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Cite as: AVS Quantum Sci. 4, 013801 (2022); https://doi.org/10.1116/5.0073075
Submitted: 27 September 2021 • Accepted: 07 December 2021 • Published Online: 25 January 2022

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ABSTRACT
We investigate how hardware specifications can impact the final run time and the required number of physical qubits to achieve a quantum advantage in the fault tolerant regime. Within a particular time frame, both the code cycle time and the number of achievable physical qubits may vary by orders of magnitude between different quantum hardware designs. We start with logical resource requirements corresponding to a quantum advantage for a particular chemistry application, simulating the FeMo-co molecule, and explore to what extent slower code cycle times can be mitigated by using additional qubits. We show that in certain situations, architectures with considerably slower code cycle times will still be able to reach desirable run times, provided enough physical qubits are available. We utilize various space and time optimization strategies that have been previously considered within the field of error-correcting surface codes. In particular, we compare two distinct methods of parallelization: Game of Surface Code’s Units and AutoCCZ factories. Finally, we calculate the number of physical qubits required to break the 256-bit elliptic curve encryption of keys in the Bitcoin network within the small available time frame in which it would actually pose a threat to do so. It would require $317 \times 10^6$ physical qubits to break the encryption within one hour using the surface code, a code cycle time of 1 μs, a reaction time of 10 μs, and a physical gate error of $10^{-3}$. To instead break the encryption within one day, it would require $13 \times 10^6$ physical qubits.

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I. INTRODUCTION
With the advent of quantum computers, the race to a quantum computational advantage has gained serious traction in both the academic and commercial sectors. Recently, quantum supremacy has been claimed on quantum devices with tens of qubits. However, the targeted problems solved were theoretical in nature, and not relevant to industrial applications. Quantum advantage, conversely, is a stronger form of supremacy that shows an industrially relevant computational problem solved in a reasonable time-frame that would be practically impossible to do using any classical supercomputer. There is a large physical qubit overhead associated with quantum error correction which is required to run some of the most powerful algorithms. There are many factors that will determine the ability for different quantum computing architectures to scale. Consequently, within a given time frame, the maximum size (qubit count) of a device could vary by orders of magnitude. The code cycle time (base unit of operation in the surface code) will also vary considerably between platforms. Therefore, it is of interest to investigate the interplay between the code cycle time and the number of achievable qubits and the resulting impact on the feasibility for a particular device to achieve a quantum advantage. We calculate the physical qubit and run time requirement for relevant problems in chemistry and cryptography with a surface code error-corrected quantum computer, comparing parameters typical to various types of hardware realizations.

Algorithms that are tailored to Noisy Intermediate Scale Quantum (NISQ) devices generally consist of a hybrid approach, where a low depth circuit is parameterized and iterated through a classical optimizer. These NISQ algorithms are more often heuristic in nature than their fault tolerant counterparts, and so, providing rigorous resource estimates can be more challenging. Many of the most powerful quantum algorithms require a circuit depth that greatly exceeds the capabilities of NISQ era devices, and for some applications, the number of required
logical qubits and operations is known. The quantum threshold theorem states that a quantum computer using error correction schemes, and a physical error below a certain threshold, can suppress the logical error rate to arbitrary low levels. Therefore, one could run an algorithm with an arbitrary long circuit depth, provided enough qubits are available to perform the required level of error correction. There is a large time overhead for performing logical operations at the error-corrected level relative to operations performed on physical qubits. For the foreseeable future, classical computers will have a clock rate that is orders of magnitude faster than error-corrected quantum computers. To determine the problem size at which a quantum computer will outperform a classical computer, one must consider both the algorithmic speed up and the relative difference between their associated clock rates. By making use of parallelization schemes, quantum computers can speed up the effective rate of logical operations at the cost of additional qubits, and so the ability to scale to large enough device sizes will also play a role in determining the feasibility of reaching desirable run times.

The surface code is the most researched error correction technique, particularly in regard to end-to-end physical resource estimation. There are many other error correction techniques available, and the best choice will likely depend on the underlying architecture’s characteristics, such as the available physical qubit–qubit connectivity. Superconducting qubits are one of the leading quantum computing platforms, and these devices generally consist of static qubits arranged on a grid where only nearest neighbor interactions are natively available. Long-distance connections must be enabled by sequences of nearest neighbor swap operations, which, in the context of a NISQ device, may limit their computational power. The limited connectivity of superconducting qubits in part motivated the continued research into the surface code, which relies on only nearest neighbor interactions between physical qubits arranged on a 2D grid. In the following, we briefly introduce some of the alternative error correction techniques to the 2D surface code. Error correction codes that rely on global interactions at the physical level have favorable encoding rates as a function of code distance, but enabling this global connectivity on large devices may be challenging, or the connectivity overheads may outweigh the benefits relative to closer-range-connectivity codes. Entanglement distribution may be a viable method of enabling distant connectivity for large-scale devices with limited physical connectivity in the limit of large reserves of quantum memory. Higher dimensional error correction codes can have access to a greater range of transversal gates, which may considerably improve run times, where transversal implies that each qubit in a code block is acted on by at most a single physical gate, and each code block is corrected independently when an error occurs. Realizing this 3D (or greater) physical connectivity could be challenging for many of the current quantum platforms; photonic-interconnected modules may be the most flexible architecture with regard to its possible connectivity graph, however, currently, achieved connection speeds would present a considerable bottleneck. A variant of the 3D surface code may still be realizable with hardware that is only scalable in two dimensions because the thickness (extra dimension) can be made relatively small and independent of code distance. In Sec. II, we highlight the surface code in more detail and include relevant considerations for physical resource estimation.

Here, we highlight some of the leading quantum computing platforms, their relevant error correction strategies, and contrast their rate of operations. The surface code is the front-running proposal for error correction in superconducting devices, and their associated code cycle times (the base sequence of hardware operations) have been estimated to be in the range of 0.2–10 μs. There is a great variety within different implementations of trapped ion architectures, particularly with regard to the method of enabling connectivity. A proposed scalable trapped ion design that relies on shuttling to enable connectivity and microwave-based gates has estimated the code cycle time to be 235 μs. For this shuttling-based design, alternative error correction protocols may be more effective than the surface code due to the variable-mid-range-connectivity that is possible, but in this work, we constrain ourselves to the surface code. Small trapped ion modules connected via photonic interconnects have been envisaged with the surface code, but due to their extremely flexible connectivity, higher dimensional error correction codes may one day be utilized. The code cycle time for trapped ions with photonic interconnects would depend on how the physical qubits are distributed across modules; one approach advocates for two tiers of encoding to minimize the use of the slower interconnects. Trapped ion architectures have achieved some of the highest gate fidelities to date, the lower the base physical error rate, the lower the code distance will need to be in the error correction protocol, and therefore, fewer physical qubits would be needed per logical qubit. A fault tolerant silicon-based architecture has been proposed using the surface code with code cycles estimated to be 1.2 ms. The error correction choice for photonic devices will depend on the underlying design; the primary candidate for a particular fault tolerant proposal is the Rausendorf-Harrington-Goyal (RHG) lattice. The degree to which an architecture is scalable will vary greatly between architecture types and within particular stages of development.

In this work, we provide physical resource estimates to achieve a quantum advantage with a quantum computer using the surface code. Using some of the latest time optimization strategies, we investigate the interplay between the code cycle time and the number of achievable physical qubits in the underlying architecture. In Secs. IA and IB, we introduce quantum algorithms, which have been hypothesized to offer a disruptive quantum advantage for industry-relevant applications. We first provide a brief overview of quantum computing for chemistry and highlight the logical resource requirements for a quantum advantage use case, simulating the ground state of the FeMo-co molecule, which we use as the starting point for our investigation. We then calculate the number of physical qubits that are required to break the elliptic curve encryption of Bitcoin keys within the time frame that it actually poses a threat to do, as a function of the code cycle time.

A. Fault tolerant quantum chemistry

When numerically simulating quantum chemistry problems, typically, a set of independent functions, known as a basis set, are introduced to describe the physical wave function of the system. This introduction does not remedy the exponential increase in parameters with system size but enables one to balance the achievable accuracy against the required computational resources. Feynman was perhaps the first to envisage a quantum computer and its application to the simulation of physics and chemistry. It is now expected that quantum computers will eventually be able to perform electronic structure calculations with a quality of solution typical to the most accurate
classical methods but with run times comparable to the approximate techniques, such as density functional theory.

The quantum phase estimation (QPE) algorithm generates eigenvalues for a general unitary operator, and it can be applied to quantum chemistry to find the eigenenergies of chemistry Hamiltonians to FCI (full configuration interaction, i.e., exact) precision. Unlike the variational quantum eigensolver (VQE)\(^\text{18}\) that involves many iterations \(O(1/e^\epsilon)\) with accuracy \(\epsilon\) of low depth circuits, the QPE algorithm requires \(O(1)\) iterations of a circuit with a depth scaling as \(O(1/\epsilon)\). The large depth required in the QPE algorithm means that it will only be possible with error corrected devices because NISQ devices would lose their coherence long before the end of the circuit.

Hamiltonian simulation is used as a subroutine in the quantum phase estimation (QPE) algorithm, and it involves constructing a quantum circuit that approximates the evolution of the input state according to the Hamiltonian. Two of the main paradigms for Hamiltonian simulation are trotterization and qubitization. Qubitization\(^\text{45,36}\) can be used to simulate the Hamiltonian evolution by using quantum signal processing,\(^\text{37}\) but more commonly, it is used to generate a quantum walk\(^\text{38}\) upon which one can directly perform phase estimation. Qubitization is perhaps the most favored method for simulating chemistry Hamiltonian dynamics because it achieves the provably optimal scaling in query complexity and approximation error albeit while requiring more logical qubits than other methods.

Previous work has investigated the potential for quantum computers to provide a quantum advantage by performing ground state energy estimations on the catalytic complex known as FeMo-co.\(^\text{39-42}\) FeMo-co is a large molecule expressed in biology, which reduces \(N_2\) from the atmosphere, and a better understanding of this process could provide a significant commercial advantage by improving the efficiency of nitrogen fixation for the production of ammonia for fertilizer. In this work, we start our investigation with the latest logical resource requirements for simulating FeMo-co and calculate the number of physical qubits required to reach a desirable run time as a function of the code cycle time of the hardware.

**B. Breaking Bitcoin’s encryption**

Bitcoin, the first decentralized cryptocurrency, is continuing to grow in popularity. Bitcoin has properties that make it desirable as a hedge against inflation, for example, the rate of supply is known, decreases with time, and is entirely independent of demand. The decentralized nature of the blockchain makes it censorship resistant, and it can operate in a trustless manner. There are two main ways in which a quantum computer may pose a threat to the Bitcoin network.\(^\text{43,44}\) The first and least likely is the threat to the proof of work mechanism (mining) for which a quantum computer may achieve a quadratic speedup on the hashing of the SHA256 protocol with the Grover’s algorithm.\(^\text{45}\) The algorithmic speedup is unlikely to make up for the considerably slower clock cycle times relative to state of the art classical computing for the foreseeable future.\(^\text{43,46}\) The second and more serious threat would be an attack on the elliptic curve encryption of signatures. Bitcoin uses the Elliptic Curve Digital Signature Algorithm (ECDSA) that relies on the hardness of the Elliptic Curve Discrete Log Problem (ECDLP), and a modified version of Shor’s algorithm\(^\text{33,47}\) can provide an exponential speedup using a quantum computer for solving this problem. Bitcoin uses ECDSA to convert between the public and private keys, which are used when performing transactions. With best practices (using new addresses for each transaction), the only point at which a public key is available and relevant to a eavesdropper is after a transaction has been broadcast to the network but prior to its acceptance within the blockchain. In this window, transactions wait in the “mem pool” for an amount of time dependent on the fee paid; the time taken for this process is on average 10 min, but it can often take much longer. Gidney and Ekerä estimated that it would require 20 \(\times 10^6\) noisy qubits and 8 h to break the 2048 Rivest-Shamir-Adleman (RSA) encryption,\(^\text{48}\) which is of a comparable difficulty to the EC encryption of Bitcoin. The maximum acceptable run time for breaking Bitcoin’s encryption makes it particularly well suited to our investigation into the cost of parallelization and the interplay between the code cycle time and scalability, which we present in Sec. III.

In Sec. II, we introduce considerations for error correction and provide an overview of the space and time optimizations within the surface code that we make use of in this work.

**II. SPACE AND TIME OPTIMIZATIONS IN THE SURFACE CODE**

In this section, we briefly introduce quantum error correction in the context of resource estimation and explain some of the available strategies within a surface code setup, which are selected based upon a preference for space (physical qubit requirement) or time (final run time of the algorithm).

**A. The available gate set**

An important consideration for quantum error correction is the available logical gate set, which is generally more restricted than the underlying physical gate set. The Clifford gates are those that map Pauli operators onto other Pauli operators, and the set can be generated by various combinations of the set \(\{H, CNOT, S\}\), where the \(S\) gate is the Pauli \(Z^{1/2}\). The Gottesman–Knill theorem\(^\text{49}\) states that any Clifford circuit of finite size can be simulated in polynomial time (efficiently) with a classical computer. The Clifford gate set in combination with any non-Clifford gate is sufficient for universal quantum computation, and two of the most commonly considered non-Clifford gates are the \(T\) gate \(Z^{1/4}\) and the Toffoli gate (control-control-not). Any arbitrary single qubit gate can be decomposed into long sequences of the fixed angle \(H\) and \(T\) gates with chain length scaling with desired precision, as per the Solovay–Kitaev theorem.\(^\text{50}\)

\[
T = \begin{pmatrix} 1 & 0 \\ 0 & e^{i\pi/4} \end{pmatrix}, \quad S = \begin{pmatrix} 1 & 0 \\ 0 & i \end{pmatrix},
\]

\[
H = \frac{1}{\sqrt{2}} \begin{pmatrix} 1 & 1 \\ 1 & -1 \end{pmatrix}, \quad CNOT = \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 \\ 0 & 0 & 1 & 0 \end{pmatrix}.
\] (1)

The surface code has transversal access to the \(CNOT\), and the \(H\) and \(S\) gates can be realized with a low overhead using other techniques.\(^\text{51}\) The \(T\) gate is not transversal in the surface code, and it must be effected using methods that incur a large space-time volume overhead relative to the other gates listed here. The \(T\) gate can be constructed by consuming a magic state, \(|m\rangle = (|0\rangle + e^{i\pi/4}|1\rangle)/\sqrt{2}\), where the magic state can be produced with an error proportional to the physical error, independent of the code distance.\(^\text{52}\) To create a
sufficiently high-quality magic state, a distillation protocol\textsuperscript{54,55} can be used, which essentially involves converting multiple low fidelity states into fewer higher fidelity states. Due to the high time cost associated with magic state distillation (the production and consumption), we can make a simplifying assumption that the time required to perform the T-gates effectively determines the final run time of an algorithm, as the relative cost of performing the Clifford gates fault-tolerantly is negligible. Some algorithms more naturally lend themselves to being expressed in terms of Toffoli gates, and there exist distinct specialized distillation factories to produce the magic states required to effectuate both of these non-Clifford operations. A Toffoli gate can be decomposed using 4 T gates,\textsuperscript{56} whereas the control-control-Z (CCZ) states normally produced for the Toffoli gate can be efficiently catalyzed into two T states.\textsuperscript{5}\

B. Error correction and logical error rate

A logical qubit in the surface code consists of data qubits, which store the quantum information, and ancillary qubits, which are used for stabilizer measurements that nondestructively extract error information from the data qubits. The distance, $d$, of a code represents the minimum size of physical error that can lead to a logical error, and in the surface code, the number of physical qubits per logical qubit scales as $2^d$. The logical error rate per logical qubit, $p_l$, per code cycle as a function of the base physical error rate, $p$, can be approximated by\textsuperscript{56}

$$p_l = 0.1(100p)^{(d+1)/2}.$$  

(2)

The efficiency of the error protection decreases as the base physical error rate approaches from below the threshold of the code (here assumed to be 1%). For feasible final resource estimates, the base physical error rate will need to be close to or below $10^{-3}$, where the necessary code distance is chosen based upon both the base physical error rate and the length of the desired computation. The physical error model here is the assumption that each physical operation has a probability $p$ of also introducing a random Pauli error. The achieved gate fidelity (the go-to metric for experimentalists) cannot be directly converted with confidence to the physical error rate, without further information. The cause of the gate infidelity, where it exists on the spectrum between the two extremes of depolarizing error (decoherent noise) and unitary error (coherent noise), will determine the corresponding gate error rate. The best-case situation is the one-to-one mapping between the gate error (one-fidelity) and depolarizing error rate, and this has often been an assumption in the experimentally focused literature. A measure of gate fidelity alone cannot determine the unitarity of the noise, i.e., the relative contribution of coherent and decoherent errors. Coherent errors, such as an over rotation of an intended gate, can positively interfere with the shuttling-based trapped ion architecture with a code cycle time of 235 $\mu$s, the quantum measurement is estimated to represent $\sim10\%$ of that total time. We include a code cycle independent additional cost of $10$ $\mu$s to represent the (somewhat) hardware independent classical processing. With our resource estimation tool, which is available upon request, one could recreate the results of this paper with a different relationship between the code cycle time and reaction time. If particular surface code set up contained only a single distillation factory, then the rate of computation would likely not be limited by the reaction time but instead by the rate of magic state production. We refer to the regime of being limited by magic state production as “tick” limited. There are then three relevant regimes for surface code strategies, which are separated by the limiting factor of computation rate. Beat limited implies that the limiting factor is the rate of magic state consumption
by the data block, tick limited, the rate of magic state production by the distillation blocks, and reaction limited, where the conjectured time-optimal limit is reached, and one reaction time is spent per measurement layer.

In this work, we utilize and compare two distinct strategies of incrementally trading qubits for run-time up to the reaction limit and introduce them later in this section.

D. Distillation and topological errors

When choosing a surface code setup, one must decide upon the acceptable final success probability, where, in principle, a success probability greater than 50% would be sufficient to reach the desired precision by repeating the computation. The acceptable success probability then allows one an additional method of trading space for time, as the lower the acceptable success probability, the lower the various code distances would need to be. There are two contributions to the probability of failure: the topological error associated with the data block and the total distillation error. The topological error is the chance for at least a single logical error within the data block across the entirety of the algorithm, which can be calculated with the product of the number of logical qubits, the number of code cycles required for the algorithm, and the logical error rate, where the logical error rate is defined in Eq. (2) by the base physical error rate and the code distance on the data block. The total distillation error corresponds to the probability of at least a single faulty magic state, which is calculated by the product of the total required number of required magic states and the error rate per state. The error rate per state is determined by the particular factory chosen and its associated code distances. We can then consider the final failure probability as the linear sum of the topological error and total distillation error.

In this work, we set the allowable total distillation error at 5%, implying that across the entire algorithm, the probability that a magic state is generated with an error is 5%. We choose the appropriate distillation protocol to achieve this error rate, by selecting between T factory protocols of Litinski and by adjusting the level 1 and level 2 code distances of the AutoCCZ factory. The choice of 5% is in part motivated by the capacity of the AutoCCZ factory to reach a sufficient fidelity, given the number of magic states required in the quantum advantage cases we address in this work. We set the final topological error to be 1% and choose the appropriate code distance to achieve this by considering the total number of code cycles that the algorithm must run for. This leads to a total final error (failure probability) of ~6%. In the Litinski’s work, a final error of 2% is chosen, whereas in the Gidney and Ekerä’s work of breaking RSA encryption, the final error is 33.4%. The best choice of final error tolerance will depend on the type of problem being solved, in the case where the result being chosen and its associated code distances. We refer to this method as entanglement swapping.

E. Routing at the error corrected level

It is necessary to move logical information from one area of the device to another, and perhaps the most common requirement is the transport of magic states from the distillation factories to the data blocks. Analogous to physical qubits in superconducting devices that use logical operations to perform swaps, one can imagine performing swap operations between logical qubits in the surface code. However, alternative techniques are more often considered for enabling long-range interactions with topological error correcting codes. Using lattice surgery-based methods, long range interactions (CNOTs) can be enabled in d code cycles (1 beat), essentially independent of the distance between the two points, so long as there is a chain of free ancillary qubits between them. We refer to this method as entanglement swapping. An ancillary logical qubit can only contribute to one routing chain at a time (per beat). When defining the layout of an error correction setup, which involves choosing the number of distillation factories and orienting them with respect to the data blocks, one must also ensure there is sufficient ancillary routing space to enable the required degree of data transfer between the relevant areas. The degree of data transfer, or alternatively stated the degree of parallelization in the execution of the algorithm, is often characterized by the number of magic states consumed per beat across the entire data block. With a data block arranged into rows with an access hallway between each row (consisting of ancillary logical qubits), there are two unique ways of touching each data qubit, if this is deemed insufficient, entangled copies of the data rows can be created to ensure there are enough unique paths between the factories and the data qubits. In Sec. IIHI, we present our choice of the routing overhead factor for entanglement swapping as a function of the degree of parallelization using AutoCCZ factories.

F. Considering physical mid-range connectivity

In this section, we briefly consider the potential impacts on the required resources when the underlying architecture has access to flexible mid-range connectivity between physical qubits. For superconducting devices, two qubit operations can only be performed between nearest neighbor physical qubits, and so long-range interactions must be enabled by sequences of costly logical swap operations. Alternative hardware may have access to low overhead long range interactions between physical qubits, for example, photons are readily transported long distances, which is relevant to both photon-only hardware and for connecting small modules of trapped ions with photonic interconnects. In the blueprint for a shuttling-based trapped ion architecture, a single system is envisaged, comprised of iterative micro-fabricated chips, connected via electric fields, which allow for physical shuttling between modules. High state fidelity adiabatic shuttling has been demonstrated with a speed of ~20 ms⁻¹, diabatic techniques can enable much greater shuttling speeds, and ~80 ms⁻¹ has been demonstrated. Lau and James calculate that the maximum speed a ⁴⁰Ca⁺ ion can be transported across a 100 μm trap without excessive error is 10 000 ms⁻¹. For NISQ size devices, all-to-all connectivity should be achievable with high fidelity (relative to two qubit operations) using the shuttling-only approach, but physically shuttling completely across a device with over a million physical qubits is unlikely to be feasible due to the associated time cost. Utilization of mid-range connectivity may still enable a reduction in the routing overhead associated with entanglement swapping.
While very long-range shuttling operations may be protected from error by periodic cooling operations and mid-circuit syndrome extraction and correction, the total time cost must be considered. With entanglement swapping, long-range interactions can be enabled between logical qubits in the surface code in a single beat, provided there are sufficient available ancilla qubits between the locations. To contrast this capability, we estimate the range at which physical shuttling may remain competitive with entanglement swapping. Assuming a code distance of 30, and logical qubits distributed across a 2D square grid, we estimate that a logical qubit could interact via physical shuttling with another logical qubit in the range of 3–30 grid spaces away within a single beat (d code cycles), depending on physical ion density and shuttling speed. While this is, indeed, unlikely to be sufficient for mediating all long-range interactions between logical qubits, the capability of low-cost mid-range physical connectivity could make the transversal CNOT preferable to the more usually considered lattice surgery-based methods. Gutiérrez et al. have investigated the experimental regimes at which the transversal CNOT may outperform the lattice surgery-based methods for trapped ions, and for particular values of error contributions, the transversal CNOT may be performed a factor $10 \times$ faster. If the transversal CNOT is expected to require less time than the lattice surgery-based CNOT, then this would directly impact the rate of magic state production of distillation factories and, therefore, could reduce the total qubit overhead in the highly parallelized regime. Furthermore, mid-range physical connectivity could considerably reduce the qubit footprint of distillation factories by eliminating the need for interior ancillary routing space for entanglement swapping. Alternative error correction strategies to the 2D surface code may be achievable on hardware with flexible mid-range connectivity. We leave a more detailed analysis of the potential benefit of mid-range connectivity as future work.

In Subsection II G, we introduce the Game of Surface Codes method of trading space for time, and in Subsection II H, the AutoCCZ method, where we also include some more detailed assumptions on the necessary routing overhead for entanglement swapping.

G. Game of surface codes

In the work of Litinski, various data blocks are presented, which vary in their rate of T-gate effectuation (magic state consumption) and the number of required physical qubits. There are numerous distillation protocols, each of which varies with regard to the output fidelity, required number of physical qubits, and the rate of production. Distillation blocks can be parallelized, and this enables further space-time trade-offs.

We make use of distillation strategies presented by Litinski, where the distance associated with the distillation blocks is fine tuned and separate from the distance associated with the data blocks. The data blocks have a required code distance set by the total number of logical qubits and the total number of code cycles required to run the entire algorithm. With the total number of T gates in the algorithm, $T_{\text{count}}$, the distillation blocks only need a code distance sufficient to produce magic states with an error lower than $1/T_{\text{count}}$. The distillation blocks use a certain number of qubits and only need to be protected for a certain number of code cycles (corresponding to one full round of distillation), and these are generally both small relative to the requirements of the data blocks. This method of individual calibration of distance for the data and distillation blocks is in contrast to a prior method, where both block types are attributed the same code distance.

In the GoSC scheme, Clifford gates are addressed explicitly via Clifford tracking, i.e., all Clifford gates are commuted to the end of the circuit and absorbed into measurements. This turns T gates into Pauli product rotations, and measurements into Pauli product measurements. In general, these Pauli product rotations can be big multi-qubit operations. To account for the generalized worst-case algorithm, which may have these multi-qubit operations, the maximum rate at which the data block can consume a magic state is defined as one state per beat (d code cycles). If the input circuit is known, then it will sometimes be possible to arrange the data blocks in such a way, so that more than one state can be consumed per beat. In this investigation, we do not consider the details of the input circuit and instead rely only on meta details, such as abstract qubit count, total T count, and the measurement depth. Our utilization of the GoSC method should then be considered an upper-bound configuration, i.e., guaranteed to be able to support any algorithm configuration given the meta details. The number of distillation factories can be chosen to match the production rate to the consumption rate of the data block, at which point, we may describe the strategy as beat limited.

In the work of Litinski, further time optimizations are presented, which can be utilized to reach reaction limit where one reaction time is spent per measurement (T) layer. The reaction limited strategy is set by the total number of measurement layers, i.e., the measurement depth ($T_{\text{depth}}$), as opposed to the total T gate count ($T_{\text{count}}$) of the beat limited strategy. The average number of parallel-executable T gates per layer ($T_{\text{layer}} = T_{\text{count}}/T_{\text{depth}}$) varies across particular algorithms, and there exist methods to optimize circuits to minimize either the $T_{\text{count}}$ and $T_{\text{depth}}$ or the circuit width.

The Litinski’s method of speeding computation up beyond the beat limited case utilizes “units,” which combine the previous constructions of data and distillation blocks. The number of units can be increased until the time optimal limit (henceforth reaction-limited) is reached, where each unit can work in parallel to address a set of measurement layers. This does not contradict the previously provided definition of measurement layers, as although the units can parallelize aspects of the work, the layers must still be stitched back together requiring one reaction time per measurement layer for corrections. The reaction limit then defines the maximum number of units that can be utilized. For an algorithm requiring $n$ abstract (logical) qubits, with an average of, $T_{\text{layer}}$, T gates per measurement layer, a single unit will consist of $4n + \sqrt{n} + 1$ tiles (logical qubits) for the data block, and $2T_{\text{layer}}$ storage tiles. Each unit requires an amount of time, $t_u$, to process a single measurement layer, where it is measured in beats and scales as $T_{\text{layer}} + \sqrt{n} + 3$. Each unit will need a number of distillation factories to match the required production rate, which is $T_{\text{layer}}$ magic states per unit completion time, $t_u$. With a linear arrangement of units as is considered here and a reaction time, $RT$, the reaction (time-optimal) limit is reached with a number of units $n_u$ equal to $t_u/RT + 1$. The GoSC scheme of going beyond the beat limited rate initially incurs a space-time overhead but then allows one to trade linearly (by increasing the number of units) until the reaction limit is reached.

H. AutoCCZ factories

In the work of Gidney and Ekerä, detailed surface code layouts along with the logical algorithmic developments for breaking RSA
encryption are provided. The surface code strategy utilizes AutoCCZ factories, which create a magic state that can be consumed to effectuate the non-Clifford Toffoli gate. The “auto” refers to the fact that these factories create auto-corrected CCZ states, meaning that the potential correction operation associated with magic state consumption is decoupled and can be performed far away from the data block.

Using this scheme with the AutoCCZ factories, the beat limited rate can be surpassed without introducing units as is performed in GoSC. Provided the routing overhead is accounted for, one can continue to add AutoCCZ factories until the production rate is equal to the number of Toffoli gates per measurement layer per reaction time, at which point the reaction limit is reached.

The AutoCCZ factory is characterized by two code distances, corresponding to the two levels of the protocol. The factory is a tiered distillation scheme where the output magic states of the first level are the responding to the two levels of the protocol. The factory is a tiered distillation scheme where the output magic states of the first level are the input states to the second level. We calculate the final output fidelity following the ancillary files of Girand and Ekerä as a function of the two code distances and the base physical error rate. To choose the optimal two code distances, we assess a wide range of possible values and select the setup that reaches the desired final distillation error rate while minimizing the space-time volume of the factory. While technically possible to maintain a reasonably low distillation error rate with base physical error rates near the threshold of the code, $10^{-2}$, the associated code distances required would result in an infeasible physical qubit overhead. Assuming any code distance is acceptable, the final distillation error per state, $p_{D}$, can be described as a function of the base logical error, $p$, by $p_{D} = 34.300\mu p$, until the threshold of the code is reached. This relationship was found by numerical fitting. A limit on the allow-able code distance breaks away from this trend before the threshold, and the lower the limit, the sooner it breaks upward. In Fig. 2(b), we investigate the impact of the base physical error rate for the final qubit overhead to reach a desirable run time with the AutoCCZ method. If the final output fidelity of the AutoCCZ factory is insufficient, given the desired length of an algorithm and the base physical error rate, then alternative T gate factories may still be viable.

To reach a particular desired run time, assuming it is below the reaction limit, the number of AutoCCZ factories is chosen as necessary. Once this footprint arrangement is settled (the combination of the data block and the number of factories), the last stage of the calculation is to determine the necessary routing overhead to account for the degree of parallelization. As described in Sec. II E, long-distance interactions between the distillation block and data block can be enabled in one beat, provided there is an available chain of ancillary logical qubits (routing space) between them. As each ancillary logical qubit can only contribute to one routing chain per beat, there may need to be additional unique paths to account for the degree of parallelization. All of the necessary routing overhead is strictly accounted for in GoSC, with the construction and arrangement of the data blocks, which are then duplicated and distributed across units. In our utilization of AutoCCZ factories, we define the degree of parallelization as the number of magic states consumed across the data block per beat. We then ensure that there is a routing hallway per data block row for every state consumed per beat, to exceed two hallways per data row; entangled copies of the data block can be made (which is comparable to the entangled copies across units in GoSC). Finally, we multiply the entire area (number of logical qubits) by a factor of 1.2 to ensure the distillation factories are surrounded by hallways and for some work space around the data block for arranging routing-chains. Our utilization of AutoCCZ factories should not be considered a true upper bound for any generalized circuit, unlike GoSC units. In Sec. III, we contrast the two methods further and state our relative contribution.

I. Problem specification

We start our investigation with the logical resource requirement set out by Lee et al. to simulate FeMoco to chemical accuracy and assess the feasibility of reaching desirable run times as a function of the code cycle time and number of achievable physical qubits. In the work of Lee et al., a detailed surface code strategy is presented with algorithm specific optimizations, such as minimizing the number of data qubits that are stored for working. In contrast, the surface code strategies we utilize do not necessarily require detailed knowledge of the input circuit and are instead only a function of the logical qubit count: the T (or Toffoli) count and the measurement depth. This approach may not yield optimal results for specific algorithms, but it enables one to effectively estimate physical resource requirements for a particular algorithm as a function of strategy (the time-space optimization spectrum) and the code cycle time. We contrast two strategies: the first that can be considered to provide upper bound resource estimates for any general circuit input and closely follows the work of Litinski’s Game of Surface Codes (GoSC). We go beyond the pedagogical examples provided by Litinski by creating an automatic tool that calculates the physical resources across the space-time optimization spectrum and apply it to the algorithmic requirements of quantum advantage use cases. Furthermore, we use the tool to calculate the number of physical qubits required to reach a specified run time as a function of the code cycle time of the hardware by utilizing the necessary degree of parallelization. We investigate the impact of varying the measurement depth with a fixed total T count, on the efficiency of the GoSC unit approach and identify the optimal value in particular regimes. In addition to the Game of Surface Codes method of parallelization, we incorporate a method that uses AutoCCZ factories. For this method, we relied upon the ancillary files of Girand and Ekerä and adapted them to be flexible enough for our broader (circuit agnostic) considerations. Our intention with the routing overhead with the AutoCCZ factories is to cover a wide range of possible circuit characteristics; while it may be an over estimation for some specific circuits, it may also represent an under estimation for some worst-case situations. We accomplish our aim of quick and general resource estimation as a function of algorithm meta information, and hardware characteristics, by contrasting the upper-bound scenario of GoSC units, with a more heuristic utilization of AutoCCZ factories. With detailed knowledge of the input circuit, further optimizations of the footprint configuration are possible, but generally, these must be performed on a case-by-case basis and are non-trivial to automate. The tool used to generate the results presented in this paper is available upon request. We use the latest logical resource requirements for breaking elliptic curve encryption and estimate the number of physical qubits required to break the encryption of Bitcoin keys in the small amount of time it would actually pose a threat to do so, as a function of both the code cycle time and base physical error rate.
III. RESULTS

To calculate the results presented in this section, we use various surface code strategies, including the Game of Surface Codes scheme, which uses units to parallelize layers of T gates and AutoCCZ factories, which are both highlighted in Sec. II.

A. Simulating FeMoco as a function of the code cycle time

There has been extensive research into both algorithmic development and resource estimation in the field of fault tolerant quantum chemistry, and one of the focus points has been the FeMo-co catalyst. An improved understanding of the FeMo-co catalyst could provide considerable efficiency improvements to nitrogen fixation, which currently represents around 2% of the world’s energy usage. 

We start with some of the latest algorithmic developments by Lee et al. and investigate the feasibility of achieving a reasonable run time for different code cycle times and different surface code strategies. The associated logical resources required are 2196 logical qubits and 6.7 $10^3$ Toffoli gates. The two scatter trends utilize AutoCCZ factories with different code cycle times of 1 $\mu$s, corresponding to a future superconducting device, and 235 $\mu$s, corresponding to a future shuttling based trapped ion device. Each trend starts with one AutoCCZ factory; from there, the number of factories is incremented, and at each step, the code distance is calibrated and the resulting run time and physical qubits are plotted. Initially, the total qubit footprint is dominated by the data blocks, whereas the run time is bottle necked by the magic state production, i.e., going from one factory to two halves the expected run time. Therefore, while this is the case, adding factories results in an improvement to the space-time volume, and this can sometimes allow for a reduction in code distance, which may result in an actual reduction in total qubit count. Each time a factory is added, the magic state consumption rate per beat is defined to determine whether the routing overhead needs to be increased as described in Sec. II H. What is considered to be a desirable run time will largely depend on the importance of the problem being solved and by the speed and quality of the classical alternatives. With the one AutoCCZ factory, the superconducting device completes in around 10 days with $7.5 \times 10^5$ qubits, whereas the trapped ion device requires 2450 days and the same number of qubits, where 10 days may be considered a quantum advantage for this use case where classical computers stand no chance of providing a meaningful answer, perhaps 2450 days would not.

The good news for hardware with slower code-cycle times is that it will often be possible to still reach desirable run times, provided enough physical qubits are available. However, the associated qubit
overhead may appear daunting and implies that hardware with slower code cycle times will have to be more scalable to compete, assuming equal error rates and physical connectivity. We plot for a range of possible measurement depths, labeled as a fraction of the total Toffoli count, as this was not provided along with the other logical requirements. In the AutoCCZ scheme, the measurement depth does not directly impact the efficiency of the approach, instead it only determines in combination with the reaction time, what the time optimal (reaction) limit is. The labels then indicate the reaction limit, the point at which the trend would end, given that measurement depth.

The assumption of a base physical error rate of $10^{-3}$ is often made in the literature and may be representative of two qubit gate fidelities that have already been achieved experimentally, depending on the potential caveats highlighted in Sec. II B. Trapped ion architectures have achieved some of the highest gate fidelities to date and, generally, exceed that of superconducting devices; for that reason, here, we provide the resource estimate for the more optimistic value of $10^{-4}$ for the base physical error rate, paired with the code cycle time that may correspond to future trapped ion devices. Instead of the $60 \times 10^6$ physical qubits that were required for a run time of 10 days, now with a base physical error rate of $10^{-4}$, only $60 \times 10^6$ physical qubits would be required. Although here the factor improvement of the base physical error rate is approximately equal to the factor improvement to the physical qubit requirement, this will not always be the case. The factor difference will depend on the proximity to the threshold of the code, and we investigate the impact of the base physical error in more detail in Fig. 2(b).

In Fig. 1, we also include the Game of Surface Codes (GoSC) approach to trading space for time, where measurement layers are parallelized with constructs called "units." Each unit contains its own copy of the data block and enough factories to produce the number of magic states within the measurement layer within the time it takes to prepare the unit, which scales with both the number of magic states per layer and the number of abstract qubits. Units can be incrementally added; each one added reduces the final run time up until the reaction limit is reached. The dashed lines in Fig. 1 use units along with improved T gate factories where it is assumed that 4 T gates are required to decompose a Toffoli gate. The GoSC approach is plotted only for the code cycle time of 235 μs, but three trends are included for the different measurement depths as a fraction of the total Toffoli count. The efficiency of the GoSC approach (in addition to the reaction limit) is dependent on the measurement depth as can be seen. In Sec. III C, we investigate the impact of the measurement depth on the final qubit requirement to reach a fixed run time for the GoSC approach.

It can be seen in the figure that the AutoCCZ approach provides more favorable final resource estimates than GoSC units in this scenario. The largest contributing factor to this difference appears to be the initial setup cost for the unit approach, which is nearly an order of magnitude increase in qubits for no appreciable speed up (with

\[ \text{FIG. 2. The number of physical qubits required to break Bitcoin’s 256 elliptic curve encryption with a fixed maximum run time as a function of the code cycle time and base physical error rate. Using the latest algorithmic development for quantum circuits for elliptic curve encryption, the depth optimized approach is chosen requiring } 5.76 \times 10^7 \text{ T gates, } 2671 \text{ logical qubits, and a T (measurement) depth of } 1.08 \times 10^7. \text{ These trends utilize AutoCCZ factories to trade space for time to reach the desired run time and assume that a CCZ state can be efficiently traded for 2 T gates. Assuming the relationship between reaction time (RT) and code cycle time (CC) of } RT = CC/4 + 10 \mu s, \text{ which is motivated in Sec. II. (a) Physical qubit requirement as a function of code cycle time for maximum run times of 10 min, 1 h, and 1 day. The trends start from the chosen code cycle time of } 10^{-3} \text{ and end at the right due to reaching the reaction limit before reaching the desired run time. We use a base physical error rate of } 10^{-3} \text{ and a final output error of } \sim 6\%. \text{ (b) Physical qubit requirement as a function of the base physical error rate for a maximum run time of 1 h and code cycle time of both 1 and 100 } \mu \text{s. The trends start from the chosen base physical error rate of } 10^{-3} \text{ and end at the right due to the AutoCCZ factory being no longer able to reach the desired distillation fidelity given the base physical error and number of required states.} \]
This is in stark contrast to the AutoCCZ approach, which initially takes very favorable space-time trades by increasing the number of factories. As the rate of parallelization increases in the AutoCCZ approach, eventually, entangled copies of the data block are made to maintain sufficient access hallways between the data block and distillation blocks. Both the AutoCCZ and GoSC approaches converge toward an equal linear trade between space and time. It should be restated that the GoSC approach can be considered a true upper-bound estimate, functional for any general circuit, whereas our utilization of the AutoCCZ factories is more heuristic and may represent an underestimation for some specific circuit inputs, see Sec. II for more discussion on this.

In Fig. 1(a), the beat limited method is included for both code cycle times as diamond and square points, i.e., no units and the computation rate are limited by the data blocks magic state consumption rate. These points use T gate factories and the fast data blocks from GoSC, which have size scaling at $2^n + \sqrt{8n} + 1$ for n logical qubits and can effectuate T gates at a maximum rate of one per beat (d code cycles). The number of T factories is chosen to match the rate of magic state consumption of the fast data blocks. The beat limited situation provides comparable run times to the single AutoCCZ factory and requires a similar number of physical qubits.

To conclude this section, it appears that the AutoCCZ factories are the favorable approach to trade space for time up to the reaction limit, but a more detailed investigation into the underlying assumptions of both methods is warranted. These resource estimates are solely a function of algorithm meta information such as total T count, measurement depth, and the number of logical qubits. The surface code configuration can be optimized when paired with detailed knowledge of the input algorithm, but this process is non-trivial to automate, as we have done in this work as a function of code cycle time. Future hardware that expects to have considerably slower code cycle times than the superconducting devices may still be able to reach desirable run times, provided enough physical qubits are available, which further emphasizes the importance of scalability. The associated qubit overhead factor will range between less than 1 (here ~0.3) and 1, times the difference in the code cycle time depending on the relative degree of parallelization in the comparison. Algorithms should be optimized by minimizing the measurement depth if the reaction limit is restrictive. The physical qubit requirement may be reduced if the underlying hardware has access to low overhead mid-range physical connectivity, as discussed in Sec. II.

**B. Breaking Bitcoin’s EC encryption**

Breaking encryption has received a lot of attention in the quantum computing community since Shor’s breakthrough algorithm, which provides a near exponential speedup for prime factoring that has direct implications for breaking RSA encryption. 1

Ekerä et al. provide algorithmic improvements in addition to the surface code strategies for breaking RSA encryption, and they estimate that \(20 \times 10^9\) qubits running for 8 h could break it with a code cycle time of 1 \(\mu s\). In a blueprint for a shuttling-based trapped ion device, which estimated the code cycle time to be 235 \(\mu s\), it was originally estimated that breaking RSA encryption would require 110 days and 2 \(\times 10^9\) qubits with a base error of \(10^{-3}\), implying a device occupying an area of \(103.5 \times 103.5\) m\(^2\). With the latest algorithmic and surface code strategy improvements, we can reduce this estimate to requiring instead a run time of 10 days (a factor 10× faster) and \(650 \times 10^9\) qubits, which would imply a device size of area 80 \times 80 m\(^2\). With a base physical error of \(10^{-4}\), the device size would reduce to \(18 \times 18\) m\(^2\), and further reductions may be possible if one were to make use of the flexible mid-range connectivity that is available. Using the same relative improvement factor from mid-range connectivity as in the design blueprint, starting from the \(10^{-4}\) case, we might expect a device of size 2.5 \times 2.5 m\(^2\) to be sufficient. This is, indeed, a rough estimate, and a more rigorous understanding of how to best make use of mid-range connectivity would be required to confidently provide the performance improvements relative to a nearest neighbor approach.

Bitcoin uses the Elliptic Curve Digital Signature Algorithm (ECDSA), which relies on the hardness of the Elliptic Curve Discrete Log Problem (ECDLP), and a modified version of Shor’s algorithm46–48 can provide an exponential speedup using a quantum computer for solving this problem. The encryption of keys in the Bitcoin network is only vulnerable for a short window of time, around 10 min to an hour depending on the fee paid, as described in more detail in Sec. 1B, and this makes it a well-suited problem for our investigation.

In the work of Aggarwal et al.43 the potential threat of a quantum attack on the Bitcoin network is investigated, and two main vulnerabilities are contrasted, i.e., proof of work (mining) and the encryption of private keys. Following the conclusions of Aggarwal et al., we highlight the differences between these two vulnerabilities and the associated quantum speed up in Sec. 1B. Here, we focus on the most likely threat, the encryption of private keys, and go beyond the physical resource estimates of Aggarwal by including the latest improved logical algorithmic requirements of Håner et al.47 and by quantifying how the number of physical qubits changes based on the desired run time and the code cycle time of the hardware. In the following, we will present our resource estimates for this problem and also contrast with the work of Aggarwal et al.43

In Fig. 2(a), we plot the number of physical qubits required to break the elliptic curve encryption of Bitcoin within a run time of 1 day, 1 hour, and 10 min, as a function of the code cycle time. We use the logical resources of the depth-optimized approach provided by Håner et al.41 for a 256 bit encryption, which corresponds to \(5.76 \times 10^9\) T gates, 2871 logical qubits, and a measurement depth (\(T_{\text{depth}}\)) of \(1.88 \times 10^7\). The measurement depth of this algorithm is low relative to the \(T_{\text{count}}\) at \(\sim 10^8\), implying that there is a lot of room for parallelization before the reaction limit is reached. In Fig. 2(a), it can be seen that it would require \(317 \times 10^9\) physical qubits to break the encryption within one hour with a code cycle time of 1 \(\mu s\). To break it within 10 min with the same code cycle time, it would require \(19 \times 10^9\) physical qubits, whereas to break it within 1 day, it would require only \(13 \times 10^9\) physical qubits. The horizontal period most evident in the dash-dotted black trend for a run time of 1 day is because the desirable run time can be reached for those code cycle times with only 1 AutoCCZ factory. Once the code cycle time increases sufficiently, it is then necessary to begin adding AutoCCZ factories to maintain the desired run time. Hardware with considerably slower code cycle times than 1 \(\mu s\) will need to be able to reach larger device sizes to break the encryption within the allotted time. Even for code cycle times of 1 \(\mu s\), this large physical qubit requirement implies that the Bitcoin network will be secure from quantum computing attacks for many years. High-value transactions are likely to pay high fees, ensuring that
they are processed with higher priority and, therefore, would require considerably more physical qubits to break the encryption in time. The Bitcoin network could nullify this threat by performing a soft fork onto an encryption method that is quantum secure, where Lamport signatures are the front-running candidate, but such a scheme would require much more memory per key. The bandwidth of Bitcoin is one of the main limiting factors in scaling the network, and so, changing the encryption method in this way could have serious drawbacks.

The logical resources provided by Häner et al.\textsuperscript{47} for breaking elliptic curve encryption improve on the prior state of the art of Roetteler et al.\textsuperscript{46} by over an order of magnitude. In the quantum threat to Bitcoin work of Aggarawal et al.,\textsuperscript{1} the older and less favorable logical resource requirements were considered.\textsuperscript{3} Aggarawal et al. estimate that it would require 6.5 days and \(1.7 \times 10^6\) physical qubits to break the encryption with a base physical error rate of \(5 \times 10^{-4}\). The code cycle time is not explicitly defined, and instead, a physical gate rate of 66.6 MHz is assumed, which we estimate would correspond to a code cycle time of approximately 0.1 \(\mu\)s. Next, we calculate the physical resources using the assumptions and logical requirements of Aggarawal et al., and we find that a device with three AutoCCZ factories would complete in seven days and require \(5 \times 10^6\) physical qubits. There is rough agreement between the final physical resources between our methods; the remaining discrepancy originates from the differing estimates of the number of physical qubits that are required per logical (abstract) qubit. The conversion factor between logical to physical qubits of Aggarawal et al. is stated to be 735.5 for this problem, which should include the overhead associated with the degree of encoding (code distance), distillation factories, and routing space. We find that a code distance of 25 is required to maintain a final failure rate below 6\%, implying at least \(2 \times d^2\), or 1250, physical qubits per logical qubit. When we include the distillation and routing overhead, our final physical to logical qubit conversion factor is 2140. While a higher final failure rate may be tolerable in a heralded problem such as breaking encryption, we estimate that a code distance of at least 22 would be required, as a code distance of 21 or lower would lead to final failure rates of \(\sim 100\%\).

Häner et al.\textsuperscript{47} provide asymptotic formulas for the resources required to break the elliptic curve encryption as a function of the input encryption bit size, \(n\) (Bitcoin uses 256-bit encryption). The total logical space-time volume (\(T\) count \times logical qubits) scales to leading order with encryption bit size as \(O(n^2/\log n)\), which can be compared to the exponential time complexity of classical methods, \(O(2^n/\log n)\). We calculated the physical resources to break the elliptic curve encryption as a function of bit length, and one may expect the results to follow the same form of dependence as the total logical space-time volume. This is because we would expect the total physical qubits required to scale linearly with the total logical space-time volume, and this is generally the case except for variations due to differing efficiencies of the error correction set up at different regimes (ratio of the logical qubits to gate requirement and the degree of parallelization required). We fit the physical resources as a function of bit length with a function containing the terms that result from multiplying the asymptotic trends of \(T\) count and logical qubit count from Table 4 in the work of Häner et al.\textsuperscript{1} The generated trend fit the data well for bit string sizes in excess of 256, with some discrepancies below this value, which is likely due to the minimal degree of parallelization that was required for lower bit strings. For a code cycle time of 1 \(\mu\)s, a base physical error rate of \(10^{-3}\), and a desired run time of 0.1 day, the total physical qubits required to break the encryption as a function of bit length, \(n\), are well described by the following equation as \(0.01 \times n^{1.91}\). The fit coefficients here are a function of the desired run time and code cycle time of the hardware, which dictate the required degree of parallelization. For 250 data points of bit length in the range of 100–2000, the average percentage error between the data points and the trend was 3.7\%. The run time was maintained below 0.1 days with these hardware assumptions up to a bit length of 1730, albeit requiring \(4 \times 10^{19}\) physical qubits, at this point, the maximum degree of parallelization was reached. Hence, we show that the required number of physical qubits to break the elliptic curve encryption in a defined fixed duration scales only polynomially, with the bit length of the encryption, in contrast to classical methods that require exponential time.

In Fig. 2(b), we plot the required number of physical qubits to break the 256-bit elliptic curve encryption within 1 h as a function of the base physical error rate. In Sec. II B, the relationship between the base physical error rate and an experimentally achieved gate fidelity is explained in more detail. We plot two trends for code cycle times equal to 1 and 100 \(\mu\)s, which begin at the selected error rate of \(10^{-5}\). The trends end at the right at a physical error rate of \(\sim 2.8 \times 10^{-3}\) when the AutoCCZ factory can no longer produce a state with sufficient fidelity given the error rate and number of required states. High code distance and multi-tiered \(T\) gate factories would be able to continue further, but at a value of the 1\% error rate, the threshold of the surface code is reached. With a physical error rate higher than this threshold, increasing the code distance actually results in a larger logical error. Three distinct distances are calibrated according to the base physical error rate; first, the final topological error is maintained below 1\% by adjusting the code distance associated with the data block. Next, there is a level 1 code distance and a level 2 code distance associated with the AutoCCZ factory, which are calibrated to ensure that the final distillation error is maintained below 5\%, as per the ancillary files of Gidney and Ekker.\textsuperscript{3} We assess a wide range of code distances for the AutoCCZ factory and choose the set that minimizes the factory volume (i.e., number of qubits \times duration per cycle) while maintaining the desired error rate. With a code cycle time of 1 \(\mu\)s, it requires \(317 \times 10^6\) physical qubits to reach the 1 h run time with a base error of \(10^{-3}\); this is reduced down to \(33 \times 10^6\) for a base error of \(10^{-4}\), i.e., a factor 10 reduction. The relative reduction in the qubit overhead, associated with an order of magnitude improvement in the base error, is greater when the comparison is performed closer to the threshold of the code. For example, from \(2.8 \times 10^{-3}\) to \(2.8 \times 10^{-4}\), the qubit reduction is instead a factor 30 (in contrast to the factor 10 of the previous comparison). This highlights the importance of reaching base physical error rates of \(10^{-3}\) and lower.

C. Finding the optimal measurement depth

Logical algorithms may be optimized for particular properties, for example, either the total \(T\) gate count, \(T_{\text{count}}\), the number of measurement layers, \(T_{\text{depth}}\), or logical qubit count may be minimized.\textsuperscript{47} In the elliptic curve encryption breaking algorithm of Häner et al.,\textsuperscript{47} logical requirements are stated for each of these three possible optimizations, where in Sec. III B, the measurement depth minimized approach was chosen. The reaction limit (conjectured to be the fastest an algorithm can be run) is determined solely by the measurement depth and reaction time (i.e., independent of total gate count and code
distance), and so, the depth optimized approaches are the most suitable when room for parallelization is desired. The ratio of the measurement depth to total gate count is the inverse of the number of $T$ gates per layer, $T_{\text{layer}}$ (when considering $T$ gates as opposed to some other non-Clifford operation). In the GoSC method of parallelization with units, all aspects of the cost depend on the number of $T$ gates per layer, including the footprint of the unit, the time it takes to prepare a unit, and the number $T$ gates that are effectuated within the preparation time.

In Fig. 3, we plot the efficiency of the GoSC method as a function of this measurement depth with a fixed total gate count, and it can be seen that there is a measurement depth that leads to a minimum physical qubit footprint. This is in contrast to the AutoCCZ method, where, instead, the measurement depth only plays a role in determining the reaction limit, and it would appear in these figures as a horizontal line. In Fig. 3(a), we plot the required number of physical qubits to reach a desired run time for the logical resources required to simulate FeMoco. We include three plots for different code cycle times and maximum run times. As expected in this regime prior to reaction-limited, it can be seen that it is the ratio of the code cycle time and maximum run time that determines the physical qubit requirements. We show that the optimal measurement depth ratio is not necessarily “as small as possible,” and for this situation, it lies between $10^{-3}$ and $10^{-2}$. The discontinuous movements result from an increase in the required number of units as the measurement depth ratio becomes larger. A less demanding run time requirement necessitates fewer units for a given measurement depth, which results in less frequent and larger relative downward movements. The two distinct discontinuous movements on the blue trend (for a code cycle time of 10 $\mu$s and run time of 1 day) are due to the topological code distance changing as the space time volume changes. The black trend with a run time of 10 days can maintain the desired run time for larger measurement depths than the dotted orange line because the proportional difference in the desired run time (a factor 10) is greater than the proportional difference in their associated reaction times. The relationship between the code cycle time and reaction time that we assume in this work is explained in Sec. II C.

In Fig. 3(b), we again investigate the efficiency of the parallelization but now for an abstract algorithmic requirement with fixed space-time volume. Three trends are shown with logical qubits, $n$, corresponding to 500, 1000, and 2000, where the total $T$ gate of the algorithm is set to maintain the fixed space time volume $(n \times T_{\text{count}})$ of $5 \times 10^{13}$. The trends end at the right when the reaction limit is reached where 2000 is the last to end because it has the lowest total $T$ gate count, which, in turn, relates to a lower reaction limit for a fixed measurement depth ratio. The optimal measurement depth ratio is larger for lower logical qubit counts; for 500, the optimal is $0.6 \times 10^{-3}$, whereas for 2000, the optimal is $0.3 \times 10^{-3}$. In the following, we investigate the relationship between the optimal measurement depth and logical qubit requirement of the algorithm in more detail.

1. Optimal measurement depth and logical qubit requirement

We have shown that the optimal measurement depth given a fixed total gate count for the GoSC approach is non-trivial and can be

![Fig. 3](image-url)
a function of both the number of logical qubits, and on how demanding the desired run time is (i.e., the number of units needed). In this section, we investigate the relationship between the optimal measurement depth and the number of logical (abstract) qubits of the algorithm. The average number of T gates per measurement layer, \( T_{\text{layer}} \), is the inverse of the previously stated ratio of measurement depth to total gate count, i.e., \( T_{\text{layer}} = \frac{T_{\text{count}}}{T_{\text{depth}}} \). In Fig. 4, for a particular value of the total T gate count, \( T_{\text{count}} \), we calculate the optimal value of T gates per layer, \( T_{\text{layer}} \), which is then converted into a ratio of the number of logical qubits. We include multiple trends for different numbers of logical qubits and plot as a function of the total T gate count. This involves identifying the value of \( T_{\text{layer}} \) at which the physical qubit requirement is minimized as in Fig. 5, for each particular logical resource requirement. We choose different qubit numbers and minimum run time values to highlight the various dependencies. We can see four distinct behavioral phases for each trend; first, at low T gate counts, the minimum run time can be reached with no parallelization, and so, we state the optimal T per layer as 0 (whereas truly it is independent). The next phase is an oscillating pattern where the optimal T per layer ratio varies widely with a decreasing amplitude as the T gate count increases. The beginning of this phase is determined by the end of the previous one, where the no-parallelization (beat limited) method can no longer reach the desired run time, which is determined by the ratio of the code cycle time and minimum run time. The amplitude of the oscillation is largest at the start of parallelization because only a small number of units are initially required, and an increase in T count requirement can mean the number of required units increases, for example, the optimal number of units may change from 3 to 4, but the associated \( T_{\text{layer}} \) for this minimum can vary widely, and the magnitude of that variation reduces as the overall number of required units increases. The large variation in the optimal ratio does not imply a large variation in the required number of physical qubits. In the next phase, the optimal T per layer ratio is relatively constant at an equilibrium value, which is solely determined by the number of logical (abstract) qubits in the algorithm, where the greater the number of logical qubits, the fewer T gates per layer (as a percentage of the logical qubits) are required for the optimal physical qubit overhead. For logical qubit numbers (N) in excess of 100, the equilibrium value of the optimal number of T gates per layer is well described by the following equation: \( T_{\text{layer}} = 1.9(5)N^{0.70(5)} \), which we estimated by taking an average of the points within the equilibrium phase and fit with linear regression. The final phase starts when the algorithmic requirement becomes too demanding for the minimum run time, and so, the \( T_{\text{layer}} \) rises despite the loss in efficiency until the number of logical qubits is reached, which, here, we define as the cutoff point, where finally even the reaction limit (one reaction time per T layer) is not sufficient.

The potential degree of control over the average number of T gates per layer during algorithm construction and optimization will determine whether it is beneficial to consider the optimal value as calculated with the techniques used in Fig. 4. There are optimization techniques that can minimize either the \( T_{\text{depth}} \) and \( T_{\text{count}} \), but they generally trade-off with one another, where minimizing one may increase the other. Furthermore, this analysis is specific to the GoSC approach of parallelization with units, and earlier, we have shown that

![Figure 4](https://via.placeholder.com/150)

**Fig. 4.** The optimal value of the number of T gates per measurement layer, shown as a fraction of the number of logical qubits, plotted as a function of the T gate count of the algorithm. Base physical error of \( 10^{-3} \) and code cycle time of 1 \( \mu s \). Different plots for varied desired run times and logical qubit count, 1 day and 100 qubits in blue, 1 day and 1000 qubits in orange, 1 day, 0.1 day, and 1000 qubits in green, and 10,000 qubits in black. The maximum value of T gates per layer is limited to the number of logical qubits for each trend. There are four distinct behavioral phases for each trend; first, the trends read 0 at the left, while the beat limited approach can reach the desired run time, and then units are introduced following the GoSC method. The next phase is an oscillating pattern where the optimal T per layer ratio varies widely with a decreasing amplitude as the T gate count increases. The amplitude of the oscillation is largest at the start of parallelization because only a small number of units are initially required, and an increase in T count requirement can mean the number of required units increases, for example, the optimal number of units may change from 3 to 4, but the associated \( T_{\text{layer}} \) for this minimum can vary widely, and the magnitude of that variation reduces as the overall number of required units increases. The large variation in the optimal ratio does not imply a large variation in the required number of physical qubits. In the next phase, the optimal T per layer ratio is relatively constant at an equilibrium value, which is solely determined by the number of logical (abstract) qubits in the algorithm, where the greater the number of logical qubits, the fewer T gates per layer (as a percentage of the logical qubits) are required for the optimal physical qubit overhead. For logical qubit numbers (N) in excess of 100, the equilibrium value of the optimal number of T gates per layer is well described by the following equation: \( T_{\text{layer}} = 1.9(5)N^{0.70(5)} \), which we estimated by taking an average of the points within the equilibrium phase and fit with linear regression. The final phase starts when the algorithmic requirement becomes too demanding for the minimum run time, and so, the \( T_{\text{layer}} \) rises despite the loss in efficiency until the number of logical qubits is reached, which, here, we define as the cutoff point, where finally even the reaction limit (one reaction time per T layer) is not sufficient.
the AutoCCZ method of parallelization produces more favorable final resource estimates. As mentioned, the AutoCCZ method does not display this rich behavior with the efficiency dependence on the measurement depth, and we believe that further research is warranted to compare the underlying assumptions of these two methods of parallelization.

**IV. CONCLUSION**

Within a particular time frame, the code cycle time and the number of achievable physical qubits may vary by orders of magnitude between hardware types. When envisaging a fault tolerant implementation, there are numerous decisions to be made based on a preference for either space or time. In this work, we compare surface code strategies of parallelization that allow one to speed up the computation until the reaction limit is reached. Most of the fault tolerant resource estimation work has focused on code cycle times corresponding to superconducting architectures. A space optimized quantum advantage case study translated for hardware with slower code cycle times may lead to run times in excess of 1000 days, and so parallelization would have to be performed to reach desirable run times. In this work, we have calculated the required number of physical qubits to reach a given desirable run time for two representative quantum advantage cases (chemistry and encryption) across a range of code cycle times. The feasibility of using these time optimization strategies will depend upon the number of physical qubits achievable within a device; therefore, the scalability of an architecture will play an important role in determining whether a quantum advantage is achievable. We contrast two methods of parallelization to simulate the FeMo-Co catalyst; first, a Game of Surface Codes approach that should be considered an upperbound, and second, a more heuristic utilization of AutoCCZ factories. We find in this situation that the AutoCCZ factories produce more favorable resource estimates, and the difference is mostly due to the high initial set up cost of parallelization with Game of Surface Code Units. With a single AutoCCZ factory, a superconducting device with a 1 μs code cycle time would require $7.5 \times 10^6$ qubits to simulate FeMo-co in ~10 days, whereas a shuttling based trapped ion device with a 235 μs code cycle time would take 2450 days. By increasing the number of factories, the space-time trade is initially favorable (as opposed to linear), and the trapped ion device can reach the same 10 day run-time with 600 $\times 10^6$ qubits. In this comparison, the factor difference between the physical qubit requirement is $\sim 3 \times$ less than the factor difference in the code cycle time. Trapped ion architectures have generally been shown to achieve higher two qubit fidelities than superconducting devices, and so we have calculated that if a base physical error rate of $10^{-4}$ was achieved, the trapped ion device would instead require $60 \times 10^6$ qubits for the 10 day run time.

We have investigated the effect of varying the ratio of the T gate count and T depth (the average T gates per layer) and identified the optimal value for a constrained run time against general algorithmic requirements. Here, we focused on the GoSC unit method of parallelization, as it was unique in displaying rich T depth dependence.

We apply our methods to the logical resources required to break the 256 elliptic curve encryption, which is used to secure public keys in the Bitcoin network. We use the logical resource requirements of the latest algorithmic developments, which improve on the previous state of the art by $\sim 2$ orders of magnitude. There is a small window of time, approximately 10–60 min, in which the public keys are available and vulnerable after the initiation of a transaction. We quantify the number of physical qubits required to break the encryption in one hour as a function of code cycle time and the base physical error rate. It would require approximately $317 \times 10^6$ physical qubits to break the encryption within one hour using the surface code and a code cycle time of 1 μs, a reaction time of 10 μs, and physical gate error of $10^{-3}$. To instead break the encryption within one day, it would require only $13 \times 10^6$ physical qubits. If the base physical error rate was instead the more optimistic value of $10^{-4}$, $33 \times 10^6$ physical qubits would be required to break the encryption in 1 h. This large physical qubit requirement implies that the Bitcoin network will be secure from quantum computing attacks for many years (potentially over a decade). Alternative error correction techniques, in particular those that benefit from a more flexible physical qubit connectivity as often found in trapped ion-based quantum computers, could potentially offer considerable improvements to the requirements, but the lower rate of logical operations must also be factored in. The Bitcoin network could nullify this threat by performing a soft fork onto an encryption method that is quantum secure, but there may be serious scaling concerns associated with the switch. We hope to motivate continued research into end-to-end resource estimation for alternative error correction schemes to the surface code, and to determine how best to make use of the available physical connectivity of different quantum hardware platforms.

**ACKNOWLEDGMENTS**

This work was supported by the U.K. Engineering and Physical Sciences Research Council via the EPSRC Hub in Quantum Computing and Simulation (No. EP/T001062/1), the U.K. Quantum Technology hub for Networked Quantum Information Technologies (No. EP/M013243/1), the European Commission’s Horizon-2020 Flagship on Quantum Technologies under Project No. 820314 (MicroQC), the U.S. Army Research Office under Contract No. W911NF-14-2-0106, the Office of Naval Research under Agreement No. N62909-19-1-2116, and the University of Sussex. We thank Craig Gidney, Daniel Litinski, and Niel de Beaudrap for helpful discussions and for their suggestions to improve the manuscript.

**AUTHOR DECLARATIONS**

**Conflict of Interest**

The authors declare no conflicts of interest.

**DATA AVAILABILITY**

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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