

Quantum Science and Technology Master's thesis

Quantum tensor networks for sampling

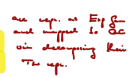
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Abstract

Efficient sampling from complex probability distributions is a cornerstone of modern machine learning, enabling critical tasks such as probabilistic inference, generative modeling, and optimization. Classical sampling methods, including Markov Chain Monte Carlo and rejection sampling, face prohibitive computational challenges as the dimensionality and logical complexity of target distributions grow. This thesis investigates the potential of quantum computing—specifically amplitude amplification algorithms—to overcome these limitations and deliver practical advantages for machine learning sampling tasks. Through a systematic methodology, logical contensor networks, which are then encoded into quantum circuits. Simulations on up to 25 qubits demonstrate the scalability of quantum amplitude amplification, while comparative analyses with classical baselines quantify speedup and robustness un-The results reveal that quantum-assisted sampling achieves quadratic speedups for structured distributions, though hardware constraints and noise pose significant challenges. This work bridges quantum computing and machine learning by formalizing a framework for integrating quantum sampling into classical workflows, offering a pathway toward scalable probabilistic inference in AI systems.



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Chapter 1

Introduction

Motivation and Background

Sampling is an essential part of a machine learning algorithm. Whenever the program makes a decision, handles predictions and uncertainty or gives solutions to a problem, it is a form of sampling dear from a vast set of information based on past data and / or knowledge. Sampling therefore is fundamental in enabling probabilistic inference, uncertainty quantification, and optimization in high-dimensional spaces. As models are increasingly sophisticated — incorporating rich logical structures or large graphical dependencies — their complexity escalates and the computational demands of sampling algorithms grow exponentially. Classical methods, while foundational, struggle to balance accuracy and efficiency in high-dimensional or multimodal distributions. Quantum computing, with its inherent parallelism and interference capabilities suggest that quantum algorithms could offer speedups for certain sampling tasks, motivating a systematic investigation of their applicability and impact for machine learning [1][2]. This thesis explores the intersection of quantum computing and machine learning, focusing on how quantum amplitude amplification can revolutionize sampling tasks in artificial intelligence. The following sections elaborate on the motivation, challenges, opportunities, objectives, and structure of this work.

Classical Sampling in Machine Learning: Challenges

Classical sampling algorithms, such as Markov Chain Monte Carlo (MCMC) and rejection sampling, face significant barriers in contemporary machine learning applications. High-dimensional distributions with complex dependencies or sharp peaks result in slow mixing times, autocorrelation, and exponential resource scaling. For example, sampling from Bayesian networks with hundreds of variables or training deep generative models with multimodal posteriors often becomes computationally intractable. These challenges are exacerbated by the "curse of dimensionality," where the volume of the sampling space grows exponentially with the number of variables. Even state-of-the-art methods like Hamiltonian Monte Carlo or variational inference

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require trade-offs between approximation accuracy and computational cost, leaving room for fundamentally new computational paradigms.

Quantum Computing: Opportunities for Sampling

Quantum computing introduces novel strategies for sampling through principles such as superposition or entanglement, enabling new computational paradigms for sampling. Recent research demonstrates that quantum algorithms can sample from complex distributions with fewer resources than classical counterparts, particularly for problems with combinatorial structure or where classical simulation is intractable or even computationally prohibitive. Quantum amplitude amplification, a generalization of Grover's search algorithm [3] enables quadratic speedups in identifying "good" solution states within unstructured search spaces. Thus by encoding probability distributions into quantum states, this technique can efficiently sample from distributions that would be unfeasible for classical systems. Also, recent advances in quantum hardware, such as improved gate fidelities and coherence times suggest that near-term devices may soon support practical implementations. However, practical deployment depends on circuit depth, noise resilience, hardware capabilities and seamless integration with classical machine learning workflows, posing as critical challenges for quantum procedures to effectively overcome classical ones [4]

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Thesis Objectives and Contributions

This thesis aims to bridge the gap between theory and practice in quantum-assisted sampling for machine learning, through the following contributions:

- A systematic methodology for mapping logical connectives and graphical models to tensor networks formalization of exponential family distributions, enabling their representation as quantum circuits.
- A framework for integrating quantum sampling into the TNREASON library, enabling hybrid quantum-classical inference for nested probabilistic models.
- Implementation and simulation of amplification-based quantum sampling routines, analyze their theoretical speedup over classical rejection sampling in structured distributions, and assess their empirical performance on real quantum hardware.
- Compare quantum sampling with classical baselines (rejection sampling, MCMC) across synthetic and structured datasets, quantifying speed, accuracy, and scalability.

Resource estimation studies for near-term quantum hardware, providing actionable insights for algorithm deployment on superconducting and trapped-ion platforms.

Thesis Structure

The remainder of this work is organized as follows:

- Chapter 2 reviews foundational concepts in logical connectives, graphical models, and exponential familiy distributions, with their tensor network representations.
- Chapter 3 introduces quantum computing, defining quantum gates and measurements, then turning to quantum amplitude amplification.
- Chapter 4 destribes my implementation of mapping a one dimensional logical formula to a quantum circuit, and performing the previously defined quantum measurement and amplitude amplification , to classical baselines.
- Chapter 5 benchmarks quantum sampling against classical methods, with analyzing scalability, and comparing theoretical speedups, and discusses practical deployment constraints.
- Chapter 6 explores broader implications, limitations, and future research directions, finally concludes with a synthesis of contributions and an outlook on quantum sampling in AI.

Chapter 2

Foundations

2.1 Logical Connectives and Probabilistic Models

Artificial Intelligence (AI) and Machine Learning (ML) inherently involve sampling problems during training, learning, and optimization. Data and knowledge can be represented through diverse models, each with distinct advantages. Probabilistic graphical models—such as Bayesian networks and Markov Random Fields—have become dominant in real-world applications due to their natural handling of uncertainty and explicit representation of variable dependencies via graph structures. Conversely, in the still well used logical approaches data is naturally encoded as logical propositions with inference traditionally retrieving deterministic conclusions, though modern extensions integrate probabilistic interpretations and sampling. With extending the practical usage of logics, the field of Statistical Relational AI bridges this gap, unifying logical relations with statistical models to handle uncertainty systematically. This synthesis, often termed neuro-symbolic AI [5], leverages the structured reasoning of logic and the uncertainty modeling of graphical models. For instance, logical rules can be embedded as soft constraints in probabilistic frameworks, enabling hybrid inference that balances symbolic precision with statistical robustness.

Both probabilistic and logical systems assume a factored structure, where system states are assignments to a set of variables. This structure admits a natural tensor representation, where variables correspond to tensor indices and their assignments to tensor entries. For example, a probability distribution over binary variables X_1, \ldots, X_n can be encoded as a rank-n tensor $\mathcal{T}[X_1, \ldots, X_n]$, with each entry storing the probability of a specific joint assignment. Similarly, logical formulas map to tensors via one-hot encodings, where entries indicate formula satisfaction (1, TRUE) or violation (0, FALSE).

Tensor methods provide a unifying formalism for reasoning algorithms across probabilistic and logical frameworks. Marginalization in Bayesian networks corresponds to tensor contraction, while logical inference reduces to multilinear operations on formula-encoded tensors. This shared mathematical foundation enables seamless in-

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tegration of probabilistic graphical models and symbolic logic, paving the way for scalable quantum sampling algorithms that exploit tensor network decompositions.

2.1.1 Propositional Logic in Al

Propositional logic is a fundamental tool in artificial intelligence for representing and manipulating knowledge in a formal, symbolic manner. In this framework, knowledge is encoded as a set of atomic propositions, each of which can be either true or false. These atomic propositions can be combined using logical connectives such as AND (\land) , OR (\lor) , NOT (\neg) , and IMPLIES (\rightarrow) to form more complex statements or formulas.

Logical reasoning in AI often involves determining the satisfiability of a set of formulas, performing inference (e.g., deducing new facts from known ones), and checking entailment between statements. For example, given a knowledge base consisting of rules and facts, an AI system can use logical inference mechanisms such as Modus Ponens or Resolution to derive new conclusions. This symbolic approach underpins many classical AI systems, including expert systems, rule-based engines, and automated theorem provers.

Importantly, logical connectives can be mapped to indicator functions or binary variables, which facilitates their integration into probabilistic models. For instance, the truth value of a conjunction of variables can be represented as the product of their indicator variables. This mapping provides a bridge between symbolic logic and statistical modeling, enabling hybrid approaches that combine the strengths of both paradigms.

2.1.2 Graphical Models: Bayesian and Markov Networks

Probabilistic graphical models (PGMs) provide a powerful framework for representing the conditional dependencies among random variables in complex systems. There are two principal types of graphical models: Bayesian networks and Markov random fields (also known as Markov networks).

Bayesian networks are directed acyclic graphs (DAGs) where each node corresponds to a random variable, and directed edges encode conditional dependencies. The joint probability distribution factorizes according to the graph structure:

$$P(X_1, ..., X_n) = \prod_{i=1}^n P(X_i \mid Pa(X_i))$$
 (2.1)

where $Pa(X_i)$ denotes the set of parent nodes of X_i . This factorization enables efficient inference and learning, especially when the graph is sparse.

Markov random fields are undirected graphs where nodes represent random variables and edges encode direct probabilistic interactions. The joint distribution is expressed as a product of potential functions over cliques (fully connected subsets of nodes):

$$P(X_1, \dots, X_n) = \frac{1}{Z} \prod_{C \in \mathcal{C}} \psi_C(X_C)$$
 (2.2)

where C is the set of cliques, ψ_C are non-negative potential functions, and Z is the partition function ensuring normalization. It also can be defined through a partition function and an energy function:

$$P(X) = \frac{1}{Z} exp\left(\sum_{i} w_{i} \phi_{i}(X)\right)$$
(2.3)

This energy based representation perfectly overlaps with the Exponential family representation.

Both Bayesian networks and Markov networks can represent high-dimensional distributions compactly, capturing complex dependencies among variables. Many logical knowledge bases and rule systems can be embedded within graphical models, providing a bridge between symbolic and probabilistic reasoning. For example, logical rules can be encoded as hard or soft constraints in the potentials of a Markov network or as conditional probability tables in a Bayesian network.

2.2 Exponential Family Distributions

2.2.1 Definition and Properties

The exponential family is a broad class of probability distributions that share a common functional (energy based) form, making them particularly amenable to statistical inference and learning [6][7]. A probability distribution $p(x \mid \eta)$ belongs to the exponential family if it can be written as:

$$p(x) = \exp\{\eta^T \phi(x) - A(\eta)\}$$
(2.4)

where h(x) is the base measure, η is the vector of natural (canonical) parameters, $\phi(x)$ is the vector of sufficient statistics, and $A(\eta)$ is the log-partition function (also called the cumulant generating function) that ensures normalization.

Key properties of exponential family distributions include:

- Sufficient statistics: The function $\phi(x)$ captures all information about the data relevant to parameter estimation.
- Conjugacy: Many exponential family distributions admit conjugate priors, simplifying Bayesian inference.

- Tractable moments: Moments and cumulants of the distribution can be computed as derivatives of $A(\eta)$.
- Generalization: Many common distributions, such as the Bernoulli, multinomial, normal, and Poisson, and Markov Logic Networks as well are members of the exponential family.

In the context of AI and machine learning, exponential familiy representations can provide a general, unified framework that encompasses many commonly used distributions, such as probabilistic graphical models and logic-based systems. By choosing appropriate sufficient statistics (such as indicator functions for logical formulas), one can encode logical constraints and probabilistic dependencies within a single, tractable mathematical formalism.

2.2.2 Examples Relevant to Logical Connectives

Many logical knowledge bases, Markov logic networks (MLNs), and rule-based systems can be naturally expressed within the exponential family framework. In MLNs, for example, each logical formula ϕ_i is associated with a weight θ_i , and the probability of a world (i.e., a complete assignment of truth values) is given by:

$$P(X) \propto \exp\left(\sum_{i} \theta_{i} \phi_{i}(X)\right)$$
 (2.5)

Here, $\phi_i(X)$ is an indicator function that evaluates to 1 if the formula is satisfied by X, and 0 otherwise. The weights θ_i determine the strength of each logical constraint: large positive values enforce hard constraints, while smaller values allow for soft, probabilistic reasoning.

This representation allows for the seamless integration of symbolic logic and statistical learning. Logical connectives can be encoded as sufficient statistics in the exponential family, and their weights can be learned from data. As the weights become large, the model approaches a purely logical system; as they decrease, the model becomes more probabilistic, capturing uncertainty and noise in the data.

2.3 Tensor Networks

2.3.1 Tensor Network Basics

A tensor is a multi-dimensional array generalizing scalars (0th order), vectors (1st order), and matrices (2nd order) to higher dimensions. Formally, an nth-order tensor

 $\mathcal{T} \in \mathbb{R}^{d_1 \times \cdots \times d_n}$ has n indices, each ranging over dimensions d_1, \ldots, d_n . Tensors enable compact representation of high-dimensional functions, such as joint probability distributions over many variables.

A tensor network is a structured factorization of a high-order tensor into a collection of lower-order tensors connected via contractions over shared indices. Graphically, nodes represent tensors, and edges denote contracted indices.

Key operations in tensor networks include:

- Contraction: Summing over shared indices, e.g., $C[i,j] = \sum_k A[i,k]B[k,j]$ for matrix multiplication.
- **Decomposition**: Approximating \mathcal{T} as a network of smaller tensors via formats (like Canonical-Polyadic (CP), Matrix Product State (MPS), or tensor train (TT)).
- **Normalization**: Enforcing constraints (e.g., non-negativity) to represent valid probability distributions.

Tensor networks excel at capturing locality and hierarchy in data. For example, a tree tensor network (TTN) mirrors the structure of a Bayesian network, with each node representing a conditional probability table and edges encoding variable dependencies.

2.3.2 Tensor Networks for Probabilistic Models

The joint probability distribution of a graphical model can be encoded as a tensor where each entry $\mathcal{T}[x_1,\ldots,x_n]$ stores $P(X_1=x_1,\ldots,X_n=x_n)$. Tensor networks factorize this high-dimensional tensor into localized components reflecting the model's conditional independence structure.

Example: Bayesian Network as a Tensor Network

Consider a Bayesian network over binary variables X_1, X_2, X_3 with edges $X_1 \to X_2$ and $X_2 \to X_3$. The joint distribution is:

$$P(X_1, X_2, X_3) = P(X_1)P(X_2|X_1)P(X_3|X_2)$$
(2.6)

Which factorizes to a tensor network:

$$\mathcal{T}[X_1, X_2, X_3] = \sum_{Y} \mathcal{A}[X_1] \mathcal{B}[X_1, X_2, Y] \mathcal{C}[X_2, X_3, Y]$$
 (2.7)

where Y is an auxiliary index connecting the two conditional tensors $\mathcal{B}[X_1, X_2, Y]$ and $\mathcal{C}[X_2, X_3, Y]$, with $\mathcal{A}[X_1]$ being the prior tensor to X_1 .

Example: Markov Random Field as a Tensor Network

Consider a Markov random field (MRF) over binary variables X_1, X_2, X_3 with cliques $\{X_1, X_2\}$ and $\{X_2, X_3\}$. Its joint distribution is:

$$P(X) = \frac{1}{Z}\psi_{12}(X_1, X_2)\psi_{23}(X_2, X_3)$$
(2.8)

This corresponds to a tensor network:

$$\mathcal{T}[X_1, X_2, X_3] = \sum_{Y} \psi_{12}[X_1, X_2, Y] \psi_{23}[Y, X_2, X_3]$$
 (2.9)

where Y is an auxiliary index connecting the two clique tensors. This meets the Tensor Network representation of Propositional Logical formulas as well, where we can define the unique Logical connectives through the $\psi_{X_n,X_m,Y}$ functions, with X_n, X_m being the variables for the connective and the Y should be the 'ancilla' connecting the connectives to have the Logical formula in the end. end Applications:

- Marginalization: Computing $P(X_i)$ reduces to contracting all indices except X_i . For a TTN, this scales linearly in n rather than exponentially.
- Sampling: Ancestral sampling in Bayesian networks corresponds to sequential contractions in directed tensor networks.
- Learning: Maximum likelihood estimation becomes tensor factorization, e.g., using alternating least squares (ALS).

2.4 Tensor Network Representation of Exponential **Families**

Exponential family distributions, due to their structured parameterization and sufficient statistics, are naturally suited for representation as tensor networks. This correspondence enables efficient manipulation, marginalization, and sampling—crucial for high-dimensional probabilistic models and for mapping to quantum circuits.

2.4.1 Slice Tensor Decomposition

A central technique for representing high-order tensors arising from exponential family models is slice tensor decomposition. In this approach, a high-dimensional tensor $\mathcal{T}[X_1,\ldots,X_n]$ encoding the joint distribution is factorized into a sum of lower-rank tensors (slices) along one or more modes:

$$\mathcal{T}[X_1, \dots, X_n] = \sum_{k=1}^r \lambda_k \,\mathcal{A}_k[X_1] \otimes \mathcal{B}_k[X_2, \dots, X_n]$$
 (2.10)

where λ_k are scalar coefficients, and \mathcal{A}_k , \mathcal{B}_k are subtensors or vectors corresponding to particular slices. This decomposition preserves the conditional independence structure of the underlying graphical or logical model and allows for efficient storage and computation, especially when the tensor admits a low-rank structure.

Slice decomposition is particularly powerful for models with factorizable sufficient statistics, such as those arising in Markov logic networks or graphical models, where each logical formula or clique potential can be associated with a tensor slice. The resulting network of slices can then be contracted (multiplied and summed over shared indices) to recover marginals or conditionals.

2.4.2 Operational Mapping: Step-by-Step Example

To illustrate, consider a simple exponential family model over three binary variables X_1, X_2, X_3 . The joint distribution can be encoded as a rank-3 tensor $\mathcal{T}[X_1, X_2, X_3]$. Suppose the model factorizes as:

$$\mathcal{T}[X_1, X_2, X_3] = \sum_{k=1}^r \mathcal{S}_k[X_1] \cdot \mathcal{U}_k[X_2] \cdot \mathcal{V}_k[X_3]$$
 (2.11)

where each $S_k, \mathcal{U}_k, \mathcal{V}_k$ is a vector (or slice) over its respective variable, and r is the decomposition rank. This format is equivalent to the canonical polyadic (CP) decomposition and is especially efficient when r is small compared to the full tensor size.

In practice, such decompositions can be obtained via algebraic methods (e.g., alternating least squares) or by exploiting the structure of the model (e.g., using the factor graph or logical formulae). The resulting tensor network can then be contracted to compute marginals, conditionals, or to generate samples.

Chapter 3

Quantum Sampling Algorithms

3.1 Quantum Gates, Circuits, and Measurement

Quantum gates, circuits, and measurements together provide the operational foundation for all quantum algorithms. Gates manipulate the amplitudes and phases of quantum states, circuits implement complex transformations, and measurement extracts classical information, enabling quantum algorithms to outperform their classical counterparts in sampling, search, and inference [8].

This section summarizes the essential elements relevant for later procedures, such as amplitude amplification and the collective quantum sampling algorithm.

3.1.1 Qubit States and Quantum Registers

A single qubit is described by a state vector in a two-dimensional Hilbert space,

$$|a\rangle = v_0|0\rangle + v_1|1\rangle,$$

where v_0 and v_1 are complex amplitudes satisfying $|v_0|^2 + |v_1|^2 = 1$. The basis states $|0\rangle$ and $|1\rangle$ are represented as column vectors:

$$|0\rangle = \begin{bmatrix} 1 \\ 0 \end{bmatrix}, \quad |1\rangle = \begin{bmatrix} 0 \\ 1 \end{bmatrix}.$$

For multi-qubit systems, the overall state is given by the tensor product of individual qubit states. For example, a two-qubit state is

$$|\psi\rangle = v_{00}|00\rangle + v_{01}|01\rangle + v_{10}|10\rangle + v_{11}|11\rangle.$$

3.1.2 Quantum Gates: Basic Building Blocks

Quantum gates are unitary operations acting on one or more qubits. They generalize classical logic gates but can create superposition and entanglement, enabling quantum parallelism.

Single-Qubit Gates:

• Pauli-X (NOT) Gate: Flips $|0\rangle$ to $|1\rangle$ and vice versa.

$$X = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix}$$

• Pauli-Y and Pauli-Z Gates: Y combines a bit and phase flip; Z applies a phase flip to $|1\rangle$.

$$Y = \begin{bmatrix} 0 & -i \\ i & 0 \end{bmatrix}, \quad Z = \begin{bmatrix} 1 & 0 \\ 0 & -1 \end{bmatrix}$$

• Hadamard (H) Gate: Creates superposition. Applied to $|0\rangle$, it yields $(|0\rangle + |1\rangle)/\sqrt{2}$.

$$H = \frac{1}{\sqrt{2}} \begin{bmatrix} 1 & 1\\ 1 & -1 \end{bmatrix}$$

• Rotation Gates: $R_x(\theta)$, $R_y(\theta)$, and $R_z(\theta)$ rotate the qubit state around the respective axes by angle θ .

Multi-Qubit Gates:

- CNOT (Controlled-NOT) Gate: A two-qubit gate that flips the target qubit if the control qubit is $|1\rangle$. Essential for generating entanglement.
- Toffoli (CCNOT) Gate: A three-qubit gate; flips the third (target) qubit if both control qubits are |1\(). Important for universal reversible computation and error correction.
- SWAP Gate: Exchanges the states of two qubits.

Quantum gates are represented as unitary matrices. A gate acting on n qubits is a $2^n \times 2^n$ unitary matrix.

3.1.3 Quantum Circuits

A quantum circuit is a sequence of quantum gates applied to qubits, typically followed by measurement. Circuits are often depicted as diagrams, with qubits as horizontal lines and gates as symbols acting on these lines. For example, the circuit for creating a Bell state applies a Hadamard gate to the first qubit, followed by a CNOT with the first as control and the second as target, as seen in the figure below 3.1.

The action of a circuit on an initial state $|\psi_0\rangle$ is a sequence of unitary transformations:

$$|\psi_{\text{final}}\rangle = U_k \cdots U_2 U_1 |\psi_0\rangle,$$

where each U_i is a quantum gate.

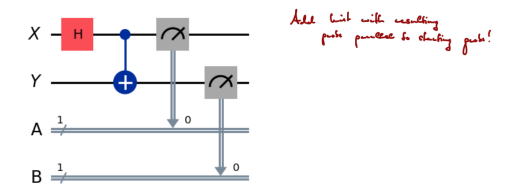


Figure 3.1: Two-qubit entangling circuit with measurement added at the end

3.1.4 Measurement in Quantum Mechanics

A measurement is the testing or manipulation of a physical system to yield a numerical result. In quantum mechanics it is when a quantum state collapses to a classical outcome. In the computational basis, measuring a qubit in state $|a\rangle = v_0|0\rangle + v_1|1\rangle$ yields $|0\rangle$ with probability $|v_0|^2$ and $|1\rangle$ with probability $|v_1|^2$.

For multi-qubit systems, measurement projects the state onto one of the basis vectors $|k\rangle$, with probability $|v_k|^2$. The outcome is inherently probabilistic, a fundamental departure from classical computation.

Entanglement and Measurement: Measurement on one qubit of an entangled pair instantaneously determines the outcome of the other, a phenomenon with no classical analog. For example, measuring one qubit of the Bell state $(|00\rangle + |11\rangle)/\sqrt{2}$ collapses the other qubit to the same value, although causality must be preserved, hence the no communication theorem should be obeyed [9].

3.2 Amplitude Amplification: Theory and Practice

Amplitude amplification is a quantum algorithmic technique that generalizes Grover's search [3], providing a quadratic speedup for the identification or sampling of "good" states in a large, unstructured search space. The power of amplitude amplification lies in its ability to systematically increase the probability amplitude associated with desirable outcomes, while keeping the state space normalized, making it a central primitive for quantum-enhanced sampling in machine learning.

3.2.1 Grover's Algorithm and Generalizations

Grover's algorithm is the canonical example of amplitude amplification. In the classical setting, searching for a marked item in an unsorted database of size N requires, on average, O(N) queries, as with random selection, each and every item would have a $\frac{1}{N}$ probability of finding. Grover's quantum approach reduces this to $O(\sqrt{N})$ by exploiting quantum superposition and entanglement.

The algorithm operates in an N-dimensional Hilbert space \mathcal{H} , where each basis state $|k\rangle$ represents a possible solution. A Boolean oracle function $\chi: \{0,1\}^n \to \{0,1\}$ identifies "good" states. The oracle operator **O** applies a phase flip to these states, while the diffusion operator **D** amplifies their amplitudes.

Brassard et al. [10] extended Grover's idea to arbitrary initial states and general quantum algorithms, formalizing the amplitude amplification operator:

$$\mathbf{Q} = -\mathcal{A}\mathbf{B}_0\mathcal{A}^{-1}\mathbf{B}_{\gamma}$$

where \mathcal{A} prepares the initial state, \mathbf{B}_{χ} flips the phase of good states, and \mathbf{B}_0 flips the phase of the all-zero state. Repeated application of \mathbf{Q} rotates the quantum state in the two-dimensional subspace spanned by the good and bad components, exponentially increasing the probability of measuring a good state.

3.2.2 Oracle and Diffusion Operator Design

The effectiveness of amplitude amplification hinges on the careful design of the oracle and diffusion operators.

Oracle Operator O: The oracle is a unitary operator that marks the set of good states by flipping their phase. For a basis state $|k\rangle$, in $\mathcal{S} := \{|k\rangle\}_{k=0}^{N-1}$,

$$\mathbf{O} |k\rangle = (-1)^{\chi(k)} |k\rangle$$

where $\chi(k)=1$ for good states and 0 otherwise. In practical terms, the oracle is implemented as a quantum circuit that evaluates the Boolean function χ and applies a controlled-Z or multi-controlled Toffoli gate to flip the phase of the target states. For simple logical connectives, such as AND or OR, the circuit construction is straightforward. For more complex constraints, the circuit depth increases, but the principle remains the same: the oracle must be a reversible, unitary operation that encodes the solution set into phase flips.

Diffusion Operator D: The diffusion operator, or "inversion about the mean," amplifies the amplitudes of the marked states. For an initial state $|\psi\rangle = \mathcal{A}|0\rangle$, the diffusion operator is

$$\mathbf{D} = 2 |\psi\rangle \langle \psi| - \mathbb{I}$$

This operator reflects the quantum state about the initial state vector. In the special case where $|\psi\rangle$ is the uniform superposition, **D** can be implemented by Hadamard gates, a phase flip on $|0\rangle$, and another round of Hadamards. For non-uniform initial states, the diffusion operator is constructed by applying the inverse of the state preparation circuit, a phase flip on $|0\rangle$, and then re-applying the state preparation.

Geometric Interpretation: The interplay between the oracle and diffusion operators can be visualized as a sequence of reflections in a two-dimensional subspace. With a projection operator emerging from the definition of **O**;

$$\mathcal{P} := \sum_{\chi(k)=1} |k\rangle \langle k|$$

defining the good and bad subspaces respectively:

$$\mathcal{H}_G := Im(\mathcal{P}) = span\{|k\rangle \in \mathcal{S}_{op} \mid \chi(k) = 1\}$$
$$\mathcal{H}_B := Ker(\mathcal{P}) = span\{|k\rangle \in \mathcal{S}_{op} \mid \chi(k) = 0\}$$

If we adopt the convention of \sum' for summation over the solution states, and \sum'' for summation over the non-solutions, we can define normalized states as such:

$$|x_G\rangle = \sqrt{\frac{1}{M}} \sum_{x}' |x\rangle$$

$$|x_B\rangle = \sqrt{\frac{1}{N-M}} \sum_{x}^{"} |x\rangle$$

where N is the number of states (2^n for n qubits) and M is the number of solution states. Thus the initial state can be realized as:

$$|\phi\rangle = \sqrt{\frac{N-M}{N}} |x_B\rangle + \sqrt{\frac{M}{N}} |x_G\rangle = \cos(\theta) |x_B\rangle + \sin(\theta) |x_G\rangle$$

This is then a two-dimensional subspace spanned by the vectors $|x_G\rangle$ and $|x_B\rangle$ is stable under the action of **Q**. The oracle reflects the state about the bad subspace, while the diffusion operator reflects about the initial state.

The composition of these two reflections is a rotation by 2θ towards the good subspace. This geometric process underlies the quadratic speedup of amplitude amplification.

3.2.3 Parameter Update Interpretation

The action of the amplitude amplification operator \mathbf{Q} can be interpreted as two reflections – once over the bad states, and then over the average amplitude of our

states (i.e. the register itself in the two-dimensional subspace) – which geometrically is a rotation in the two-dimensional subspace of the Hilbert space spanned by the projections of the initial state onto the good and bad subspaces. The initial state can be written as

$$|\psi\rangle = \sin(\theta) |\psi_q\rangle + \cos(\theta) |\psi_b\rangle$$

where $|\psi_q\rangle$ and $|\psi_b\rangle$ are normalized projections onto the good and bad subspaces, respectively. As the state is stable under the action of \mathbf{Q} , the rotaion angle of θ is given. Thus, Each application of **Q** rotates the state by 2θ , such that after k iterations,

$$\mathbf{Q}^{k} | \psi \rangle = \sin((2k+1)\theta) | \psi_{q} \rangle + \cos((2k+1)\theta) | \psi_{b} \rangle$$

The probability of measuring a good state thus increases quadratically with the number of iterations, reaching a maximum when $(2k+1)\theta \approx \pi/2$. add informative picture about the rotation has the sampling which are the second of the sampling which are the sampling which are the sampling which are the sampling which are the sampling with the s

findings In the context of exponential family distributions, amplitude amplification can be leveraged to sample from distributions where the "good" states correspond to assignments satisfying logical constraints or high-probability regions. The initial quantum state encodes the base distribution (e.g., uniform or prior), and the oracle marks the desired subset. Each amplification step updates the effective parameters of the distribution, concentrating probability mass on the target set.

Mathematically, if the initial probability of a good state is P_0 , then after k applications of amplitude amplification, the probability becomes $P_k = \sin^2((2k+1)\theta)$, where $\theta = \arcsin(\sqrt{P_0})$. In exponential family terms, this corresponds to a shift in the canonical parameter:

$$\theta_k = \ln\left(\frac{(1 - P_0)P_k}{P_0(1 - P_k)}\right)$$

This provides a direct link between quantum amplitude amplification and classical parameter updates in probabilistic models.

Circuit Construction: In practice, the amplitude amplification circuit is constructed as follows:

- 1. Prepare the initial state $|\psi\rangle$ using the unitary \mathcal{A} .
- 2. Apply the oracle **O**.
- 3. Apply the diffusion operator **D**.

- 4. Repeat the sequence $\mathbf{Q} = \mathbf{DO}$ for the optimal number of iterations.
- 5. Measure the final state in the computational basis.

The oracle and diffusion operators are implemented as modular subroutines, allowing for flexibility in defining different logical connectives or constraints.

3.2.5 Scalability and Simulator Limitations

The scalability of amplitude amplification is determined by the complexity of the oracle and diffusion operators, as well as the available quantum resources. For simple logical connectives and small models, both operators can be implemented with shallow circuits, and the algorithm can be simulated efficiently on classical hardware. For more complex logical formulas or high-dimensional models, the circuit depth and qubit count increase, and classical simulation becomes intractable.

In this work, quantum simulations were performed for circuits up to 25 qubits using the Qiskit Aer simulator, balancing accuracy and computational feasibility. For larger problem sizes, efficient simulation may require tensor network-based methods or access to real quantum hardware. The quadratic speedup of amplitude amplification remains, but practical implementation is constrained by current hardware and software limitations.

3.3 From Tensor Networks to Quantum Circuits

Tensor networks unify the representation of probabilistic models with quantum state encoding, as probabilistic models can be 2.3.2, and quantum states, such as matrix product states (MPS) and projected entangled pair states (PEPS) are inherently represented as tensor networks [TNforQC]. Therefore tensor networks provide a unified framework for representing and manipulating probabilistic models, as inference and Amplitude amplification in quantum algorithms operates on these networks, providing a pathway to quantum-enhanced sampling.

3.3.1 Quantum Circuit Construction

visualize maybe?

To map a tensor network to a quantum circuit, each tensor (or slice) is encoded as a quantum state or as a set of parameterized quantum gates. The contraction of tensors, which in the classical setting corresponds to summing over shared indices, is implemented in the quantum setting by entangling qubits or applying multi-qubit gates. For example, in a network where each variable X_i is binary, a qubit is allocated for each variable, and the entries of the tensor slices determine the amplitudes of basis states after the application of parameterized rotations.

The construction proceeds as follows:

- 1. **Initialization**: Prepare the quantum register in a reference state (e.g., $|0\rangle^{\otimes n}$).
- 2. **State Preparation**: Apply a sequence of single- and multi-qubit gates to encode the tensor slices, such that the resulting quantum state amplitudes correspond to the (normalized) entries of the target tensor network. For instance, parameterized rotation gates (R_X, R_Y, R_Z) can be used to set the amplitudes according to the tensor entries, while Hadamard gates may be used to create initial superpositions
- 3. **Entanglement**: Use CNOT and controlled gates to implement contractions between slices, reflecting dependencies in the original probabilistic model. In my model later, entanglement will be inherently present due to the nature of the encoding, with Toffoli gates being frequently utilized over ancillas (i.e. shared indices of the Tensor Network).

This mapping is particularly efficient when the tensor network has low rank or sparse structure, as is often the case for models with strong conditional independence. In such scenarios, the number of required gates and the circuit depth can be kept polynomial in the number of variables, making the approach feasible for near-term quantum devices [Ran 2020].

3.3.2 Gate Decomposition and Resource Estimates

The quantum circuit depth and resource requirements depend on the structure and rank of the underlying tensor network:

- Qubit Count: Each variable in the model typically requires one qubit for binary variables (or $\lceil \log_2 d_i \rceil$ qubits for d_i -ary variables) as well as one auxiliary qubit (i.e. ancilla qubit) for connecting the cliques of probabilistic model representations 2.3.2.
- Gate Complexity: The number of gates scales with the number of tensor slices and the connectivity of the network. For a CP decomposition of rank r over n variables, the circuit requires $\mathcal{O}(rn)$ parameterized gates [11].
- Circuit Depth: For chain-like (MPS) or tree-like (TTN) tensor networks, the depth is O(n) or $O(\log n)$, respectively.

In summary, the mapping from exponential family distributions to tensor networks, and subsequently to quantum circuits, enables scalable quantum sampling for complex probabilistic models. This approach exploits both the structure of the underlying model and the computational power of quantum devices, providing a pathway to quantum advantage in probabilistic inference and machine learning.

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