Error characterization in quantum information processing: A protocol for analyzing spatial correlations and its experimental implementation

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We present a protocol for error characterization and its experimental implementation with four qubits in liquid state NMR. The method is designed to retrieve information about spatial correlations and scales as $O(n^w)$, where $w$ is the maximum number of qubits that have non-negligible interaction. We discuss the practical aspects regarding accuracy and implementation.

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I. INTRODUCTION

Precise and reliable control of the system constituting a quantum information processor (QIP) remains one of the biggest challenges in the quantum information field. In order to assess the reliability of a device and to tailor quantum error correction schemes to a faulty one, we need to characterize the errors occurring in the system. Quantum process tomography (QPT) [1] presented a first answer to this problem, providing full characterization of the process under analysis. Experimental implementations of QPT have been already conducted in a variety of small systems [2–5]. Nevertheless, QPT becomes impractical beyond a few qubits, as it requires $O(2^m)$ experiments for a system of $n$ qubits. In recent years, the idea of getting less information at a lower cost has become a popular strategy in tackling error characterization, and several works have been devoted to the subject [6–12]. Our proposal fits in this context, providing a subset of information yet still using scalable resources.

The scheme we present selectively keeps information about the spatial correlations of the errors occurring in the process under study (a gate, a noisy channel, etc.). Both the magnitude and the structure of the errors are relevant to evaluate fault-tolerance. In particular, fault-tolerance threshold theorems are designed for certain conditions of spatial correlation (also termed range or locality) [13]. So even when it is experimentally determined that only up to $w$ qubits are involved in an error process, we have to further establish in which way the $\binom{n}{w}$ possible sets are being affected.

We have implemented our protocol in a liquid state NMR four-qubit QIP. The core mathematical work for this protocol was introduced in [7]. Here we extend our proposal to a more general setting and include an experimental realization. Our basic method, like others proposed [1,6–8,10,11], assumed error-free implementation stages. This idea is of course unrealistic in practice, and implementation errors complicate the task of reliable error characterization. Thus here we have included an analysis of their effect.

II. THEORY BEHIND THE PROTOCOL

We start by describing the action of a general map on the state of an $n$-qubit system ($D=2^n$), described by an initial state $\rho_0$, as

$$S(\rho_0) = \int p(\vec{\eta})E(\vec{\eta})\rho_0E^\dagger(\vec{\eta})d\vec{\eta}. \quad (1)$$

The vector $\vec{\eta}$ denotes $D^2$ complex coefficients $\{\eta_0, \eta_1, \ldots, \eta_{D^2-1}\}$ that parametrize $E$, an arbitrary operator in the Hilbert space $\mathcal{H}_D$, as

$$E = \eta_0 I + \sum_{l=1}^{D^2-1} \eta_l O_l, \quad O_l = \bigotimes_{j=1}^n O_l^{(j)}, \quad (2)$$

where each $O_l^{(j)}$ is an element of the Pauli group $\{I, \sigma_x, \sigma_y, \sigma_z\}$ but at least one factor in each $O_l$ is a Pauli matrix. Notice that $\text{Tr}[O_l O_{l'}] = D(\delta_{ll'} - \frac{1}{D})$. When $p(\vec{\eta})$ is a (real) non-negative distribution, Eq. (1) describes an arbitrary completely positive (CP) map, and the condition $\int p(\vec{\eta})|\vec{\eta}|^2d\vec{\eta} = 1$ guarantees the preservation of $\text{Tr}[\rho]$. This representation of a CP map is an operator-sum representation with continuous parameters. We prefer this form as it is more suitable to describe nonunitary dynamics arising from stochastic Hamiltonians. Our parameters, the $\vec{\eta}$, are trivially related to the parametrizations of the operator-sum representations used in other works [8,10]. Also, for small $\eta_l$ it is possible to relate Eq. (1) to a description of the noise in terms of generators rather than propagators [7].

When necessary, we shall denote the $\eta_l$ in more detail as $\eta_{j,k}^{x,y,z}$, where $j > k > \ldots$ label qubits, and $p, q, \ldots = x, y, z$. Therefore $\eta_{j,k}^{x,y,z}$ labels a term in Eq. (2) that is a product of $\sigma_p$ for qubit $j$, $\sigma_q$ for qubit $k$, etc., and $I$ for the qubits absent in the subscript. Notice that the number of qubits in the subscript gives the Pauli weight (also Hamming weight) of the term [14].

In our protocol we measure a subset of $m$ qubits at a time, which will allow us to extract the magnitude of the errors involving that subset. So we now break the system in two: $m$ qubits belonging to the Hilbert space $\mathcal{H}_M$, which will be the ones measured, and $\bar{m}=n-m$ qubits belonging to the complementary space $\mathcal{H}_{\bar{M}}$, $\mathcal{H}_M = \mathcal{H}_\bar{M} \otimes \mathcal{H}_{\bar{M}}$. We require the initial state to be separable in these two spaces, and within $\mathcal{H}_M$, thus

$$\rho_0 = \rho_0^{(M)} \otimes \rho_0^{(\bar{M})} \quad \text{and} \quad \rho_0^{(M)} = \bigotimes_{j \in M} \rho_0^{(j)}. \quad (3)$$

We now perform a $U(2)^m$ twirl on the target map $S$, producing the effective map $S^{\mathcal{H}_M}$.
where the $C_j$ are $m$-fold tensor products of the twirl operators on one qubit, for the $m$ qubits in $H_q$. For the task we propose, a subset of $K=6$ operators from the Clifford group will suffice. We refer to this minimum set of operators required to apply the twirl [Eq. (4)] as the 6$^m$-Clifford element pool (see Appendix A for references and some mathematical details on this twirling).

The reduced density matrix $\rho_1^{(M)} = \text{Tr}_q[\rho_1]$ is the state of the $m$ qubits we measure. In general, the fidelity decay, which gives a measure of how $\rho_1$ differs from the initial $\rho_0$, encodes information about the map $S$ we are characterizing [6,7]. In particular, considering only the qubits we will measure and expressing $\rho_0^{(M)} = (I^{\otimes m} + \rho_{\text{dev}}^{(M)})/2^m$, we obtain

$$\text{Tr}[(\rho_1^{(M)})^2] - \text{Tr}_m[\rho_0^{(M)}\rho_1^{(M)}] = \gamma^{(M)} - \Omega^{(M)}_{\text{dev}},$$

and $\Omega^{(M)}_{\text{dev}} = 0$ if $\rho_{\text{dev}}^{(M)} = 0$ ($\Omega^{(M)}_{\text{dev}}$, for $\rho_{\text{dev}}^{(M)} \neq 0$ can be exactly computed if desired). The quantity defined in the left-hand side of Eq. (5) is the fidelity decay rate [6,7] we will analyze in this work. In Eq. (6) we have denoted $\langle \ldots \rangle = \int d\tilde{\eta} \ldots d\tilde{\eta}$, $\mathcal{P}_j = \text{Tr}[(\rho_j^{(i)})^2]$ the initial purity of each of the $M$-qubits, and $\mathcal{C}_j(l) = \left\{ \begin{array}{ll} (2/3)(1 - \mathcal{P}_j/2) & \text{if } O_j^l = \sigma_x, \sigma_y, \sigma_z \\ \mathcal{P}_j & \text{if } O_j^l = I \end{array} \right.$

For the derivation of Eqs. (5) and (6) we use the equivalence between a Clifford twirl and a Haar twirl [15], and apply the tools developed in [16]. See also Appendix A.

In sum (6), a $t$-term vanishes when $O_t$ is the identity operator for the $m$ qubits being measured. So by choosing different sets of $M$-qubits, it is possible to leave out certain $\eta_j$ in a given $\gamma$. Note however that $C_j$ does not distinguish the direction of the Pauli matrices, so there is an implicit coarse-graining of all the $O_t$ that have the identity $I$ for the same subset of $m$ qubits. This is how we get the “collective coefficients” $\eta_{j}^{\text{col}}$,

$$|\eta_j^{\text{col}}|^2 = \sum_{p=x,y,z} |\eta_j^p|^2, \quad |\eta_j^{\text{col}}|^2 = \sum_{p=x,y,z} |\eta_j^{p,\text{col}}|^2, \quad \text{etc.} \tag{8}$$

For example, the terms $\sigma_x(1)\sigma_x(3)$ and $\sigma_x(1)\sigma_z(3)$ contribute collectively to $\eta_{j}^{\text{col}}$.

By combining the $\gamma$’s from different $M$-sets it is possible to further isolate the collective coefficients. If we prepare $a$ and $b$ in a pure state so $\mathcal{P}_a = \mathcal{P}_b = 1$, we obtain

$$\frac{9}{4} (\gamma^{(a)} + \gamma^{(b)} - \gamma^{(a,b)}) = \langle \eta_{a,b}^{\text{col}}|^2 \rangle + \sum_j \langle \eta_{a,b,j}^{\text{col}}|^2 \rangle + \cdots. \tag{9}$$

(See Table B for the formulas of the $\gamma$’s involved in this example.) If three-body and higher multibody terms can be neglected, Eq. (9) gives $\langle \eta_{a,b}^{\text{col}}|^2 \rangle$. Similarly, the combination of the seven $\gamma^{(a)}$, $\gamma^{(b)}$, and $\gamma^{(a,b)}$ for a set of three qubits $a,b,c$ would return $\langle \eta_{a,b,c}^{\text{col}}|^2 \rangle$, and so on. The collective coefficients then report on the spatial correlations of errors.

### III. PROTOCOL

We present now a systematic protocol for measuring the collective coefficients involving the $m$ qubits of a particular subset.

(i) Prepare each of the qubits to be measured in the initial state $|0\rangle$. Prepare each of the other $m$ qubits in the maximally mixed state $I/2$.

(ii) Apply one of the $m$-fold Clifford operators from the 6$^m$-Clifford element pool.

(iii) Implement the target gate or noise under study.

(iv) Invert the Clifford operator applied in (ii).

(v) Measure the projection of the resulting state on the initial state $|0\rangle$, for each of the qubits being measured.

To implement the twirl, we repeat (i)–(v) each time taking a different operator from the 6$^m$-Clifford element pool, and average the results.

In a canonical QIP, the implementation of this protocol to measure the decay rates involving $m$ qubits will require $N$ realizations. This will take care of: (1) preparing the desired initial state as in step (i) (starting from the $|0\rangle^{\otimes m}$ state, and randomly flipping the $\bar{m}$ qubits we do not measure); (2) measuring the $\gamma$’s through repeated projective measurements of the $m$ qubits as prescribed in step (v) (notice Eq. (5) becomes $1 - \text{Tr}_m[\rho_0^{(M)}\rho_1^{(M)}] = \gamma^{(M)}$ for the proposed initial state); (3) implementing the twirl approximately, by randomly drawing the twirl operators for steps (ii) and (iv) from the corresponding 6$^m$-Clifford element pool (or from an infinite pool of one-qubit random rotations).

With this strategy, the outcome of the measurement step (v) is a Bernoulli variable, and thus $N$ can be estimated from usual statistics. More precisely, the standard error in our estimation of the decay rate will be $\sigma_N \leq 1/\sqrt{N}$ (following the central limit theorem) so for a desired precision $\delta$ we must have $N \geq \delta^2$. On the other hand, the Chernoff bound tells us that, for a desired precision $\delta$ and an error probability $\epsilon_N \ll 1$, we must have $N = \log(2/\epsilon_N)/(2\delta^2)$, which is a stronger requirement when $\delta < 2e^{-2}$. In any case, $N$ is independent of the number of qubits; thus the efficiency of the protocol is independent of the size of the system.

In the case of liquid state NMR ensemble QIP, pseudopure state preparation allows for initialization in the $I/2$ state over the ensemble of molecules, and ensemble measurements avoid the need of repeated realizations in order to perform step (v) by quantum state tomography (QST). This is the case in our experiment.

The measurement step (v) retrieves the information to calculate the decay rates for the $m$ qubits and for any smaller subset of qubits within those. For example, twirling qubits $a$ and $b$ only ($m=2$), we can obtain $\gamma^{(a)}$, $\gamma^{(b)}$, and $\gamma^{(a,b)}$ in one shot, and calculate $\langle \eta_{a,b}^{\text{col}}|^2 \rangle$ as in Eq. (9) neglecting the higher multibody terms. This procedure can be repeated for the $\binom{m}{2} = m(m-1)/2$ pairs of qubits, and by doing so all the collective coefficients for one- and two-body terms can be extracted.

The scalability of the method goes as follows. If we can neglect the multi-body terms above a certain Pauli weight $w$,
and \( N \) is the number of realizations required to measure the fidelity \( \gamma \) for \( w \) qubits, then with \( N(C_w) \leq Nw^w/w! \) experiments we can estimate all the non-negligible coefficients. This should be compared against \( N2^w \), the overhead in the number of experiments required for QPT. We emphasize here that our proposal seeks to characterize the correlations among up to \( w \) qubits in order to establish the range of the noise, not to characterize the process fully.

To use our protocol, the negligibility of multi-body terms above a certain Pauli weight \( w \) must be established \textit{a priori}. In a canonical QIP we could apply our protocol to measure all the \( n \) qubits, obtain all the decay rates \( \gamma(j), \gamma(k), \ldots, \gamma(1,\ldots,n) \), and extract all the collective coefficients after only \( N \) experiments (independently of \( n \)). In this way we can handle all the Pauli weights, from zero to \( n \). But the error in the decay rates \( \sigma_j \) propagates into the \( \eta_{\text{col}}^{i,j} \) for \( m \) qubits inefficiently, roughly as \( \sigma_j \chi = \sum_{j=1}^{w} (\xi_j)^{i,j} \sigma_j \). Therefore our strategy is to look for spatial correlations after establishing a cutoff Pauli weight \( w \) to our knowledge, neither QPT nor other proposals so far are able to make use of the negligibility of high order correlations in order to gain further insight. In general, even after it has been established that a subset of parameters is null, it is not trivial to direct the measurement procedure to target the non-null ones exclusively.

An approach to establishing this cutoff \( w \), demanding the same resources as our protocol, is to apply the method developed by Emerson et al. \cite{8} which gives the probability of errors happening, distinguishing them only by Pauli weight (this is an average of all the \( \xi_j^{i,j} \) coefficients having Pauli weight \( w \)). It is worth pointing out that both methods require the same experimental work: the algorithms retrieve different information because the measurement and processing of the data is different, but actually both protocols can be implemented simultaneously and used complementarily.

### IV. EXPERIMENTAL RESULTS

We implemented our protocol in a liquid state NMR QIP using the four \( ^{13}\text{C} \)-labeled carbons of crotonic acid \cite{17} in a 400 MHz Bruker spectrometer. The initial state preparation and all the gates required by the protocol were implemented with rf pulse sequences engineered using either gradient ascent pulse engineering (GRAPE) \cite{18} or strongly modulating pulses (SMP) \cite{17(c)} methods. Their simulated gate fidelities \( F_g \) were on average 0.98. The typical experimental performance of one-qubit gates on the spectrometer is 1%–2% below their simulated fidelities. See \cite{2} for details on the model used in the simulation.

We studied the following processes (see Table I for more details): (i) A time suspension sequence \( I_G \) since it is important to study our ability to “do nothing” in a system with a natural Hamiltonian that is always on. (ii) An engineered error creating a coupling between qubits 1 and 2, of the form \( C_{12}(\beta) = \exp(-i\beta \sigma_z^1 \sigma_z^2) \). We chose \( \beta = 0.1 \), and \( \beta = 0.4 \) (the previous one applied consecutively four times). (iii) \( \sigma_z^1 \)-controlled-\text{NOT} (CNOT) gate between qubits 1 and 2: CNOT \( = 0.5(I + \sigma_z^{(1)} + \sigma_z^{(2)} - \sigma_z^{(1)} \sigma_z^{(2)}) \). Also, this gate applied twice: CNOT\( ^2 = I \). These gates are more complex than one-qubit operations (which are typically less than 1 ms long) and they all involve refocusing idle times (periods of free evolution under the internal Hamiltonian) in their pulse sequences.

The results on the measurement of the collective coefficients for the qubit pairs (1,2), (2,3), and (1,4) are presented in Table I. The pair (1,2) is the one targeted by the \( C_{12} \) and CNOT gates while the other two are the pairs involving qubits 1 or 2 with the highest \( J \)-coupling \cite{17}. We expect the errors for the three chosen pairs to be larger (due to internal evolution that is not perfectly refocused). See Appendix C for more details on the experiment and simulations using liquid state NMR QIP.

| Gate   | \( \langle |w|_{i,j}^{\text{col}} \rangle \) | \( \langle |w|_{j,3}^{\text{col}} \rangle \) | \( \langle |w|_{j,4}^{\text{col}} \rangle \) |
|--------|---------------------------------|---------------------------------|---------------------------------|
| \( I_2 \) | meas 0.02                      | 0.02                            | 0.01                            |
|        | theo 0.00                       | 0.00                            | 0.00                            |
| \( F_g = 0.96 \) | simG 0.01                       | 0.02                            | 0.00                            |
|        | simE 0.01                       | 0.03                            | 0.00                            |
| \( F_g = 0.99 \) | simG 0.02                       | 0.00                            | 0.00                            |
|        | simE 0.02                       | 0.00                            | 0.00                            |
| \( C_{12}(0.1) \) | meas 0.26                       | 0.03                            | 0.03                            |
| \( C_{12}(0.4) \) | meas 0.23                       | 0.01                            | 0.00                            |
|        | simG 0.24                       | 0.02                            | 0.00                            |
| \( CNOT \) | meas 0.32                       | 0.01                            | 0.03                            |
| \( CNOT^2 \) | meas 0.07                       | 0.05                            | 0.04                            |
| \( CNOT^4 \) | meas 0.01                       | 0.01                            | 0.00                            |
|        | simG 0.01                       | 0.01                            | 0.00                            |
|        | simE 0.01                       | 0.02                            | 0.00                            |

The results \( \langle |w|_{i,j}^{\text{col}} \rangle \) for the various pairs of qubits shown in Table I exhibit good agreement with the predicted ones. Notice that the largest differences between measured and predicted appear on the pair (1,2), and on the most complex gates: \( CNOT^2 \), CNOT, and \( C_{12}(0.4) \). This indicates that these deviations are due to the errors expected from spurious processes in our QIP, particularly an imperfect refocusing during the gate sequence rather than from an imperfect implementation of the protocol.
The simulations account for well-known sources of error in liquid state NMR QIP (imperfect pulse design [17(c)], rf field inhomogeneities [2], and the presence of the magnetically active hydrogens in crotonic acid [17]). It is worth mentioning that the main errors occurring in one-qubit gates are correlated one-body errors (this is, a one-qubit rotation that is slightly off), and they do not introduce two-body errors, which are the main target of a spatial correlation analysis. This can be noticed also in the fact that the sin$G$ values are similar to the sin$E$ values (a change of 0.00–0.01, except in two cases where we found 0.02 and 0.03). Moreover, there is a contribution arising from $T_2$ relaxation. However, a calculation over the theoretical propagator and the numerics over the simulated one show a change on the order of 0.01.

Elements outside our model of the system, which would explain further the gap between theory and experiment, are $B_0$ (static) field inhomogeneities, the presence of transients and residual nonlinearities in the spectrometer circuitry, and an imperfect spectral fitting of the measured signal. These are well-known issues in liquid state NMR QIP, whose effect falls within the 1%–2% error.

We must differentiate between the implementation errors in the protocol (initial state preparation, one-qubit twirling, and readout), and the errors in the gate under study. The former ones affect the accuracy of protocol, which accounts for the measured non-null coefficients that are expected to be zero. As discussed, rf field inhomogeneities, the presence of hydrogen, and $T_2$ relaxation already give an error bar $\sigma_{x} \approx 0.03$. There are still other sources of error mentioned above that could make $\sigma_{x}$ larger, but within that order.

Moreover, a fiducial initial state preparation is critical to the success of the algorithm. We can quantify this as follows. If we call $\epsilon_i$ the error in the initial state preparation, and similarly we call $\epsilon_1$ the error in the implementation of the Clifford gates, an error propagation in the formula for the $\gamma$'s gives $\sigma_\gamma^2 \leq \frac{1}{4}(1+4\gamma)+\epsilon_1^2$, and then it is simple to propagate this into the formulas for the $\langle \eta_l \rangle$: for example, for a pair of qubits $a$ and $b$, we follow Eq. (9) and obtain $\sigma_{\eta}^2 = \frac{1}{3} \sqrt{\sigma_{\eta_a}^2 + \sigma_{\eta_b}^2 + \sigma_{\eta_{ab}}^2}$. These $\epsilon$'s account for nonstatistical errors (typically correlated one-body errors) and set the accuracy of the method. Given the low complexity of initial state preparation and one-qubit operators, these $\epsilon$'s are smaller than the errors in target operations (a fact reflected, for example, in the gate fidelities). This is why even though the theory was developed for error-free initial state preparation and twirl, the actual implementation can still retrieve information about the target operations.

V. DISCUSSION ON THIS AND OTHER PROPOSALS

Our protocol belongs to the family of characterization strategies based on the use of twirling to coarse-grain the original $O(2^n)$ complex parameters to a poly$(n)$ number of parameters (cf. the work by Emerson and co-workers in [6,8]). In our case, we gain detail about the process under study (spatial correlations). Note also that the required resources in our proposal are within the minimal performance expected from a functional QIP: fiducial state preparation and readout, and a set of $6n$ one-qubit gates.

One of the protocols presented by Bendersky et al. in [10] also returns similar information (although our method performs a coarse graining of the directions). Nevertheless, their proposal requires more demanding resources [although still scaling as poly$(n)$]: a full twirl on $U(D)$ and the implementation of the $O_l$ operators.

Other characterization methods include the ancilla-assisted ones in [10,11] but unfortunately they have a rather strong requirement: one or more clean error-free qubits within the system. Contrasting with all the proposals discussed so far, the method developed by Knill et al. in [9] does not require error-free stages, allowing for certain type of errors to occur during the whole computation. Unfortunately it is not yet evident how to take their scheme beyond one-qubit QIP [12].

On a different note, we would like to point out a particular feature of the variables $\langle \eta_l \rangle$, which are the diagonal elements of the so-called $\chi$ matrix in the $O_l$ basis (cf. the $a_i$ in [8], or the $x_{\text{sm}}$ in [10]). When the error $E$ acts on a short time we can expect the coefficients to be small and Eq. (2) can be taken instead as a first-order Taylor expansion of $E$, where the $\eta$ with $l \geq 1$ play the role of a generator of the error. This idea was originally developed in [7] under the same setting but aiming to study the generators directly led to limitations in the error model. Nevertheless, drawing the connection between the two opens the possibility of identifying a generator, which is essentially a Hamiltonian with varying parameters $\eta$ whose dynamics we can only observe on average through the $\langle \eta_l \rangle$. This interpretation allows a different insight into the dynamics of the system, as many physical processes are better described by the action of a stochastic Hamiltonian rather than by an operator-sum representation arising from a system+bath picture.

In conclusion, we have presented a method to characterize the spatial correlations occurring in a gate or process under study, showing its potential through a liquid state NMR QIP experiment. Second, we have pointed out the need of experimental feedback in order to arrive to a not only scalable but also feasible protocol, as the one we introduced. Finally, we have analyzed the relevance of implementation errors, showing the need for strategies that are not only scalable but also robust.

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APPENDIX A: ON THE CLIFFORD TWIRL

Consider the twirling of the map $S$ for $m$ qubits in the Hilbert space $\mathcal{H}_m$, of dimension $2^m$. As shown in [15], a Haar twirl is equivalent to Clifford twirl as follows

$$\rho_1 = \int_{U(M)} dUU^\dagger S(U\rho_0U^\dagger)U$$  \hspace{1cm} (A1)
\[
\gamma^{(a)} = A_a \left( \langle |\eta^a_1|^2 \rangle + \sum_{j \neq a} \langle |\eta^a_{j1}|^2 \rangle + \sum_{k > j \neq a} \langle |\eta^a_{j,k1}|^2 \rangle + \cdots \right) \quad \text{and similarly} \quad \gamma^{(b)},
\]

\[
\gamma^{(a,b)} = \frac{8P_aP_b + 2(P_a + P_b) - 4P_aP_b}{9} \left( \langle |\eta^{col}_a|^2 \rangle + \sum_{j \neq a} \langle |\eta^{col}_{j,a}|^2 \rangle + \cdots \right) + A_a P_b \left( \langle |\eta^{col}_a|^2 \rangle + \sum_{j \neq a} \langle |\eta^{col}_{j,a}|^2 \rangle + \sum_{k > j \neq a} \langle |\eta^{col}_{j,k,a}|^2 \rangle + \cdots \right),
\]

where \ldots denote the corresponding higher order multi-body terms. If we prepare the qubits in a pure state so \(P_a = P_b = 1\), the combination

\[
\gamma^{(a)} + \gamma^{(b)} - \gamma^{(a,b)} = \frac{4}{9} \left( \langle |\eta^{col}_{a,b}|^2 \rangle + \sum_{j \neq a,b} \langle |\eta^{col}_{j,a,b}|^2 \rangle + \cdots \right)
\]

leaves only the collective coefficients involving both the qubits \(a\) and \(b\).

**APPENDIX C: DETAILS ON THE EXPERIMENTAL IMPLEMENTATION**

The internal Hamiltonian of the system in the rotating frame is given by

\[
H_{\text{int}} = \hbar \sum_{j=1}^{4} \frac{\omega_{\delta,j}}{2} \sigma^{(j)}_z + \hbar \sum_{k > j=1}^{4} \pi J_{\delta,k} \sigma^{(j)}_+ \sigma^{(k)}_-, \quad \text{(C1)}
\]

where the chemical shifts, at our 9.4 T spectrometer, are of the order of kHz: \(\omega_{\delta,1} = 6650.6\) Hz, \(\omega_{\delta,2} = 1695.8\) Hz, and \(\omega_{\delta,3} = 4210.0\) Hz. The \(J\)-couplings are \(J_{12} = 72.6\) Hz, \(J_{13} = 69.8\) Hz, \(J_{14} = 7.1\) Hz while \(J_{24} = 1.6\) Hz, \(J_{13} = 1.3\) Hz, and \(J_{14} = 41.6\) Hz (according to the characterization of the sample we used; see also [17]).

The experimental initial state preparation over the four qubits reported a correlation with the theoretical one that was on average 0.99 (0.98 the lowest). The correlation for the targeted qubits (a pair of qubits) was, in each case, between 1.00–0.99 (the pseudopure state preparation was designed to optimize this correlation).
We implemented the twirl of pairs of qubits exactly using 36 Clifford operators. We had to pick one of the eight available six-element subsets of Clifford operators for each qubit, and we chose the subset that performed best experimentally: we applied each candidate to the thermal equilibrium state and compared the experimental performance with the theoretical one. This criterion coincided with choosing the subset that best took the equilibrium state to the I/2 state for the qubit being twirled.

To perform QST on four qubits with a liquid state NMR QIP, we used a set of 18 readout pulses. Therefore the number of experiments required to measure one collective coefficient $|\eta_{q,ij}^{\text{col}}|^2$ for a given pair of targeted qubits $(a,b)$ for a particular gate under study was 648, plus 18 experiments to characterize the initial state $|00\rangle(00)(I/2)^{\otimes 2}$ corresponding to preparing that pair in a pseudopure state. We performed QST of the full system therefore having a broader knowledge of the experimental performance but this is not required by the protocol: only the target qubits must be measured.

The negligibility of higher order multibody terms in the gates under study is to be expected in liquid state NMR QIP. In a simple model, these gates consist basically of periods of free evolution of length $\tau$ [the corresponding propagator is $U_\tau = \exp[-i H_{\text{int}} \tau / \hbar]$ separated by $\pi$-pulses on some of the qubits (so $U_\tau = \exp[-i(\pi/2 + \epsilon) \sigma^z_j]$, already accounting for some error $\epsilon$). For example, the sequence for the gate $I_E$ is

$$U = \frac{1}{\sqrt{2}} \begin{pmatrix} 1 & 0 & 0 & 1 \\ 0 & 1 & 1 & 0 \\ 0 & 1 & 1 & 0 \\ 1 & 0 & 0 & 1 \end{pmatrix},$$

where $\tau$ denotes free evolution for a time $\tau$, and $\pi^x_j$ denotes a $\pi$ pulse (180° rotation) around the $p$ axis for the qubits $j$.

Using the Baker-Campbell-Hausdorff (BCH) formula [19] is straightforward to see that in the building block $U_i U_{\pi^x_j}$ three-body- and higher order terms will appear with a factor at least $\frac{\pi^2}{2}$ smaller with respect to any possible one-body and two-body terms. For the values of $J_{i,k} \tau$ involved in our experiment, we have $\frac{J_{i,k} \tau}{\pi^2} \leq 0.14$. This means that any possible three-body and four-body terms would appear with a coefficient ten times smaller than the ones for one-body and two-body terms.

On the other hand, the simulation of the engineered pulse sequences used in the experiment showed that all the three-body and four-body terms appear with collective coefficients $|\eta_{i,k,\tau}^{\text{coll}}, 3,4 − |^2 < 0.005$ for the CNOT and $C_{12}(0.4)$ gates, and $<0.002$ for the rest. These are much smaller than the differences between measured and predicted values of $|\eta_{q,ij}^{\text{col}}|^2$, which can be better explained as implementation errors in the protocol or genuine gate errors arising from imperfect refocusing.

[14] The Pauli weight (also Hamming weight) of a given $O_j$ is the number of factors in it that are not the identity $I$. In a quantum information processor, an operator $O_j$ with Pauli weight $w$ can cause error in up to $w$ of the $n$ qubits of the register.