau) = h\tilde{H}. \quad (8.36)$$

This Hamiltonian fulfils  $\hat{H}^3 = h^3\mathbb{I}$ . In this case, due to the existence of three independent operators  $\{\mathbb{I}, \tilde{H}, \tilde{H}^2\}$ , we can have at most 9 possible operators contributing to the evolution of the state  $\tilde{\rho}(t)$ . Unlike the  $H^2 \propto \mathbb{I}$  case, where the terms of odd order in  $h$  vanish, in the present case, all the 9 terms contribute. For simplicity, we focus on Gaussian disorder, for which we can write the final disorder-averaged state  $\tilde{\rho}(t)$  as

$$\begin{aligned} \tilde{\rho}(t) = & \frac{1}{9} \left[ 3 \left( \mathbb{I} \otimes \mathbb{I} + \tilde{H}^{*2} \otimes \tilde{H} + \tilde{H}^* \otimes \tilde{H}^2 \right) \right. \\ & + \underbrace{\vec{v}(t) \cdot (1, 1, 1)^T}_{G_1(t)} \left( 2\mathbb{I} \otimes \mathbb{I} - \tilde{H}^{*2} \otimes \tilde{H} - \tilde{H}^* \otimes \tilde{H}^2 \right) \\ & + \underbrace{\vec{v}(t) \cdot (1, \omega^2, \omega)^T}_{G_2(t)} \left( 2\tilde{H}^* \otimes \tilde{H} - \tilde{H}^{*2} \otimes \mathbb{I} - \mathbb{I} \otimes \tilde{H}^2 \right) \\ & \left. + \underbrace{\vec{v}(t) \cdot (1, \omega, \omega^2)^T}_{G_3(t)} \left( 2\tilde{H}^{*2} \otimes \tilde{H}^2 - \tilde{H}^* \otimes \mathbb{I} - \mathbb{I} \otimes \tilde{H} \right) \right] \tilde{\rho}(0), \end{aligned} \quad (8.37)$$

where for convenience we have defined the vector  $\vec{v}(t) \equiv (e^{\frac{3}{2}\sigma^2 t^2}, e^{\frac{3}{2}\omega\sigma^2 t^2}, e^{\frac{3}{2}\omega^2\sigma^2 t^2})$ . Note, for a non-Hermitian system, the state  $\tilde{\rho}$  has to be normalized by  $\text{Tr}(\tilde{\rho})$  in order to obtain normalized probabilities. See the derivation below for a concrete understanding.

To see how this analytical equation regulates the dynamics, we first look at  $t = 0$ , where  $\vec{v}(0) = (1, 1, 1)$ . In this limit, only the first two lines of Eq. (8.37) survive and add up to give  $\mathbb{I} \otimes \mathbb{I} \tilde{\rho}(0)$ , as expected. Similar to the qubit case, we can also identify the time-dependent functions  $G_1(t), G_2(t), G_3(t)$ , which govern the full dynamics at  $t > 0$  and which differ only by the combinations of coefficients  $\{1, \omega, \omega^2\}$ .

**Derivation.** For the case  $\hat{H}^3 = h^3\mathbb{I}$ , we can write more generally  $H^k = h^k \tilde{H}^{k \pmod{3}}$ . As before, the terms in Eq. (8.10) that are odd in  $h$  vanish, and thus we are left with

$$\begin{aligned} & \left[ \sum_{k=0}^{2n} \binom{2n}{k} (-H^*)^{2n-k} \otimes H^k \tilde{\rho}(0) \right] \\ & = \left[ \sum_{k=0}^{2n} \binom{2n}{k} h^{2n} (-1)^{2n-k} \tilde{H}^{*(2n-k) \pmod{3}} \otimes \tilde{H}^{k \pmod{3}} \tilde{\rho}(0) \right]. \end{aligned} \quad (8.38)$$

As  $\tilde{H}^3 \propto \mathbb{I}$ , the dynamics is generated by  $\{\mathbb{I}, \tilde{H}, \tilde{H}^2\}$  and its complex conjugates, thus when written in super-operator formalism like  $B^* \otimes A$ , where  $A, B \in \{\mathbb{I}, \tilde{H}, \tilde{H}^2\}$ , there are 9 possible terms. Now, we can use  $\omega^3 = 1$ , to have a closed form of the combinatorial terms,

$$\begin{aligned}
\binom{n}{0} - \binom{n}{3} + \binom{n}{6} \dots &= \frac{0^n + (1-\omega)^n + (1-\omega^2)^n}{3} \equiv W_{(n,1)}, \\
-\binom{n}{1} + \binom{n}{4} - \binom{n}{7} \dots &= \frac{1}{3} \left( \frac{\delta_{n0}}{1} + \frac{(1-\omega)^n}{\omega} + \frac{(1-\omega^2)^n}{\omega^2} \right) \equiv W_{(n,\omega)}, \\
\binom{n}{2} - \binom{n}{5} + \binom{n}{8} \dots &= \frac{1}{3} \left( \frac{\delta_{n0}}{1} + \frac{(1-\omega)^n}{\omega^2} + \frac{(1-\omega^2)^n}{\omega} \right) \equiv W_{(n,\omega^2)}.
\end{aligned} \tag{8.39}$$

Using the above simplifications, the disorder-averaged density matrix is

$$\begin{aligned}
\tilde{\rho}(t) = & \sum_{n=0}^{\infty} \left[ \frac{(-it)^{6n}}{(6n)!} \mathbb{E}[h^{6n}] \left\{ W_{(6n,1)} \mathbb{I} \otimes \mathbb{I} + W_{(6n,\omega)} \tilde{H}^{*2} \otimes \tilde{H} + W_{(6n,\omega^2)} \tilde{H}^* \otimes \tilde{H}^2 \right\} \right. \\
& + \frac{(-it)^{6n+2}}{(6n+2)!} \mathbb{E}[h^{6n+2}] \left\{ W_{(6n+2,1)} \tilde{H}^{*2} \otimes \mathbb{I} + W_{(6n+2,\omega)} \tilde{H}^* \otimes \tilde{H} + W_{(6n+2,\omega^2)} \mathbb{I} \otimes \tilde{H}^2 \right\} \\
& \left. + \frac{(-it)^{6n+4}}{(6n+4)!} \mathbb{E}[h^{6n+4}] \left\{ W_{(6n+4,1)} \tilde{H}^* \otimes \mathbb{I} + W_{(6n+4,\omega)} \mathbb{I} \otimes \tilde{H} + W_{(6n+4,\omega^2)} \tilde{H}^{*2} \otimes \tilde{H}^2 \right\} \right] \tilde{\rho}(0).
\end{aligned} \tag{8.40}$$

Using  $(1-\omega)^{6n} = (-3)^{3n}$  in Eq. (8.39), we can further simplify the  $W(6n, \cdot)$  terms,

$$\begin{aligned}
W_{(6n,1)} &= \frac{\delta_{n0} + 2(-3)^{3n}}{3}, \quad W_{(6n,\omega)} = W_{(6n,\omega^2)} = \frac{\delta_{n0} - (-3)^{3n}}{3}, \\
W_{(6n+2,1)} &= W_{(6n+2,\omega^2)} = \frac{-(-3)^{3n+1}}{3}, \quad W_{(6n+2,\omega)} = \frac{2(-3)^{3n+1}}{3}, \\
W_{(6n+4,1)} &= W_{(6n+4,\omega)} = \frac{-(-3)^{3n+2}}{3}, \quad W_{(6n+4,\omega^2)} = \frac{2(-3)^{3n+2}}{3}.
\end{aligned} \tag{8.41}$$

Note, the  $\delta_{n0}$  terms in the coefficients corresponding to the operators  $\mathbb{I} \otimes \mathbb{I}$ ,  $\tilde{H}^{*2} \otimes \tilde{H}$ ,  $\tilde{H}^* \otimes \tilde{H}^2$  are responsible for the time-independent terms in Eq. (8.37). The factor of 2 in front of  $\mathbb{I} \otimes \mathbb{I}$ ,  $\tilde{H}^* \otimes \tilde{H}$  and  $\tilde{H}^{*2} \otimes \tilde{H}^2$ , and  $-1$  in all the other terms of the time-dependent part are also due to the above relation in Eq. (8.41). Now, we need to simplify the time-dependent term along with disorder-averaging  $\mathbb{E}[h^\#]$ . In the case of uniform disorder, the result can be expressed in terms of a hypergeometric function. For Gaussian disorder, using  $\mathbb{E}[h^{2n}] = \sigma^{2n}(2n-1)!!$ , one obtains rather compact expressions such as

$$\begin{aligned}
\sum_{n=0}^{\infty} \frac{(-it)^{6n}}{(6n)!} \mathbb{E}[h^{6n}] &= \sum_{n=0}^{\infty} \frac{(-it)^{6n}}{(6n)!} \sigma^{6n} (6n-1)!! \\
&= \sum_{n=0}^{\infty} \frac{(-i\sigma t)^{6n}}{(6n)!!} = \sum_{n=0}^{\infty} \frac{(-\sigma^2 t^2)^{3n}}{2^{3n} (3n)!} = \sum_{n=0}^{\infty} \frac{(-\frac{1}{2} \sigma^2 t^2)^{3n}}{(3n)!},
\end{aligned} \tag{8.42}$$

and similarly for the terms with  $3n$  replaced by  $3n+1$  and  $3n+2$ . The resulting infinite series can be expressed as the sum of three exponentials having permutations of  $(1, \omega, \omega^2)$  in the exponents,

$$\sum_{n=0}^{\infty} \frac{x^{3n+j}}{(3n+j)!} = \frac{e^x + \omega^{2j} e^{\omega x} + \omega^{4j} e^{\omega^2 x}}{3}, \quad \text{for } j = 0, 1, 2, \tag{8.43}$$

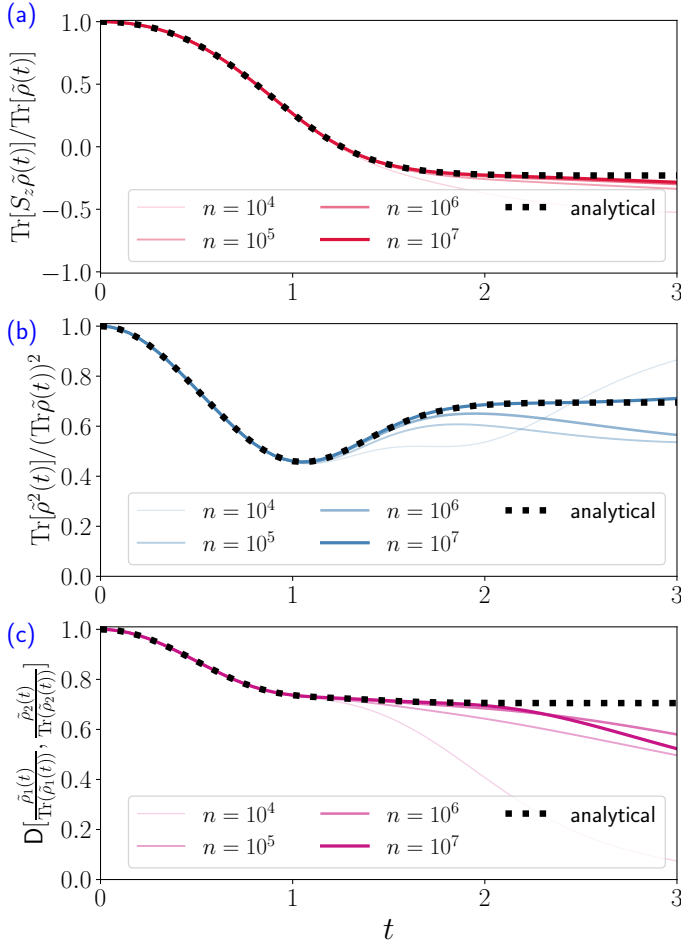
These relations allow us to express the three time-dependent functions  $G_1, G_2, G_3$  in terms of  $\vec{v} = (e^x, e^{\omega x}, e^{\omega^2 x})$  and the three vectors  $(1, 1, 1)^T$ ,  $(1, \omega^2, \omega)^T$ ,  $(1, \omega, \omega^2)^T$ , leading to the Eq. (8.37) describing the disorder-averaged time evolution of a qutrit.



In Fig. 8.4a, we plot the expectation value of a Hermitian observable, the spin-1 magnetization  $S_z = \text{Diag}(1, 0, -1)$  [see Eq. (8.47)] with an initial statevector  $|\psi_0\rangle = (1, 0, 0)^T$ . Its time evolution can be analytically expressed as

$$\frac{\text{Tr}(S_z \tilde{\rho}(t))}{\text{Tr}(\tilde{\rho}(t))} = \frac{12 + 2G_1 - 2^{5/3}G_3}{15 + e^{\frac{3}{2}\sigma^2 t^2} + 2^{4/3}G_2 + 2^{2/3}G_3}, \quad (8.44)$$

which can be compared to the evolution of  $\sigma_z$  in the Hermitian qubit case, see Eq. (8.25). Again, the evolution averaged over the Gaussian distribution decreases monotonically. The numerical average over  $10^4$  trajectories agrees qualitatively with the analytical prediction for the average, but even at  $10^7$  independent samples, slight deviations are still visible at the longest times of  $t \sim 3$  considered.



**Figure 8.4.** (a) The magnetization of the disorder-averaged qudit state reaches a plateau for Gaussian disorder, whereas (b) the purity reaches a dip around  $t \simeq 1$ , followed by an increase and a final plateau. However, the dynamical map being non-unital, it does not witness non-Markovianity. (c) The trace distance does not witness non-Markovianity, as it decays monotonically, without any revival. Beyond very early times, for all the quantities of magnetization, purity, and trace distance, the numerical average converges to the analytical solution only when using a large number of disorder realizations.

Moreover, we calculate the purity of the normalized state [642],

$$\begin{aligned} \frac{\text{Tr}[\tilde{\rho}^2(t)]}{(\text{Tr}\tilde{\rho}(t))^2} = & 3 \left( 81 - 6G_1 + 6 \cdot 2^{1/3}G_2 + 5G_1^2 + 4 \cdot 2^{2/3}G_2^2 \right. \\ & \left. + 6 \cdot 2^{1/3}G_3^2 - 2^{4/3}G_1G_2 - 4G_2G_3 - 2^{8/3}G_1G_3 \right) \\ & / \left( 15 + e^{\frac{3}{2}\sigma^2 t^2} + 2^{4/3}G_2 + 2^{2/3}G_3 \right)^2. \end{aligned} \quad (8.45)$$

As we can see in Fig. 8.4b, the purity has a dip around  $t \simeq 1$ , after which it rises again to finally saturate in a plateau (we have checked that the dip does not derive



from the normalization, which monotonically increases in time as an exponential). This behavior is unlike the qubit case, where the purity of the averaged state monotonically decays. In contrast to the qubit case, the dynamical map generated by the non-Hermitian qutrit Hamiltonian ensemble is non-unital (see Section 8.6), meaning that the non-monotonic behavior of the purity in this case is not a good witness for non-Markovianity. To study a potential non-Markovianity, we also plot the trace distance in Fig. 8.4c. Its monotonous decay with time is compatible with a Markovian evolution.

These quantities do, however, illustrate an interesting aspect: Even at  $10^6$  independent disorder samples, considerable deviations become visible already at around  $\sim 1.75 - 2$ . With  $10^7$  samples, the purity reaches good convergence during the time frame considered, while the trace distance continues to display significant deviations. As this example shows, a vast number of realizations can be needed to match the numerical ensemble average to the analytical predictions, even for relatively short times. Thus, even in simple systems, it can be a considerable asset to be able to analytically extract converged results without the need for extensive numerical averaging.

## 8.5 Case III: $\hat{H}^3 = h^2 \hat{H}$

In the examples above, we have considered cases where finite powers of the Hamiltonian return to  $\mathbb{I}$ . As mentioned in Sec. 8.2.2, we can derive similar analytical predictions also in the case when  $\hat{H}^p = h^q \hat{H}^q$ , with  $q < p$ . In this section, we illustrate this possibility for  $\hat{H}^3 = h^2 \hat{H}$ , i.e.,  $p = 3$  and  $q = 1$ . The simplest instance of this case is given by  $3 \times 3$  matrices describing spin-1 operators,  $S_\alpha$ ,  $\alpha = x, y, z$ . They follow the commutation relations  $[S_\alpha, S_\beta] = i\hbar\epsilon_{\alpha\beta\gamma}S_\gamma$  [643], and we choose the matrix representation

$$S_x = \frac{1}{\sqrt{2}} \begin{pmatrix} 0 & 1 & 0 \\ 1 & 0 & 1 \\ 0 & 1 & 0 \end{pmatrix}, S_y = \frac{1}{\sqrt{2}} \begin{pmatrix} 0 & -i & 0 \\ i & 0 & -i \\ 0 & i & 0 \end{pmatrix}, S_z = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & -1 \end{pmatrix}. \quad (8.46)$$

Interestingly, these spin-1 operators also form a universal basis for qutrit computation [490, 497, 644]. However, they follow a different symmetry than the qutrit or Potts operators. In particular, they are Hermitian, and they obey  $(S_\alpha)^3 = S_\alpha$ . It is, therefore, interesting to investigate the difference in the time-evolution these operators generate after disorder averaging.

In similarity to the qubit case, Eq. (8.13), we choose a Hamiltonian describing a rotation around a specific axis,

$$\hat{H} = \frac{h}{\sqrt{3}}(S_x + S_y + S_z) \Rightarrow \hat{H}^3 = h^2 \hat{H}. \quad (8.47)$$

Using similar techniques to the previous cases, we can write down the final equation,

$$\begin{aligned} \tilde{\rho}(t) = & \left[ \mathbb{I} \otimes \mathbb{I} + (G' - 1) \left\{ \tilde{H}^{*2} \otimes \mathbb{I} + \mathbb{I} \otimes \tilde{H}^2 \right\} \right. \\ & \left. + \frac{1}{2} (1 - G) \tilde{H}^* \otimes \tilde{H} + \frac{1}{2} (3 + G - 4G') \tilde{H}^{*2} \otimes \tilde{H}^2 \right] \tilde{\rho}(0), \end{aligned} \quad (8.48)$$

where  $G$  is identical to the qubit case (Tab. 8.1), and  $G'$  has a similar form as  $G$  but with  $t \rightarrow t/2$ .

**Derivation.** As in the previous cases, terms of odd order in  $h$  vanish. Thus, starting from the Eq. (8.10), we are left with

$$\sum_{n=0}^{\infty} \frac{(-iht)^{2n}}{(2n)!} \left[ \sum_{k=0}^{2n} \binom{2n}{k} (-\tilde{H}^*)^{2n-k} \otimes \tilde{H}^k \right]. \quad (8.49)$$

One can consider the terms  $n = 0, 1, 2, \dots$  one by one and, by putting the terms together, can find the following equation:

$$\begin{aligned} \mathbb{I} \otimes \mathbb{I} &+ \sum_{n=1}^{\infty} \frac{(-iht)^{2n}}{(2n)!} \underbrace{\binom{2n}{0}}_{=1} \tilde{H}^{*2} \otimes \mathbb{I} - \sum_{n=1}^{\infty} \frac{(-iht)^{2n}}{(2n)!} \underbrace{\sum_{k=0}^{n-1} \binom{2n}{2k+1}}_{=2^{2n-1}} \tilde{H}^* \otimes \tilde{H} \\ &+ \sum_{n=2}^{\infty} \frac{(-iht)^{2n}}{(2n)!} \underbrace{\sum_{k=1}^{n-1} \binom{2n}{2k}}_{=2^{2n-1}-2} \tilde{H}^{*2} \otimes \tilde{H}^2 + \sum_{n=1}^{\infty} \frac{(-iht)^{2n}}{(2n)!} \underbrace{\binom{2n}{2n}}_{=1} \mathbb{I} \otimes \tilde{H}^2. \end{aligned} \quad (8.50)$$

For the Gaussian disorder case, using Eq. (8.18), disorder averaging yields

$$\begin{aligned} \mathbb{I} \otimes \mathbb{I} &+ \left( e^{-\sigma^2 t^2/2} - 1 \right) \tilde{H}^{*2} \otimes \mathbb{I} - \frac{1}{2} \left( e^{-2\sigma^2 t^2} - 1 \right) \tilde{H}^* \otimes \tilde{H} \\ &+ \frac{1}{2} \left( 3 + e^{-2\sigma^2 t^2} - 4e^{-\sigma^2 t^2/2} \right) \tilde{H}^{*2} \otimes \tilde{H}^2 + \left( e^{-\sigma^2 t^2/2} - 1 \right) \mathbb{I} \otimes \tilde{H}^2. \end{aligned} \quad (8.51)$$

For the uniform disorder case, using Eq. (8.22), we obtain

$$\begin{aligned} \mathbb{I} \otimes \mathbb{I} &+ \left( \frac{\sin bt}{bt} - 1 \right) \tilde{H}^{*2} \otimes \mathbb{I} - \frac{1}{2} \left( \frac{\sin 2bt}{2bt} - 1 \right) \tilde{H}^* \otimes \tilde{H} \\ &+ \frac{1}{2} \left( 3 + \frac{\sin 2bt}{2bt} - 4 \frac{\sin bt}{bt} \right) \tilde{H}^{*2} \otimes \tilde{H}^2 + \left( \frac{\sin bt}{bt} - 1 \right) \mathbb{I} \otimes \tilde{H}^2. \end{aligned} \quad (8.52)$$

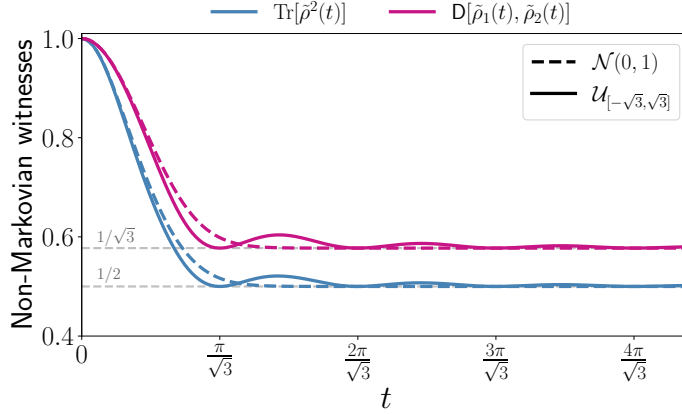
Thus, we obtain the Eq. (8.48) with two time-dependent functions  $G(t)$  and  $G'(t)$ .

Choosing the initial statevector  $|\psi_0\rangle = (1, 0, 0)^T$ , the expectation value of  $S_z$  and the purity take the following forms:

$$\begin{aligned} \text{Tr}(S_z \tilde{\rho}(t)) &= \frac{1}{3} (1 + 2G'(t)), \\ \text{Tr}(\tilde{\rho}^2(t)) &= \frac{1}{18} (9 + G^2(t) + 8G'^2(t)). \end{aligned} \quad (8.53)$$

Figures 8.1 and 8.5 respectively compare the time-evolution for both quantities for the Gaussian and uniform distribution. For the Gaussian distribution, both observables monotonously decay to a plateau. The one of the magnetization,  $\text{Tr}(S_z \tilde{\rho}(t \rightarrow \infty)) = 1/3$ , matches with the long-time plateau obtained in the qubit case, see Eq. (8.25), although the approach towards it is delayed, as it is governed by  $G'(t)$  in the case of

spin-1 instead of  $G(t)$  as was the case for qubits. For the uniform distribution, both the magnetization and purity of the disorder-averaged state oscillate, with a period that is twice that of the qubit case (see Fig. 8.1 for magnetization, and Fig. 8.2 and 8.5 for purity). In the long-time limit, the oscillation dies out, and the dynamics approach the plateau obtained in the case of Gaussian disorder. As discussed in Sec. 8.2.3, as the map in the present case is unital (see Section 8.6), the oscillations of the purity indicate an emergent non-Markovianity of the averaged time evolution.



**Figure 8.5.** Time evolution of purity and trace distance of the disorder-averaged state for Gaussian (dashed line) and uniform (solid line) distribution. The time scales governing the evolution are twice those of the corresponding qubit dynamics (Fig. 8.2).

## 8.6 Unitality of the dynamical maps

A dynamical map is called unital when  $\Lambda_t[\mathbb{I}] = \mathbb{I}$ , i.e., a fully mixed state remains fully mixed. Here,  $\Lambda_t$  acts on the density matrix, not on its vectorized form. We now show that the map for the qubit and spin-1 cases is unital, while it is non-unital for the non-Hermitian qutrit case considered.

**Case I:** For the qubit case, from Eq. (8.15) we have

$$\Lambda_t[\mathbb{I}] = \left( \frac{1 + G(t)}{2} \right) \mathbb{I} + \left( \frac{1 - G(t)}{2} \right) \tilde{H} \mathbb{I} \tilde{H}^\dagger = \mathbb{I}, \quad (8.54)$$

where we used the fact that  $\tilde{H}^\dagger = \tilde{H}$  and  $\tilde{H}^2 = \mathbb{I}$ , showing that the dynamical map for the qubit system is unital. Hence, purity can act as a non-Markovian witness (see Fig. 8.2 and behaves similarly to trace-distance).

**Case II:** For the qutrit ( $\mathbb{Z}_3$ ) case, from Eq. (8.37) we have

$$\begin{aligned} \Lambda[\mathbb{I}] = & \frac{1}{3} (\mathbb{I} + \tilde{H} \tilde{H}^{\dagger 2} + \tilde{H}^2 \tilde{H}^\dagger) + \frac{G_1(t)}{9} (2\mathbb{I} - \tilde{H} \tilde{H}^{\dagger 2} - \tilde{H}^2 \tilde{H}^\dagger) \\ & + \frac{G_2(t)}{9} (2\tilde{H} \tilde{H}^\dagger - \tilde{H}^{\dagger 2} - \tilde{H}^2) + \frac{G_3(t)}{9} (2\tilde{H}^2 \tilde{H}^{\dagger 2} - \tilde{H}^\dagger - \tilde{H}). \end{aligned} \quad (8.55)$$

As all the coefficients of  $G_i(t)$  (in the bracket) remain non-zero, the dynamical map corresponding to a qutrit is non-unital. Hence, purity can not act as a non-Markovian witness (see Fig. 8.4 that purity behaves differently from trace-distance).

**Case III:** For the spin-1 case, from Eq. (8.48) we have

$$\begin{aligned}\Lambda_t[\mathbb{I}] &= \mathbb{I} + (G' - 1)(\tilde{H}^{\dagger 2} + \tilde{H}^2) \\ &\quad + \frac{1}{2}(1 - G)\tilde{H}\tilde{H}^\dagger + \frac{1}{2}(3 + G - 4G')\tilde{H}^2\tilde{H}^{\dagger 2}.\end{aligned}\tag{8.56}$$

Note that  $\tilde{H}^{\dagger 2} + \tilde{H}^2 - 2\tilde{H}^2\tilde{H}^{\dagger 2} = 0$  and  $\tilde{H}^2\tilde{H}^{\dagger 2} - \tilde{H}\tilde{H}^\dagger = 0$  as  $\tilde{H}^\dagger = \tilde{H}$  and  $\tilde{H}^3 = \tilde{H}$ . Thus, we can rewrite the above equation as

$$\Lambda_t[\mathbb{I}] = \mathbb{I} - (\tilde{H}^{\dagger 2} + \tilde{H}^2) + \frac{1}{2}\tilde{H}\tilde{H}^\dagger + \frac{3}{2}\tilde{H}^2\tilde{H}^{\dagger 2} = \mathbb{I},\tag{8.57}$$

and thus the dynamical map for the spin-1 system is unital. Hence, purity can act as a non-Markovian witness (see Fig. 8.5 that purity behaves similarly to trace-distance).

## 8.7 Summary

In this chapter, we have derived an exact equation for the disorder-averaged dynamics of periodic Hamiltonians, independent of the initial states, applicable for Hermitian and non-Hermitian systems, and valid for arbitrary evolution times. These equations rely not on the specific representation of the Hamiltonians but rather only on their  $(q, p)$ -potency class as defined by the relation  $\hat{H}^p \propto \hat{H}^q$ . The effect of the disorder is then entirely captured in a few analytic functions derived from the disorder moments, which can be easily exchanged in the final equations according to the disorder at hand. This analytic approach significantly reduces the numerical overhead typically associated with disorder averaging. As often occurs in such a scenario [200, 201, 577–583], the disorder averaging induces open-system-like dynamics even in inherently closed systems. In cases where the equation defining the time-evolved state can be inverted, we have also derived a master equation governing the disorder-averaged system dynamics. Moreover, even without a formal master equation, we have shown how non-Markovian characteristics emerge as a function of the disorder distribution and its strength. These factors determine the periodicity of revivals observed in non-Markovian witnesses. Such phenomena could be leveraged when using an ensemble of disordered Hamiltonians to simulate open system dynamics [582, 645] as well as for applications in quantum-information processing [200, 201, 625–628]. It is an interesting question to ask what it means to witness non-Markovianity in a closed system without a genuine bath, and how the associated backflow of information should be interpreted.

The analytic expressions may help form a better understanding of errors in quantum computing by enabling insights into noise modeling or aiding in reverse-engineering disorder distributions tailored for specific applications. Another application can be in qudit-based quantum computations, where the effect of the disorder depends not only on the hardware implementation but also on the specific representation of the qudit operator. It thus becomes crucial to investigate which combinations of qudit dimension  $d$ , related to  $q, p$ , and disorder distributions  $P(h)$  offer better resilience against decay caused by averaging over uncontrolled disorder in the qudit gates. Finally, in the context of optimal control, the optimal gate pulses for noisy hardware can be modified, knowing the decay rate caused by specific errors [646]. Beyond an intrinsic interest in analytic solutions to disorder problems, such investigations could thus guide the development of more robust quantum computing systems.

## Chapter 9

# Complexity transitions in chaotic quantum systems

*complexity or resources are not one-size-fits-all*

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## 9.1 Introduction

The interplay between interactions and disorder [564] in quantum many-body systems gives rise to complex systems with extremely rich phenomenology, characterized by varying degrees of chaoticity and ergodicity [647, 648, 648–651]. Although analytical approaches are available in certain limits of ergodic [652–655] and localized regimes [656–659], and while quantum simulations [75, 90, 318, 660] have explored various regimes of disordered interacting systems [661–668], their treatment remains a significant challenge both for theory and experiment. In order to translate that difficulty into a measurable quantity, various notions have been proposed [91, 669–671]. Notable examples of such “complexity markers” include entanglement entropy [672] and stabilizer Rényi entropy [169], which have clear implications both for the difficulty of numerically treating [91, 154, 167, 542, 673–677] the system under study as well as for the quantum resources [47] an experiment needs in order to generate the target state [678, 679]. Such markers are also deeply connected to the question of what actually defines a quantum system as complex [172, 176, 680–682]. As a disordered quantum many-body system enters into respectively less ergodic and more localized regimes, the number of basis states involved is reduced (see Fig. 9.1b), suggesting intuitively a suppression of quantum complexity. As of yet, it remains, however, an open question how precisely ergodicity transitions are related to changes in complexity markers, and in particular, whether these define qualitatively different complexity regimes.

In this chapter, we map out complexity quantifiers (see Fig. 9.1) in prototypical disordered models that exhibit multiple distinct regimes: the Rosenzweig–Porter (RP) model [683, 684], the Power-Law Random Banded Matrix (PLRBM) model [685], and the Sachdev–Ye–Kitaev (SYK) [686, 687] model coupled to an Ising chain (see Fig. 9.3). The RP and PLRBM are paradigmatic models of chaotic systems that interpolate between ergodic and non-ergodic regimes, and they are central to the understanding of Anderson localization [558, 688] and (multi)fractal states [656, 689, 690]. The SYK model is a prototypical example of a maximally chaotic quantum many-body system [691], which plays a key role in the phenomenological description of non-Fermi liquid behavior [686, 692] and in the study of quantum black holes through the holographic principle [693, 694]. We scrutinize these models using three different markers of complexity: (i) The fractal dimension [695–699], a key observable for identifying

localized regions and mobility edges, quantifies how much a state is spread over a computational basis. (ii) The entanglement entropy across half–half bipartitions estimates a state’s quantumness in terms of its inability to be expressed as a direct product state [91]. However, not all entangled states are computationally challenging. An outstanding example is the class of stabilizer states, which contain highly-entangled states but can be tackled efficiently classically thanks to the Gottesman–Knill theorem [149, 150]. In contrast, non-stabilizer states fall outside this framework and are exponentially hard to represent on a classical computer. To capture this distinction, we use as the third measure (iii) the nonstabilizerness [85] (or “magic”) of a state, quantified through the stabilizer Rényi entropy (SRE) [169]. While the fractal dimension and entanglement entropy have been instrumental in characterizing non-ergodic quantum systems already for some time [685, 700–702], magic is currently emerging as a powerful tool in various contexts, including the study of ergodicity [168, 521, 524, 703–706], quantum chaos [525, 707, 708], and as a sensitive probe for identifying quantum phase transitions [518, 526, 528, 709–712].

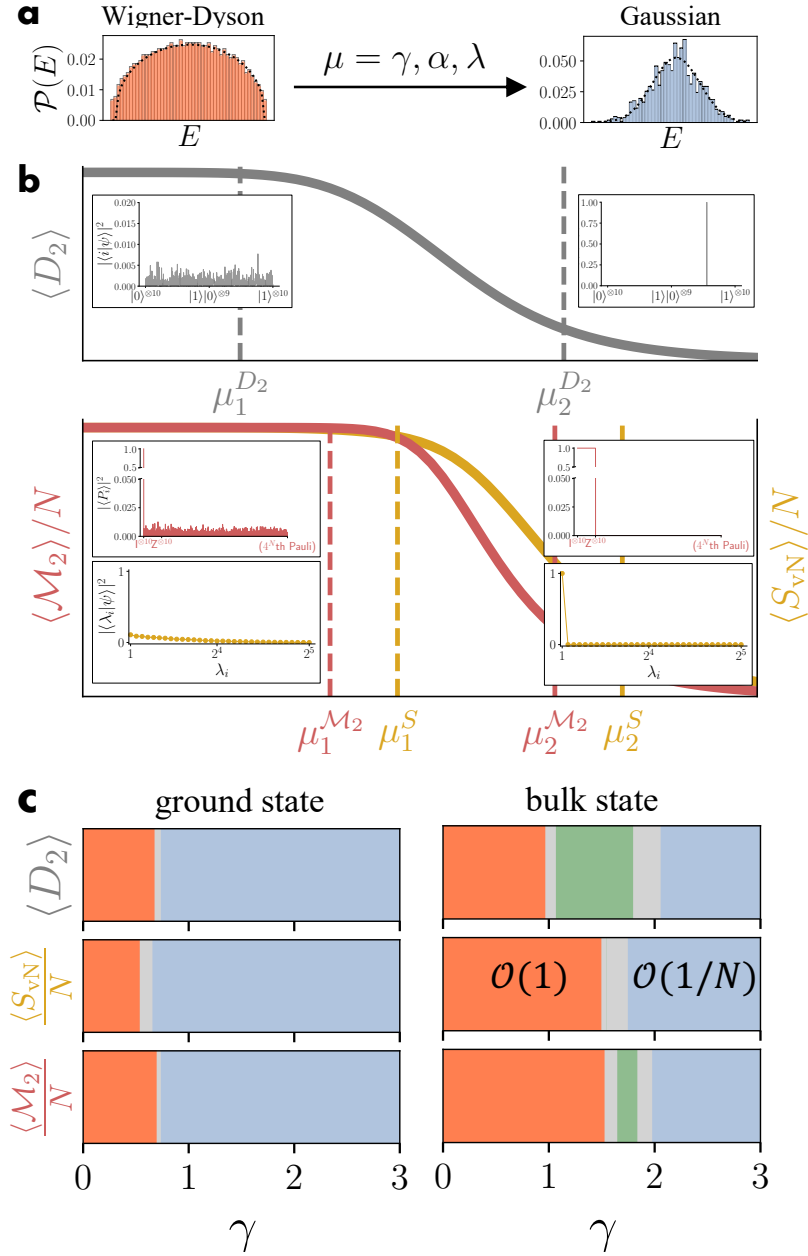
In our work, we find that all three complexity markers show sharp phase transitions, separating regimes of different levels of complexity. Notably, the phase transition points are distinct when an extended fractal phase is present, as in the bulk of the RP spectrum. In contrast, when such a phase is absent, as in the ground state of the RP model, all considered markers detect the transition points close to each other, see Fig. 9.1c. Furthermore, an analysis of the SYK<sub>4</sub>+Ising model reveals that the SRE exhibits a richer structure than the other markers and, in particular, is significantly more sensitive to underlying symmetries of the many-body systems, such as fermion parity and time reversal. Overall, this comparative analysis suggests that no single marker can independently describe all relevant regimes of the system, which may undergo distinct complexity transitions. These results not only yield insights into the physics of disordered quantum many-body systems but also help identify the complexity marker best suited to capture the computational hardness across different phases [676, 677, 682, 713, 714], guiding the selection of appropriate simulation methods.

Finally, our studies complement ongoing research on several complexity markers in ergodic and integrable/localized systems, including Krylov complexity [715–719], logarithmic multifractality [640], operator spreading [720], holographic complexity [721], or quantum-computational complexity [722].

The remainder of the chapter is structured as follows. In Sec. 9.2, we present the complexity markers used to characterize the various quantum phases explored in this study. In Secs. 9.3 and 9.4, we apply these markers to analyze random matrix models, specifically the Rosenzweig–Porter model and the power-law random banded matrix model. In Sec. 9.5, we investigate the Sachdev–Ye–Kitaev model interpolated towards the integrable Ising model. Our main findings and possible future research directions are presented in Sec. 9.8. We also find that the self-averaging properties of the considered complexity markers (Section 9.6.2) can also detect similar phase transitions.

## 9.2 Complexity markers

A plethora of different approaches exists to simulate quantum states, each with varying regimes of validity and efficiency, making it essential to scrutinize the complex-



**Figure 9.1.** Complexity markers across a transition from a chaotic (ergodic) to an integrable (localized) phase. **(a)** As the control parameter  $\mu$  drives a random matrix model towards localization, the eigenvalue distribution evolves from the Wigner–Dyson semicircle to a Gaussian profile. **(b)** Comparison of three complexity markers (data for the RP model). A high (low) fractal dimension  $D_2$  indicates delocalization (localization) in the computational basis. High (low) magic  $\mathcal{M}_2$  implies support over an extensive set of  $4^N$  Pauli strings (concentration on a limited subset). Strong (weak) entanglement  $S_{\text{vN}}$  reflects high (low) Schmidt rank and the presence of many (few) computational basis states in the superposition. While all three markers feature a high-complexity regime, the transition towards intermediate- or low-complexity regimes can happen at distinct points ( $\mu_{1,2}^{D_2} \neq \mu_{1,2}^{\mathcal{M}_2} \neq \mu_{1,2}^S$ ). Regimes of different complexity thus overlap but do not coincide. **(c)** The phase transition points in the RP model, obtained through finite-size scaling analysis, differ across the three complexity markers (orange: high [ $\mathcal{O}(1)$ ] complexity; green: intermediate; blue: low complexity [ $\mathcal{O}(1/N)$ ]). Faded regions indicate numerical uncertainty in determining regime boundaries. Notably, the ground state exhibits a sharp transition directly from high to low complexity, bypassing the intermediate regime. In contrast, bulk eigenstates display a well-defined intermediate phase, with a significant variation in transition points across the considered markers.



ity of quantum states through multiple markers. This section introduces the main markers we will employ to analyze how the system evolves from an ergodic to an integrable/localized regime, the fractal dimension, the maximal entanglement entropy, and the stabilizer Rényi entropy (SRE).

### 9.2.1 Fractal dimension

The fractal dimension can be derived from the inverse participation ratio (IPR), and quantifies the spread of the wavefunction in a given basis, which in our case will be provided by the computational basis states  $\{|i\rangle\}_{i=1}^L$ ,  $L$  being the Hilbert space dimension. Fractal dimension and IPR are commonly used markers to investigate many-body localization [651, 688, 690, 695, 699], as a state that has support on just a few elements of the basis will be in the localized phase, while an ergodic wavefunction will be spread over many states of any generic basis, as shown in Fig. 9.1b. In computational terms, a highly localized state implies a smaller part of the Hilbert space needs to be handled, making it easier to simulate classically [723, 724].

For a wavefunction  $|\psi\rangle$  expressed in terms of the computational basis with coefficients  $\psi(i) = \langle i|\psi\rangle$ , the  $q^{\text{th}}$  IPR is defined as

$$I_q(\psi) = \sum_{i=1}^L |\psi(i)|^{2q}, \quad q > 1, \quad (9.1)$$

where  $L = 2^N$  is the Hilbert space dimension for a space of  $N$  qubits. The fractal dimension  $D_q$  is defined from the IPR as [696–698]

$$D_q = \lim_{L \rightarrow \infty} \frac{\log_L(I_q)}{(1-q)}. \quad (9.2)$$

Typically, one considers  $L \gg 1$  and the limit is dropped, such that the fractal dimension is given by

$$D_q = \frac{\log_2(I_q)}{N(1-q)}. \quad (9.3)$$

When the wavefunction  $\psi$  is confined to a small region of the Hilbert space, the system is considered to be in the localized phase, marked by  $D_q = 0$  in the thermodynamic limit. For a fully extended state, in contrast, where the wavefunction spreads uniformly over all basis states, one obtains  $D_q = 1$ . States in the ergodic phase, however, are usually not entirely uniformly distributed. Instead, they can be approximated by Haar-random states<sup>1</sup>, which do not saturate the fractal dimension for finite system sizes. For  $q = 2$ , the corresponding fractal dimension is given by [725]

$$D_2^{\text{Haar}} = -\frac{1}{N} \log_2 \left( \frac{2}{2^N + 1} \right). \quad (9.4)$$

For  $N \rightarrow \infty$ , one recovers  $D_2^{\text{Haar}} \rightarrow 1$ . For non-extended states, as are expected to occur in non-ergodic but delocalized phases, just a fraction of the volume will be occupied, and we will have  $D_q < 1$ . If  $D_q$  is a constant across all values of  $q$ , the state

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<sup>1</sup>A Haar random state refers to a quantum state that is sampled uniformly at random from the set of all possible pure quantum states in a given Hilbert space, according to the Haar measure.



is called single-fractal (or just fractal), which is the case for the Rosenzweig–Porter model in the non-ergodic extended regime [689]. In the general case,  $D_q$  is a function of  $q$ . Such a state is called multi-fractal. An example occurs in the power-law random banded model at the Anderson transition [685, 688]. Here, we focus on  $q = 2$  only.

## 9.2.2 Entanglement entropy

The second marker we consider is the entanglement entropy maximized with respect to all possible equal-sized (half–half) bipartitions of the state in the computational basis. The maximal entanglement entropy is a key measure of quantum correlations between subsystems. The presence of entanglement has been identified as one of the first limitations for efficient classical simulations of quantum states [154, 726, 727], and entanglement entropy provides an estimate of the bond dimension required in tensor-network methods [91, 542, 673, 674]. Moreover, it is widely used to probe many-body localization and quantum phase transitions [728–732].

Given a bipartition of the Hilbert space  $\mathcal{H} = \mathcal{H}_A \otimes \mathcal{H}_B$  and a pure state  $|\psi\rangle \in \mathcal{H}$ , the von Neumann entropy is defined as  $S(\rho_A) = -\text{Tr}(\rho_A \log \rho_A)$ , with  $\rho_A = \text{Tr}_B |\psi\rangle \langle \psi|$  the reduced density matrix for subsystem  $A$ . The maximal entanglement entropy is then defined as the von Neumann entropy maximized over all possible half–half bipartitions of the system,

$$S_{\max} = \max_{L_A = \lfloor N/2 \rfloor} S(\rho_A). \quad (9.5)$$

For Haar-random states, the entanglement entropy follows the Page curve [733], which for  $L_A \leq L_B$  reads

$$S_{L_A, L_B}^{\text{page}} = \sum_{k=L_B+1}^{L_A L_B} \left( \frac{1}{k} \right) - \frac{L_A - 1}{2L_B}, \quad (9.6)$$

where  $L_{A,B}$  is the Hilbert space dimension of  $\mathcal{H}_{A,B}$ . In this chapter, we work in the convention where logarithms are in base 2, so that  $S_{L_A, L_B}^{\text{page}}/N \approx \frac{1}{2}$  in the large- $N$  limit.

## 9.2.3 Stabilizer Rényi entropy (SRE)

The last complexity marker we consider is the stabilizer Rényi entropy (SRE). Recall from Sec. 2.7.5 that the SRE captures how a state is distributed when expressed in the basis of Pauli operators  $P \in \mathcal{P}_N$ , where  $\mathcal{P}_N$  represents the Pauli group for  $n$ -qubits. Stabilizer states, which are common eigenstates of a maximal set of mutually commuting Pauli operators, exhibit a highly concentrated probability distribution of the Pauli expectation values, confined to a subset of  $2^N$  operators, and thus are easily simulable [148, 149, 151, 158, 166, 167]. In contrast, a Haar-random state has approximately equal weight on all  $P$ 's [168]. Thus, a natural way to quantify the nonstabilizerness of a state  $|\psi\rangle$  is by assessing the spread of this distribution using the  $q$ -th order stabilizer Rényi entropy,

$$\mathcal{M}_q(\psi) = \frac{1}{1-q} \log_2 \sum_{P \in \mathcal{P}_N} \frac{|\langle \psi | P | \psi \rangle|^{2q}}{2^N}. \quad (9.7)$$

In this work, we will use the second-order Rényi entropy,  $q = 2$ . The SRE vanishes for a stabilizer state and grows as the state exhibits increasing degrees of magic.

### SRE in Haar-random states

For unitary-invariant Haar-random states, the expectation value of Pauli strings can be easily calculated analytically by averaging over Haar-random states. Using the formula [734]

$$\int |\psi\rangle \langle \psi|^{\otimes k} d\psi = \frac{(L-1)!}{(L+k-1)} \sum_{\pi \in S_k} W_\pi, \quad (9.8)$$

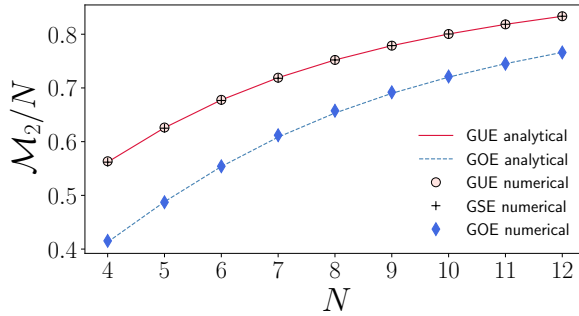
where  $S_k$  is the permutation group and  $W_\pi$  is the replica-permutation operator and  $L = 2^N$ , we find

$$\int |\langle \psi | P_i | \psi \rangle|^2 d\psi = \begin{cases} 1 & i = 0, \\ \frac{1}{2^{N+1}} & i \neq 0. \end{cases} \quad (9.9)$$

The stabilizer Rényi entropy (SRE) can be calculated using similar techniques [168] and takes the following analytical forms depending on the ensemble with respect to which the Haar measure is invariant:

$$\begin{aligned} \mathcal{M}_2^{\text{Haar}}(\text{GUE}) &= \mathcal{M}_2^{\text{Haar}}(\text{GSE}) = -\log_2 \left( \frac{4}{3 + 2^N} \right), \\ \mathcal{M}_2^{\text{Haar}}(\text{GOE}) &= -\log_2 \left( \frac{7}{6 + 2^N} \right). \end{aligned} \quad (9.10)$$

This value approaches the maximal value of 1 in the thermodynamic limit.



**Figure 9.2.** The magic computed for GUE and GSE ensembles exhibits similar behavior, while the GOE shows a distinct trend. The numerically obtained values are in excellent agreement with the corresponding analytical predictions.

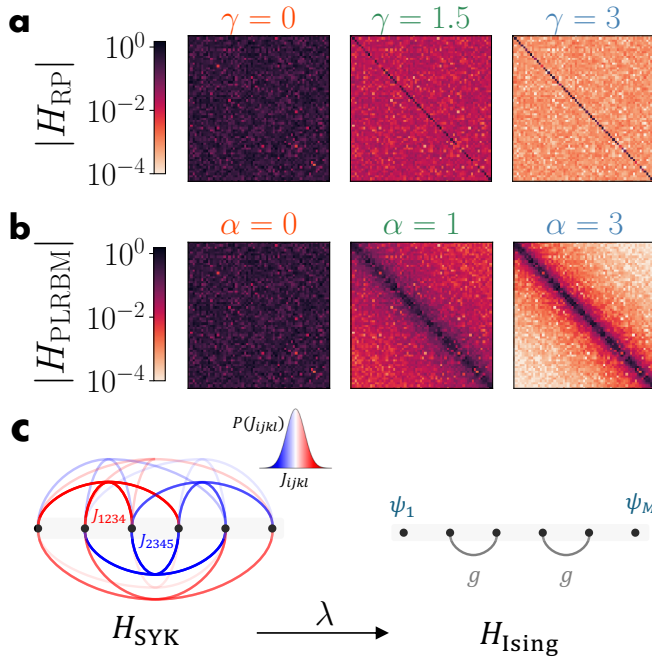
In random matrix theory, the Gaussian ensembles—GOE, GUE, and GSE—are distinguished by their symmetry classes and the associated Dyson index  $\beta \in \{1, 2, 4\}$ , corresponding to real symmetric, complex Hermitian, and quaternionic self-dual Hermitian matrices, respectively [735]. Let  $H$  be an  $N \times N$  Hermitian matrix. The probability distribution over the ensemble is given by  $P(H) \propto \exp\left(-\frac{\beta}{4\sigma^2} \text{Tr}(H^2)\right)$ , where  $\beta = 1$  corresponds to GOE,  $\beta = 2$  to GUE, and  $\beta = 4$  to GSE. The choice of  $\beta$  determines the symmetry class of the matrices:

- GOE ( $\beta = 1$ ):  $H \in \mathbb{R}^{N \times N}$ , with  $H = H^T$  (real symmetric).
- GUE ( $\beta = 2$ ):  $H \in \mathbb{C}^{N \times N}$ , with  $H = H^\dagger$  (complex Hermitian).
- GSE ( $\beta = 4$ ):  $H$  is quaternionic Hermitian (self-dual), which can be represented by  $2N \times 2N$  complex matrices with symplectic structure.

In what follows, we investigate how the three complexity markers discussed above evolve as a system transitions from a chaotic random matrix regime to a localized phase. We consider three representative models: the generalized Rosenzweig–Porter (RP) model (Sec. 9.3), the Power-Law Random Banded Matrix (PLRBM) model (Sec. 9.4), and the Sachdev–Ye–Kitaev (SYK<sub>4</sub>) model coupled to an Ising chain

(Sec. 9.5). These models are characterized by a single tunable parameter, which controls the interpolation between chaotic and localized behavior. Moreover, the RP and PLRBM models exhibit Anderson criticality and support multifractal eigenstates [656, 689], providing a fertile ground for analyzing the distinct responses of the complexity markers.

## 9.3 Rosenzweig-Porter model



**Figure 9.3.** Three model Hamiltonians exhibiting a transition from chaotic (ergodic) to localized phases, controlled by tunable parameters. **(a)** Rosenzweig–Porter (RP) model: A random matrix ensemble where the off-diagonal elements have decreasing weight as the parameter  $\gamma$  increases. **(b)** Power-Law Random Banded Matrix (PLRBM) model: Characterized by off-diagonal elements that decay with a power-law determined by the exponent  $\alpha$ . **(c)** SYK<sub>4</sub>+Ising hybrid model: The coupling strength  $\lambda$  interpolates between the Sachdev–Ye–Kitaev (SYK<sub>4</sub>) model with all-to-all random four-body interactions drawn from a Gaussian distribution (left,  $\lambda = 0$ ) and an Ising chain, expressed in Majorana fermions (right,  $\lambda = 1$ ).

In this section, we study the above complexity markers for the Rosenzweig–Porter (RP) model, both for bulk and ground states. By numerically analyzing varying system sizes, we identify sharp transitions that occur at different points for different markers, and we find distinct behavior in the bulk and ground state.

### 9.3.1 Model

The RP model is an ensemble of  $L \times L$  random Hermitian matrices defined as [683]

$$H_{ij} = \delta_{ij} h_i + \frac{1}{L^{\gamma/2}} V_{ij} (1 - \delta_{ij}). \quad (9.11)$$

Here,  $h_i$  as well as  $V_{ij}$  are independent random Gaussian variables (real for  $h_i$  and complex for  $V_{ij}$ ) with mean zero and standard deviation 1. At  $\gamma = 0$ , the RP model recovers a Random Matrix Theory (RMT), which in the case of the above choices is given by the Gaussian unitary ensemble (GUE). Upon increasing  $\gamma$ , the model interpolates between this RMT and a diagonal Hamiltonian reached at  $\gamma \rightarrow \infty$ , whose eigenvalues are completely uncorrelated.

The relative simplicity of the RP model enables an analytical treatment of both its spectral properties [689, 736] and eigenvector statistics [737–739], allowing for a

comprehensive characterization of the model in the bulk of the energy spectrum. In the ergodic phase, which spans the range  $0 \leq \gamma < 1$ , the model behaves like an RMT, with Wigner–Dyson level spacing and Porter–Thomas statistics for the eigenvector probability distribution, signaling quantum chaos and ergodicity in Hilbert space. At  $\gamma = 1$ , a transition to non-ergodic behavior occurs, and at  $\gamma = 2$  Anderson localization sets in. In the regime  $2 < \gamma < \infty$ , wavefunctions are localized within a small region of Hilbert space, and the level spacing distribution exhibits Poissonian statistics. A key feature of the RP model is the presence of an intermediate regime between the ergodic and localized phases, in the range  $1 < \gamma < 2$ , where wavefunctions remain extended but occupy only a fraction of the full Hilbert space, exhibiting fractal behavior that deviates from conventional ergodicity without being fully localized. The fractal dimension defined in Eq. (9.3) is a particularly effective marker for these transitions [689], leading to the characterization

$$\langle D_q \rangle = \begin{cases} 1 & \gamma < 1, \\ 2 - \gamma & 1 \leq \gamma \leq 2, \\ 0 & \gamma > 2. \end{cases} \quad (9.12)$$

The same phase transitions have been confirmed not only through eigenvalue and eigenstate statistics but also via dynamical properties, such as diffusion characteristics [740, 741], as well as survival and return probabilities [742, 743]. Given the presence of different well-established regimes, the RP model serves as an ideal benchmark for evaluating how the behavior of the various complexity markers across different phases, as we discuss in detail in the following section.

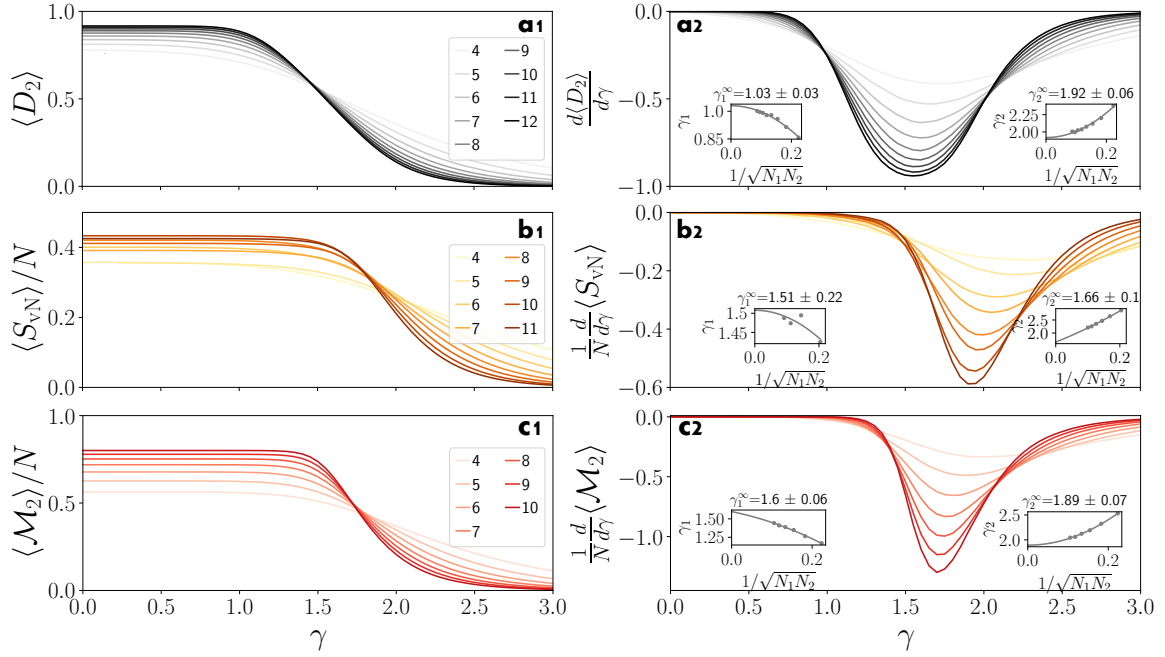
### 9.3.2 Results

In this section, we comparatively study the behavior of eigenvectors—both ground state and bulk states—undergoing a localization transition using the complexity markers defined in Sec. 9.2: fractal dimension  $\langle D_2 \rangle$ , maximum half-chain von-Neumann entanglement entropy  $\langle S_{\text{vN}} \rangle$ , and stabilizer Rényi entropy  $\langle \mathcal{M}_2 \rangle$ , where the angular brackets denote ensemble averaging. We first analyze the bulk eigenstates of the RP model in detail before extending our discussion to the ground state. We focus on the parameter range  $\gamma \in [0, 3]$  for varying system size  $N = 4$  to 12 (corresponding to matrix dimension  $2^N \times 2^N$ ), averaged over multiple samples.

**Bulk state analysis:** To characterize the bulk behavior of the RP model, we compute each complexity marker by averaging over the central 20%<sup>2</sup> of the ordered eigenstates. At  $\gamma = 0$ , the system effectively behaves like a Haar-random ensemble, with all three markers saturating their respective analytical values, as described in Sec. 9.2. In the opposite limit, as  $\gamma \rightarrow \infty$ , the Hamiltonian becomes fully diagonal, leading to eigenstates that are localized in the computational basis. In this regime, both the fractal dimension  $D_2$  and the entanglement entropy  $S_{\text{vN}}$  vanish. Furthermore, since the diagonal Hamiltonian commutes with all  $2^N$  Pauli strings composed of only  $I$  and  $Z$ , these are the only ones contributing to the Pauli spectrum. Hence,

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<sup>2</sup>We obtain the same result if more than 20% of the bulk eigenstates are chosen instead, see Section 9.3.3.



**Figure 9.4.** (a1-c1) Complexity markers (normalized with respect to  $N$ ), averaged over the central 20% of the bulk eigenstates of the RP model for different system sizes from  $N = 4$  up to  $N = 12$ , and averaged over samples from 20000 for  $N = 4$  to 500 for  $N = 12$ . All markers plateau at the maximum value in an extended regime at small  $\gamma$ , overlapping with the model's ergodic regime. The  $N$ -dependent value matches with predictions from the GUE ensemble that is exactly recovered at  $\gamma = 0$ . The plateau's extent in terms of  $\gamma$  depends on the marker: the plateau is largest for  $S_{vN}$  and smallest for  $D_2$ . At large  $\gamma$ , the model enters a localized regime, characterized by a low value of all markers. The slope at which the model goes from the ergodic to the localized regime is different for all markers, as becomes clear from the derivatives (a2-c2). The crossings in the derivatives (insets in a2-c2) determine two different transition points  $\gamma_1$  and  $\gamma_2$ . There are, therefore, three different regimes of high, intermediate, and low complexity, which do not coincide for the different complexity markers used.

every eigenstate is a stabilizer state, resulting in vanishing magic. The Pauli spectra for different regimes are shown in Section 9.7.

For the fractal dimension, two critical points are known within  $\gamma \in (0, 3)$ , corresponding to transitions from ergodic to non-ergodic at  $\gamma_1^c = 1$  and then to the localized phase at  $\gamma_2^c = 2$  [Eq. (9.12)]. However, it is not a priori clear that entanglement and magic should exhibit the identical behavior. As can be seen in Fig. 9.4b1,c1, at small  $\gamma$  they exhibit  $\mathcal{O}(1)$  scaling, indicating a high-complexity regime that one may expect in an ergodic phase. At a marker-dependent value  $\gamma_1$ , the complexity begins to decay, signaling entry into the non-ergodic regime. This decay continues until a second point  $\gamma_2$ , beyond which the markers eventually stabilize to a  $\mathcal{O}(1/N)$  plateau as  $N \rightarrow \infty$ , indicating a low-complexity regime consistent with a large degree of localization. Within the intermediate regime  $\gamma_1 < \gamma < \gamma_2$ , all markers display fractal-like scaling of the form  $\mathcal{O}(N^\delta)$  with  $-1 < \delta < 0$ .

Notably, the points  $\gamma_1$  and  $\gamma_2$ , and therefore the parameter ranges corresponding to high- and low-complexity behavior, differ significantly between markers. For example, as shown in Fig. 9.4, the entanglement entropy features the broadest high-complexity plateau (panel b1), while the fractal dimension shows the narrowest (panel a1). We can make the separation into distinct regimes more precise by examining the first derivative of each marker with respect to  $\gamma$ . The intersection of these derivatives for

different system sizes  $N$  provides a reliable signature of emerging non-analytic behavior, indicating the onset of a sharp transition in the thermodynamic limit. We extract the transition points by extrapolating the crossing positions between two consecutive system sizes,  $N_1$  and  $N_2$ , as a function of the inverse geometric mean,  $1/\sqrt{N_1 N_2}$  [744]. These extrapolations are shown in the insets of Fig. 9.4a2-c2.

In the case of  $\frac{d\langle D_2 \rangle}{d\gamma}$ , the crossing points for consecutive system sizes lie within a relatively narrow range, see the insets of Fig. 9.4a2. By applying a quadratic fit to the crossing locations as a function of  $1/\sqrt{N_1 N_2}$ <sup>3</sup>, we extract the transition points in the thermodynamic limit as  $\gamma_1^\infty[D_2] = 1.03 \pm 0.03$  and  $\gamma_2^\infty[D_2] = 1.92 \pm 0.06$ . These values are in excellent agreement with the analytically predicted transition points from Eq. (9.12), thereby validating our numerical method. The quoted uncertainties are derived from the covariance matrix of the fit parameters.

In the half-system entanglement entropy, an even-odd behavior is observed in the  $\mathcal{O}(1)$  regime: In this regime, the system assumes a volume-law entanglement determined by the subsystem size  $\lfloor N/2 \rfloor$ , leading to nearly identical half-system entanglement densities for  $N$  and  $N + 1$ . However, once the system departs from the  $\mathcal{O}(1)$  regime, this pairing breaks down, as the volume law no longer holds (see Fig. 9.4b1). Considering this effect, we use only even  $N$  when extracting  $\gamma_1^\infty$  via finite-size scaling, while we include all available  $N$  for determining  $\gamma_2^\infty$ . Given the limited number of even-sized systems, the estimate for  $\gamma_1^\infty[S_{\text{vN}}] = 1.51 \pm 0.22$  has a comparatively larger uncertainty. Even when taking these error bars into account, we have that  $\gamma_1^\infty[S_{\text{vN}}] > \gamma_1^\infty[D_2]$ . Conversely, we find  $\gamma_2^\infty[S_{\text{vN}}] = 1.66 \pm 0.10$ , which is smaller than  $\gamma_1^\infty[D_2]$ , indicating that the intermediate regime identified by the entanglement entropy is significantly narrower than the non-ergodic fractal regime.

The behavior of magic is smoother across system sizes. The scaling of its crossing points yields a transition point  $\gamma_1^\infty[\mathcal{M}_2] = 1.60 \pm 0.06$  in the same range as that of the entanglement entropy. However, the transition to the low-complexity  $\mathcal{O}(1/N)$ -regime occurs at a higher value,  $\gamma_2^\infty[\mathcal{M}_2] = 1.89 \pm 0.07$ , rather more closely aligned with that of the fractal dimension  $\gamma_2^\infty[D_2]$ .

Among all the transition points, the most notable contrast lies in  $\gamma_1^\infty$ , where  $S_{\text{vN}}$  and  $\mathcal{M}_2$  display a new phase transition that occurs significantly later than  $\gamma = 1$  and that is undetected by  $D_2$ . A similar trend is evident in their self-averaging behavior (see Section 9.6.2), which undergoes abrupt changes that reinforce this observation. This behavior may stem from the fact that fractal states, while non-ergodic, can still exhibit high entanglement and high magic [706], causing these markers to remain large even after the system departs from full ergodicity. Consequently, the extended regime between  $\gamma_1^\infty[D_2]$  and  $\gamma_1^\infty[\mathcal{M}_2]$  represents a distinct non-ergodic phase, where the states remain highly entangled and highly magical, yet are no longer fully ergodic. This aligns with the intuition that the non-ergodic fractal phase can host highly complex states.

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<sup>3</sup>To find the transition points in the thermodynamic limit, we fit a second-order polynomial curve to the crossing point with respect to  $1/\sqrt{N_i N_{i+1}}$ , of the form

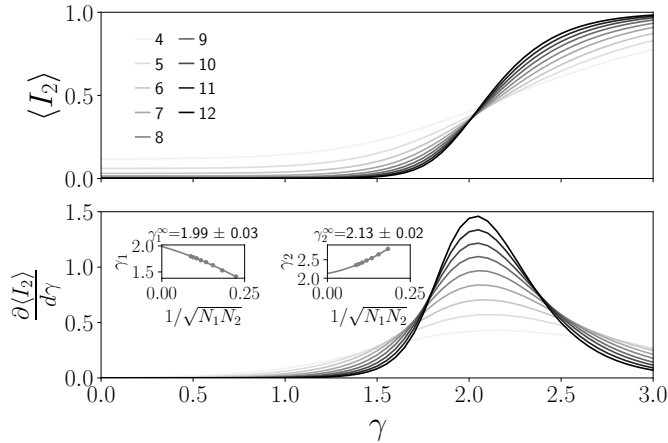
$$\gamma = a + b \frac{1}{\sqrt{N_i N_{i+1}}} + c \frac{1}{N_i N_{i+1}}. \quad (9.13)$$

By extrapolating to  $N \rightarrow \infty$ , we obtain the transition points  $\gamma^\infty = a$ . To obtain the correct points after scaling, we found it crucial to set the acceptable parameter range of  $b$  and  $c$  while fitting the curve. For obtaining  $\gamma_1$ , the fitted curve should move to higher  $\gamma$  with increasing  $N$ , i.e.,  $b, c \leq 0$ , while for  $\gamma_2$ , it should move to smaller  $\gamma$ , i.e.,  $b, c \geq 0$ .



Finally, we highlight an additional noteworthy observation. As discussed in Section 9.6.2, among the various complexity markers, the SRE exhibits superior self-averaging properties across nearly all regimes, except in the interval  $(\gamma_1^\infty[D_2], \gamma_1^\infty[\mathcal{M}_2])$ , where the entanglement entropy shows faster convergence. This enhanced self-averaging behavior makes the SRE a particularly reliable probe for identifying transitions in finite-size systems, where statistical fluctuations are significant. At the same time, the fact that entanglement entropy outperforms SRE in a specific regime underscores the complementary nature of these markers in capturing the structure of complex quantum states.

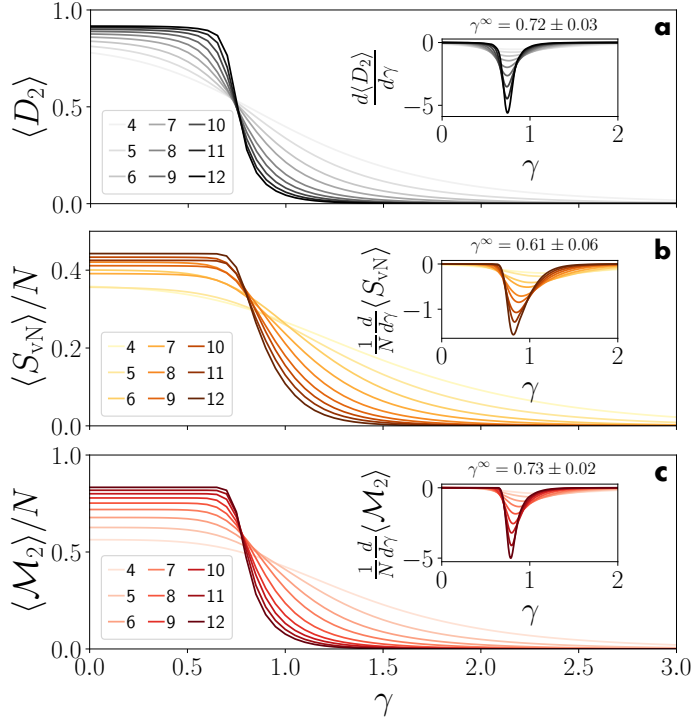
**Comparison between IPR and  $D_2$ :** Although the fractal dimension  $D_2$  can be formally related to the inverse participation ratio (IPR)  $I_2$ , defined in Eq. (9.1), via the expression  $D_q = \frac{\log_2(I_q)}{N(1-q)}$ , relying solely on  $I_q$  to detect phase transitions in such models can be challenging. This is because the multifractal nature of the wavefunctions manifests through the scaling relation  $I_q \propto 1/L^{D_q(q-1)}$  [685, 688, 690, 695, 699], which can be captured in  $D_q$  instead of  $I_q$ . As illustrated in Fig. 9.5, the averaged IPR  $\langle I_2 \rangle$  reveals only the transition near  $\gamma \simeq 2$ , corresponding to the non-ergodic to localized transition, while it fails to resolve the ergodic to non-ergodic transition. This limitation highlights the advantage of using the fractal dimension as a more sensitive diagnostic of intermediate, multifractal phases.



**Figure 9.5.** Unlike the fractal dimension, the average IPR does not resolve two distinct transition points in the bulk phase diagram of the RP model (Fig. 9.4). Instead, the derivative crossings yield nearly coincident transition points, suggesting a single transition point at  $\gamma_c \simeq 2$ .

**Ground state analysis:** The analysis of random matrix models is traditionally focused on the properties of the full eigenspectrum. However, their ground state tends to be more sensitive to ergodicity breaking, giving rise to a mobility edge [649, 745–750] and motivating a dedicated analysis.

As shown in Fig. 9.6, again all complexity markers approach their Haar-random values in the small- $\gamma$  regime, with complexity scaling as  $\mathcal{O}(1)$ . However, they begin to deviate from this value at a smaller  $\gamma$  than their bulk counterparts. Such an earlier transition point in the ground state compared to the bulk indicates the presence of a mobility edge (see Section 9.3.3). Furthermore, all markers transition more rapidly to a  $\mathcal{O}(1/N)$  scaling than was the case in the bulk. As a result, in the large- $N$  limit, the intermediate region with scaling  $\mathcal{O}(1/N^\delta)$  appears to vanish entirely, giving rise to a sharp, first-order-like transition directly from  $\mathcal{O}(1)$  to  $\mathcal{O}(1/N)$  scaling. These abrupt



**Figure 9.6.** Complexity markers for the ground state of the RP model. Inset: first derivative with respect to  $\gamma$ . A direct transition from an ergodic to a localized phase is observed. The transition points detected through fractal dimension  $D_2$  (a) and SRE density  $\mathcal{M}_2/N$  (c) coincide within error bars. The entanglement entropy  $S_{vN}$  (b) indicates the transition at a slightly smaller value of  $\gamma$ , but with a larger error bar due to the even-odd effect.

transitions are identified by extrema in the derivatives of the complexity markers<sup>4</sup>, signaling non-analytic behavior in the thermodynamic limit (see insets of Fig. 9.6). After finite-size scaling, we find that both the fractal dimension and magic detect the ground-state transition at around the same value,  $\gamma^\infty[D_2] = 0.72 \pm 0.03$  and  $\gamma^\infty[\mathcal{M}_2] = 0.73 \pm 0.02$ , respectively. In contrast, for entanglement it appears somewhat earlier, at  $\gamma^\infty[S_{vN}] = 0.61 \pm 0.06$ , but again with a larger error bar. Given the relative vicinity between the different transition points, we cannot exclude the possibility that they identify the same phase transition. Above, we found that the transitions in  $S_{vN}$  and  $\mathcal{M}_2$  occurred well within the fractal regime. One may thus conjecture that all transition points agree when an intermediate fractal regime is missing.

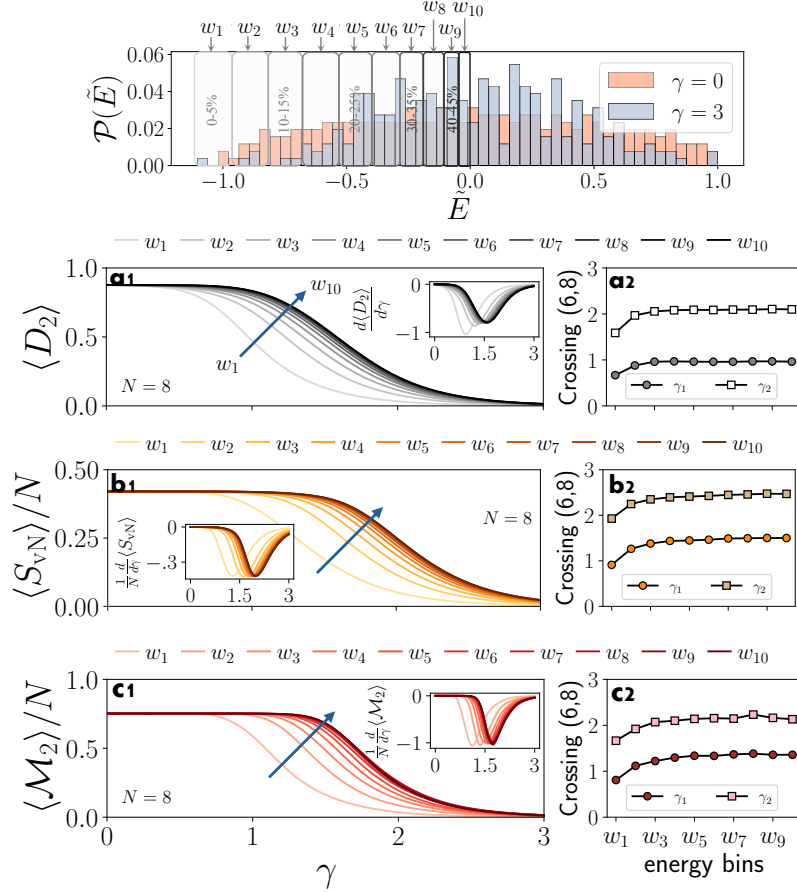
### 9.3.3 Mobility edge in the energy spectrum

The transition to the localized phase can be energy-dependent, a phenomenon referred to as a mobility edge [649, 746–750]. For instance, low-energy states can become localized at lower values of the parameter regime compared to states near the middle of the spectrum. Random banded models are known to exhibit such a mobility edge [745]. A similar trend is observed in our numerical analysis of the RP model (see Fig. 9.1c). As a result, for certain values of the control parameter, low-energy states may already be localized while high-energy states remain ergodic. Given this sensitivity to the energy window, it is crucial to investigate how different complexity markers respond to variations in the chosen spectral region.

In Fig. 9.7a1–c1, we analyze consecutive 5% segments of the energy spectrum. Since the spectrum is symmetric on average, we focus only on the lower half. As we move toward the center of the spectrum, the transition curves systematically shift to

<sup>4</sup>Similar to bulk states, the derivatives of the ground state’s complexity marker for systems of size  $N$  and  $N + 1$  intersect at two distinct points. However, after finite-size scaling, these crossings converge to a single transition point, which is consistently identified by the extrema of the derivative.





**Figure 9.7.** Dependence of the ergodic regime on energy window in the RP model. Top panel: The eigenenergies span a broad range, with densely packed, nearly degenerate states located in the spectral bulk, while the edges are populated by relatively few states. In the ergodic phase ( $\gamma = 0$ ), the energy distribution is flatter as compared to the localized phase ( $\gamma = 3$ ). The eigenstates are ordered in ascending energy, and the lower half of the spectrum is divided into 10 equal windows, labeled  $w_1$  to  $w_{10}$ , each containing 5% of all eigenstates. As the energy window shifts toward the spectral center (thin to thick lines), the  $\mathcal{O}(1)$  ergodic plateau and corresponding transition points  $\gamma_1$  and  $\gamma_2$  move to higher  $\gamma$  values for all complexity markers (**a1–c1**). This shift saturates beyond approximately  $w_3$  (**a2–c2**), indicating that the crossing points remain essentially unchanged when averaging over bulk regions ranging from as narrow as the central 10% to as broad as 90% of the spectrum. Data shown for  $N = 6$  (10000 samples) and  $N = 8$  (4000 samples). Insets of (**a1–c1**) show the derivative of the complexity markers, from which the crossing points are calculated.

the right, indicating that the transition point occurs at higher values of the control parameter, consistent with the presence of a mobility edge. Additionally, the curves corresponding to higher-energy windows quickly collapse onto one another, suggesting rapid convergence. This behavior supports our choice of using the central 20% of the spectrum for the bulk calculations in section 9.3.2, as this range already displays stable and converged behavior.

The rapid convergence becomes even clearer when examining the evolution of the crossing points of the derivatives of the complexity markers, which we used to pinpoint the transitions. As shown in Fig. 9.7a2–c2, these crossing points quickly stabilize to a constant value above the energy window 10%–15%, confirming the robustness of our choice.

## 9.4 Power-Law Random Banded Matrix model

To further deepen our understanding of the complexity transitions in random matrix models, in this section we examine another paradigmatic model—the Power-Law Random Banded Matrix (PLRBM) model.

### 9.4.1 Model

The PLRBM model [685] is conceptually similar to the RP model, as it is a random matrix model with suppressed off-diagonal elements. Nevertheless, its physics and phase diagram have distinct features from the RP model. The PLRBM model is a random matrix ensemble whose elements are given by

$$H_{ij} = G_{ij}a(|i - j|). \quad (9.14)$$

In our case,  $G$  is chosen as a GUE matrix. The coefficients  $a(r)$  implement a polynomial decay of the off-diagonal elements beyond a certain range,

$$a(r) = \begin{cases} 1 & r < b, \\ (r/b)^{-\alpha} & r \geq b, \end{cases} \quad (9.15)$$

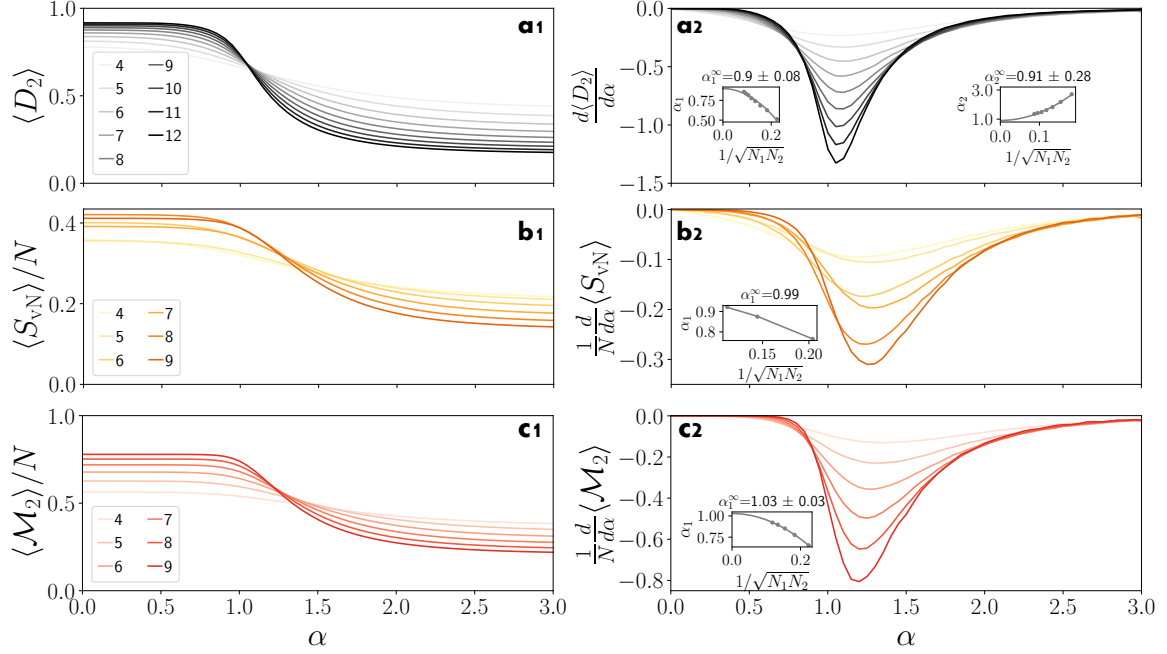
where  $b$  is called the bandwidth parameter and  $\alpha$  is the decay exponent. In this work, we set  $b = 1$ , i.e., only terms adjacent to the diagonal remain intact, and  $\alpha$  remains the single tunable parameter.

The model is known to behave like a random matrix theory for  $\alpha < \frac{1}{2}$  [685, 751]. In the regime  $\frac{1}{2} < \alpha < 1$ , the eigenvectors depart from the Porter–Thomas distribution and follow a generalized hyperbolic distribution [752], signaling the presence of a weakly-ergodic regime distinct from that of the RP model. A recent study on the variable-range SYK<sub>2</sub> model reveals clear signatures of this non-ergodic transition at  $\alpha = \frac{1}{2}$  in the many-body spectral statistics [753]. There is an Anderson critical point at  $\alpha = 1$  [685, 754], where the eigenvectors display proper multifractal behavior that has been intensively studied in the literature [656, 688, 690]. In the range  $1 < \alpha < \frac{3}{2}$ , the PLRBM enters a localized yet superdiffusive phase, while it becomes fully localized for  $\alpha > \frac{3}{2}$ .

The differences between the PLRBM and RP models provide a valuable framework for a comparative analysis of how the complexity markers behave in the different regimes, as we will see in what follows. In this section, we focus on the bulk, again considering the central 20% of the energy eigenstates.

### 9.4.2 Results

The numerical results for the three complexity markers as a function of  $\alpha \in [0, 3]$  are plotted in Fig. 9.8. Note that compared to the RP model, the PLRBM model has more significant fluctuations across realizations due to the persistence of some long-range correlations, which do not vanish in the large  $N$  limit (see Fig. 9.3b). For instance, while 1000 samples suffice for the RP model, we use 10000 samples for the PLRBM at  $N = 8$  (see Section 9.6.1) when computing the SRE—the most self-averaging of the three markers—yet noticeable fluctuations persist.



**Figure 9.8.** (a1-c1) Complexity markers (normalized with respect to  $N$ ), averaged over the central 20% of bulk eigenstates of the PLRBM model for  $N = 4$  up to  $N = 9$  ( $N = 12$  for the fractal dimension) averaged over different sample sizes of 35000 to 3000, respectively. Throughout an extended regime at small  $\alpha$ , all markers plateau at the value matching predictions from the GUE ensemble (recovered by the model at  $\alpha = 0$ ). At large  $\alpha$ , the model transitions to a localized regime, characterized by a lower value of all the markers. The transition point can be deduced from the derivatives of the complexity markers (a2-c2). The crossings in the derivatives (insets a2-c2) effectively find only one distinct transition point. Therefore, in contrast to the RP model, there are only two different regimes of high and low complexity that coincide with the ergodic and localized phases, respectively.

At  $\alpha = 0$ , the state behaves again equal to a Haar-random state and reaches the maximal value for all complexity markers, similar to the RP model. This value extends up to around  $\alpha = 1$ , creating a plateau with value  $\mathcal{O}(1)$  and indicating the extended ergodic regime of the model. Depending on the marker, the complexity begins to decay at a specific value  $\alpha_1$ , signaling the entry of the system into the non-ergodic regime. In the limit for  $\alpha \rightarrow \infty$ , the PLRBM model becomes a tridiagonal matrix. Hence, it does not commute with all the  $2^N$  diagonal Pauli strings and, consequently, the magic is nonzero (see Section 9.7). For the same reason, the eigenstates cannot completely localize in the computational basis. Indeed, in Fig. 9.8 we see that for large values of  $\alpha$  the markers do not vanish and still show a finite  $N$ -dependence. As before, we identify the phase transition points for each complexity marker by considering the positions of the crossing points of the first derivatives in the thermodynamic limit.

For the fractal dimension, we find two crossing points  $\alpha_1^\infty[D_2] = 0.90 \pm 0.08$  and  $\alpha_2^\infty[D_2] = 0.91 \pm 0.28$ , indicating that, in the large  $N$  limit, they will collapse to the same value. The curves for entanglement entropy and magic at different system sizes cross just at one transition point  $\alpha_1^\infty[S_{vN}] = 0.99^5$  and  $\alpha_1^\infty[\mathcal{M}_2] = 1.03 \pm 0.03$ .

<sup>5</sup>For  $S_{vN}$ , the entanglement entropy exhibits even-odd effects even in the  $\mathcal{O}(1)$  regime, prompting us to focus on even  $N$  values to locate the transition. To improve resolution near the critical point, we compute additional data for  $N = 10$ , resulting in three crossing points—(4,6), (6,8), and (8,10). These enable a quadratic fit that recovers the Anderson localization transition at  $\alpha = 1$ , although the limited number of points precludes a reliable error estimate.

Note there might be a second crossing point that remains undetected due to the more sizable fluctuations in the localized regime for large  $\alpha$ . Thus, in contrast to the RP model, our analysis does not detect an intermediate regime that is non-ergodic but delocalized. Rather, the available data suggest a direct transition from the delocalized to the localized phase close to the Anderson critical point at  $\alpha = 1$ .

Although the analysis of the PLRBM model is restricted to smaller system sizes and exhibits larger fluctuations, it supports what we conjectured for the RP model: the complexity markers assume distinct values across the ergodic-to-non-ergodic transition only in the presence of a genuine fractal (or multifractal) phase. Indeed, within error bars, in the PLRBM model, the transition points identified by all three complexity markers coincide (or almost coincide) with the Anderson critical point, the only transition captured by the fractal dimension in the thermodynamic limit. We do not report here the results for the ground state behavior of the PLRBM model since they qualitatively resemble that of the RP model considered in Sec. 9.3.2: the transitions occur earlier than in the bulk [755], yet no intermediate phase is observed. This is consistent with expectations, as the markers used do not reveal a clear extended non-ergodic regime even in the bulk of the PLRBM model.

## 9.5 SYK<sub>4</sub>+Ising model

In this section, we consider a quantum many-body model subject to a chaotic–integrable transition to contrast with the random-matrix behavior. For this, we choose a hybrid model consisting of the SYK model interpolating towards the Ising model. The SYK model, originally introduced as a phenomenological model for non-Fermi liquids [686], is a quantum mechanical system consisting of  $M$  Majorana fermions with random  $q$ -body interactions. It has accumulated significant attention due to the combination of strong interactions and solvability in the large- $M$  limit [687, 756], making it a paradigmatic system to study quantum chaos beyond RMT. Moreover, randomly interacting many-body systems, like the SYK model, are known to behave drastically differently from RMT near the edge of their energy spectrum [757]. This makes it particularly interesting to compare the complexity markers for the ground state of this model to those of the RP model of Sec. 9.3.2. Further, the SYK model is a prototypical system in the study of black holes and quantum gravity [693, 694] through the AdS/CFT correspondence [758, 759]. Indeed, at low energies, it exhibits an emergent conformal symmetry and is effectively described by a Schwarzian action [756], mirroring the behavior of Jackiw–Teitelboim gravity [760]. Since the duality becomes manifest in the infrared, it is of particular interest to understand the ground-state properties of the model in order to assess the stability of the holographic phase under perturbations.

### 9.5.1 Model

The system under study is composed of  $M$  interacting Majorana fermions, described by operators  $\psi_i$  fulfilling the commutation relations  $\{\psi_i, \psi_j\} = \delta_{ij}$ . The Hilbert space of the model is isomorphic to that of  $N = M/2$  qubits. The interactions between the fermions are chosen to interpolate between a chaotic Hamiltonian at  $\lambda = 0$ , provided

by the SYK model, and a fully localized, integrable Ising chain reached at  $\lambda = 1$ ,

$$H = (1 - \lambda)H_{\text{SYK}_4} + \lambda H_{\text{Ising}}. \quad (9.16)$$

The SYK<sub>*q*</sub> model is a system with  $M$  all-to-all interacting Majorana fermions with random  $q$ -body interactions, (see Fig. 9.3c where  $q = 4$ ), defined as

$$H_{\text{SYK}_q} = (i)^{q/2} \sum_{1 \leq i_1 < \dots < i_q \leq M} J_{i_1 \dots i_q} \psi_{i_1} \dots \psi_{i_q}, \quad (9.17)$$

where the couplings  $J_{i_1 \dots i_q}$  are independent and identically distributed random variables drawn from a Gaussian distribution with zero mean and variance given by  $\langle J_{i_1 \dots i_q}^2 \rangle = J^2(q-1)!/M^{q-1}$ . In this article, we work with  $q = 4$ .

The Ising Hamiltonian is given by an anti-ferromagnetic ZZ spin chain without a magnetic field,  $H_{\text{Ising}} = \sum_{i=1}^{N-1} g \sigma_i^z \sigma_{i+1}^z$ . It can be written in terms of Majorana operators using the Jordan–Wigner transformation, which maps each Majorana operator to a Pauli string consisting of  $N$  Pauli matrices [761],

$$\psi_{2i-1} = \sigma_1^x \otimes \dots \otimes \sigma_{i-1}^x \otimes \sigma_i^z \otimes \mathbb{I}_{i+1} \otimes \dots \otimes \mathbb{I}_N, \quad (9.18)$$

$$\psi_{2i} = \sigma_1^x \otimes \dots \otimes \sigma_{i-1}^x \otimes \sigma_i^y \otimes \mathbb{I}_{i+1} \otimes \dots \otimes \mathbb{I}_N. \quad (9.19)$$

Using  $\psi_{2i} \psi_{2i+1} = \mathbb{I}_1 \otimes \dots \otimes \mathbb{I}_{i-1} \otimes -i \sigma_i^z \otimes \sigma_{i+1}^z \otimes \mathbb{I}_{i+2} \otimes \dots \otimes \mathbb{I}_N$ , we obtain

$$H_{\text{Ising}} = i \sum_{i=1}^{M/2-1} g \psi_{2i} \psi_{2i+1}, \quad (9.20)$$

where only interactions between even-odd Majorana fermions remain, see Fig. 9.3c.

The Ising contribution to the model in Eq. (9.20) introduces purely neighboring and equal-strength interactions between the Majorana fermions. Thus, an increasing  $\lambda$  gradually suppresses the disordered all-to-all interactions characteristic of the ergodic SYK<sub>4</sub> regime. The effect is reminiscent of the suppression of off-diagonal terms in the RP model in Eq. (9.11) considered above.

The symmetries of this model play an important role in understanding the behavior of complexity markers, as will be discussed in detail in the following section. Both the SYK<sub>4</sub> and Ising model Hamiltonian commute with the fermion parity operator

$$(-1)^F = (i)^M \psi_1 \dots \psi_M = (\sigma^x)^{\otimes M}. \quad (9.21)$$

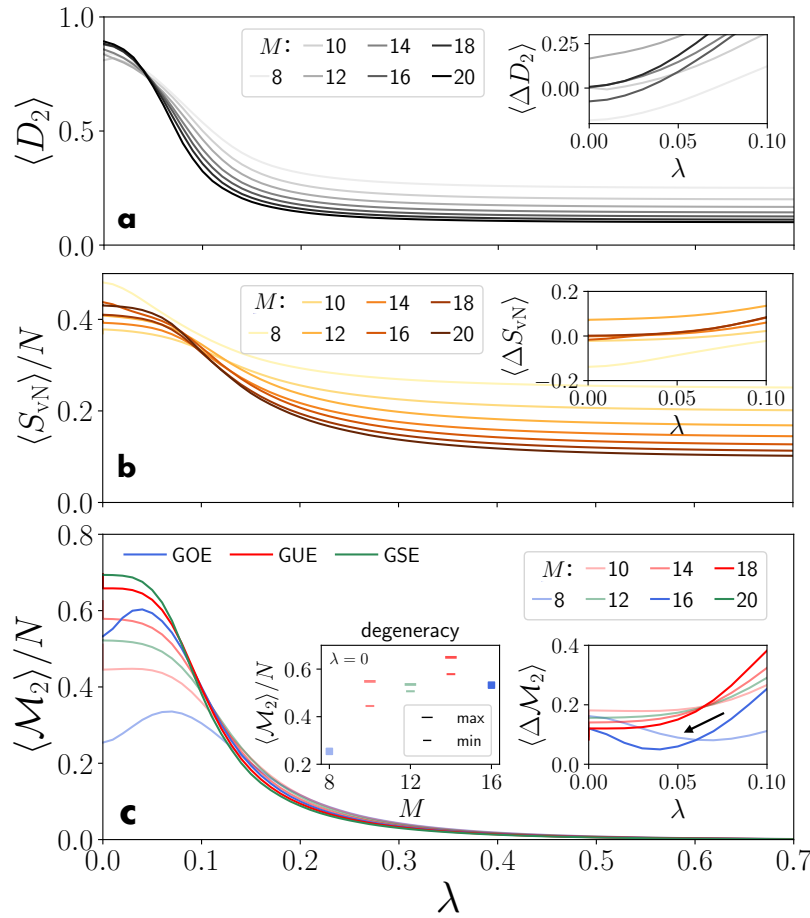
In addition,  $H_{\text{SYK}_4}$  commutes with the anti-unitary time reversal operator  $\mathcal{T}$ . The interplay between  $(-1)^F$  and  $\mathcal{T}$  determines the RMT ensemble and degeneracy of the ground state of the SYK model [762]: for  $M \bmod 4 = 2$  the ensemble corresponds to GUE, for  $M \bmod 8 = 0$  the corresponding ensemble is GOE, and for  $M \bmod 8 = 4$  it is GSE<sup>6</sup>. In contrast,  $H_{\text{Ising}}$  as given in Eq. (9.20) anticommutes with  $\mathcal{T}$  due to the presence of the imaginary unit in the Hamiltonian, lifting for  $\lambda \neq 0$  the symmetry class to GUE for all  $M$ .

<sup>6</sup>The Gaussian symplectic ensemble describes the random  $L \times L$  Hermitian quaternionic matrices. The numerical results were obtained by representing the GSE matrices as  $2L \times 2L$  complex matrices. For consistency, when comparing the SYK<sub>4</sub> results falling in the GSE symmetry class with the corresponding Haar-random value, we take an ensemble of dimension  $L^{N-1}$  such that the Hilbert space dimension of SYK<sub>4</sub> matches the dimension of the complex GSE representation.

### 9.5.2 Results

Similarly to the RP and PLRBM models, we carry out a comparative analysis of three complexity markers for the SYK<sub>4</sub> + Ising model. Additionally, we present a more detailed study of magic across different  $M$ , which turns out to be sensitive to the model's symmetries introduced above. To be consistent with the rest of the chapter, we consider the number of qubits  $N = M/2$  instead of the number of Majorana fermions when calculating the density of the markers.

In the integrable Ising model at  $\lambda = 1$ , the ground state is doubly degenerate. However, introducing a small perturbation from the SYK Hamiltonian,  $\lambda \lesssim 1$ , lifts this degeneracy. Due to the fermion parity symmetry of the full Hamiltonian, the system selects a specific ground state: an equal superposition of the two antiferromagnetic



**Figure 9.9.** Ensemble averaged complexity markers for the groundstate of the SYK<sub>4</sub> + Ising model for different number of Majorana fermions  $M$ . (a) The fractal dimension  $D_2$  starts to decrease for any non-zero  $\lambda$  and approaches the asymptotic value of the Ising model relatively quickly. Inset: Difference of  $D_2$  to the Haar-random value, indicating that for sufficiently large  $M$ ,  $D_2$  saturates this value at  $\lambda = 0$ . (b) The  $S_{vN}$ -density shows a small remnant of a plateau around the SYK<sub>4</sub> limit ( $\lambda = 0$ ). Inset: for large  $M$  also the  $S_{vN}$ -density approaches the Haar-random value at small  $\lambda$ . (c) For  $\lambda \gtrsim 0.5$ , the average magic remains zero, while it reaches its maximum near  $\lambda \sim 0$ , except for states corresponding to the GOE ensemble (which reflects the symmetry of the SYK model at  $\lambda = 0$ ), where the peak occurs away from zero. Left inset: At  $\lambda = 0$ , the ground states corresponding to the GUE and GSE ensembles are degenerate, where a range of SRE values is possible; however, as soon as  $\lambda$  deviates from zero, it chooses the minimum. Right inset: The difference to the Haar-random value is larger than for  $D_2$  and  $S_{vN}$ .



spin configurations, with a relative phase of  $\pm 1$ . As a result, in this limit, the fractal dimension and maximal von-Neumann entropy density approach the values of  $1/N$ . In contrast, the magic tends to zero since this superposition is a stabilizer state (see Section 9.5.3). All considered markers approach the values of the  $\lambda \sim 1$  limit relatively rapidly without appreciable changes beyond  $\lambda \gtrsim 0.5$ , see Fig. 9.9.

In the chaotic regime for  $\lambda \gtrsim 0$ , unlike in the RP ground state, we cannot a priori know if the complexity markers saturate the Haar-random values since we are now considering a many-body system. The deviation from the Haar-random value is quantified by

$$\langle \Delta \mathcal{C} \rangle = \frac{\langle \mathcal{C}_{\text{Haar}} \rangle}{N} - \frac{\langle \mathcal{C} \rangle}{N}, \quad (9.22)$$

where  $\mathcal{C}$  denotes one of the three markers  $D_2$ ,  $S_{\text{vN}}$ , or  $\mathcal{M}_2$ . As shown in the insets of Fig. 9.9, for sufficiently large  $M$ ,  $\langle \Delta D_2 \rangle$  and  $\langle \Delta S_{\text{vN}} \rangle$  tend to zero as  $\lambda \rightarrow 0$ , whereas  $\langle \Delta \mathcal{M}_2 \rangle$  is strictly larger than zero. This behavior can be understood by the fact that, similarly to what we have seen for  $\lambda \sim 1$ , the ground state for small  $\lambda$  has fixed fermion parity, which means that only Pauli strings composed of the product of an even number of Majorana operators (those that preserve the symmetry) contribute to a non-zero  $\mathcal{M}_2$ . Since these comprise only half of all possible Pauli strings, the overall  $\mathcal{M}_2$  is smaller than the Haar-random value. The situation is, however, a bit more subtle in the strict  $\lambda = 0$  case since in pure SYK<sub>4</sub> the ground state is not always unique. For  $M = 0 \bmod 8$  (GOE), there is just one ground state with fixed fermion parity. In contrast, for GUE and GSE,  $M \neq 0 \bmod 8$ , the presence of degeneracy allows for the superposition of the two ground states with opposite fermion parity, which can enhance the SRE. We numerically determine the maximum and minimum values of this linear combination<sup>7</sup>, as shown in the left inset of Fig. 9.9. Notably, the minima coincide with the values shown in the main plot (c) at small but non-zero  $\lambda$ , confirming that the Ising Hamiltonian breaks the degeneracy and selects one specific parity.

Moreover, the SRE exhibits a peculiar  $M$ -dependent behavior at small  $\lambda$ , which can be attributed to the characteristics of the underlying RMT ensembles [694, 763]. Specifically, when  $M \bmod 8 = 0$ ,  $\mathcal{M}_2$  begins at a lower value, and then rises to a peak at some small nonzero  $\lambda$ , followed by a monotonic decrease. This behavior can be comprehended in the following way: for finite-size systems, the GOE ensemble yields lower magic than GUE or GSE for Haar-random states (see Eq. (9.10) and Fig. 9.2 in Section 9.2.3). As contributions from the Ising Hamiltonian kick in at  $\lambda \neq 0$ , the symmetry class is lifted from GOE to GUE, leading to a rise of the value for the SRE in the case of  $M \bmod 8 = 0$ . In contrast, for  $M \bmod 4 = 2$  and  $M \bmod 8 = 4$ , the SRE starts at its maximal value, and the addition of the Ising contribution only decreases it.

As we move further away from  $\lambda \sim 0$ , all three markers start deviating from their ergodic value without exhibiting a long plateau as in the RP model—indicating that the ground state is rather susceptible to perturbations by the Ising contribution (see Fig. 9.9), unlike the  $\lambda \gtrsim 0.5$  regime. Such fragility of the SYK<sub>4</sub> ground state resembles

<sup>7</sup>Given the two ground states  $\text{GS}_1$  and  $\text{GS}_2$  with opposite fermion parity and a generic linear combination

$$|\psi\rangle = \sin \theta |\text{GS}_1\rangle + e^{i\phi} \cos \theta |\text{GS}_2\rangle, \quad (9.23)$$

we numerically optimize  $\langle \mathcal{M}_2(\psi) \rangle$  over a  $200 \times 200$  grid in the  $(\theta, \phi)$  plane to determine the maximum and minimum values.

recent observations in a related model interpolating between SYK<sub>4</sub> and SYK<sub>2</sub> [711], which implements a crossover from many-body chaos to single-body chaos, while always maintaining volume-law magic in the ground state. In our model, the perturbed Ising ground state is a stabilizer state; therefore, the transition in magic appears sharper, reminiscent of the transition observed in the RP ground state (Fig. 9.6).

We expect the ensemble-dependent features discussed above to disappear in the  $M \rightarrow \infty$  limit. Indeed, the  $M \bmod 8 = 0$  peak seems to progressively move to smaller values of  $\lambda$  and  $\Delta\mathcal{M}_2$  progressively decreases if we compare the same ensembles. Nonetheless, such finite-size effects are particularly relevant for characterizing realistic many-body chaotic systems beyond random matrix universality [757, 764]. The analysis highlights the utility of the SRE as a diagnostic tool due to its sensitivity to the model's symmetry structure, distinguishing it from other markers such as the fractal dimension and entanglement entropy.

### 9.5.3 Complexity markers for superposition of two computational basis states

It is instructive to examine the behavior of the three complexity markers for a simple superposition of two computational basis states. Consider the following  $N$ -qubit state:

$$|\psi\rangle = \sin\theta |a\rangle + e^{i\phi} \cos\theta |\bar{a}\rangle, \quad (9.24)$$

where  $|a\rangle$  and  $|\bar{a}\rangle$  are  $N$ -qubit orthogonal computational basis states differing by a Hamming distance of  $N$ , i.e., they are locally orthogonal at each qubit site. For such a state, the complexity markers are given by:

$$D_2 = -\frac{1}{N} \log_2 (\sin^4\theta + \cos^4\theta), \quad (9.25)$$

$$S_{vN} = -[\cos^2\theta \log_2 (\cos^2\theta) + \sin^2\theta \log_2 (\sin^2\theta)]. \quad (9.26)$$

The SRE is given by [765]:

$$\mathcal{M}_2 = -\log_2 \left( \frac{1 + \cos^4 2\theta + \sin^4 2\theta (\cos^4 \phi + \sin^4 \phi)}{2} \right). \quad (9.27)$$

Among the three markers, only the SRE is sensitive to the relative phase  $\phi$ . In the special case of an equal superposition with  $\pm 1$  difference, i.e.,  $\sin\theta = \cos\theta = \frac{1}{\sqrt{2}}$  and  $\phi = 0$ , or  $\pi$ , the complexity markers simplify to

$$D_2 = \frac{1}{N}, \quad \frac{S_{vN}}{N} = \frac{1}{N}, \quad \frac{\mathcal{M}_2}{N} = 0. \quad (9.28)$$

Such a superposition state appears in the  $\lambda \rightarrow 1$  limit of the SYK+Ising model. Due to the fermion parity symmetry of the full Hamiltonian, the ground state has double degeneracy and forms a superposition of two degenerate states connected by  $(-1)^F$ :

$$(-1)^F |\text{GS}_1\rangle = + |\text{GS}_2\rangle. \quad (9.29)$$

Moreover, the symmetry ensures that they form equal superpositions without any complex relative phase, as explained in the following way. Consider the generic su-



perposition

$$|\psi\rangle = \sin \theta |\text{GS}_1\rangle + e^{i\phi} \cos \theta |\text{GS}_2\rangle . \quad (9.30)$$

As  $|\psi\rangle$  should also be symmetric under  $(-1)^F$  with possible eigenvalues  $\pm 1$ ,

$$(-1)^F |\psi\rangle \stackrel{!}{=} \pm |\psi\rangle \Rightarrow \theta = \pi/4, \phi = 0, \pi. \quad (9.31)$$

Now, close to  $\lambda \rightarrow 1$ , as both ground states are computational basis states, we can use the above Eq. (9.28). That is, we expect fractal dimension and von-Neumann entropy density to scale as  $1/N$  and SRE to vanish, which is exactly what we observe in Fig. 9.9c.

## 9.6 Effects of samples

Analyzing disordered systems requires averaging over several random disorder instances to eliminate instance dependence, i.e., reduce the variance of results. However, it is often a model, its phase, and an observable dependent. For example, a highly ergodic phase with a self-averaging observable requires fewer samples compared to a localized phase to obtain the same order of variance. In this section, first, we discuss the number of samples we use for each model and complexity markers. Then we discuss how the self-averaging property of the complexity markers also contains the information of the phase transition.

### 9.6.1 Details on number of samples

For numerical reproducibility, we provide detailed tables listing the exact number of disorder realizations used for the results for different models (see Tables 9.1-9.3). To keep the numerical effort manageable under the exponential growth of the Hilbert-space dimension with system size  $N$ , we progressively reduced the number of realizations with increasing  $N$ . However, this reduction does not compromise the accuracy of our results, as the observables become increasingly self-averaging with larger  $N$  (see Fig. 9.10a1,b1,c1).

Moreover, we adapted the number of samples for the marker under consideration. The computational complexity for computing the maximum entanglement grows as  $\binom{N}{N/2} \times \text{construction}(\rho_A) \times \text{diag}(\rho_A) = \frac{2^{5/2N}}{\sqrt{N}}$ . Similarly, the complexity for computing the Stabilizer Rényi Entropy (SRE) scales as  $\mathcal{O}(2^{2N})$ , due to the size of the Pauli group. In comparison, the computational complexity for calculating the fractal dimension is more modest, scaling as  $\mathcal{O}(2^N)$ , permitting us to increase the number of samples to even further improve the quality of the results.

Among the considered models, we found the most substantial sample-to-sample fluctuations in the PLRBM model, making a larger number of realizations necessary to achieve satisfactory statistical accuracy. In contrast, due to the large density of states in the bulk, a comparatively smaller number of realizations was sufficient for the RP bulk model as compared to studies on its ground state. For the SYK+Ising model, we maintained a comparable sample size to that of the RP bulk case across all markers.

Model / N	4	5	6	7	8	9	10	11	12
RP bulk	20000	15000	10000	5000	4000	3000	2000	1000	500
RP gs	20000	15000	10000	5000	4000	3000	2000	1000	500
PLRBM bulk	20000	15000	10000	5000	4000	3000	2000	1000	500
SYK+Ising	5000	4000	3000	2000	1000	500	100	–	–

**Table 9.1.** Number of samples used for fractal dimension estimates across different models and system sizes  $N$ .

Model / N	4	5	6	7	8	9	10	11	12
RP bulk	5000	4000	3000	2000	1000	800	100	30	–
RP gs	20000	15000	10000	5000	4000	3000	500	100	50
PLRBM bulk	35000	24000	40000	16000	10000	3000	500	–	–
SYK+Ising	5000	4000	3000	2000	1000	500	100	–	–

**Table 9.2.** Number of samples used for maximum entanglement entropy estimates across different models and system sizes  $N$ .

Model / N	4	5	6	7	8	9	10	11	12
RP bulk	5000	4000	3000	2000	1000	500	80	–	–
RP gs	20000	15000	10000	5000	4000	1000	200	100	50
PLRBM bulk	35000	24000	20000	20000	10000	1500	–	–	–
SYK+Ising	5000	4000	3000	2000	1000	500	100	–	–

**Table 9.3.** Number of samples used for Stabilizer Rényi Entropy estimates across different models and system sizes  $N$ .

## 9.6.2 Self-averaging property

In disordered quantum many-body systems, the self-averaging property refers to the tendency of observables to converge to their ensemble-averaged (disorder-averaged) values as the system size increases. A physical quantity  $\mathcal{C}$  is said to be self-averaging if its relative variance vanishes in the thermodynamic limit. This can be quantified by the dimensionless ratio

$$R_{\mathcal{C}} = \frac{\text{Var}(\mathcal{C})}{\langle \mathcal{C} \rangle^2}, \quad (9.32)$$

where  $\langle \cdot \rangle$  denotes the disorder average. In ergodic phases, where the system uniformly explores its Hilbert space,  $R_{\mathcal{C}} \rightarrow 0$  as the system size increases, indicating strong self-averaging. In contrast, in non-ergodic or localized regimes,  $R_{\mathcal{C}}$  can remain finite even in the thermodynamic limit, reflecting strong sample-to-sample fluctuations and the breakdown of ergodicity.

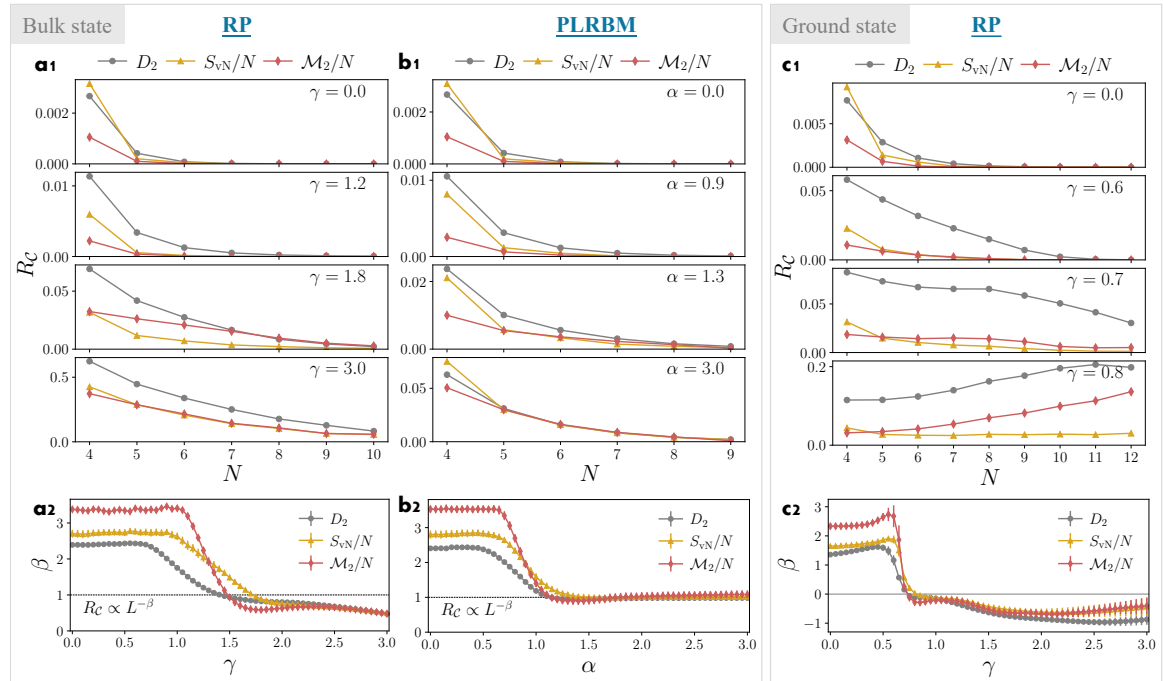
**RP bulk states.** In Fig. 9.10a1, we find that the complexity markers exhibit self-averaging behavior exclusively within the ergodic regime ( $\gamma \leq 1$ ). Beyond this point, deviations begin to emerge. Notably, at  $\gamma = 1.2$ , the fractal dimension  $D_2$  starts losing its self-averaging character due to its scaling as  $\mathcal{O}(N^\delta)$ , whereas both the normalized von Neumann entropy  $S_{\text{vN}}/N$  and the magic  $\mathcal{M}_2/N$ , which remain  $\mathcal{O}(1)$ , continue

to exhibit self-averaging. At stronger disorder strengths, such as  $\gamma = 1.8$ , the entanglement remains more strongly self-averaging, consistent with a larger plateau in Fig. 9.4b1 of the  $\mathcal{O}(1)$  regime. At  $\gamma = 3$ , none of the markers remain self-averaging, in accordance with their  $\mathcal{O}(N^\delta)$  and  $\mathcal{O}(1/N)$  scaling behaviors, respectively.

Furthermore, the strength of self-averaging can be inferred from the scaling behavior of  $R_C$  with system size  $L = 2^N$ , i.e.,

$$R_C \propto L^{-\beta} = 2^{-N\beta}. \quad (9.33)$$

A system is *strongly self-averaging* if  $\beta \geq 1$ , and *weakly self-averaging* if  $0 < \beta < 1$ . Notably, Fig. 9.10a2 shows that up to  $\gamma = 1.5$ , even in the non-ergodic phase, the system remains strongly self-averaging. Beyond  $\gamma = 2.5$ , all markers exhibit the same exponents. Additionally,  $\mathcal{M}$  demonstrates stronger self-averaging in the ergodic regime and undergoes the most pronounced change around the transition points.



**Figure 9.10.** Self-averaging behavior of different complexity markers and their scaling exponents. **(a1)** For the bulk of the RP model, in the ergodic regime ( $\gamma = 0$ ), all markers exhibit strong self-averaging, indicated by a vanishing relative variance. At intermediate disorder ( $\gamma = 1.2$ ),  $D_2$  already begins to deviate, showing non-self-averaging behavior, while others remain stable. At  $\gamma = 1.8$  and  $\gamma = 3$ , none of the markers are self-averaging, consistent with the breakdown of ergodicity and the emergence of strong sample-to-sample fluctuations. **(a2)** Scaling exponent  $\beta$  as a function of  $\gamma$  for RP bulk states. All three markers exhibit strong self-averaging ( $\beta > 1$ ) up to  $\gamma = 1.5$ , and after  $\gamma = 2.5$ , they show same  $\beta$ . Notably,  $\mathcal{M}_2$  displays the steepest variation in  $\beta$ , indicating a more pronounced change in self-averaging across the transition. The deviation from the plateau region of constant  $\beta$  occurs in the same order, i.e.,  $D_2$  then  $S_{vN}$  and  $\mathcal{M}_2$  as observed in Fig. 9.4. **(b1)** In the bulk states of the PLRBM model, complexity markers exhibit behavior analogous to the RP model, with magic showing the highest degree of self-averaging, followed by entanglement entropy, and fractal dimension displaying the least self-averaging in the ergodic phase. **(b2)** The scaling exponent of all the markers becomes weakly self-averaging ( $\beta \sim 1$ ) close to  $\alpha \sim 1$ , and the  $\alpha > 1$  regime is comparatively more self-averaging than the bulk states of the RP model. **(c1)** In the ground state of the RP model, all the markers stop exhibiting the self-averaging behavior after crossing the phase transition point  $\gamma^\infty \sim 0.75$ , leading to  $\beta < 0$  as seen in **(c2)**.

**RP ground states.** A key distinction between the bulk and ground state phases is the absence of a fractal regime in the latter, where only ergodic and localized phases are detected via complexity markers. Notably, the ground state exhibits a first-order phase transition, raising the question of whether self-averaging behaves differently compared to the bulk. As shown in Fig. 9.10c1, self-averaging holds at  $\gamma = 0$ , but breaks down beyond the transition point at  $\gamma = 0.8$ , the marker even increases with  $N$ . This behavior is reminiscent of the vanishing marker values at large  $\gamma$  (see Fig. 9.6).

**PLRBM bulk states.** Similar to the RP bulk states, the PLRBM bulk state also shows the strongly self-averaging behavior, and near the Anderson critical point, at  $\alpha = 1$ , all the complexity markers still remain strongly self-averaging, i.e.,  $\beta = 1$  (see Fig. 9.10b2), which is a behavior quite different from that in the RP bulk states.

Since the scaling behavior of the self-averaging property in complexity markers reflects the ergodicity transitions observed in the markers themselves, one might speculate that in disordered systems where fluctuations play a crucial role, the self-averaging property could serve as a diagnostic tool for phase transitions.

## 9.7 Pauli spectrum for RP and PLRBM bulk states

The stabilizer Rényi entropy is defined as the Rényi entropy of a probability distribution constructed from the expectation values of Pauli strings on a given quantum state. Since a probability distribution inherently contains more information than any of its finite-order entropies, additional structure—particularly relevant in complex phases—is revealed by examining higher-order moments  $\mathcal{M}_q$ . Moreover, non-ergodic behavior is often captured by higher-order entropies, such as the fractal dimension  $D_q$ .

It is therefore worth investigating if relevant signatures can emerge from the *Pauli spectrum*, defined as the set of squared Pauli expectation values on the state [168, 766]:

$$\text{spec}(|\psi\rangle) = \left\{ |\langle\psi|P_i|\psi\rangle|^2, P_i \in \mathcal{P}_N; i = 0, 1, \dots, 4^N - 1 \right\}. \quad (9.34)$$

The support of the Pauli spectrum,  $i \in [0, 4^N - 1]$ , changes depending on  $N$ . The *frequency distribution* of these expectation values,

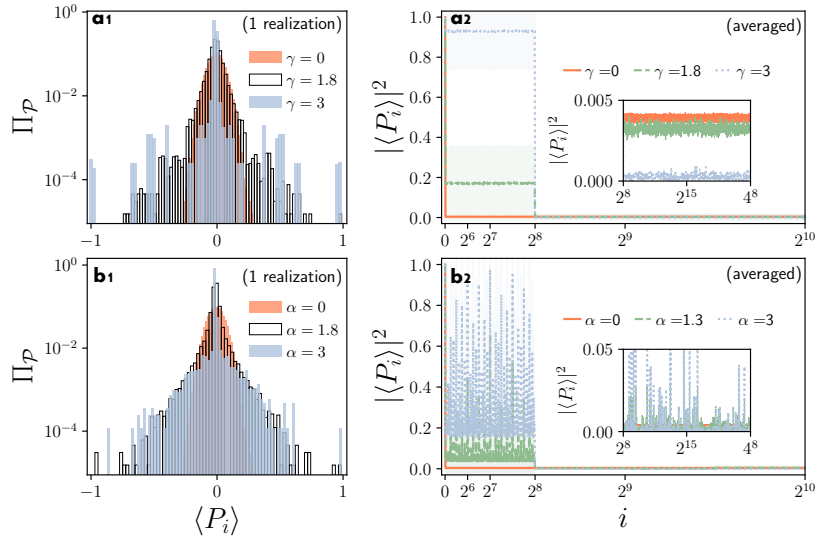
$$\Pi_{\mathcal{P}}(|\psi\rangle) = \sum_i \frac{\delta(x - x_{P_i})}{4^N}, \quad x_{P_i} = \langle\psi|P_i|\psi\rangle, \quad (9.35)$$

offers a diagnostic tool for understanding the ergodicity behavior independent of  $N$ . However, individual realizations would have different  $\Pi_{\mathcal{P}}$ , which can average away relevant features. Instead, we focus on the averaged Pauli spectrum, where the expectation value for each  $P_i$  is averaged over states from each realization. To better interpret its structure, we organize the Pauli strings by indexing  $i$  in a specific order: the first  $2^N$  indices  $i \in [0, 2^N - 1]$  correspond to diagonal operators  $\{I, Z\}^{\otimes N}$ , followed by all other (non-diagonal) strings. The overall ordering is then refined by sorting the values of  $\langle P_i \rangle$  in descending order, using the localized state at  $\alpha = 10$  or  $\gamma = 10$  as a reference. This choice is motivated by the fact that localized states exhibit significant weight on the diagonal strings (see Fig. 9.11 a2, b2), and thus provide a natural order.

Ergodic states follow the Haar-random prediction (see Section 9.2.3) and exhibit an almost uniform distribution over Pauli strings around the value  $\frac{1}{2^N+1}$ , except for the identity component  $I^{\otimes N}$ , which saturates to 1, resulting in  $\Pi_{\mathcal{P}}$  sharply peaked around zero (orange bars in Fig. 9.11 **a1**, **b1**) and low variance in the frequency distribution (orange shaded region in **a2**, **b2**). In contrast, localized states have only a few significant Pauli components (**a2**, **b2**), leading to larger individual  $\langle P_i \rangle$  values and a broadened spectrum (blue bars in **a1**, **b1**).

Notably, exact stabilizer states yield Pauli expectation values of either +1 for all  $2^N$  stabilizers, or a balanced set of +1 and -1 values [168], as can be observed in the RP bulk case (blue lines in **a2**). In the PLRBM model, however, bulk localized states do not align with the computational basis  $\{I, Z\}^{\otimes N}$ , resulting in a more dispersed Pauli spectrum (**b1**) compared to the RP model (**a1**). Even after averaging over random realizations, the PLRBM bulk state retains high variance in its Pauli spectrum across the non-ergodic and localized regimes, except in the ergodic phase where the distribution sharply concentrates [767].

In summary, the Pauli spectrum offers insight into why an ergodic state, which contains high magic, requires a larger number of basis states and is thus considered more complex, while a localized state, with non-zero contributing terms limited to a few Pauli strings, is considered less complex (see Fig. 9.1b). Furthermore, the difference between the RP and PLRBM models is clearly captured in the Pauli spectrum, where the less suppressed off-diagonal terms of the PLRBM model yield strong fluctuations even in the localized regime, thus providing a direct interpretation of the relatively high magic in the localized regime of PLRBM bulk states (see Fig. 9.8c1), compared to its RP counterpart.



**Figure 9.11.** Frequency distribution for a single realization (left) and averaged Pauli-spectrum for a bulk state in the middle of the spectrum (right) of the RP (top) and PLRBM models (bottom). Data for  $N = 8$  with 4000 and 8000 realizations, respectively. In the ergodic phase (orange), expectation values are nearly uniformly distributed over all Pauli strings with minimal variance (shaded regions in **a2**, **b2**), resulting in a distribution with small variance around  $\langle P_i \rangle = 0$  in  $\Pi_{\mathcal{P}}$  (**a1**, **b1**). In contrast, localized states (blue) exhibit pronounced peaks in  $\Pi_{\mathcal{P}}$ , reflecting non-zero expectation values aligned with the stabilizer basis (notably within  $i = [0, 2^N]$  in **a2**, **b2**). The intermediate regime shows partial delocalization with mixed features. These distinct distribution patterns are reflected in the magic values. Compared to RP, localization in PLRBM bulk states is weaker (**b1**) and exhibits reduced self-averaging behavior (**b2**).

## 9.8 Summary and outlook

In this work, we have conducted a comparative study of various complexity markers across families of Hamiltonians that interpolate between ergodic, extended non-ergodic, and localized phases. When a fractal region in the parameter space is present, such as for states in the bulk of the Rosenzweig–Porter model spectrum, the different markers identify high and low complexity regimes in overlapping but distinct parameter regions. Both entanglement entropy and stabilizer Rényi entropy undergo a transition within the fractal phase that is not detected by the fractal dimension, indicating the existence of a non-ergodic yet complex regime with respect to these two markers. Moreover, the stabilizer Rényi entropy reveals an additional extended intermediate regime with non-maximal magic, before eventually dropping to low complexity in correspondence to the localization point. The presence of an extended fractal region seems to be crucial to the appearance of new complexity regimes: when such a region is absent, as in the ground state of the Rosenzweig–Porter model and the bulk of the power-law random banded model, the transition points identified by all complexity markers remain consistent. The behavior of the stabilizer Rényi entropy suggests that it can uncover subtle features in the quantum phase diagram that might otherwise go unnoticed using conventional localization markers. It would be interesting to corroborate these findings through analytical studies, e.g., of the Rosenzweig–Porter model akin to those employed for fractal dimensions [689].

Moreover, our analysis of the ergodic to integrable transition of the  $\text{SYK}_4$  + Ising model suggests that stabilizer Rényi entropy may be a more sensitive marker to the underlying symmetries of the model, as indicated by its non-monotonic behavior as compared to the fractal dimension and the entanglement entropy. This observation motivates a more systematic investigation of the relationship between nonstabilizer-ness and symmetry classes in SYK-type models, extending beyond the cases considered in Refs. [762, 763]. Additionally, it would be valuable to study other quadratic perturbations of the SYK model to determine whether the complexity markers studied here can consistently capture the observed fragility of ground-state ergodicity. This is especially relevant for assessing the stability of the holographic phase, which has been argued to remain robust under deformations only within a vanishingly small region in the thermodynamic limit [768].

Investigating the presence of non-ergodic but highly complex regimes would not only shed light on the structure of the quantum phase diagram but also help assess the sensitivity of non-stabilizer magic to non-ergodic behavior and complexity transitions. This could be further pursued by analyzing many-body systems undergoing a many-body localization transition, where multifractal phases are expected to arise, such as disordered spin systems [769, 770] and the Bose–Hubbard model [748, 771]. Localization also plays a significant role in quantum optimization [243, 570, 571, 574, 772], whose target problems are often described by spin-glass models [236, 529, 538]. In these contexts, our work offers a promising perspective for probing complex characteristics of physical systems approaching the many-body localization transition.

The presence of distinct crossover values at finite system sizes is crucial for both classical and quantum simulations, as the feasible techniques are often constrained by available quantum resources. For instance, in noisy intermediate-scale quantum (NISQ) devices, the bottleneck is the ability to build up entanglement across the device [773], while in tensor networks, the reachable bond dimension constrains the

accessible entanglement entropy [91, 542, 673, 674]. In contrast, the efficiency of classical calculations using the Gottesmann–Knill framework [148, 149, 151, 158, 166] as well as the cost of broad classes of fault-tolerant quantum computers [774, 775] is determined by the number of  $T$ -gates [167, 545], which is related to the magic in a quantum state [165]. In this context, our analysis could guide the identification of the most efficient simulation strategies for each regime.





## Chapter 10

# Conclusions and Outlook

In this thesis, we demonstrated how various quantum resources enhance our understanding of quantum optimization and simulation, providing critical insights into the performance and capabilities of quantum technologies. After introducing the concepts of quantum resources and quantum optimization in Part I, we dedicated Part II entirely to quantum optimization and Part III to disordered systems.

We explored the quantumness of Quantum Approximate Optimization Algorithm (QAOA) and Quantum Annealing (QA) through squeezing, multipartite entanglement, and nonstabilizerness (magic), as well as experimentally measured entanglement on an IBM quantum device. We shed light on the role of quantum correlation in QAOA by connecting two fields of quantum technology: quantum computation and quantum metrology. The goal of metrology is to establish measurement protocols with increased sensitivity, and quantum computation aims to obtain the solution state with reduced uncertainty, both requiring a reduction in variance. By expressing the combinatorial MaxCut problem in terms of a collective spin basis, we are able to generate spin-squeezed states by finding optimal solutions to the MaxCut problem. We showed this both numerically and on the IBM Quantum chip. Then we connected this squeezing to quantum Fisher information and multipartite entanglement. Furthermore, squeezing generated by QAOA enabled us to propose a benchmark, where more squeezing is associated with a higher probability of the solution state, conventionally referred to as the approximation ratio of the quantum algorithm. Moreover, such squeezing can also detect errors in the circuit. We also extended this connection between the two fields to their benefits: using QAOA to generate arbitrary Dicke states, and using squeezed states to warm-start QAOA.

To understand the presence of multipartite entanglement in quantum optimization, we utilized the generalized geometric measure, a measure of genuine multipartite entanglement, in quantum annealing and QAOA. Using numerical benchmarks, we analyzed its occurrence in the annealing schedule in detail. We observed that entanglement increases with anneal time, and for successful instances, it decreases again, creating a multipartite entanglement barrier. We explored how such a barrier relates to the algorithm's success. We also proved how multipartite entanglement provides an upper bound to the overlap of the instantaneous state with an exact solution. Vice versa, the overlaps to the initial and final product states, which can be easily measured experimentally, offer upper bounds for the multipartite entanglement during the entire schedule.

Noisy devices raise the question of whether entanglement persists with longer circuit depth, as its absence would render the algorithm's success entirely dependent on classical algorithms. To validate such quantum effects in real devices, we chose

the new Heron processor from IBM Quantum to test entanglement generation in the QAOA layers, using the SWAP circuit. We first measured the purity of the subsystems, then extracted the maximum Schmidt coefficients, which served as a proxy for entanglement. We found that such entanglement persists even for large systems and significant depth, showing on the one hand that quantum effects play a role in the QAOA, as well as that current hardware is mature enough to run quantum optimization.

We also analyzed the presence of nonstabilizerness (a proxy for classical simulability) in QAOA. We found that both stabilizer Rényi entropy (SRE) and mana initially grow with circuit depth, peaking and then decreasing as the algorithm converges, creating a barrier that limits the algorithm’s capability for shallow circuits. We found curves corresponding to different depths collapse under a simple rescaling, and we revealed a non-trivial relationship between the final SRE and the fidelity of final states. Finally, we identified a similar nonstabilizerness barrier also in quantum annealing.

In Part III, we turned to disordered systems. Since disorder mimics noise, understanding its effects helps model noisy devices. Averaging over disorder is computationally costly, so we developed an exact framework for disorder-averaged dynamics generated by periodic Hamiltonians (including (p, q)-potent ones). These results are valid for arbitrary initial states, evolution times, and even non-Hermitian systems. The ensemble dynamics resemble open-system evolution, with non-Markovian features depending on the disorder distribution and Hamiltonian periodicity. We illustrated this for systems built from spin-1/2, spin-1, and clock operators.

We also investigated complexity transitions in paradigmatic models such as the power-law random banded matrix model, the Rosenzweig–Porter model, and a hybrid SYK+Ising model. Using fractal dimension, von Neumann entropy, and SRE as markers, we identified sharp complexity transitions through finite-size scaling. These markers, while aligned in ergodic and localized regimes, diverged in intermediate phases. SRE in particular was highly sensitive to symmetries like fermion parity and time reversal. Our results show that no single marker suffices: combining different resources offers a better understanding of phase behavior and classical simulability.

Throughout this thesis, we explored various aspects of quantum resources and their exploitation in quantum optimization and many-body physics scenarios. Such quantum resources define what is easy and hard, given the free and resourceful operations within the resource theory. In quantum optimization, we observed that squeezing, multipartite entanglement, and nonstabilizerness all grow with the quantum run of the algorithm, validating that quantum algorithms are indeed quantum. Moreover, when asked for classical optimization problems, one is looking for a classical solution. In that case, all such resources must be reduced to the classical level, creating a resource barrier. The existence of such barriers makes the resource necessary for quantum algorithms, and understanding them is essential, as success is closely related to it. In many-body physics, such peaks can be related to phase transitions or crossovers, and it becomes interesting to understand if these barriers or transitions appear at the same transition points for different resources. As we discovered, this is not the case, suggesting that there may be multiple phases that require combining different resources for a comprehensive understanding. Our results provided more profound insights into how quantum resources influence quantum optimization, enabling a better understanding of the success of such algorithms and the recognition of new phases behind the wall of combined complexity markers arising from resource theories

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in many-body physics.

We now turn to a discussion of possible future directions and outlooks based on the findings presented in this thesis. Since we had obtained squeezing for eight-qubit QAOA on IBM hardware, the quantum hardware architectures have evolved into over 100 qubits with substantially improved gate fidelities [460, 776]. Moreover, platforms based on trapped ions and neutral atoms have become increasingly accessible, making it feasible to perform cross-platform [777], large-scale benchmarks using squeezing as a diagnostic tool. Such an approach could potentially realize squeezed states with unprecedented precision [332]. This would not only validate the quantum nature of QAOA but also help identify the breakdown points of classical tensor-network simulations, offering a compelling case for quantum computational advantage. In parallel, growing attention has been devoted to understanding the behavior of quantum annealers [778] such as those developed by D-Wave [779, 780]. Exploring how different practical annealing schedules influence entanglement generation, and whether this entanglement correlates with the success of those schedules, could offer valuable insights into the behavior of quantum devices. While our study of nonstabilizerness in QAOA was limited to small system sizes due to computational overhead, recent advancements in tensor-network simulations of QAOA [265, 544] now open the door to probing larger system sizes. This would allow us to test the universality of the magic barrier observed across scales. Additionally, since SRE is, in principle, experimentally measurable [530], there is a clear opportunity to extend our entanglement-measurement pipeline to include nonstabilizerness; thereby concluding the empirical investigation of quantum resources explored in this thesis. A particularly intriguing avenue concerns the relationship between squeezing and non-classicality, specifically through the appearance of negative Wigner functions [781], which are also directly related to mana, a magic monotone. This raises the fundamental question of whether entanglement and magic are generated in a correlated manner within the QAOA framework. While several counterexamples suggest these resources can vary independently in other scenarios [522, 782, 783], if QAOA does exhibit correlated resource generation, it would represent a distinctive and possibly unique feature of the QAOA ansatz. This warrants deeper exploration, particularly in light of the discrete nature of the Clifford group and the challenge of designing QAOA evolutions that avoid magic generation entirely, allowing simulation through stabilizer formalism [151].

In the context of disorder-averaged dynamics, our focus was on exact analytical results, which led us to consider a specific type of Hamiltonian. Extending such analysis to other Hamiltonian classes, especially non-Hermitian systems [784], presents an exciting theoretical direction. In such regimes, fundamental questions arise: What constitutes a quantum resource under non-Hermitian dynamics? And do traditional criteria for non-Markovianity still apply? Furthermore, it is worth exploring whether experimental observation of non-Markovian features can facilitate a modelling method for the underlying disorder, which would be of interest for quantum control theory [785]. Our resource-based complexity analysis in random matrix models, encompassing entanglement, nonstabilizerness, and fractal dimensions, provides a comprehensive view of quantum complexity. Extending this multi-marker framework to other many-body systems and quantum optimization problems could uncover hidden structures in the non-ergodic regime, potentially leading to new classifications of quantum phases and a deeper understanding of quantum information structure in complex systems.



# List of Publications

## Peer-reviewed articles.

- **G. C. Santra**, F. Jendrzejewski, P. Hauke, D.J. Egger (2024). Squeezing and quantum approximate optimization.  
*Physical Review A*, 109(1), 012413 (**Editor's suggestion**) [[325](#)].  
Chapter [4](#)  
Contribution: This project explores the connection between quantum squeezing and the Quantum Approximate Optimization Algorithm (QAOA), and uses this link to study the entanglement structure within QAOA. My contributions included performing numerical simulations of QAOA and squeezing, calculating the quantum Fisher information and multipartite entanglement, identifying the connection between squeezing and the proposed entanglement benchmark, and extending the analysis to arbitrary Dicke states. I also worked on implementing a warm-start approach for QAOA, contributed to the interpretation of the results, and participated in writing the manuscript.
- **G. C. Santra**, S.S. Roy, D.J. Egger, P. Hauke (2025). Genuine multipartite entanglement in quantum optimization.  
*Physical Review A*, 111(2), 022434. [[503](#)]  
Chapter [5](#)  
Contribution: This project investigates the role of genuine multipartite entanglement in quantum optimization algorithms and its connection to algorithmic performance. My contributions included writing numerical code for entanglement measures and quantum annealing, generating and analyzing data, interpreting the results, and providing analytical proofs for the entanglement bounds. I also contributed to the project design and manuscript writing.

## Preprints.

- **G. C. Santra**, P. Hauke (2024). Disorder-averaged Qudit Dynamics.  
*arXiv preprint* arXiv:2412.17519. [[786](#)]  
Chapter [8](#)  
Contribution: This project aims to understand how open quantum system dynamics can emerge from disorder-averaged closed quantum systems. My contributions included deriving exact analytical calculations, executing numerical validations, and carrying out detailed calculations for all illustrative examples. I also contributed to the project design and manuscript writing.
- **G. C. Santra**, A. Windey, S. Bandyopadhyay, A. Legramandi, P. Hauke (2025). Complexity transitions in chaotic quantum systems.

*arXiv preprint* arXiv:2505.09707. [522]

Chapter 9

Contribution: This project investigates various complexity markers in disordered many-body quantum systems. My contributions included developing the numerical code for all complexity markers and random matrix models, running simulations for the Rosenzweig–Porter model in both the bulk and ground-state regimes, and preparing all figures for the manuscript. I also contributed to the project design and manuscript writing.

- C. Capecci, **G.C. Santra**, A. Bottarelli, E. Tirrito, P. Hauke (2025). Role of Nonstabilizerness in Quantum Optimization.

*arXiv preprint* arXiv:2505.17185. [765]

Chapter 7

Contribution. This project explores the role of nonstabilizerness in the Quantum Approximate Optimization Algorithm (QAOA). Together with the first author, I contributed to the development of the numerical code for evaluating nonstabilizerness markers in qudit systems. I contributed to the analytical calculations used to interpret the data, project design, and wrote parts of the manuscript.

### Under preparation

- **G.C. Santra**, P. Hauke, D.J. Egger (2025). Validating quantum effects in quantum optimization

*to be arXived*

Chapter 6

Contribution. This project focuses on identifying and validating multipartite entanglement in quantum optimization algorithms on IBM hardware. My contributions included setting up the numerical framework for the experiments, gathering experimental data, conducting exact and noisy simulations, and analyzing data. I also contributed to the project design, interpretation of the results, and the writing of the manuscript.

### Others (not included in the thesis)

- A.N. Madhusudan<sup>1</sup>, **G.C. Santra**<sup>1</sup>, I. Kaur, W. Li, R. Nath. (2024). Commensurate supersolids and re-entrant transitions in an extended Bose-Hubbard ladder.

*arXiv preprint* arXiv:2407.20107. [787]

- **G.C. Santra**, E. Ballini, A. Bottarelli, J. Mildenerberger, M.M. Wauters, P. Hauke. Quantum Resource in Non-Abelian Lattice Gauge Theory

*under preparation*

- **G.C. Santra**, J. Mildenerberger, P. Hauke, R.C. Almeida. Stabilizer Fisher Information

*under preparation*

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<sup>1</sup>These authors contributed equally.

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