arXiv:2411.08955v1 [quant-ph] 13 Nov 2024

Fermion-qubit fault-tolerant quantum computing

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(Dated: November 15, 2024)

Simulating the dynamics of electrons and other fermionic particles in quantum chemistry [1], material science [2], and high-energy physics [3] is one of the most promising applications of fault-tolerant quantum computers. However, the overhead in mapping time evolution under fermionic Hamiltonians to qubit gates [4-7] renders this endeavor challenging [8, 9]. We introduce fermion-qubit fault-tolerant quantum computing, a framework which removes this overhead altogether. Using native fermionic operations we first construct a repetition code which corrects phase errors only. We then engineer a fermionic color code which corrects for both phase and loss errors. We show how to realize a universal fermionic gate set in this code, including transversal Clifford gates. Interfacing with qubit color codes we realize qubit-fermion fault-tolerant computation, which allows for qubit-controlled fermionic time evolution, a crucial subroutine in state-of-the-art quantum algorithms for simulating fermions [10–13]. We show how our framework can be implemented in neutral atoms, overcoming the apparent inability of neutral atoms to implement nonnumber-conserving gates by introducing a neutral-atom braiding gate using photodissociation of bosonic molecules. As an application, we consider the fermionic fast Fourier transform, an important subroutine for simulating crystalline materials [12], finding an exponential improvement in circuit depth from $\mathcal{O}(N)$ to $\mathcal{O}(\log(N))$ with respect to lattice site number N and a linear improvement from $\mathcal{O}(N^2)$ to $\mathcal{O}(N \log(N))$ in Clifford gate complexity compared to state-of-the-art qubit-only approaches [14]. Our work opens the door to fermion-qubit fault-tolerant quantum computation in platforms with native fermions such as neutral atoms, quantum dots and donors in silicon, with applications in quantum chemistry, material science, and high-energy physics.

Solving the dynamics of many strongly-interacting fermionic quantum particles is central to numerous scientific and technological challenges, including explaining the dynamics of quarks in the early universe, understanding the reaction dynamics of molecules, and elucidating high-temperature superconductivity. While qubit quantum computers promise to help solve these open problems, simulating the evolution of fermions encoded in qubits encounters a large overhead in both the number of

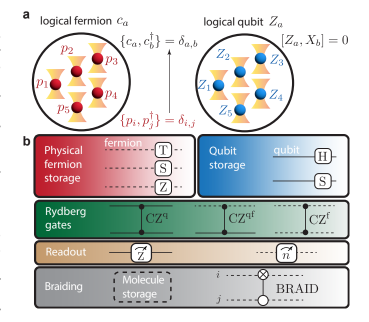


FIG. 1. Logical fermions and architecture. a, A logical qubit is encoded in a codeblock formed by many physical qubits. Similarly, we encode a logical fermion into a codeblock formed by many physical fermions. While logical qubit operators commute, logical fermionic operators anti-commute between two codeblocks. Physical fermions can be encoded in fermionic neutral atoms trapped in optical tweezers or lattices (indicated by hour-glass shapes) or in electrons trapped by electric fields in quantum dots or donors in silicon (Methods). b, Neutral-atom architecture, circuit notation, and universal qubitfermion gate set. Single-fermion gates are performed by detuning the tweezer potential; single-qubit gates by Raman lasers; CZ gates by Rydberg lasers; braiding gates by photodissociation of molecules as well as inter-tweezer tunneling. In the braiding gate, the site i, c.f. definition in Eq. (4), is indicated by a cross. We distinguish fermion sites from qubits by using dashed lines and drop the ^f superscript in circuits.

two-qubit gates and the circuit depth [5–7], which means that a solution of fermionic problems with near-term [8, 9] and even fault-tolerant quantum computers [12] is extremely challenging. It is possible to circumvent this issue by using quantum simulation platforms which host controllable fermionic degrees of freedom, such as quantum dots [15], donors in silicon [16], and light-trapped neutral atoms [17–20]. Using those platforms, analog simulations of condensed-matter models have been achieved [21–25] and proposals for studying quantum

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chemistry in an analog [26] or digital [27] way have been made. However, turning these fermionic platforms from a tool of scientific exploration into a computational tool requires error-correction and fault-tolerance techniques, for which no practical scheme has been found so far.

Here, we close this gap by introducing a framework for fermion-to-fermion fault-tolerant quantum computation. We first prove a theorem that fermionic error correction is not possible using only number-conserving gates, which are most easily implemented in experimental platforms. We then show how to circumvent this theorem by using a non-number-conserving gate and concepts from Majorana error correction. This enables us to introduce error-correcting codes for which we explicitly show how to perform a universal gate set including fault-tolerant transversal gates. To enable usage of state-of-theart quantum algorithms requiring qubit-controlled gates [10], we introduce qubit-fermion fault-tolerant computation. We apply our framework to time evolution under the Hamiltonian of a crystalline material and study the advantages of our framework compared to all-qubit quantum computation. Finally, we show how to realize these codes and their operations in neutral atoms by removing the previously assumed restriction to number-conserving gates [27] in neutral atoms and introducing the physics behind the implementation of a non-number-conserving braiding gate.

FERMION-TO-FERMION ERROR CORRECTION GATE SET

Our central goal is the construction of an error correcting code that encodes a logical fermion, which we define as a creation operator c_a^{\dagger} anti-commuting between different codeblocks a and b, i.e. $\{c_a^{\dagger}, c_b\} = \delta_{ab}$, see Fig. 1a. This is by contrast to fermionic operators encoded into a single codeblock of a topological error correcting code [7, 28, 29], which are difficult to address via transversal gates [30] and whose efficiency degrades when applied to fermion models that are not local in 2D. Constructing logical operators that anticommute between different codeblocks is impossible using physical qubits because no qubit operator can be constructed which anticommutes between separate codeblocks. Instead, we use physical fermionic degrees of freedom p_i^{\dagger} acting on physical sites *i* and fulfilling $\{p_i^{\dagger}, p_j\} = \delta_{ij}$ to encode logical fermions. A physical fermion can be realized for example by a fermionic atom trapped with light or an electron in a quantum dot. For concreteness, we will focus on the former implementation and comment on a quantum dot implementation in the Methods. As a second crucial ingredient, we interface these fermions with qubits, which are encoded in two internal energy levels of an atom.

We first consider a fermion-number-conserving gate set acting on many fermionic and qubit sites labeled by indices i, j,

$$\mathbf{T}_{i}^{\mathrm{f}} = \exp\left(i(\pi/4)n_{i}\right),\tag{1}$$

$$CZ_{ij}^{f} = \exp\left(i\pi n_{i}n_{j}\right), \qquad (2)$$

$$CZ_{ij}^{qf} = \exp\left(i\pi n_i^q n_j\right),\tag{3}$$

along with the qubit Hadamard gate H, and the qubit phase gate S. For this gate set to be universal (see Methods for a proof), we also need a gate which moves fermions between sites, $\sqrt{iSWAP}_{ij}^{f} = \exp\left(i(\pi/4)\left(p_{i}^{\dagger}p_{j} + h.c.\right)\right)$. For convenience, we also define the gates $S_{i}^{f} = (T_{i}^{f})^{2}$ and $Z_{i}^{f} = (S_{i}^{f})^{2}$ not necessary for universality. CZ^{af} is a qubit-fermion gate dependent on the fermionic number operator $n_{j} = p_{j}^{\dagger}p_{j}$ and its qubit analogue $n_{i}^{q} = |1\rangle \langle 1|_{i}$, i.e. the projector on the $|1\rangle$ state of the qubit. Note that, in principle, we do not require qubits for fermion universality [31], but as we will show, being able to couple fermions to qubits will lead to key advantages. Neutral-atom platforms are considered to be constrained to this number-conserving gate set [27] as usually the atom number is conserved during gates and each tweezer is initially prepared in a number eigenstate.

The first question is therefore whether these numberconserving operations suffice to encode logical fermions, which is equivalent to demanding that the codestates are eigenstates of the total fermion number operator. Unfortunately, such states are not sufficient:

Theorem. An error-correcting code using eigenstates of the total fermion number operator as codestates does not have logical fermion operators.

The proof (Methods) relies on the fact that codestates that are eigenstates of the total fermion number operator need to be eigenstates with the same number eigenvalue due to the Knill-Laflamme conditions. This means that logical operators cannot change the total fermion number and therefore need to be even weight in fermionic creation/annihilation operators. Hence, logical operators are bosonic.

To circumvent this theorem, we use fermion parity eigenstates as codestates, i.e. states that are superpositions of states that contain either only an odd or only an even number of fermions. A universal gate set on such states requires a non-numberconserving gate. In analogy to the braiding operation performed in Majorana nanowires [32], we define the braiding gate

$$BRAID_{ij} = \exp\left(i\frac{\pi}{4}\left(p_i^{\dagger} - p_i\right)\left(p_j^{\dagger} + p_j\right)\right), \quad (4)$$

which will also turn out to be a natural logical gate in our codes. We propose a scheme to implement this gate in neutral atoms at the end of this work.

We envision a zoned architecture in which the gates are implemented in separate physical regions of the quantum processor, which has been experimentally realized for qubitonly neutral atom quantum computing in Ref. [33]; we specify the zones in which each of the gates is performed in Fig. 1b, and introduce our circuit notation.

To define our codes, it is convenient to introduce Majorana fermion operators, which are the real and imaginary parts γ_i , $\tilde{\gamma}_i$ of complex fermionic operators, $p_i^{\dagger} = \frac{1}{2}(\gamma_i + i\tilde{\gamma}_i)$, fulfilling $\{\gamma_i, \gamma_j\} = 2\delta_{ij} = \{\tilde{\gamma}_i, \tilde{\gamma}_j\}$ and $\{\gamma_i, \tilde{\gamma}_j\} = 0$. We construct codes in which we encode two logical Majorana fermions $\gamma^{\rm L}$ and $\tilde{\gamma}^{\rm L}$. These are then combined into a logical complex fermion $c^{\dagger} = \frac{1}{2}(\gamma^{\rm L} + i\tilde{\gamma}^{\rm L})$. In order for logical operators

between different codeblocks to fulfill Majorana fermion anticommutation relations, they each must be a product of an odd number of γ_i and $\tilde{\gamma}_i$, i.e. they must have odd operator weight.

REPETITION CODE FOR PHASE ERRORS

We start with a repetition code which only corrects for phase errors $E = n_i$. This is the dominant error source in neutral atoms [27]. As an introductory example, consider the two-site code

$$\left|0\right\rangle_{\rm L} = \frac{1}{\sqrt{2}} \left(\left|00\right\rangle - \left|11\right\rangle\right),\tag{5}$$

$$\left|1\right\rangle_{\rm L} = \frac{1}{\sqrt{2}} \left(\left|10\right\rangle + \left|01\right\rangle\right),\tag{6}$$

where $|11\rangle = p_2^{\dagger} p_1^{\dagger} |00\rangle$ and $p_1^{\dagger} |00\rangle = |01\rangle$ and $|0\rangle, |1\rangle$ denotes the absence, presence of a fermion, respectively. The above code can detect a single phase error, but does not detect loss errors $E = p_i$. The code has two logical Majorana fermion operators $\gamma^{\rm L} = \gamma_1$ and $\tilde{\gamma}^{\rm L} = \tilde{\gamma}_2$, from which we form a logical complex fermion operator by $c^{\dagger} = \frac{1}{2} (\gamma_1 + i\tilde{\gamma}_2)$. This operator acts as $c^{\dagger} |0\rangle_{\rm L} = |1\rangle_{\rm L}$, $c^{\dagger} |1\rangle_{\rm L} = 0$. We therefore interpret $|0\rangle_{\rm L}$ and $|1\rangle_{\rm L}$ as logical fermionic Fock states. Two codeblocks aand b each encode such an operator and they fulfill $\{c_a^{\dagger}, c_b^{\dagger}\} = 0$. Hence, c^{\dagger} is indeed a logical fermion operator. This simple twosite code is also a stabilizer code, see Fig. 2a: its codestates are defined by the operator $i\tilde{\gamma}_1\gamma_2$ which acts as the unity operator on both states, $i\tilde{\gamma}_1\gamma_2 |0\rangle_{\rm L} = |0\rangle_{\rm L}$ and $i\tilde{\gamma}_1\gamma_2 |1\rangle_{\rm L} = |1\rangle_{\rm L}$. This enables generalization to detecting N-1 phase errors by using N complex physical sites and the stabilizers $i\tilde{\gamma}_i\gamma_{i+1}$ for 1 < i < N. For $N \ge 3$, we can also correct phase errors as we show below. This code can also be used as a purely error-detecting code for all errors except atom-loss errors on the edges. In fact, this is the code defined by the ground space of the Kitaev chain, and the two logical Majorana operators γ_1 and $\tilde{\gamma}_N$ are its well-known edge states used as qubits in Majorana nanowires [32]. However, by contrast to nanowires, we do not implement the Kitaev Hamiltonian in our system. Hence, we cannot rely on the gap between the ground space of the Hamiltonian and the excited states in order to suppress errors and keep the computation in the ground space of the Hamiltonian. Instead, we require active error correction.

For implementing active error correction, we need to measure where errors happened and correct for them, which we show how to do in Fig. 2b. We use our qubit-fermion gate to map a measurement of the fermion density *n* to a measurement of the phase of the qubit, which we transform to a measurement of the population of the qubit using Hadamard gates. To measure the stabilizer instead of *n* on the fermions we conjugate the qubitfermion gate with two braiding gates as well as phase gates. An ancilla qubit is used for each of the stabilizers. Dependent on the measurement outcome, Z^f gates are performed on the fermions to correct for the errors. This scheme shows the first advantage of using qubits: destructive qubit readout is faster than a non-destructive *n* measurement [35]. To prepare a logical codestate $|0\rangle_{L}$, we use a chain of braiding gates—the S[†]

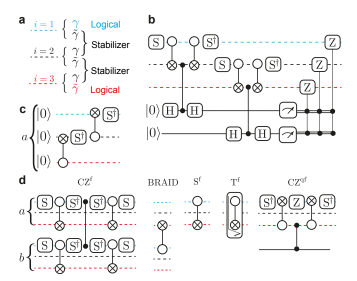


FIG. 2. Logical operations for a repetition code protecting against phase errors. a, Illustration of splitting physical fermions for N = 3 into virtual Majorana fermions. Stabilizers are given by $i\tilde{\gamma}_1\gamma_2$, $i\tilde{\gamma}_2\gamma_3$, and the logical fermionic creation operator is given by $c^{\dagger} = \frac{1}{2}(\gamma_1 + i\tilde{\gamma}_3)$. b, Syndrome measurement and correction. The two stabilizers are measured by using a Hadamard test on qubits. Conditioned on the measurement outcome, gates are performed to correct for errors. c, Encoding circuit, translated from Ref. [34], realized as a chain of braidings. Note that $(S^{f})^{\dagger}$ can be interpreted as intra-site braiding. d, Logical gates. $\sqrt{\text{BRAID}}_{31} = \exp(-\frac{\pi}{8}\tilde{\gamma}_3\gamma_1)$, which only acts on the two edge atoms. See Methods for a derivation of these circuits.

gates effectively braid the two Majoranas on the same physical site, Fig. 2c. Note that we do not error correct the qubit ancillas, which is reasonable because their dephasing time is much longer than that of the fermions and we only act with a handful of gates on them before measuring.

Next, we discuss how to implement logical gates. We show the circuits in Fig. 2d, see Methods for a proof. The basic building block for the logical CZ^f is the physical CZ^f . Similar to the stabilizer measurement, conjugation with braiding gates "spreads" the entanglement induced by the physical CZ^f to the correct site, with S^f gates selecting whether $\tilde{\gamma}$ or γ is acted upon. The braiding gate is simple as it amounts to a physical braiding gate between γ_1 of one block and $\tilde{\gamma}_N$ of the other. The S^f gate amounts to a braiding gate between the first and Nth site of a single codeblock. Similarly, the T^f gate is implemented as a braiding operation with half the duration of the S^f gate. Finally, the qubit-fermion CZ^{qf} is implemented similarly to the CZ^f

The utility of this code is based on the fact that, in neutral atoms, phase errors are orders of magnitude more frequent than atom-loss errors. However, because braiding gates convert a single phase error into two correlated atom-loss errors, which cannot be detected, the depth of the circuits which can be performed in practice is still limited by the phase coherence time. Hence, this code is not fault-tolerant even in the absence of atom-loss errors. The fact that logical gates only act on the edge fermions on the other hand reduces the depth of the circuits, as the gate overhead for implementing logical gates is independent of N. It might be beneficial to reduce gate depth further by replacing the CZ^{f} gates and BRAIDs in Fig. 2d with their arbitrary-angle versions, which turns our gates to arbitraryangle logical gates. However, for large-scale computations, a truly fault-tolerant code is necessary.

FERMION-QUBIT FAULT TOLERANCE

We now show how to construct an error-correcting code correcting for both phase errors E = n and atom-loss errors E = p and show how to perform fault-tolerant gadgets. In the context of encoding qubits in Majorana nanowire platforms, Bravyi, Terhal, and Leemhuis [31] noted that a particular class of qubit error correcting codes, so-called weakly self-dual Calderbank-Shor-Steane (CSS) codes, can be converted into a Majorana stabilizer code by replacing qubit operators in the stabilizers by Majorana fermions. Motivated by the prospect of high error-resilience for codes with odd-weight logical operators, they noted that for codes with an odd number of physical Majorana fermions, a single odd-weight logical operator can be encoded. This operator is given by the total parity, i.e. the product of all Majorana operators. Our key observation is that, because there is an odd number of physical Majorana fermions, this logical operator is in fact a logical Majorana fermion operator. Even more importantly, our framework using complex physical fermions enables the encoding of two such Majorana codes in the same set of physical fermions. It turns out that a general prescription to constructing complex fermion codes from weakly self-dual CSS codes is to replace the Z operators in the qubit code stabilizers by γ operators and the qubit X operators by $\tilde{\gamma}$ operators, see Fig. 3a. The logical Majorana operators are then given by the two parities $\gamma^{\mathrm{L}} = \prod_{i=1}^{N} \gamma_i$ and $\tilde{\gamma}^{L} = \prod_{i=1}^{N} \tilde{\gamma}_{i}$, from which we construct a single logical fermion operator $c^{\dagger} = \frac{1}{2} (\gamma^{L} + i \tilde{\gamma}^{L})$. As an example, we consider the family of triangular honeycomb Majorana color codes, which using our procedure above, encodes a single logical fermion with odd distance d into $N = (3d^2 + 1)/4$ physical fermions. In particular, d = 3 is the fermionic version of the Steane code [36], Fig. 3a. Such codes can correct d-2loss or phase errors.

Magically, this fermion Steane code yields highly intuitive and easy-to-implement fault-tolerant logical gates. Of central utility is the fact that we can use qubits for controlled operations. We use this fact to measure stabilizers using the Hadamard test, as shown in Fig. 3b. From these stabilizers, we can reconstruct whether a $\tilde{\gamma}$ or γ error (or both) happened on a fermion. Such single-Majorana errors are corrected by coupling an ancilla site a prepared in $|0\rangle$ to the site with the error, acting with $(BRAID_{ai})^2 = \gamma_i \tilde{\gamma}_a$ and then measuring the ancilla. The measurement circuit in Fig. 3b is made fault-tolerant by for example using the Shor scheme, i.e. replacing the single ancillary qubit with a set of qubits in a cat state and repeatedly measuring the stabilizers. This stabilizer measurement is also used for fault-tolerant preparation of the logical codestates from a trivial starting state. Note that the correction of stabilizers is optional for both state preparation and error correctioninstead of enforcing stabilizers to be +1, their value can instead

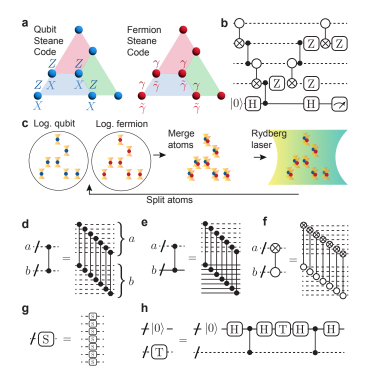


FIG. 3. Fermion-qubit fault-tolerant quantum computation. a, Construction of the fermionic Steane code, the smallest member of the family of color codes we consider. Stabilizers are obtained by replacing products of X(Z) operators in the qubit code with products of γ ($\tilde{\gamma}$), respectively. Corners on areas with the same color are part of a stabilizer as shown for the bottom left area. b, Mid-circuit measurement of $-\gamma_1\gamma_2\gamma_3\gamma_4$ (counting sites from bottom to top). By inverting the BRAIDs (i.e. $BRAID_{12} \rightarrow BRAID_{21}$), we measure $-\tilde{\gamma}_1\tilde{\gamma}_2\tilde{\gamma}_3\tilde{\gamma}_4$, the other stabilizer. c, Neutral-atom implementation of transversal CZ^{qf} gates, performed by interleaving a logical qubit and fermion and applying a global Rydberg laser. This implements the circuit shown in subfigure e. \mathbf{d} , Transversal CZ^f. This is implemented by the logical qubit in subfigure c with a logical fermion. e, Transversal qubit-fermion CZ^{qf} . Transversal **f**, BRAID and **g**, S^{f} gates are similarly implemented by parallel operations. h, T^f gate. Implemented by using a T gate applied on an ancilla qubit, which is then mapped onto the fermion. All the operations shown here are straightforwardly generalized to codes with larger (odd) d (see Methods).

be tracked [37].

Transversal logical gates are implemented, similar to qubit Steane codes, by applying the corresponding physical gate in parallel on all atoms, c.f. Fig. 3c-g (Methods). This set of gates turns out to be the Majorana Clifford gates BRAID, CZ^f, S^f, which in analogy to qubit Clifford gates map a single Majorana operator string to another single Majorana string. These transversal Clifford gates are inherently fault-tolerant as an error on one physical codeblock can only spread to one physical fermion in the other codeblock.

One of the key tools enabled by our approach is the interfacing of our fermionic Steane code with a qubit Steane code (with matching N) using the qubit-fermion CZ^{qf} gate: by applying physical CZ^{qf} in parallel (see Fig. 3e), we realize a logical, transversal CZ^{qf} . This enables fault-tolerant qubit-fermion quantum computation. In particular, we will discuss in the next

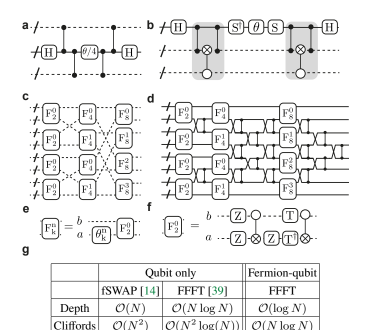


FIG. 4. Fermion-qubit logical quantum simulation primitives and comparison to the qubit-only approach. a, Qubit-controlled arbitrary-angle interaction gate. The box with label " $\theta/4$ " is a qubit gate $\exp(-i(\theta'/2)X)$ with $\theta' = \theta/4$, synthesised from discrete gates using the Solovay-Kitaev algorithm. b, Qubit-controlled arbitraryangle braiding gate. The gray box is the qubit-controlled braiding gate $\exp(-(\pi/4)Z\tilde{\gamma}_i\gamma_j)$. c, FFFT circuit implemented using logical fermions. Crossing lines indicate swapping, implemented by movement of the atoms. d, Corresponding FFFT circuit implemented using logical qubits, c.f. Ref. [40]. e, Decomposition of the n-mode FFFT into an arbitrary-angle phase gate $\exp(-i\theta n)$ with $\theta = \theta_k^n = 2\pi k/n$ and a two-mode FFFT. f, Compilation of the two-mode FFFT into fermionic fault-tolerant gates. g, Depth and Clifford complexity of qubit-only and fermion-qubit algorithms for simulating the kinetic energy term in crystalline materials, i.e. the first term in Eq. (7).

session that this has immense utility for implementing modern quantum algorithms for simulating fermions.

This gate set of BRAID, CZ^{qf} , CZ^{f} , S^{f} , together with H and S on qubits, forms the "qubit-fermion Clifford group" as it maps a Majorana-qubit string to a single other Majorana-qubit string. This gate set needs to be combined with a T^{f} gate or a qubit T gate to enable universal qubit-fermion computation. While Majorana fermion schemes for *T*-gate synthesis have been proposed [38], they are more involved than qubit schemes due to the requirement to create two-fermion magic states. We instead propose using a qubit T gate applied on a qubit Steane code and transferring it to the fermions using our transversal logical CZ^{f} gate, Fig. 3h. This therefore enables a significant simplification compared to Majorana T-gate synthesis. electrons are cast into a discrete lattice model through a choice of basis, which leads to a Hamiltonian coupling fermionic operators as $H_{qchem} = \sum_{ij} J_{ij} c_i^{\dagger} c_j + \sum_{ijkl} V_{ijkl} c_i^{\dagger} c_j^{\dagger} c_k c_l$, with J_{ij} and V_{ijkl} constants that depend on the choice of basis. This generally leads to a number of Hamiltonian terms scaling as $\sim N^4$ with the basis size N. For a Trotterized simulation, this implies the same scaling for the number of gates per Trotter step. However, for crystalline materials, the plane-wave-dual basis [12] enables writing the chemistry Hamiltonian as

$$H = \text{FFFT}^{\dagger} \sum_{i} \epsilon_{i} n_{i} \text{FFFT} + \frac{1}{2} \sum_{ij} U_{ij} \left(n_{i} - \frac{1}{2} \right) \left(n_{j} - \frac{1}{2} \right), \quad (7)$$

where ϵ_i , μ_i , and U_{ij} are classically calculable coefficients and FFFT denotes the fermionic fast Fourier transform [40]. This formulation hence reduces the scaling of the number of terms to $\sim N^2$. Efficient algorithms for dynamics, as well as for ground and finite-temperature state preparation, in addition require controlling this Hamiltonian on a qubit [10, 11].

Our next goal is therefore to implement qubit-controlled arbitrary-angle evolution under each of the terms in Eq. (7) individually. Time evolution under the Hamiltonian $\epsilon_i n_i$ until time t is implemented by replacing the T gate in Fig. 3h with an approximate arbitrary-angle rotation of angle $\epsilon_i t$, where t corresponds to time, synthesized from discrete gates with e.g. the Solovay-Kitaev algorithm. The interaction gate exp $\left(-iZ\frac{\theta}{2}(n_i-\frac{1}{2})(n_j-\frac{1}{2})\right)$ is implemented by the circuit shown in Fig. 4a, which maps the arbitraryangle single-qubit gate onto the fermions using CZ^{qf} gates. A related gadget shown in Fig. 3b realizes an arbitraryangle qubit-controlled braiding gate $\exp(-\frac{\theta}{2}Z\tilde{\gamma}_i\gamma_j)$, where again CZ^{qf} gates map the phase gate from the qubit to a fermion. The BRAIDs then "spread" the gate onto the second fermion. Two such gadgets are combined to yield the qubitcontrolled hopping gate via $\exp\left(iJ_{ij}\Delta tZ\left(c_{i}^{\dagger}c_{j}+h.c.\right)\right) =$ $\exp(-\frac{J\Delta t}{2}Z\tilde{\gamma}_j\gamma_i)\exp(-\frac{J\Delta t}{2}Z\tilde{\gamma}_i\gamma_j)$. These arbitrary-angle gates also enable time evolution under the more general Hamiltonian H_{qchem} by using the compilations in Ref. [27]. Finally, we need to implement the FFFT, which may be reduced to the 1D FFFT circuit shown in Fig. 4c, c.f. Ref. [40]. In analogy to the classical fast Fourier transform, it uses a divide-andconquer reduction to two-site Fourier transforms F_N^k (Fig. 4e, f) whose phases depend on the number of sites N and an index k. Despite the fact that it is a 1D transformation, it requires non-local connectivity, which we can simply implement by moving the tweezers.

ADVANTAGES OVER QUBIT QUANTUM SIMULATION

LOGICAL QUBIT-FERMION QUANTUM SIMULATION

A prime area of application for fermionic quantum computing is simulating electrons in molecules and materials. In second quantized approaches, the continuous-space wavefunctions of We now show that our approach yields an exponential depth advantage over the most efficient qubit algorithm for simulating trotterized time evolution under the kinetic energy term in crystalline materials (first term in Eq. (7)). The, to our knowledge, most efficient qubit algorithm uses fermionic-SWAP (fSWAP) networks, leading to $\mathcal{O}(N)$ depth and $\mathcal{O}(N^2)$ Clifford gates per Trotter step, where N is the number of fermionic sites in the model [14]. Local encodings of qubits into fermions [5–7] yield little advantage for such all-to-all connected hoppings. The FFFT described above can also used in qubits, where atom rearrangements are replaced by layers of fSWAP=SWAP·CZ gates, Fig. 4d. These layers cannot be contracted even in all-to-all connected hardware such as tweezer arrays or trapped ions [9]. This leads to a gate depth of $\mathcal{O}(N \log N)$ and $\mathcal{O}(N^2 \log(N))$ Clifford gates, which is worse than the fSWAP network implementation.

By contrast, in our fermionic platform, the fact that fSWAP gates are implemented by atom rearrangements and can therefore be contracted (Fig. 4c), leads to a Clifford complexity of $\mathcal{O}(N \log(N))$ and a depth of $\mathcal{O}(\log N)$, yielding an exponential improvement for the latter. Moreover, our compilation of the constituent FFFT subroutines (see Fig. 4e, f) only takes two braiding and three single-fermion gates and T-gate depth 1, which is more gate-efficient than currently known qubit decompositions taking three CNOTs, nine single-qubit gates and T-gate depth 2 [13].

For the more general H_{qchem} , at first sight, no depth advantage is possible because fermi-SWAP networks achieve the optimal depth $\mathcal{O}(N^3)$ [14], which our approach reproduces with a smaller prefactor. However, asymptotically, there are only $\mathcal{O}(N^2)$ non-negligible terms in the often-used Gaussian basis [41]. We therefore expect that, for large N, we will again yield at least an $\sim N$ depth advantage by the fact that our fermion approach always achieves the optimal depth by parallelization, whereas in the qubit approach, most of the fermi-SWAP network would still have to be traversed.

Within our fault-tolerant gate set, no advantage in the number of T gates is possible if the T-gate count is the only parameter which is optimised for. This can be directly seen by mapping the fermionic gates to qubit gates using the Jordan-Wigner transform and noting that only the T^{f} gate requires T gates (Methods). We note that steady progress in T-gate synthesis means they do not completely dominate the simulation cost over Clifford gates [30].

IMPLEMENTATION IN NEUTRAL ATOMS

Finally, we describe how to implement our framework in neutral atoms, which requires the implementation of the univeral qubit-fermion gateset shown in Fig. 1. CZ^{qf} and CZ^{f} are implemented using a high-lying Rydberg state of the atom, similarly to the routinely implemented qubit-qubit CZ^{q} [27, 42]. \sqrt{iSWAP}^{f} is implemented by bringing two tweezers close to each other, inducing tunneling of atoms between the sites [27] or by placing the atoms into an optical lattice [43] and controllably performing tunneling by using superlattices [44]. T^f, S^f, Z^f are implemented by detuning the trapping frequency of the target tweezer or optical lattice site. What is left to show is how to implement BRAID. We propose its realisation as a product BRAID_{ij} = PAIR'_{ij} \sqrt{iSWAP}^{f}_{ij} , where PAIR'_{ij} is

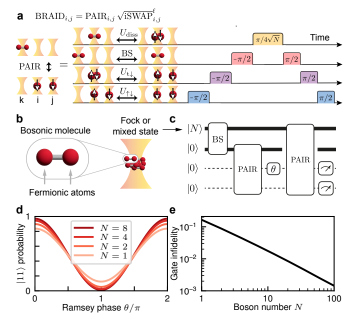


FIG. 5. Implementation of the braiding gate in neutral atoms. a, Illustration of the sequence of operations to implement an effective braiding gate. Braiding is a product of pairing and the tunneling gate $\sqrt{\text{iSWAP}}^{\text{f}} = U_{t\downarrow}(\pi/4)$. $U_{t\downarrow}(\theta)$ and $U_{\uparrow\downarrow}(\theta)$ act only on fermions, while the beamsplitter acts only on molecules, and $U_{\rm diss}$ is a fermionmolecule gate. The pairing gate dissociates a bosonic molecule into two fermions on two separate sites. The pulse duration of the dissociation interaction depends on the average number $N = \langle b^{\dagger}b \rangle$ of molecules in the tweezer. b, The molecules are formed from the same atomic species as the fermionic atoms used. c, Ramsey sequence to probe the effective coherence of a "Bell-pair" $(|00\rangle + i |11\rangle)/\sqrt{2}$ created by the effective pairing gate. The phase gate is defined as $\exp(-i\theta n)$. The beamsplitter gate, defined above Eq. (9), has phase $\theta = \pi/4$. **d**, Ramsey fringes as a function of initial molecule Fock state number N. e, Simulated pairing gate infidelity as a function of molecule Fock state number N. See Fig. S1f for the circuit used to determine the infidelity.

the pairing gate

$$\text{PAIR}'_{ij} = \exp\left(i\frac{\pi}{4}\left(p_i^{\dagger}p_j^{\dagger} + h.c.\right)\right). \tag{8}$$

To implement pairing, we propose the pulse-sequence in Fig. 5a. It employs photodissociation of homonuclear diatomic bosonic molecules consisting of the same atomic species as the fermions [45], see Fig. 5b. Photodissociation realizes the bosonfermion gate $U_{\text{diss}} = \exp\left(-i\frac{\pi}{4\sqrt{N}}\left(b_ip_{i\uparrow}^{\dagger}p_{i\downarrow}^{\dagger} + b_i^{\dagger}p_{i\downarrow}p_{i\uparrow}\right)\right)$, where *b* is the bosonic annihilation operator of molecules (we assume there are *N* of them) residing in the ground state of the tweezer and obeying commutation relations $[b_i, b_j^{\dagger}] = \delta_{ij}$. \uparrow and \downarrow label two internal states of the atom, where so far we assumed that all atoms reside in \downarrow , i.e. $p_i = p_{i\downarrow}$. In order to convert this same-site gate into a non-local gate, we conjugate it with a fermion spin-selective tunneling $U_{t\downarrow}(\theta) = \exp\left(-i\theta\left(p_{i\uparrow}^{\dagger}p_{j\uparrow} + h.c.\right)\right)$ and a fermionic spin-flip $U_{\uparrow\downarrow}(\theta) = \exp\left(-i\theta\left(p_{i\downarrow}^{\dagger}p_{i\uparrow} + h.c.\right)\right)$. To suppress inter-

actions between the molecules and the fermions, the molecules reside in a separate tweezer when not in use for gates and are tunneled into the target tweezer by moving the two tweezers close to each other, implementing a beam-splitter operation $BS(\theta) = \exp\left(\theta\left(b_i^{\dagger}b_j - h.c.\right)\right)$. Altogether, the sequence in Fig. 5a implements

$$\text{PAIR}_{ij} = \exp\left(i\frac{\pi}{4\sqrt{N}}\left(b_k p_i^{\dagger} p_j^{\dagger} + h.c.\right)\right). \tag{9}$$

We now discuss how we use Eq. (9) to effectively implement Eq. (8). If the molecules were in a coherent state with large amplitude \sqrt{N} , we could replace $b \rightarrow \langle b \rangle = \sqrt{N}$ and the two gates would be equal. However, in the case of atoms trapped in dipole traps, Fock states or mixtures thereof are more naturally prepared than coherent states. To overcome this conceptual challenge, we use the little-known fact that large-NFock states cannot be distinguished from coherent states by measurements [46].

To show that we can indeed effectively implement Eq. (8)employing a large-N Fock state of molecules, consider the thought experiment shown in Fig. 5c. The first pairing gate is supposed to create a "fermionic Bell-pair" $(|00\rangle + i |11\rangle)/\sqrt{2}$. To probe whether there is coherence, a phase gate with phase θ is applied and a second PAIR gate rotates the measurement basis into the effective Bell basis. This realizes a Ramseytype experiment. Simulating this experiment numerically, we find Ramsey fringes with a contrast that approaches unity as $N \to \infty$. This shows that, after the first pairing gate, there was indeed effective coherence between states $|00\rangle$ and $|11\rangle$. An intuitive way to understand this observation is that the molecule number distribution in each mode after the beamsplitter is approximately Poissonian-just like in a coherent state. The entanglement between both modes then leads to an effective phase coherence between the modes. This effective coherence is then revealed by the second pulse, which disentangles the fermions from the molecules. We say "effective" coherence because the reduced density matrix of the fermions is always diagonal; indeed, by measuring the total number of molecules, one projects the fermions onto $|00\rangle$ or $|11\rangle$. Therefore, it is important that the molecules are not measured while the fermions are entangled with them. In Fig. S1, we additionally perform state tomography numerically, showing that the state after the first pairing gate is indeed effectively a Bell pair.

We probe the gate fidelity of our effective pairing gate in Fig. 5e using Choi state tomography, which we generalize to fermionic gates, see circuit shown in Fig. S1. We find that only 100 molecules suffice to create a gate with > 99.8 % fidelity. In the Methods, we also show that even a mixed state with a Poissonian number distributation can be used instead of $|N\rangle$. Non-trivially, Fig. 5c and Fig. 5d show that our scheme works even if the second pairing gate is not performed on the same molecule site, which enables full parallelizability of pairing operations. For that to be the case, the molecule tweezers need to be effectively coherent with respect to each other, just like lasers manipulating different qubits within a quantum computer need to be phase-locked. The cross-coherence of molecule tweezers is guaranteed by creating them from one common Fock

state with a beamsplitter as done in Fig. 5c. To parallelize this further, many molecule tweezers can be created by repeatedly applying beamsplitters starting from one large Fock state, for

OUTLOOK

example by tunneling from a single large dipole trap.

Our work opens the field of fermion-qubit error correction and fault-tolerant algorithms. Of prime importance is the error threshold of our codes for the strongly biased errors present in experiments. Moreover, fermionic versions of quantum low-density-parity check codes may be developed to enable a finite rate of logical fermions per codeblock. Due to our reliance on transversal gates, algorithmic fault tolerance can be explored within our approach [30]. Finally, we assumed that the bosonic molecule mode does not need to be error-corrected, which might require a more restrictive lower bound on the mode occupation N.

Our present scheme employs single-species fermions. While two-species models, for example spin up and spin down electrons in quantum chemistry, can be mapped to single-species computation, an architecture involving two species, as partially employed in our pairing gate, might make the implementation more efficient.

We have so far mainly used the presence of qubits as a helper to implement logical operations. In high-energy physics, the possibility to directly simulate qubit-fermion models enables simulation of fermions coupled to large-spin degrees of freedom in lattice gauge theories [27, 47]. For this purpose, qudits could be directly implemented with large-spin atomic species [48, 49], rotational levels of molecules [50], or an oscillator mode [51]. Relatedly, the depth-efficiency of our approach might enable simulation of quantum gravity models [52].

Boson-fermion models could be studied by combining our schemes with oscillator-qubit quantum simulation [47, 53] using either motional modes of neutral atoms in tweezers [47, 54] or bosonic atoms. Suitable boson-to-boson error correction schemes would need to be found [55, 56]. This way, the longstanding problems of the influence of phonon-electron interactions on thermal, structural, and vibrational material properties [57] as well as the interplay of molecular vibrations with the electronic wavefunction in chemistry [58] could be investigated using potentially vastly shallower circuits than with qubit-only platforms.

Finally, our architecture might offer an advantageous approach not only for simulating fermions but also for quantum computations that are natively qubit-only. We encoded logical fermion operators in odd-weight fermion operators—instead, logical qubits can be encoded in even-weight logical operators, for example Majorana surface codes which have been shown to have high thresholds [59]. One advantage of this approach is that lost atoms are replenished by coupling to the molecule bath, offering an alternative to continuous atom reloading [43], which is one of the challenging aspects of neutral-atom quantum computing.

ACKNOWLEDGMENTS

We acknowledge discussions with Christopher Kang, Paul Julienne, Robert Ott, Shraddha Singh, Maryam Mudassar, Nathan Wiebe, Torsten Zache, Johannes Zeiher, and Hengyun Zhou.

Funding. This material is based upon work supported in part by the U.S. Department of Energy, Office of Science, National Quantum Information Science Research Centers, Quantum Systems Accelerator. Additional support is acknowledged from the following agencies. This work was supported in part by the Department of Energy (DOE), Office of Science, Office of Nuclear Physics, via the program on Quantum Horizons: QIS Research and Innovation for Nuclear Science (award no. DE-SC0023710) and the National Science Foundation's Quantum Leap Challenge Institute in Robust Quantum Simulation (award no. OMA-2120757). A.V.G. was also supported in part by the DoE ASCR Accelerated Research in Quantum Computing program (award No. DE-SC0020312), AFOSR MURI, DoE ASCR Quantum Testbed Pathfinder program (awards No. DE-SC0019040 and No. DE-SC0024220), NSF STAQ program, and DARPA SAVaNT ADVENT. E.C. was also supported in part by ARO MURI (award No. SCON-00005095), and DoE (BNL contract No. 433702). Support is acknowledged from the U.S. Department of Energy, Office of Science, National Quantum Information Science Research Centers, Co-design Center for Quantum Advantage (C2QA) under contract number DE-SC0012704. C2QA participated in this research.

Competing interests. All authors declare no competing interests.

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Methods

PROOFS

Proof of Theorem

In this section, we present the details behind the proof of the Theorem.

We first discuss some preliminaries. To construct a valid code, we need to satisfy the Knill-Laflamme condition [60]. It ensures that errors do not send codestates to states that have an overlap with other codestates. If this condition were not fulfilled, then the detection of an error could lead to a projection onto the wrong codestate, which is an uncorrectable logical error. Formally, the condition states that, for an error operator E to be detectable, $PEP \propto P$ needs to be fulfilled, where P is the projector on the codespace, i.e. the space spanned by the logical codestates $|0\rangle_{\rm L}$ and $|1\rangle_{\rm L}$, representing, respectively, the absence and presence of a logical fermion.

Lemma 1. Codestates that are eigenstates of the total number operator in a single codeblock, $N_{tot} = \sum_j n_j$, must have the same number eigenvalue.

Proof. The Knill-Laflamme condition for phase errors $E = n_j$ implies $PN_{\text{tot}}P \propto P$. Furthermore, having codestates that are number eigenstates implies $(1 - P)N_{\text{tot}}P = 0$. Taking these two conditions together, we find $N_{\text{tot}}P \propto P$. This implies that all codestates must have the same number eigenvalue.

Definition. Fermionic logical operators are operators that anti-commute between two codeblocks.

We note that this definition does not exclude fermionic logical operators that anti-commute within one codeblock [29]. However, our definition is more restrictive - logical operators that anti-commute within one codeblock may commute between two codeblocks. This is for instance the case when encoding two anti-commuting logical operators in a qubit surface code [28].

Lemma 2. Fermionic logical operators have odd weight in physical Majorana operators.

Proof. First, we define new Majorana operators η such that $\eta_{2i} = \gamma_i$ and $\eta_{2i+1} = \tilde{\gamma}_i$. We then write the logical operators O_a and O_b on two codeblocks as a product of η Majorana operators. Two such products of Majorana operators fulfill [31]

$$O_a O_b = (-1)^{|A| \cdot |B| + |A \cap B|} O_b O_a,$$
(S1)

where A and B are the supports on which O_a and O_b are defined (i.e. on the lattice on which the η operators are defined). Because O_a and O_b are defined on two codeblocks, their support does not overlap, $|A \cap B| = 0$. Therefore, they can only fulfill fermionic anti-commutation relations if |A| and |B| are odd, i.e. if the two operators have odd weight.

Theorem. An error correcting code using number eigenstates as codestates does not have fermionic logical operators.

Proof. First, consider stabilizer codes using number eigenstates. In this case, by Lemma 1, the total fermion parity $\exp(i\pi N_{\rm tot})$ is a stabilizer because all the codestates have the same number eigenvalue, which is an integer. As noted in proposition 6.1 in Ref. [31], this implies that there cannot be an odd-weight logical operator. To repeat the argument in Ref. [31], this is because all logical operators need to commute with the stabilizers and hence the total fermion parity. This is only possible if they are even weight in Majorana operators, which means they are not fermionic logical operators by Lemma 2.

More generally, consider two logical operators $O_a^{\rm L}$ and $O_b^{\rm L}$ acting on two codeblocks a and b. For them to be logical fermionic operators, they need to fulfill anti-commutation relations when acting on the codespace, i.e. $\{O_a^L, O_b^L\}P_aP_b =$ 0, where P_a and P_b are the projectors on the code spaces of aand b. To see whether this is the case, consider $O_a^{\rm L}O_b^{\rm L}P_aP_b$. Because of Lemma 1, the total fermion parity in codeblock a commutes with P_a . Hence, the operator decomposition of P_a in terms of Majorana operators must only contain operator strings with even weight. Eq. (S1) then implies that P_a commutes with any fermion operator in codeblock b, in particular the logical ones, $O_a^{\rm L} O_b^{\rm L} P_a P_b = O_a^{\rm L} P_a O_b^{\rm L} P_b$. Note that for $O_a^{\rm L}$ to be a logical operator, $(1 - P_a) O_a^{\rm L} P_a = 0$ must hold [61], implying $O_a^{\rm L} P_a = P_a O_a^{\rm L} P_a$. Because $P_a O_a^{\rm L} P_a$ only acts on the code space, it commutes with the fermion parity. Hence, its operator decomposition only contains even-weight operators and therefore commutes with any operator in codeblock b. This implies $O_a^{\rm L} P_a O_b^{\rm L} P_b = O_b^{\rm L} O_a^{\rm L} P_a P_b$. Hence, $[O_a^{\rm L}, O_b^{\rm L}] P_a P_b =$ 0 and logical operators are not fermionic.

Proof of absence of T-gate advantage

In this subsection, we present a proposition proving that there is no T-gate advantage.

Proposition 1. Fermion-to-fermion error correcting codes with transversal BRAID, CZ^{f} , S^{f} gates and non-transversal T^{f} gate do not have a lower minimal T^{f} -gate count than the *T*-gate count of the equivalent qubit circuit.

Proof. We show this by converting the fermionic circuit to a qubit circuit using the Jordan-Wigner encoding. The CZ^f gate maps to a qubit CZ, which can be compiled to qubit Clifford gates, whereas S^f and T^f gates directly map to S and T gates, respectively. BRAID gates map to multi-qubit gates, but their compilation is possible using only Clifford gates. Hence, it is always possible to construct a qubit circuit with exactly the same number of T gates as the fermion circuit.

Proof of universality of the discrete qubit-fermion gate set

In this subsection, we show the universality of our discrete qubit-fermion gate set.

Proposition 2. BRAID, CZ^{f} , CZ^{qf} , H, S and T (or T^{f}), are universal for qubit-fermion quantum computation.

Proof. We would like to do universal qubit-fermion quantum computation on a system with L_q qubits and L_f fermions. To do this, we introduce $2L_q$ fermionic ancillas and encode L_q qubits into dual-rail fermion qubits, i.e. $|0\rangle \rightarrow |01\rangle$, $|1\rangle \rightarrow |10\rangle$. We then swap the initial qubit state into those dual-rail fermion qubits (we discuss how to do so below). We then perform the qubit-fermion circuit entirely using the universal fermionic gate set BRAID, CZ^f , and T^f [31] on the $L_f + 2L_q$ fermions (where T^f gates are implemented using additional qubit ancillas and T, H and CZ^{qf} gates as shown in Fig. 3h). At the end of the circuit, we swap the information back from the fermionic ancillas into the qubits.

It remains to show how to implement dual-rail-qubit SWAP gates. A SWAP gate is decomposed into three CNOT gates of two types, one where the qubit is the control and one where the dual-rail qubit is the control. If the dual-rail qubit is the control, a CNOT is simply $HCZ^{qf}H$, where the CZ^{qf} is acting on the first fermion site (because it is in $|1\rangle$ when the dual-rail qubit is in $|1\rangle$). If the qubit is the control, we decompose the CNOT as

$$e^{i\frac{\pi}{4}\left(1-X^{\mathrm{DR}}\right)\left(1-Z\right)}.$$
 (S2)

Noting that $X^{\text{DR}} = c_1^{\dagger}c_2 + c_2^{\dagger}c_1$, we can decompose this gate into a qubit phase gate, a $\pi/4$ hopping gate, and a $\pi/4$ qubit-controlled hopping gate, whose compilations are shown in Fig. 4b in terms of the claimed gate set.

COMPILATION OF LOGICAL OPERATIONS

We show that the circuits shown in Fig. 2 and Fig. 3 indeed implement the target logical operations.

Useful identities for fermion gates

In this subsection, we provide useful identities for proving circuit identities.

First, we summarize in Table S1 how Majorana and qubit operators transform under our gate set.

BRAID gate identities

In our discrete gate set, the only braiding operation we consider is BRAID_{*ij*} = exp $\left(-\frac{\pi}{4}\tilde{\gamma}_i\gamma_j\right)$. For the compilations of logical gates, the braidings exp $\left(-\frac{\pi}{4}\tilde{\gamma}_i\tilde{\gamma}_j\right)$ and exp $\left(-\frac{\pi}{4}\gamma_i\gamma_j\right)$ are also useful. These can be engineered by

$$\exp\left(-\frac{\pi}{4}\tilde{\gamma}_{i}\tilde{\gamma}_{j}\right) = (\mathbf{S}_{j}^{\mathbf{f}})^{\dagger}\mathbf{BRAID}_{ij}\mathbf{S}_{j}^{\mathbf{f}},$$
$$\exp\left(-\frac{\pi}{4}\gamma_{i}\gamma_{j}\right) = \mathbf{S}_{i}^{\mathbf{f}}\mathbf{BRAID}_{ij}(\mathbf{S}_{i}^{\mathbf{f}})^{\dagger},$$
$$\mathbf{BRAID}_{ij}^{\dagger} = \mathbf{Z}_{j}^{\mathbf{f}}\mathbf{BRAID}_{ij}\mathbf{Z}_{j}^{\mathbf{f}},$$
(S3)

which can be proven by rewriting $\text{BRAID}_{ij} = \frac{1}{\sqrt{2}} (1 - \tilde{\gamma}_i \gamma_j)$ and using the transformations in Table S1. Note that $(\text{BRAID}_{ij})^{\dagger} \neq \text{BRAID}_{ji}$.

	γ_i	$ ilde{\gamma}_i$	γ_j	$ ilde{\gamma}_j$	\mathbf{Z}_i	\mathbf{X}_i
T_i^{f}	$\frac{1}{\sqrt{2}}(\tilde{\gamma}_i + \gamma_i)$	$\frac{1}{\sqrt{2}}(\tilde{\gamma}_i - \gamma_i)$				
$\mathbf{S}_{i}^{\mathrm{f}}$	$ ilde{\gamma}_i$	$-\gamma_i$				
$\mathbf{Z}_{i}^{\mathrm{f}}$	$-\gamma_i$	$- ilde{\gamma}_i$				
H_i					\mathbf{X}_i	\mathbf{Z}_i
Z_i						$-X_i$
BRAID _{ij}		$-\gamma_j$	$\tilde{\gamma}_i$			
CZ_{ij}^{f}	$\gamma_i \mathrm{Z}_j^\mathrm{f}$	$ ilde{\gamma}_i \mathrm{Z}_j^\mathrm{f}$	$\gamma_j \mathbf{Z}_i^{\mathrm{f}}$	$\tilde{\gamma}_j \mathbf{Z}_i^{\mathrm{f}}$		
$\begin{array}{c c} & \text{CZ}_{ij}^{\text{f}} \\ \hline & \text{CZ}_{ij}^{\text{qf}} \end{array}$			$Z_i \gamma_j$	$\mathbf{Z}_i \tilde{\gamma}_j$	\mathbf{Z}_i	$\mathbf{Z}_{j}^{\mathrm{f}}\mathbf{X}_{i}$

TABLE S1. Transformation of Majorana and qubit operators under our gate set. We show $U^{\dagger}AU$, where U is specified in the first column and A in the first row. The indices are defined such that braiding is given by BRAID_{ij} = exp $\left(-\frac{\pi}{4}\tilde{\gamma}_i\gamma_j\right)$ and qubit-fermion controlled-Z as $CZ_{ij}^{af} = exp \left(-i\pi n_i^q n_j\right)$. An empty field means that the operator is invariant. Note that $Z_i^f = -i\tilde{\gamma}_i\gamma_i$.

CZ gate identities

From $(n^{q})^{2} = n^{q}$ and $n^{2} = n$ it follows that

$$\exp(-i\pi n_i^{q} n_j) = 1 - 2n_i^{q} n_j, \exp(-i\pi n_i n_j) = 1 - 2n_i n_j.$$
(S4)

Repetition code logical operations

In this subsection, we show explicitly that the circuits shown in Fig. 2 realize logical gates in the repetition code. Logical gates need to leave the stabilizers invariant. For the repetition code, the logical Majorana fermions are distinct from those on which the stabilizers act (c.f. Fig. 2a). The logical gate compilations below do not act on the Majorana fermions which are part of the stabilizers and therefore leave stabilizers invariant. What is left to show is that our compilations act as they should on the logical Majoranas. To do so, recall that the two logical Majoranas of a single codeblock are given by single physical Majorana fermion operators on the edges, $\gamma^{\rm L} = \gamma_1$ and $\tilde{\gamma}^{\rm L} =$ $\tilde{\gamma}_N$, and, in particular, $n = \frac{1}{2}(1 + i\tilde{\gamma}^{\rm L}\gamma^{\rm L}) = \frac{1}{2}(1 + i\tilde{\gamma}_N\gamma_1)$. In the following, we will drop the "L" superscripts and instead denote the codeblock in the superscript. For example, $\tilde{\gamma}^a_N$ is the $\tilde{\gamma}^{\rm L}$ logical Majorana operator of codeblock *a*.

Logical CZ^{f} gate

Writing the logical CZ^{f} gate between two codeblocks a and b in terms of logical Majoranas, we get

$$\exp\left(-i\pi n^a n^b\right) = \frac{1}{2} \left(1 + i\tilde{\gamma}_N^a \gamma_1^a + i\tilde{\gamma}_N^b \gamma_1^b - \tilde{\gamma}_N^a \gamma_1^a \tilde{\gamma}_N^b \gamma_1^b\right).$$
(S5)

We construct this gate from a physical CZ^{f} gate between the two edge sites, which in terms of the Majoranas is written as

$$\exp\left(-i\pi n_1^a n_1^b\right) = \frac{1}{2} \left(1 + i\tilde{\gamma}_1^a \gamma_1^a + i\tilde{\gamma}_1^b \gamma_1^b - \tilde{\gamma}_1^a \gamma_1^a \tilde{\gamma}_1^b \gamma_1^b\right), \qquad (S6)$$

where n_1^a is the number operator acting on the first site of codeblock *a*. While the general form is already correct, we need to replace $\tilde{\gamma}_1^b \rightarrow \tilde{\gamma}_N^b$ and $\tilde{\gamma}_1^a \rightarrow \tilde{\gamma}_N^a$. Such a "replacement operation" is performed by conjugating with braidings (see Table S1):

$$\exp\left(-i\pi n^a n^b\right) = e^{\frac{\pi}{4}\tilde{\gamma}^b_N\tilde{\gamma}^b_1}e^{\frac{\pi}{4}\tilde{\gamma}^a_N\tilde{\gamma}^a_1}e^{-i\pi n^a_N n^b_1}e^{-\frac{\pi}{4}\tilde{\gamma}^a_N\tilde{\gamma}^a_1}e^{-\frac{\pi}{4}\tilde{\gamma}^b_N\tilde{\gamma}^b_1}.$$
 (S7)

Using Eq. (S3) to express this sequence in terms of BRAID and S^f gates, we find the circuit shown in Fig. 2d. Note that this compilation also works in the same way for an arbitraryangle gate $\exp(-i\theta n^a n^b)$ by replacing the physical CZ^f with $e^{-i\theta n_N^a n_1^b}$.

Logical braiding gate gate.

A logical braiding gate is given in terms of the Majorana fermions as

$$BRAID_{ab} = \exp\left(-\frac{\pi}{4}\tilde{\gamma}_N^a\gamma_1^b\right).$$
 (S8)

This is in fact a physical braiding gate between the edges of two codeblocks, as depicted in Fig. 2d.

Logical qubit-fermion gate.

Similarly, we decompose the logical qubit-fermion gate as

$$\exp\left(-i\pi n^{q}n^{a}\right) = e^{\frac{\pi}{4}\gamma_{1}^{a}\gamma_{N}^{a}}e^{-i\pi n^{q}n_{N}^{a}}e^{-\frac{\pi}{4}\gamma_{1}^{a}\gamma_{N}^{a}}.$$
 (S9)

Again using Eq. (S3), we find the circuit shown in Fig. 2d.

Logical phase gate.

A logical fermion phase gate $\exp(i\theta n^a)$ is written in terms of the Majorana fermions as

$$\exp\left(i\frac{\theta}{2}(1+i\tilde{\gamma}_N^a\gamma_1^a)\right).$$
 (S10)

Therefore, this gate is, up to a global phase, equivalent to an arbitrary-angle braiding between the two edge Majoranas. In particular, the S^f gate is equivalent to BRAID and the T^f gate is a $\pi/8$ braiding.

Syndrome measurement.

For the measurement of the syndromes $i\tilde{\gamma}_i\gamma_{i+1}$, we study the following transformation of the final Z measurement of the ancilla qubits: $U^{\dagger}Z_{\rm anc}U$ where we use $U = H_{\rm anc}CZ_{{\rm anc},i+1}^{\rm qf}BRAID_{i+1,i}S_i^{\rm f}H_{\rm anc}$. Using the transformation rules in Table S1, we find $U^{\dagger}Z_{\rm anc}U = Z_{\rm anc}i\tilde{\gamma}_i\gamma_{i+1}$, and, therefore, initializing the ancilla in $|0\rangle$, a measurement of the ancilla yields a measurement of $i\tilde{\gamma}_i\gamma_{i+1}$. The additional gates in Fig. 2b compared to the circuit defined here are needed to not disturb the logical information. This enables true mid-circuit error correction.

Color code logical operations

In this subsection, we show that the circuits in Fig. 3 for CZ^{t} , BRAID, and CZ^{qf} are indeed logical gates.

We show that stabilizers transform to products of stabilizers and that the gates transform the logical Majorana operators in exactly the same way as the physical gates transform physical Majorana operators (c.f. Table S1). Therefore, we study the unitary transformation $U^{\dagger}\gamma^{L}U$ and $U^{\dagger}\tilde{\gamma}^{L}U$ of the logical Majorana operators under the circuits shown in Fig. 3. All of the gates are transversal because each physical gate acts only on a single physical fermion in each codeblock, similarly to the qubit-only architecture in Ref. [33].

Similar to the notation in the previous Methods subsection on repetition code logical operations, we attach a superscript to physical Majorana operators to indicate the codeblock and let the physical site index *i* only run from 1 to the number of physical sites within the codeblock. For example, γ_i^a indicates physical lattice site *i* in codeblock *a*, where $i \in \{1, 7\}$ for the seven-site triangular color code. The below calculations hold for all triangular color codes.

BRAID gate

The gate

$$BRAID_{ab} = \prod_{i} \exp\left(-\frac{\pi}{4}\tilde{\gamma}_{i}^{a}\gamma_{i}^{b}\right)$$
(S11)

is a logical braiding gate. To see this, first note that the individual braiding gates commute with Majorana operators on sites $j \neq i$ because they are quadratic. Therefore, this gate transforms the logical operators $\tilde{\gamma}^a = \prod_i \tilde{\gamma}_i^a$ and $\gamma^b = \prod_i \gamma_i^b$

$$BRAID_{ab}^{\dagger}\tilde{\gamma}_{b}^{L}BRAID_{ab}$$

$$=\prod_{i}\exp\left(\frac{\pi}{4}\tilde{\gamma}_{i}^{a}\gamma_{i}^{b}\right)\gamma_{i}^{b}\exp\left(-\frac{\pi}{4}\tilde{\gamma}_{i}^{a}\gamma_{i}^{b}\right)$$

$$=\prod_{i}\tilde{\gamma}_{i}^{a}=\tilde{\gamma}_{a}^{L},$$

$$BRAID^{\dagger},\tilde{\gamma}_{i}^{L}BRAID_{ab}$$
(S12)

$$= \prod_{i} \exp\left(\frac{\pi}{4}\tilde{\gamma}_{i}^{a}\gamma_{i}^{b}\right)\tilde{\gamma}_{i}^{a}\exp\left(-\frac{\pi}{4}\tilde{\gamma}_{i}^{a}\gamma_{i}^{b}\right)$$
$$= \prod_{i} \left(-\gamma_{i}^{b}\right) = -\gamma_{b}^{L}, \tag{S13}$$

where in the very last step we used the fact that the triangular color codes have an odd number of physical fermions.

Stabilizers transform to stabilizers because they are products of an even number of Majorana fermions consisting of only either γ or $\tilde{\gamma}$, but not both. Braiding will transform them to products of the other type residing on the same lattice sites of the other codeblock. Because of the self-duality (in the sense of the symmetry $\gamma \leftrightarrow \tilde{\gamma}$ of the stabilizers) of our code, the resulting product is also a stabilizer. Because stabilizers consist of an even product of Majorana operators, there is also no minus sign resulting from the transformation.

 S^{f} gate

The transversal gate

$$\mathbf{S}_{a}^{\mathrm{f}} = \prod_{i} \mathbf{S}_{i}^{\mathrm{f}} \tag{S14}$$

is shown to be a logical phase gate S^{f} in the same way as the BRAID gate.

The CZ^f gate is given by

$$CZ_{ab}^{f} = \prod_{i} \exp\left(-i\pi n_{i}^{a} n_{i}^{b}\right).$$
(S15)

To check the transformation of the logical operators, consider

$$\left(\mathrm{CZ}_{ab}^{\mathrm{f}}\right)^{\dagger} \gamma^{a} \mathrm{CZ}_{ab}^{\mathrm{f}} = \prod_{i} \exp\left(i\pi n_{i}^{a} n_{i}^{b}\right) \gamma_{i}^{a} \exp\left(-i\pi n_{i}^{a} n_{i}^{b}\right)$$
(S16)

$$= \prod (-i\gamma_i^a \tilde{\gamma}_i^b \gamma_i^b) \tag{S17}$$

$$= -i\gamma_a^{\rm L}\tilde{\gamma}_b^{\rm L}\gamma_b^{\rm L}.$$
 (S18)

The last step is non-trivial: the global sign is a combination of the overall $(-i)^n$ and a minus sign resulting from reordering

the physical Majorana fermions. The triangular Honeycomb color codes have $n = (3d^2 + 1)/4$ physical fermions for odd d [62]. For convenience, we define a \tilde{d} as $d = 2\tilde{d} + 1$, and write $n = 3\tilde{d}^2 + 3\tilde{d} + 1$. One can show that $\prod_{i=1}^n (\gamma_i^a \tilde{\gamma}_i^b \gamma_i^b) = (-1)^{\lfloor n/2 \rfloor} \prod_{i=1}^n \gamma_i^a \prod_{i=1}^n \tilde{\gamma}_i^b \prod_{i=1}^n \gamma_i^b$. Therefore, the overall prefactor is given by $(-i)^{3d^2+3\tilde{d}+1}(-1)^{\lfloor (3\tilde{d}^2+3\tilde{d}+1)/2 \rfloor} = -i$ for all integer \tilde{d} .

Similarly, stabilizers transform to products of stabilizers after the transformation. To see the global sign, note that, in the triangular color codes, stabilizers are products of either four or six Majorana operators. For four operators, $(-i)^4 = 1$ and the sign from reordering is $(-1)^{\lfloor 4/2 \rfloor} = 1$. For six operators, $(-i)^6 = -1$ and the sign from reordering is $(-1)^{\lfloor 6/2 \rfloor} = -1$, again yielding an overall +1.

CZ^{qf} gate

Our qubit-fermion gate CZqf is implemented as

$$CZ_{ab}^{qf} = \prod_{i} \exp\left(-\pi n_i^{q,a} n_j^b\right).$$
(S19)

This is shown in the same manner as the previous gates. In particular, the fact that $\prod_{i=1}^{n} (\tilde{\gamma}_{i}^{a} \gamma_{i}^{a}) = (-1)^{\lfloor n/2 \rfloor} \prod_{i=1}^{n} \tilde{\gamma}_{i}^{a} \prod_{i=1}^{n} \gamma_{i}^{a}$ leads to the correct minus signs. Stabilizers map onto products of fermion and qubit stabilizers under the action of this gate. Therefore, it is a valid logical qubit-fermion gate.

PAIRING GATE

In this section, we present some details of the neutral-atom scheme for implementing pairing presented in the main text.

Non-local pairing from on-site pairing

We demonstrate that the pulse sequence shown in Fig. 5a of the main text indeed implements the target inter-site braiding gate

$$\text{PAIR}_{ij} = \exp\left(i\frac{\pi}{4\sqrt{N}}(e^{i\varphi}b_k p_i^{\dagger}p_j^{\dagger} + h.c.)\right), \quad (S20)$$

where we generalized the definition in the main text to include a phase φ . The pulse sequence is arranged such that the dissociation gate

$$U_{\rm dis} = \exp\left(-i\frac{\pi}{4\sqrt{N}} (e^{i\varphi}b_i p_{i\uparrow}^{\dagger} p_{i\downarrow}^{\dagger} + h.c.)\right)$$
(S21)

is conjugated with a series of unitaries. We therefore have to show that these unitaries transform bosonic and fermionic creation/annihilation operators in such a way that U_{dis} is transformed into PAIR_{*ij*}. First, the beamsplitter gate

$$BS = \exp\left(\frac{\pi}{4}(b_i^{\dagger}b_k - h.c.)\right)$$
(S22)

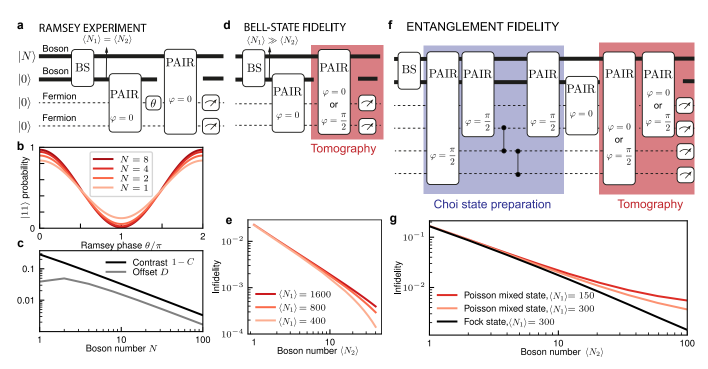


FIG. S1. Benchmarking the pairing gate. Pairing gate definition in Eq. (S20), with the occupation number entering its phase given by $\langle N_i \rangle$ of the mode *i* it acts on. **a**, Circuit for implementing the Ramsey circuit, reproduced from Fig. 5c of the main text. The beamsplitter phase is $\pi/4$. **b**, Ramsey fringes, reproduced from Fig. 5d of the main text. **c**, Contrast *C* and offset *D* obtained from fitting $\frac{1}{2} + \frac{1}{2} (C \cos(\theta) - D)$ to the Ramsey fringes in subfigure b. **d**, Circuit for performing tomography within the $|00\rangle$ and $|11\rangle$ manifold. The beamsplitter phase is $\arccos\left(\sqrt{\langle N_2 \rangle/(\langle N_1 \rangle + \langle N_2 \rangle)}\right)$, chosen such that the average occupation number of the first, second bosonic mode (counted from the top) is $\langle N_1 \rangle$, $\langle N_2 \rangle$ after the beamsplitter, respectively. **e**, Infidelity $1 - \langle \psi | \rho | \psi \rangle$, where $|\psi\rangle = (|00\rangle + i |11\rangle)/\sqrt{2}$ and ρ is the density matrix reconstructed from the numerical tomography. Initial state is the same is in a. **f**, Circuit for numerically measuring the average gate fidelity for all possible input states of the form $\alpha |00\rangle + \beta |11\rangle$. The Choi state is defined as $|\psi_{\text{Choi}}\rangle = (|0000\rangle + |1111\rangle)\sqrt{2}$. The tomography is performed within the subspace spanned by the states $|0000\rangle$, $|0011\rangle$, $|1100\rangle$, $|1111\rangle$. **g**, Infidelity $1 - \langle \psi_{\text{Choi}} | \rho | \psi_{\text{Choi}} \rangle$ for either a Fock state input $|N\rangle$ on mode 1 (black line) or a mixed state on mode 1 with a Poissonian number distribution with average boson number $\langle N_1 \rangle + \langle N_2 \rangle$ (colored lines). The black line is shown in Fig. 5e of the main text.

acts as $(BS^{\dagger})^{2}b_{i}^{\dagger}BS^{2} = b_{k}^{\dagger}$. Hence, $(BS^{\dagger})^{2}U_{dis}BS^{2} = \exp\left(-i\frac{\pi}{4\sqrt{N}}(b_{k}p_{i\uparrow}^{\dagger}p_{i\downarrow}^{\dagger} + h.c.)\right)$. Similarly,

$$U_{t\downarrow} = \exp\left(-i\frac{\pi}{2}(p_{i\downarrow}^{\dagger}p_{j\downarrow} + h.c.)\right)$$
(S23)

acts as $U_{\mathrm{t}\downarrow}^{\dagger}p_{i\downarrow}^{\dagger}U_{\mathrm{t}\downarrow}=ip_{j\downarrow}^{\dagger}$ and

$$U_{\uparrow\downarrow} = \exp\left(-i\frac{\pi}{2}(p_{i\uparrow}^{\dagger}p_{i\downarrow} + h.c.)\right)$$
(S24)

acts as $U^{\dagger}_{\uparrow\downarrow}p^{\dagger}_{i\uparrow}U_{\uparrow\downarrow}=ip^{\dagger}_{i\downarrow}.$ Therefore,

$$\mathrm{PAIR}_{ij} = U_{\uparrow\downarrow}^{\dagger} U_{\mathrm{t}\downarrow}^{\dagger} (\mathrm{BS}^{\dagger})^2 U_{\mathrm{dis}} \mathrm{BS}^2 U_{\mathrm{t}\downarrow} U_{\uparrow\downarrow}, \qquad (S25)$$

which is the sought-after pulse sequence when identifying spinless fermions with fermions in the \downarrow state. The Hermitian conjugate of each gate is performed by conjugating with phase gates, e.g. $U_{\uparrow\downarrow}^{\dagger} = Z_{\downarrow}^{f} U_{\uparrow\downarrow} Z_{\downarrow}^{f}$.

Benchmarking the pairing gate

Here we show that Eq. (S20) indeed effectively implements a pairing gate as defined in Eq. (8) when the molecule/boson mode on which it acts has large occupation. To do so, we discuss three different standard benchmark experiments and study them numerically: Ramsey spectroscopy, state tomography of a Bell state, and process/gate tomography based on measuring the entanglement fidelity.

Ramsey experiment

First, we want to test whether there is effective coherence created after applying the pairing gate. As for qubits, this is done by performing a Ramsey-type experiment, i.e. applying two $\pi/2$ pairing pulses, with a phase gate in-between. At the same time as testing whether this is the case, we also test whether two separate molecule/boson modes can be used for the two pulses. It is clear that these gates need to be effectively phase coherent. We achieve this by creating two modes with Poissonian number distributions, mimicking a coherent state, by applying a 50/50 beamsplitter on the state $|N0\rangle$, see Fig. S1a (also shown in Fig. 5c of the main text). Being able to perform both gates using separate bosonic modes is important—this tests whether ultimately we can parallelize pairing operations. In Fig. S1b (also shown in Fig. 5d of the main text), we show the result of the Ramsey sequence. We indeed find a Ramsey fringe, verifying that effective coherence has been created in the fermions. The contrast increases with N as shown in Fig. S1c, and a slight offset of the oscillations is visible. More quantitatively, we fit an oscillatory function to the fringe (see figure caption for its definition), finding that the contrast approaches unity as $\propto 1/N$. At the same time, the offset decreases with the same functional dependence. Hence, in the limit $N \rightarrow \infty$, perfect Ramsey oscillations are approached.

Bell-state fidelity

In order to probe that the state after the pairing gate is not only an effective coherent superposition, but exactly the state that we aim to prepare, in Fig. S1d we construct a circuit effectively performing Bell-state tomography. To do so, we perform three separate circuits: we always first apply a pairing gate to effectively prepare the Bell state $|00\rangle + i |11\rangle$ and then perform either a pairing gate with $\varphi = 0$ or $\varphi = \pi/2$ or no pairing gate at all. From these three measurements, we reconstruct the density matrix in the $|00\rangle$ and $|11\rangle$ manifold. In order to separate the boson-number dependence of the preparation step from the tomography step, we choose the beamsplitter phase such that the boson number N_1 of the mode we use to perform the second gate is much larger than the boson number N_2 of the mode we use to perform the first gate. Having reconstructed the density matrix, we show the infidelity of the Bell-state preparation as a function of N_2 , for different values of N_1 . We find in Fig. Fig. S1e that the curves collapse as $N_1 \rightarrow \infty$, indicating again a power-law dependence of $\propto 1/N.$ Crucially, already for $N_2 \approx 20$, we find a Bell-state fidelity of approx. 99.9%.

Gate fidelity

Finally, we would like to benchmark our gate for all possible initial states. To do so, we perform gate/process tomography by measuring the entanglement fidelity F_{e} , relying on the relation

$$F_{\rm avg} = 1 - \frac{dF_{\rm e} + 1}{d+1},$$
 (S26)

where d is the Hilbert-space dimension and

$$F_{\rm avg} = \int \mathrm{d}\psi \, \langle \psi | U^{\dagger} \mathcal{E}(|\psi\rangle \, \langle \psi |) U | \psi \rangle \tag{S27}$$

is the state-averaged fidelity between the target unitary U and the realized channel \mathcal{E} . The entanglement fidelity is measured by first preparing the Choi state $|\psi_{\text{Choi}}\rangle = \frac{1}{\sqrt{d}} \sum_{i=1}^{d} |i\rangle |i\rangle$, constructed using an ancillary system of the same size as the target system, applying the target channel $\mathcal{E} \otimes \mathbb{1}$, and finally performing state tomography on the resulting state. In our case, \mathcal{E} is the pairing gate and hence all gates stay in the even parity sector such that d = 2, $|\psi_{\text{Choi}}\rangle = \frac{1}{\sqrt{2}} (|00\rangle |00\rangle + |11\rangle |11\rangle)$, and the tomography is only performed in that sector.

We show the entanglement fidelity in Fig. S1g, black line (same as Fig. 5e), as a function of the boson number N_2 in the second mode, where the result is converged with respect to N_1 . We find that > 99.8 % gate fidelity is achieved for $N_2 = 100$.

Finally, in experiment, it is much easier to prepare a mixed state with Poissonian number distribution with average number $\overline{N} = N_1 + N_2$ rather than a Fock state, for example by preparing a Bose-Einstein condensate [63] in that tweezer (\bar{N} is the number of condensed molecules in the ground state of the tweezer). In that case, there will be a shot-to-shot miscalibration of the angle of the pairing gate, leading to a further source of infidelity. To check the robustness of our gate with respect to this effect, we replace the initial state $|N0\rangle$ of the molecules with a state $\rho_{\bar{N}} \otimes |0\rangle \langle 0|$, where $\rho_{\bar{N}} = \sum_{n} p_{\bar{N}}(n) |n\rangle \langle n|$ and $p_{\bar{N}}(n) = \frac{\bar{N}^n e^{-\bar{N}}}{n!}$ is a Poisson distribution with mean \bar{N} . The phase of the pairing gate is fixed to $\pi/(4\sqrt{N})$. We show the resulting average infidelity as red lines in Fig. S1g. We find that, for small N_2 , the result is indistinguishable from the Fock-state result. For larger N_2 , deviations occur. These deviations become smaller as N_1 increases. Hence, we can expect that as \overline{N} and N_2 increase, arbitrarily high fidelities can be achieved even when the initial molecule state is a mixed state with a Poissonian number distribution.

Experimental considerations

We discuss more detailed considerations for the parameter regimes that need to be fulfilled to implement the pairing scheme.

For spin-selective tunneling $U_{t\downarrow}$, a magnetic field gradient needs to be applied to suppress tunneling of the \uparrow spin species. Similarly, to guarantee that no fermionic atoms tunnel from the target tweezer into the BEC tweezer during the application of the beamsplitter (BS), we apply a magnetic field gradient and detune the two trap frequencies such that the detuning induced by the magnetic field gradient is compensated by the tweezer detuning for the molecules. Because the molecules carry twice the magnetic moment of atoms, the atom tunneling is still suppressed by the residual gradient. For dissociation, the rf frequency $\nu_{\rm RF}$ is chosen to bridge the bare atom energy $u_{\uparrow} - \nu_{\downarrow}$ as well as the binding energy $E_{\rm B}$ of the molecule and the trap depth $\omega_{tweezer}$ of the target tweezer (which again enters due to the differing magnetic moments of atoms and molecule), $\nu_{\rm RF} = \nu_{\uparrow} - \nu_{\downarrow} + E_{\rm B} + \omega_{\rm tweezer}$. This can be easily fulfilled: typical trap frequencies are in the tens of kHz, and binding energies in ⁶Li₂ are in the hundreds of kHz, for ⁸⁴Sr₂ even in the hundreds of MHz [45, 64, 65]. Binding energies may be tuned by a Feshbach resonance, however this needs to be counterbalanced with the interactions between bosons and fermions as well as between the \downarrow and \uparrow fermions, which need to be small for the duration of the gate. The coupling q also needs to be much smaller than the tweezer trap frequency in order to guarantee that the atom pair is created in the motional ground state of the tweezer.

Implementation in dots and donors in silicon

In a solid-state implementation in dots or donors, spinpolarized electrons encode fermions. Qubits with long coherence times can be encoded in nuclear spins, in which case a CZ^{qf} gate is implemented using the hyperfine interaction [66]. Alternatively, a qubit can be encoded in the electron spin. Both qubit-qubit and qubit-fermion entangling gates are then implemented by Heisenberg exchange [67, 68]. Single-fermion gates may be realized using electric-field gates, and the pairing gate using the proximity effect to a superconductor [69], where Cooper pairs take the role of molecules.