

# Quantum simulation of a space-time atom in loop quantum gravity

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## I. INTRODUCTION

A major open problem in NISQ is whether modern quantum computers can practically solve classically intractable problems [1]. We had access Google's Weber QPU, a 53 qubit superconducting quantum computer. It is the same computer used in [2]. The following is a report of using the Weber QPU to simulate spin-foam amplitudes in Loop Quantum Gravity (LQG). It also serves to analyze whether NISQ computers are fit to simulate LQG.

Quantum gravity is a field of theoretical physics attempting to find a quantum mechanical description of gravity. LQG is a theory based on the quantization of space-time, where entangled quantum tetrahedra give rise to 3+1 dimensional space-time in a way that unifies quantum mechanics and general relativity [3]. Because of the complexity of the Planck scale's degrees of freedom, plus the exponential growth in the dimension of a Hilbert space, quantum gravitational systems are too difficult for classical supercomputers [4]. However, quantum computation gives an exponential speedup to problems and a dramatic increase in spatial efficiency; two space-time atoms have a wavefunction of  $2^{40}$  states. This requires  $\sim 8$  TB on a classical computer, but only 40 qubits on a quantum computer.

## II. LOOP QUANTUM GRAVITY

We analyze covariant LQG using the spin-foam model. In the spin-foam model, 3-dimensional space is described by a 3-dimensional quantum space state. Evolution from initial 3-dimensional space states to final 3-dimensional space states creates the 4-dimensional quantum spacetime regions [5]. Spin-foam amplitudes are the probability amplitudes of the spin-foam model after one such evolution.

Calculation of the spin-foam amplitudes is an essential step to understand the spin-foam model in LQG. Unfortunately, spin-foam amplitudes are exponentially difficult to calculate, making non-trivial spin-foam models impossible for classical computers. That is why quantum computers are the most convenient way to calculate these amplitudes especially superconducting circuits provide a competitive result while building practical quantum computers.

### A. Constraints

There are 3 constraints in LQG. First is the Gauss constraint, which is used for constraining the spin-networks with the equation  $0 = P_G |T\rangle$ , where  $|T\rangle$  is the tensor product state of the spin-network. The second constraint is spatial diffeomorphism, which is used for characterizing the topology between all spin networks. The third one is the Hamiltonian constrain which is the most difficult to satisfied, is used for the encodes temporal dynamics of the LQG [6].

### B. Quantum Tetrahedra and Spin-Networks

In any  $n$ -dimensional space, an  $n$ -simplex is the shape with the fewest possible number of faces; a 2-simplex is a triangle, a 3-simplex is a tetrahedron, and so on to higher dimensions. One central tenet of LQG is quantized space-time, and the tetrahedron is the simplest 3 dimensional geometry that can be used for this quantization. Therefore, to get discrete 3-dimensional space

we will choose the 3-simplex, or tetrahedron, to be our discrete unit of space [7]. One tetrahedron is defined by the equation:

$$\vec{J}_1 + \vec{J}_2 + \vec{J}_3 + \vec{J}_4 = 0 \quad (1)$$

where  $\vec{J}_i = (J_x, J_y, J_z)$  is the angular momentum vector of the  $i$ th face [5]. We can then define a quantum tetrahedron, or qubit of space, with the following [8], where  $\theta$  and  $\phi$  are angles on the Bloch sphere:

$$|t\rangle = \cos\left(\frac{\theta}{2}\right) |0_L\rangle + e^{i\phi} \sin\left(\frac{\theta}{2}\right) |1_L\rangle \quad (2)$$

$$|0_L\rangle = \frac{1}{2}(|01\rangle - |10\rangle)(|01\rangle - |10\rangle) \quad (3)$$

$$|1_L\rangle = \frac{1}{\sqrt{3}}[|1100\rangle + |0011\rangle - \frac{1}{2}(|01\rangle + |10\rangle)(|01\rangle + |10\rangle)] \quad (4)$$

This quantum tetrahedron is the discrete unit of 3 dimensional quantum space. A collection of connected quantum tetrahedra gives rise to a spin-network graph, where each node in the graph is a tetrahedron and links are formed by gluing adjacent faces [9]. A spin-network graph is given by the tensor product:

$$|T\rangle = \otimes_{i=1}^N |t_i\rangle \quad (5)$$

where  $N$  is the number of tetrahedra.

The 4-simplex is a geometric object used to create discrete 3+1 dimensional space-time [7], so in order to properly simulate LQG spin-foam amplitudes we need to create a 4-simplex with our quantum tetrahedra. An  $n$ -simplex is created by gluing  $n+1$  simplices from the  $n-1$  dimension; a 2 dimension triangle is created by “gluing” 3 lines together. To create the 4-simplex, we will glue 5 quantum tetrahedra together with entanglement between faces. A space-time atom is then the process where  $m$  quantum tetrahedra glue to form  $5-m$  quantum tetrahedra [5]. Two faces glued together is equivalent to a “link” in the spin-network, given by a collection of Bell states:

$$|W\rangle = \otimes_{l=1}^{2N} |\epsilon_l\rangle, \quad |\epsilon_l\rangle = \frac{|01\rangle - |10\rangle}{\sqrt{2}} \quad (6)$$

where  $|\epsilon_l\rangle$  is a link  $l$  in the graph. A 4-simplex forms a maximally entangled spin-network graph of five quantum tetrahedra.

### C. Transition Amplitudes

Evolution of 3-dimensional space states corresponding to 3-dimensional boundaries of space-time encode the spin-foam amplitudes, or transition amplitudes, between the initial and final spin-networks. The transition amplitude represents local dynamics of a spin-network [10] by showing the change in probability amplitudes. To satisfy the Gaussian constraint, a spin-foam amplitude is given by:

$$A = \langle\psi| P_G |\psi\rangle \quad (7)$$

## III. QUANTUM SIMULATION

We performed quantum simulations of spin-networks of two and five quantum tetrahedra to get the spin-foam amplitude of topologically inequivalent spin-networks. In order to minimize

the errors due to noise and coherence times, we only consider the  $\theta = 0$  case where  $|t\rangle = |0_L\rangle$ . One qubit of space is created with the operator  $U$  such that  $|t\rangle = U|0000\rangle$ , and the Bell state is created with operator  $B$  such that  $|\epsilon\rangle = B|00\rangle$ .

The spin-foam amplitude of a spin-network of quantum tetrahedra is given by the inner-product:

$$A = \langle W|T \rangle \quad (8)$$

To simulate this, we first prepare the system in the all zero state,  $|0\dots 0\rangle$ . We then create  $N$  qubits of space by acting with  $U$  on sets of 4 qubits. A spin-network is then created by maximally entangling the qubits of space with inverse Bells states  $B^\dagger$ . Finally, we find our amplitude by measuring the final amplitude of the  $|0\dots 0\rangle$  state. This algorithm is equivalent to the inner-product:

$$A = \langle 0\dots 0|B^\dagger U|0\dots 0\rangle \quad (9)$$

While the amplitude is given by the inner product, the results from the quantum simulation will be the probability obtained from the amplitude.  $\langle W|T \rangle$  is the spin-foam amplitude, while  $|\langle W|T \rangle|^2$  is the probability.

### A. Circuit

Our circuit must satisfy as many of the constraints in LQG as possible. The Hamiltonian constraint is currently an active area of research, and has been simulated using adiabatic quantum computing [11], but it is not considered here. The diffeomorphism constraint is satisfied implicitly in the structure of the circuit [4], and we satisfy the Gauss constraint with the operators  $P_G = B^\dagger U$ .

The operators mentioned above are equivalent to the following quantum circuits:

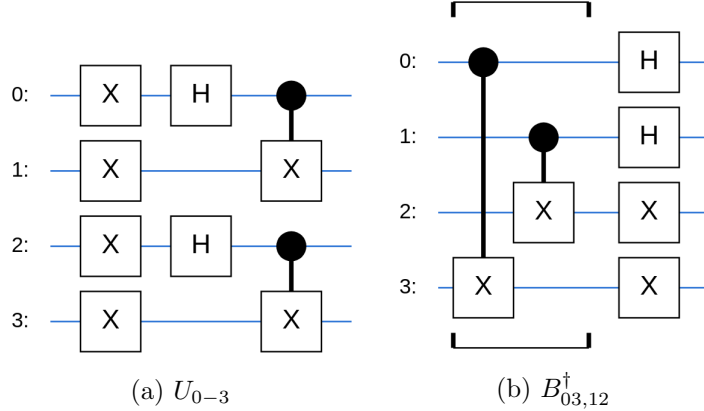


Figure 1: Circuit diagrams [15]

Notice how in the  $|0_L\rangle$  state, the entanglement is only within the pairs (0,1) and (2,3). This is part of what makes the  $|0_L\rangle$  state trivial; the quantum tetrahedron's faces are each only entangled with one other faces. This is not the case for any  $\theta \neq 0$  in the  $|t\rangle$  state.

### B. Topology

Spin-networks have different transition amplitudes based on the configuration of links in the graph. In the  $|0_L\rangle$  state, when all amplitudes of the wavefunction are uniform, this transition

amplitude is a dependent variable of, mostly, the topology of the quantum circuit. In this case, the final wavefunction has uniform amplitudes, so the amplitude of the  $|0\dots 0\rangle$  state is just a measure of the possible states of the system, like if a card just had  $\frac{1}{52}$  on it if it was picked from a full deck, or  $\frac{1}{13}$  if it was just picked from a suite of cards. The topology of each of our circuits can be defined by the number of independent “chains” of qubits that emerge and form shapes. In our observation, amplitudes of different spin-networks from the same topology can be different up to a sign change. Formal proofs of this will be presented in future work.

### C. Dipole Spin Network

This can be thought of as the gluing of two quantum tetrahedra, with each face paired to another face. It is the minimal triangulation of the 3-sphere, also known as the dipole cosmology [12]. We can organize expected values into two topological groups: one chain and two chains. Expected values are shown in the table below.

Group	$ \langle W 00_L\rangle ^2$
Single	0.0625
Double	0.015625

Table I: Spin-foam amplitude of dipole topological groups

The different circuits are shown in figure 2, brown is the connection in state preparation, blue is the connection in the Bell state.

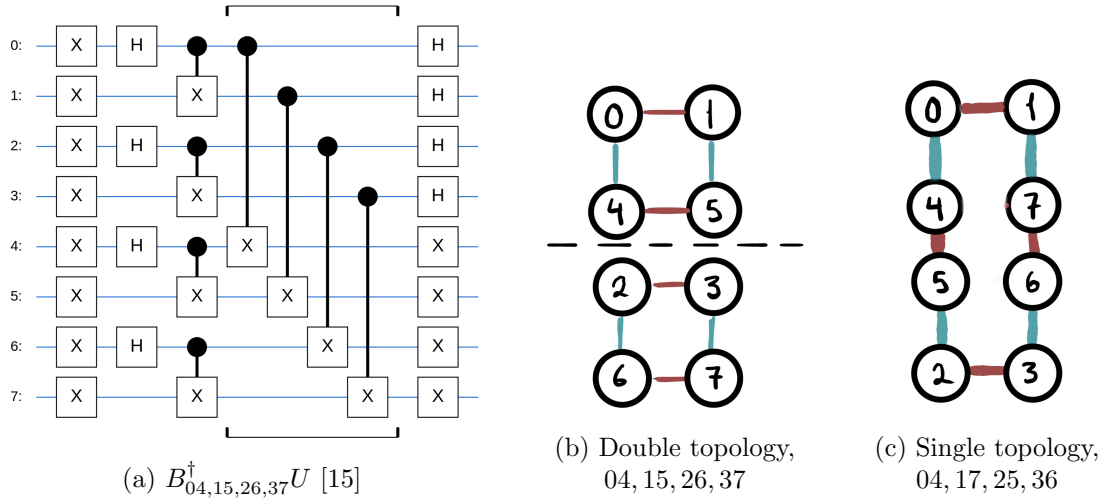


Figure 2: Dipole circuit and topology

### D. 4-Simplex Spin Network

To simulate a 4-simplex, we simulate the creation of a spin-network with maximum entanglement between the vertices. This is the gluing of five quantum tetrahedra to form a space-time atom. It has topology of one, two, and three chains. Expected values are shown in the table below.

The topological groups correspond to the following circuits

Group	$\langle W 00000_L\rangle$
Single	$\pm 2^{-9}$
Double	$\pm 2^{-8}$
Triple	$\pm 2^{-7}$

Table II: Spin-foam amplitude of 4-simplex topological groups

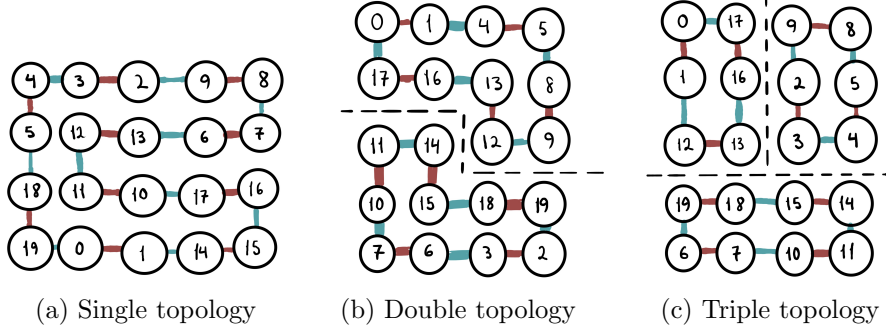


Figure 3: 4-simplex topology

## IV. RESULTS

### A. State Preparation

$\langle 1_L 1_{Lsim}\rangle$	$\langle 0_L 0_{Lsim}\rangle$
0.8618	0.9938

Table III: State preparation fidelity

The  $|1_L\rangle$  and  $|0_L\rangle$  state were each prepared 1000 times and their wavefunctions calculated. Above is the dot product with the Hermetian conjugate with the numerically calculated expected wavefunction. This fidelity is much higher than anything seen in the other literature [4],[8].

### B. Submission One

As of now, we have only received experiment results from one experiment with the QPU, to varying degrees of success.

#### 1. QPU Qubit Malfunction

Unfortunately, there was an error on Weber's (4, 8) qubit. Every time this qubit was used as a control, it would reset the target qubit to zero. The target qubit would then encounter an  $X$  flip in the inverse Bell state phase, resulting in a qubit with a final state of 1 for every run. We used the qubit on every experiment except for the state preparation and the single-chain dipole network, meaning those experiments are the only ones with useful results.

## 2. Single-Chain Dipole

Averaged over one-hundred thousand runs, the results of the single-chain dipole, figure 2.c, are shown below. The expected amplitude is 0.015625. The final state is always expected to have an even parity, so that can be used for post selection.

	Results	Post-selected
$ \langle W 00_L\rangle ^2$	0.01151	0.01869
Percent error	26.336%	16.616%

Table IV:  $\langle 0 \dots 0 | B_{04,17,25,36}^\dagger U | 0 \dots 0 \rangle$

Note that if the post-selected and non-post-selected results were averaged, we would get  $|\langle W|0_L0_L\rangle|^2 = 0.01510$ , which is percent error = 03.365%.

The 1 state is much more unstable than the 0 state, which means that many error channels, such as photon loss, will bias the 0 state[13]. Therefore, when we post-select on the even parity, only errors where at least one 1 state is present are removed. However, many of our errors are likely causing extra occurrences of the  $|0 \dots 0\rangle$  state, which will not be removed. In order to combat this on the second submission, the  $X$  gates at the end of the inverse Bell state were changed such that we are looking for states of half zeros and half ones to count the amplitude of  $|0 \dots 0\rangle$ . This way, the amplitude would not be overinflated by post-selection.

## C. Discussion and Future Work

Our simulations for the first submission were relatively simple, and closer to a trivial case. They form a simple 3-dimensional boundary for 3+1 dimensional space-time, and the simulated spin-foam amplitudes are a measure of the dynamics of quantum geometries in that space-time.

Even though the first submission did not return many results, the work is promising. As can be seen in the single-chain dipole network, results directly from the QPU are very close to the theoretically expected results. This problem is especially promising to be successful on a quantum computer because it has a short runtime— the circuit has less than ten moments when converted to the  $\sqrt{i}$ SWAP gate set. This gives the algorithm high resistance to the low coherence times that frequently occur in superconductors, giving it promising prospects for NISQ computing.

There is still a substantial amount of work to be further done. Next, we will create some formal proofs the topological properties of the circuits. A state preparation circuit with better fidelity for  $|t(\theta \neq 0)\rangle$  will be found, which will hopefully lead to high-fidelity simulations involving a qubit of space in superposition of the  $|0_L\rangle$  and  $|1_L\rangle$  state. This would allow us to run the above algorithm with spacetime tetrahedra which are eigenstates of the volume operator,  $|V_\pm\rangle = \frac{|0_L\rangle \mp i|1_L\rangle}{\sqrt{2}}$  [5]. We will also attempt to glue two space-time atoms together, as can be seen in [14], but with the larger 40 qubit system.

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