Quantum process tomography of the quantum Fourier transform

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The results of quantum process tomography on a three-qubit nuclear magnetic resonance quantum information processor are presented and shown to be consistent with a detailed model of the system-plus-apparatus used for the experiments. The quantum operation studied was the quantum Fourier transform, which is important in several quantum algorithms and poses a rigorous test for the precision of our recently developed strongly modulating control fields. The results were analyzed in an attempt to decompose the implementation errors into coherent (overall systematic), incoherent (microscopically deterministic), and decoherent (microscopically random) components. This analysis yielded a superoperator consisting of a unitary part that was strongly correlated with the theoretically expected unitary superoperator of the quantum Fourier transform, an overall attenuation consistent with decoherence, and a residual portion that was not completely positive—although complete positivity is required for any quantum operation. By comparison with the results of computer simulations, the lack of complete positivity was shown to be largely a consequence of the incoherent errors which occurred over the full quantum process tomography procedure. These simulations further showed that coherent, incoherent, and decoherent errors can often be identified by their distinctive effects on the spectrum of the overall superoperator. The gate fidelity of the experimentally determined superoperator was 0.64, while the correlation coefficient between experimentally determined superoperator and the simulated superoperator was 0.79; most of the discrepancies with the simulations could be explained by the cumulative effect of small errors in the single qubit gates. © 2004 American Institute of Physics. [DOI: 10.1063/1.1785151]

I. INTRODUCTION

In order to develop larger and more powerful quantum information processing devices, it is essential to quantify the precision with which they can be controlled. This information is generally reported as a single number, the fidelity of the operation. 1–3 Although fidelity is a reasonable measure of control, it gives experimentalists little useful information about what went wrong or how to improve their control over the quantum system. Quantum process tomography5–7 (QPT) provides additional information that may be useful in this regard, by yielding an estimate of the quantum operation that was actually implemented. This in turn provides a stringent check on the completeness and accuracy of the system-plus-apparatus model used to design the implementation. It is nevertheless a challenging task to interpret the deviations of this estimate from the superoperator implied by the model in terms of specific defects in the model. Additional errors introduced during the QPT procedure itself further complicate the analysis of the QPT results. In this paper we explore these issues via a concrete case study, in which QPT is performed on a previously reported three-qubit quantum Fourier transform (QFT) implemented using a nuclear magnetic resonance (NMR) quantum information processor.8

The dynamics of an isolated quantum system are described by the Schrödinger equation, which gives rise to an $N \times N$ unitary operator, where $N$ is the dimension of system’s Hilbert space. Open quantum systems, however, generally interact with an inaccessible environment and thereby undergo decoherence.3 Furthermore, in the case of expectation value measurements as in liquid-state NMR, each element of the statistical ensemble may undergo a slightly different unitary operation which, though nonrandom, is difficult to distinguish from decoherence.9,10 The statistics of measurements on open quantum systems are generally described by an $N \times N$ density operator, and the evolution of a density operator under an incoherent distribution of Hamiltonians and/or interactions with an environment, although nonuni-
tary, remains linear and is described by a superoperator. The goal of quantum process tomography is to determine this superoperator.

Methods for implementing QPT have been presented in Refs. 5–7, 12, and 13, and two-qubit NMR implementations of QPT have previously been reported.14,15 In the present paper, QPT is carried out with the aim of validating the mathematical model of the system-plus-apparatus used to design the NMR implementation of a multiqubit, entangling unitary operation, and to identify the types and strengths of the errors that occurred. We have found it useful to classify the errors as coherent, incoherent, and decoherent, because each class is related to specific shortcomings in the experimental implementation. Coherent errors are systematic errors in the net unitary operation that was actually implemented. Incoherent errors refer to unwanted unitary evolution which is not uniform across the ensemble of spin systems in the NMR sample so that even though each member of the ensemble undergoes strictly unitary evolution, the evolution of the ensemble averages appears nonunitary. The effects of incoherence are reversible, at least in principle, and knowledge of the coherent and incoherent errors can be used to design better quantum gates (unitary operations). In contrast, decoherent errors are due to unknown interactions with an inaccessible environment, so they are not reversible and can be eliminated only by relatively costly changes to the apparatus or the way in which information is encoded within it.

An important benefit of QPT is that it poses a rigorous test of the accuracy of the mathematical model of the system-plus-apparatus used to design and interpret the experiments. This is done by comparing the experimental results of QPT to the results of computer simulations of the complete QPT procedure, based upon the same model. Simulations based on the model used here were also used in designing the strongly modulating control fields by which the desired unitary operation as well as all the unitary operations needed for the QPT procedure were implemented.10 This was done by minimizing the difference between the desired qubit rotation operator and the quantum operation obtained by simulating the effect of the control fields on the spins in the molecule used for the experiments. As a result, any incorrectness in the model directly affects the reliability of the experiments, but in ways that, by definition, differ from the simulations. This can suggest ways to improve the model, after which further simulations will pinpoint the remaining experimental errors.

The operation on which QPT was performed is the QFT. The QFT constitutes a key subroutine in several quantum algorithms6,17 as well as in certain methods for simulating quantum dynamics on a quantum computer. In algorithms such as Shor’s factoring algorithm, the QFT is used to extract periodic features of wave functions, while in simulations of quantum dynamics it is used to move between the position and momentum representations. The QFT is defined as follows:

\[
U_{\text{QFT}}|x\rangle = \frac{1}{\sqrt{N}} \sum_{x'} e^{2\pi i xx'/N} |x'\rangle.
\]

The QFT has been expressed20,21 as a sequence of one-qubit

![FIG. 1. Circuit diagram for implementation of the quantum Fourier transform for three qubits. For each qubit \(j\), starting with the most significant, a series of conditional phase gates are implemented between qubit \(j\) and all qubits more significant than \(j\), followed by a Hadamard (H) on \(j\). The amount of phase added is \(\theta^j = \pi 2^{-j}\). A bit reversal (two-headed arrow) completes the QFT.](image)

Hadamard gates \(H^j\), which rotate the \(j\)th qubit from a computational basis state to an equal superposition of computational basis states, and two-qubit conditional phase gates \(B^{jk}(\theta)\), which rotate the phase of qubit \(k\) by \(\theta\) if qubit \(j\) is in the state \(|1\rangle\). In this notation, the complete gate sequence of the three-qubit QFT is (reading from right to left)

\[
\text{Swap}^{13} \times B^{23} \times \left( \frac{\pi}{2} H \right) \times B^{13} \times \left( \frac{\pi}{4} H \right) \times B^{12} \times \left( \frac{\pi}{2} H \right).
\]

where \(\text{Swap}^{jk}\) is a swap gate between qubits \(j\) and \(k\) (see Fig. 1). This gate sequence has been implemented via NMR to demonstrate the ability of the QFT to extract periodicity,8 and as part of the quantum baker’s map.22

The remainder of this paper is organized as follows. Section II gives an overview of the experimental and computational procedures used for QPT, together with the metrics by which the results were compared to those of the theoretical model. Section II describes incoherent errors and their effects on QPT, in particular the apparent lack of complete positivity of the results, while Sec. III describes the experimental system and procedures used to implement QPT in detail. Section IV presents a complete description of the system-plus-apparatus model by which the results of QPT were interpreted. This is followed in Sec. V by an overview of the experimental results, and in Sec. VI by a detailed comparison of the results with the model’s predictions. Finally, Sec. VII contains an analysis of the discrepancies between the experimental and simulated results, with the goal of determining their probable origins. The paper concludes with brief discussion of the implications of our findings for the field of quantum information processing as a whole.

II. QUANTUM PROCESS TOMOGRAPHY

There are several methods of performing QPT outlined in the literature. Some of these methods7,12 require increasing the Hilbert space size beyond that of the system whose dynamics are to be studied. This is unappealing for current experimental studies of quantum information processing where qubits are at a premium. The procedure used in this work (see Fig. 2) is similar to those given in Refs. 5 and 6 and results in an \(N^2 \times N^2\) complex-valued matrix, hereafter referred to as a “supermatrix.” Because of the inevitable experimental errors made in the QPT procedure, this supermatrix will not, in general, correspond to any completely

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be calculated as
to be linearly independent, the experimental supermatrix can
determined, and given that the input states have been chosen
experimental input and output states have been completely
output states via quantum
input states, followed by determination of all of the resulting
the desired unitary operation on a complete set of
errors committed during the readout steps.

completely positive part of the superoperator is largely due
better estimate of the superoperator that was actually imple-
to such a quantum operation. This is expected to yield a
recently introduced by Havel\textsuperscript{13,15} to obtain the best least-
positive and trace-preserving superoperator
\begin{equation}
M_{\text{obs}} = \mathcal{R}_{\text{out}}^{-1} \mathcal{R}_{\text{in}},
\end{equation}

\begin{equation}
C(\rho_{\text{th}}, \rho_{\text{op}}) = \frac{\text{tr}(\rho_{\text{op}}\rho_{\text{th}})}{\sqrt{\text{tr}(\rho_{\text{th}}^2)\text{tr}(\rho_{\text{op}}^2)}},
\end{equation}

where \( \rho_{\text{th}} \) and \( \rho_{\text{op}} \) are the traceless parts of the density matrices of the desired and measured input states, respectively. The input states were prepared from the equilibrium spin state by means of suitable nonunitary operations \( S_{\text{in}} \) and the magnitudes of their traceless parts taken as the reference against which all subsequent losses of coherence (or magnetization in NMR) due to nonunitary evolution were measured. This was done by scaling the correlation of the output states by a factor which measures the loss of coherence,\textsuperscript{3} yielding the so-called attenuated correlation

\begin{equation}
\tilde{C}_A(S_{\text{th}}, S_{\text{op}}) = \frac{\text{tr}(S_{\text{th}}^\dagger S_{\text{op}})}{\sqrt{\text{tr}(S_{\text{th}}^\dagger S_{\text{th}})\text{tr}(S_{\text{op}}^\dagger S_{\text{op}})}},
\end{equation}

where \( \rho_{\text{th}} \) and \( \rho_{\text{op}} \) are the traceless parts of the density matrices of the theoretical and measured output states, respectively, and \( \rho_{\text{in}} \) that of the corresponding input state from above.

The average state correlation and average attenuated state correlation are reasonable measures of the overall fidelity of the implemented quantum operation, but a measure that is more clearly independent of the choice of initial states is the correlation between the superoperator matrices, i.e.,

\begin{equation}
C(S_{\text{th}}, S_{\text{op}}) = \frac{\text{tr}(S_{\text{th}}^\dagger S_{\text{op}})}{\sqrt{\text{tr}(S_{\text{th}}^\dagger S_{\text{th}})\text{tr}(S_{\text{op}}^\dagger S_{\text{op}})}},
\end{equation}

Since this quantity is not sensitive to the overall loss of magnetization, we will also use the attenuated correlation between supermatrices, namely,

\begin{equation}
\tilde{C}_A(S_{\text{th}}, S_{\text{op}}) = \frac{\text{tr}(S_{\text{th}}^\dagger S_{\text{op}})}{\sqrt{\text{tr}(S_{\text{th}}^\dagger S_{\text{th}})\text{tr}(S_{\text{op}}^\dagger S_{\text{op}})}},
\end{equation}

where \( F_c \) is the entanglement fidelity defined by Schumacher.\textsuperscript{1} Accordingly, we will refer to this quantity as the gate fidelity. It can be shown that this fidelity satisfies

\begin{equation}
\tilde{C}_A(\rho_{\text{th}}, \rho_{\text{op}}) \geq C_A(S_{\text{th}}, S_{\text{op}}),
\end{equation}

where the overbar indicates an average over any orthonormal operator basis, with equality if \( \text{tr}(\rho_{\text{in}}^2) = \text{tr}(\rho_{\text{th}}^2) \) for all the input states.\textsuperscript{3}

Our analysis utilizes two different representations of quantum operations, the supermatrix representation and the Kraus operator sum representation. The supermatrix \( S \) operates on the density matrix as

\[ m_{\text{op}} \equiv \text{tr}(U_{\text{ro}}(\rho_{\text{x}}) \sigma_-) \]

FIG. 2. A schematic of the QPT implementation used in this paper, together with two equations describing how the readout operations \( U_{\text{ro}} \) and measurements \( M_{\text{obs}} \) are performed on the ensemble of quantum systems described by the density matrix \( \rho \) [where \( \sigma_- = \frac{1}{\sqrt{2}} (\sigma_x - i \sigma_y) \)]. A set \( \{S_{\text{in}}\} \) of (not necessarily unitary) operations is performed, each in a separate experiment, on the equilibrium state \( \rho_{\text{eq}} \) of the spin system to create a complete set \( \{\rho_{\text{in}}\} \) of input states. Each input state is then determined by quantum state tomography, i.e., by repetition of the experiment with different readout pulses, \( \{U_{\text{ro}}\} \), appended to each repetition. The readout pulses rotate unobservable components of the density matrix into observable components, the mean values \( \{m_{\text{in}}\} \) of which are equal to the unobservables’ mean values before rotation. Measurement of these mean values thus allows for the reconstruction of the input state that was actually created. Next, the set of input states is recreated one state at a time, and the operation \( S_{\text{op}} \) of interest (in this paper the QFT) applied to each in turn. This gives a complete set of output states, \( \{\rho_{\text{op}}\} \). Once again, readout of each \( \rho_{\text{op}} \) requires that the experiment be repeated followed by different readout operations \( \{U_{\text{ro}}\} \). The mean values of the complete set of observables \( \{m_{\text{op}}\} \) that is measured allows reconstruction of the set of output states actually created. Finally, these estimates of \( \{\rho_{\text{op}}\} \), together with the earlier estimates of \( \{\rho_{\text{in}}\} \), are used to reconstruct an estimate of the operation \( S_{\text{op}} \) via Eq. (3). This reconstructed operation is not expected to be unitary due to decoherence during the application of \( S_{\text{op}} \) as well as errors in the state tomography procedure. Also, due to incoherent errors across the ensemble involved in NMR experiments, the supermatrix \( M_{\text{obs}} \) estimated via this QPT procedure may not correspond to a completely positive, trace-preserving superoperator.
col(ρ_fin) = S col(ρ_int),
\end{equation}
where ρ_int and ρ_fin are the initial and final density matrices, and the “col” operation stacks the columns of the density matrices on top of one another in left to right order. This results in an $N \times N$ matrix becoming a column vector of length $N^2$. We will primarily work in two different supermatrix bases, the computational (or Zeeman) basis and the product operator basis. In the computational basis the rows and columns of the $2^n \times 2^n$ density matrix are labeled by the binary expansion of their indices from $[0 \cdots 0]$ to $[1 \cdots 1]$. The product operator basis is related to the structure of the NMR Hamiltonian and the rows and columns of the supermatrix in this basis are labeled as $I_2 \otimes \cdots \otimes I_2$, $I_2 \otimes \cdots \otimes I_2 \otimes \sigma_x$, to $\sigma_z \otimes \cdots \otimes \sigma_z$, where “$\otimes$” denotes tensor multiplication, the $\sigma$’s are the standard Pauli matrices, $I_2$ is the $2 \times 2$ identity matrix, and each term has $n$ factors (see Ref. 23 for details).

The superoperator of a quantum operation can also be expressed as a Kraus operator sum.\cite{11} This sum involves a set of $N \times N$ matrices, $A_k$ (the Kraus operators) such that
\begin{equation}
\rho_{\text{fin}} = \sum_k A_k \rho_{\text{int}} A_k^\dagger,
\end{equation}
where the sum may require as many as $N^2$ terms. It is easily seen that a Kraus operator sum preserves the Hermiticity of $\rho_{\text{int}}$, and that it preserves the trace if and only if
\begin{equation}
\sum_k A_k^\dagger A_k = 1.
\end{equation}
If the operation is purely unitary there is only one Kraus operator, which is just the unitary operator in question. In general, however, there are an infinite number of equivalent Kraus operator representations of a given quantum operation. The least-squares fitting procedure described below yields a Kraus operator sum representation in which the Kraus operators are both mutually orthogonal, minimum in number, and sorted by the size of the contribution each makes to $\rho_{\text{fin}}$.

### III. INCOHERENT ERRORS

A superoperator is completely positive if it admits a Kraus operator sum representation. Although this condition holds for the physical processes usually studied in the quantum mechanics of open systems, including unitary operations and decoherence via weak interactions with a Markovian environment, it is not difficult to construct situations in which not only complete positivity, but even the distinction between a state and its transformations breaks down.\cite{24–27} Such situations can only arise when the initial state of the system is not pure, but can be represented by a probability distribution over an ensemble of pure states with density matrix $\rho_{\text{int}}$. Suppose the ensemble’s probability distribution depends on some classical parameter $c$ (usually space or time) and that $\rho_{\text{int}}$ also represents an average over $c$ (see below for a concrete example). Then if the applied transformation $K$ likewise depends on $c$, so that the transformation is correlated with the states in the ensemble, then the final density matrix $\rho_{\text{fin}}$ will generally not be equal to the result of applying the average of $K$ over $c$ to $\rho_{\text{int}}$. Indeed there will usually be no superoperator which maps every possible $\rho_{\text{int}}$ to the correct $\rho_{\text{fin}}$. In principle one could define a nonlinear transformation that produces this mapping, but this is not very useful in practice because the large amount of information needed to define a general nonlinear transformation is neither readily available nor easy to work with.

Incoherent errors are precisely those which depend upon some classical parameter that labels the members of an ensemble, and so generate a correlation between the states in the ensemble and this parameter. In the NMR experiments described below, the main source of incoherent errors is the spatial inhomogeneity in the rf (radio-frequency) field over the sample volume. This may be expressed by writing the Hamiltonian for the interaction of the spins with the rf field in the form
\begin{equation}
H_{\text{rf}}(t; \vec{r}) = \alpha(t) \gamma B_1(\vec{r}) e^{-i\Delta \omega t} \sigma_z e^{i\Delta \omega t},
\end{equation}
where $\alpha(t)$ is the envelope of the field, $\gamma$ is the gyromagnetic ratio of the spins, $B_1(\vec{r})$ is the magnitude of the field at the point $\vec{r}$, $\Delta \omega$ is the difference between the frequency of the rf field and the resonance frequency of the spins in the static magnetic field $B_0$, and the $\sigma$’s are the usual Pauli matrices. Thus, the unitary operation $U = \exp(\int_0^t dt H_{\text{rf}})$ depends upon the position $\vec{r}$ in the sample as well as on the time, i.e., $U = U(t; \vec{r})$, as does the resulting spin state $\rho(t; \vec{r}) = \rho(0; \vec{r}) U(t; \vec{r})^\dagger U(t; \vec{r})$. The equilibrium state of the spins in the static field $B_0$ is independent of $\vec{r}$ over the sample volume, so we may take $\rho(0; \vec{r}) = \rho(0)$ to be constant. The density operator that is observed is the integral over the sample volume $V$, i.e.,
\begin{equation}
\rho(t) = \int_V d\vec{r} \rho(t; \vec{r}) = \int_V d\vec{r} U(t; \vec{r}) \rho(0) U^\dagger(t; \vec{r}) = S(t) \rho(0),
\end{equation}
where $S(t)$ denotes the net superoperator of the actual quantum operation implemented. We can express this in matrix form as\cite{13}
\begin{equation}
\text{col}(S(t) \rho(0)) = \left[ \int_V d\vec{r} U(t; \vec{r}) \otimes U(t; \vec{r}) \right] \text{col}(\rho(0)),
\end{equation}
where $U$ is the complex conjugate of $U$ and the “col” operator maps $N \times N$ matrices to $(N^2)$-dimensional column vectors, as described above.

Describing the evolution of an ensemble by integrating over its spatial degrees of freedom works well for a single operation. When a second operation $S_2$ is applied following the first $S_1$, however, we must take the spatial correlations produced by the first into account in computing the correct overall superoperator. In terms of the foregoing equations, this may be expressed as
To go the other direction, i.e., to take a supermatrix $S$ and convert it into an equivalent Kraus operator sum, we first rearrange its elements into a Hermitian supermatrix called the Choi supermatrix,

$$T = \sum_{ij=0}^{N-1} \left( E_{ij} \otimes I_N \right) S(I_N \otimes E_{ij}),$$  \hspace{1cm} (16)$$

where $I_N$ is the $N \times N$ identity matrix and $E_{ij}$ is a matrix of zeros except for a 1 in the $(i,j)$th position. Then, if $T = \sum_{k} \lambda_k v_k^\dagger v_k$ is the spectral decomposition of $T$ and we assume that all the eigenvalues $\lambda_k \geq 0$, our Kraus operators $A_k$ may be shown to be

$$\text{col}(A_k) = \sqrt{\lambda_k} v_k \quad (k=1, \ldots, N).$$  \hspace{1cm} (17)$$

It follows from the above that a superoperator $S$ is completely positive if and only if its Choi supermatrix is positive semidefinite. In the event that an experimentally determined supermatrix $S_{\text{obs}}$ does not have a positive semidefinite Choi supermatrix, it has been shown$^{13}$ that the completely positive superoperator $S'$ closest to it in the least-squares sense may be obtained simply by setting the negative eigenvalues of its Choi supermatrix to zero to get a new matrix $T'$ and mapping it back to $S'$ via Eq. (16) with $S$ and $T$ swapped. This procedure, however, will result in a supermatrix $S'$ that does not preserve the trace of the density matrix. That condition can be reimposed by subtracting

$$\Delta T' = S_{\text{obs}} \text{col}(I_N) \text{col}(I_N) T' - I_{N \times N}$$  \hspace{1cm} (18)$$

from $T'$, with the result that $T'$ is no longer positive semidefinite. It has further been shown, however, that iterating on these two procedures generates a sequence of supermatrices which converges to the Choi supermatrix corresponding to the superoperator that is closest to $S$ and is both trace preserving and completely positive. This is described for the experimental results in the Sec. VIII.

In order to quantify the extent to which a supermatrix fails to be completely positive, we define the positivity as the ratio of the sum of eigenvalues of the Choi matrix corresponding to the supermatrix over the sum of positive eigenvalues of the said Choi matrix. For a completely positive superoperator the positivity is equal to 1, while the presence of negative Choi eigenvalues causes the positivity to be less than 1.

**IV. THE SPIN SYSTEM AND THE EXPERIMENTS**

The experiments were implemented on a three-qubit NMR quantum information processor.$^{28,29}$ The three qubits used were the three carbon spins in molecules of $^{13}$C-enriched alanine in an aqueous solution. The internal Hamiltonian of this system has the form

$$H_s = \pi \sum_{i=1}^{3} \nu_i \sigma_{i,z} + \frac{\pi}{2} \sum_{j \neq i=1}^{3} J_{i,j}^{ij} \sigma_{i}^{\dagger} \sigma_{j},$$  \hspace{1cm} (19)$$

where $\nu_i$ are the Larmor frequencies of the spins and $J_{i,j}^{ij}$ are the strengths of the couplings between them, both in frequency units. In our indexing scheme, a superscript label labels the carbonyl carbon of alanine, 2 the alpha carbon and 3 the methyl carbon. A separate time-dependent external

**FIG. 3**. Eigenvalues of simulated supermatrices in the complex plane. The main figure shows the location of the eigenvalues with respect to the unit circle, while the subplot shows the eigenvalue locations in greater detail. This simulation was done for alanine (see text), and includes both coherent errors due to imperfect pulses and incoherent errors due to the inhomogeneous rf power across the sample. The eigenvalues indicated by circles (○) are of the product $S_{\text{obs}} S_{\text{ini}}$ of the supermatrix describing an evolution $S_{\text{ini}}$ of a 90° pulse on spin 1 about the negative y axis times a second supermatrix $S_{\text{obs}}$ for a 180° rotation of spins 1 and 2 about the x axis. The eigenvalues indicated by crosses (×) are for the supermatrix $S_{\text{obs}}$ describing the net evolution after concatenating these pulses together in the simulator. Due to the presence of incoherence the eigenvalues of $S_{\text{obs}}$ are not the same. In particular, the presence of incoherence leads to an apparent decoherence, as evidenced by a reduction in the eigenvalue magnitudes. The average eigenvalue reduction is 1.2% for $S_{\text{ini}}$ and 0.9% for $S_{\text{obs}}$.**
Hamiltonian, shown in Eq. (11), must be added whenever a rf pulse is applied to rotate qubits. In the 7 T magnet used for the experiments, the resonant frequency of carbon-13 is 75.468 MHz. Frequency changes, also known as chemical shifts, among the spins introduce differences \( \nu_x^C - \nu_z^C = 9456.5 \text{ Hz}, \nu_y^C - \nu_z^C = 2594.3 \text{ Hz}, \) and \( \nu_x^C - \nu_y^C = 12050.8 \text{ Hz}. \) The coupling constants between the three spins are \( J_{12}^{C,C} = 54.2 \text{ Hz}, J_{23}^{C,C} = 35.1 \text{ Hz}, \) and \( J_{13}^{C,C} = -1.2 \text{ Hz}. \) In the absence of rf, the \( T_1 \) relaxation times of the three spins are all longer than 1.5 s, while the \( T_2 \) relaxation times are longer than 400 ms (see Table I for exact numbers).

To rotate qubits we used the strongly modulating pulses introduced in Refs. 3 and 10. The first generation of strongly modulating pulses was designed to perform the desired propagator while refocusing the spins’ known internal Hamiltonian;\(^3\) the second generation was designed to also compensate for rf field inhomogeneity.\(^10\) This was done by including the rf inhomogeneity profile in the simulations and, hence, in the target function minimized to design the compensated pulses. Pulses designed in this way performed nearly optimally over the range of rf powers experienced by the ensemble of spins. While the simulated peak fidelities were lower than with the uncompensated pulses, the actual performance of the compensated pulses in the spectrometer was greatly enhanced.

The method of implementing the QFT via NMR is the same as in our previous QFT implementations,\(^8,12\) with the added benefit of the rf pulses that compensated for rf inhomogeneity. The pulse sequences for the Hadamard and conditional phase gates are derived from an idempotent or correlation operator description of the propagators.\(^30\) The Hadamard gate pulse sequence is

\[
H' = \frac{\pi}{2} y - \left( \frac{\pi}{2} \right)_x.
\]

This pulse program reads as follows: apply a pulse that rotates spin \( j \) by 90° about the \( y \) axis, followed by a pulse that rotates \( j \) by 180° about the \( x \) axis. The \( B^{12} \) gate can be implemented using the coupling between qubits and the following pulse sequence:

\[
\left( \frac{\pi}{2} \right)_x - \left( \frac{\pi}{2} \right)_y - \left( \frac{\pi}{2} \right)_x - \left( \frac{\pi}{2} \right)_y - \left( \frac{\pi}{2} \right)_x - \left( \frac{\pi}{2} \right)_y - \left( \frac{\pi}{2} \right)_x - \left( \frac{\pi}{2} \right)_y
\]

where \( \phi \) is an arbitrary phase. The notation \( \theta^{jk}/2\pi j^{ik} \) represents a time interval during which \( \sigma_j^x \sigma_z^x \) evolution occurs while chemical shifts and all other qubit couplings are refocused, while the superscript \( j,k \) denotes a pulse which rotates only spins \( j \) and \( k \). The final three pulses in the above sequence perform a rotation around the \( z \) axis. In our previous work\(^8\) the bit reversal was implemented experimentally. Here the bit reversal is achieved by simply renaming the bits.

We stress that the various building blocks of the QFT implementation, i.e., the individual pulses and evolution periods, remain unchanged for all experiments. After creation of each initial state a fixed sequence is applied that is independent of the initial state.

The input states \( \{ \rho_{in} \} \) used were the 64 product operator states (neglecting the large but undetectable identity component). These are \( (\sigma_x^1 \sigma_x^2 \sigma_x^3) \) and states such as \( \sigma_x^1 \sigma^3_x \) and \( \sigma^3_x \sigma^3_x \) where the superscripts represent the spin indices, as above. These states were chosen because they are orthogonal and easy to create on a liquid-state NMR system. We order the states lexicographically starting with 0, followed by \( \sigma_x^1 \sigma_x^2 \sigma_x^3, \sigma_x^2 \sigma_x^3 \sigma_x^1, \) and so on until \( \sigma_x^1 \sigma_z^1 \sigma_z^3 = \sigma_z^1 \sigma_z^3 \sigma_z^1 \) (the complete ordering may be found in Fig. 8). The QFT gate sequence described above is applied to each of these 64 states to obtain the corresponding set of output states \( \{ \rho_{out} \} \).

In any given three-spin spectrum only 24 of the 64 product operators are observable. Thus, to completely reconstruct the state of the system, each experiment is repeated seven times with different readout pulses appended to the experiment. The real amplitudes of the peaks in the spectrum were then used as the coefficients of the product operators which were transformed into the corresponding observable product operators by the readout pulses. This procedure, known as state tomography,\(^3\) was done for all input and output states.

V. THE SYSTEM-PLUS-APPARATUS MODEL

Quantum process tomography serves experimentalists in two ways. First, it provides a thorough test of the model of
includes, in addition to the carbons, the four 

5

The larger system’s Hamiltonian has the form 

12

because it stands for “control.” This is much faster than diagonalizing the Hamiltonian with the four hydrogens included (16 times longer as opposed to \( \sim 16^3 \)).

During the free evolution periods between pulses, on the other hand, we applied additional (hard) \( \pi \) pulses to the carbons to refocus this unwanted phase evolution, in accord with the free-evolution Hamiltonian used for these (much less demanding) simulations,

\[
H_\text{S,f} = H_\text{s} + \pi \sum_{j=1}^{4} J_{H,C,j} \sigma_\text{H,i} \sigma_\text{C,j},
\]

where \( f \) stands for “free.” The frequency evolution of the hydrogens can be neglected because they are left along the \( z \) axis, thereby also avoiding population disturbances that would lead to nuclear Overhauser cross relaxation and thereby memory effects.

The search for strongly modulating control sequences was further simplified by assuming that the rf phase, amplitude, and frequency were piecewise constant, so the total unitary transformation was given by the product

\[
U_\text{tot} = \prod_{k=1}^{\text{max}} U_k = \prod_{k=1}^{\text{max}} T \exp \left[ -i \int_{t_{k-1}}^{t_k} dt H_{\text{rf,k}}(t) \right].
\]

Here, “\( T \)” denotes the usual time-ordering operator and the external rf Hamiltonian during the \( k \)th interval is given by

\[
H_{\text{rf,k}}(t) = \alpha_k \gamma_C B_1 \sum_{j=1}^{3} \left( e^{-i (\pi \nu_k + \phi_k) \sigma_{C,j}^{z}} e^{i (\pi \nu_k + \phi_k) \sigma_{C,j}^{z}} \right),
\]

where \( \alpha_k \) is the relative amplitude of the field during the \( k \)th interval [cf. Eq. (11)]. This assumption allows the unitaries \( U_k \) for each interval to be calculated exactly by transforming to an interaction frame in which \( H_{\text{rf,k}} \) becomes time independent and diagonalizing the net Hamiltonian in that frame.
The spectrometer generates the control fields by applying a time-dependent voltage to a tuned resonator. The limitations on both the control circuitry and the tuned resonator introduce time-dependent distortions of this modulation at the discontinuities between intervals. To account for this limitation, we monitor the field generated in the control coil (the antenna that interacts with the spins) and preweight the time dependent wave form to provide a close approximation to the desired shape. In addition, we use the measured modulation sequence in the simulator to follow the dependence of the propagator for small distortions.

To achieve usable sensitivity, NMR is carried out on an ensemble of spatially distributed spins. The control field thus varies over the spatial extent of the sample, which is termed rf field inhomogeneity. This variation could be reduced by using a larger coil, a smaller sample, or by selecting a particularly homogeneous region of the sample via a magnetic field gradient. All of these options, however, reduce the signal-to-noise ratio. Fortunately, the spatial variation of the rf field is constant over time, and can be measured very accurately. This allows us to perform the simulations for each member of a histogram of the variations in rf amplitude across the sample, and to combine the results as an incoherent sum just as was described for the variations in the hydrogen spin states above. A total of 33 values was included in this rf inhomogeneity histogram (see Fig. 4).

The procedure used to find a modulation sequence that correctly implements any desired unitary operation seeks to maximize the fidelity between the desired unitary superoperator $\hat{U}_{\text{th}} \otimes U_{\text{th}}$ and the simulated superoperator $S_{\text{op}}$. The latter was obtained as an incoherent or Kraus operator over the 33 rf field strengths in the experimentally measured rf inhomogeneity histogram, i.e.,

$$S_{\text{op}}(\rho) = \sum_{\ell=1}^{33} p_{\ell} U_{\ell} \rho U_{\ell}^\dagger = \sum_{\ell=1}^{33} A_{\ell} \rho A_{\ell}^\dagger.$$  \hspace{1cm} (27)

Thus, the gate fidelity as defined in Eq. (7) can be calculated directly from the Kraus operators $A_{m}$ as

$$F_{Q}(\hat{U}_{\text{th}} \otimes U_{\text{th}}) S_{\text{op}} = \frac{\text{tr}[ (\hat{U}_{\text{th}} \otimes U_{\text{th}})^\dagger S_{\text{op}} ]}{\text{tr}[ (\hat{U}_{\text{th}} \otimes U_{\text{th}})^\dagger (\hat{U}_{\text{th}} \otimes U_{\text{th}}) ]}$$

$$= \frac{1}{64} \sum_{m=1}^{33} \text{tr}[ (\hat{U}_{\text{th}} \otimes U_{\text{th}})^\dagger (A_{m} \otimes \bar{A}_{m}) ]$$

$$= \frac{1}{64} \sum_{m=1}^{33} \text{tr}[ \hat{U}_{\text{th}} A_{m} \otimes (U_{\text{th}}^\dagger A_{m}) ]$$

$$= \frac{1}{64} \sum_{m=1}^{33} |\text{tr}(U_{\text{th}}^\dagger A_{m})|^2.$$  \hspace{1cm} (28)

In simulations including the four hydrogen spins, the sum on the right-hand side of this formula must be increased by another factor of 16 in taking the partial trace over the hydrogen spins.

The relaxation superoperator for the carbons in alanine was measured in the absence of rf fields, and found to have 98.5% of its norm along the diagonal in the product operator basis. This means that the various product operator components decay monoeXponentially, without cross relaxation, so that it can be described by an $8 \times 8$ “Hadamard relaxation matrix.” This is shown pictorially in Fig. 5, while precise values for the various types of rates seen in the figure are given in Table I.

To be complete, the relaxation superoperator should also be measured as a function of the applied rf fields (and include memory effects due to nuclear Overhauser effects with the hydrogens). The QFT, however, efficiently mixes the states of the three carbon spins, so that on average over its implementation the decoherence is indistinguishable from a uniform, isotropic attenuation of all the product operator components. That is to say, QPT on the QFT is not able to

FIG. 4. The 33 point rf profile shows how much of the sample sees what fraction of the desired power (so 1.0 on the horizontal axis is the desired power). This profile was measured experimentally and used for the simulations.

FIG. 5. (Color) Measured Hadamard relaxation operator vs the product operator basis for the three carbons of alanine in the absence of rf fields, with rates color coded as indicated by the legend. The labels on the axes are a short hand for the product operators associated with x and y directions on the three carbons, with e.g., $X_1 X\rightarrow \sigma_x^1 \sigma_x$. The labels for each entry of the matrix are obtained by analogy with the product rules for the Pauli operators; for example, the matrix entry in the row labeled $1 X Y$ is $Y X Z \rightarrow \sigma_y^1 \sigma_y \sigma_z^3$. See Table I for the numerical values of the various entries.
FIG. 7. The bar graph on the left shows the amplitudes used QPT to learn about the coherent and incoherent errors processes operative in alanine, save for an average overall rate committed during implementation of a complex unitary transformation, the QFT. Therefore, this simple effective relaxation superoperator was assumed to accelerate the simulations of the overall QFT (no relaxation was assumed during the pulse design simulations, since they are not intended to correct for such effects).

VI. AN OVERVIEW OF THE EXPERIMENTAL RESULTS

Complete QPT was performed twice using different sets of strongly modulating rf control sequences to implement single spin rotations (Sec. IV). The first such iteration was done with the sequences described in Ref. 3, which refo-cussed the evolution under the alanine molecule’s internal spin Hamiltonian \( H_s \) in Eq. (19) during the sequence. The second iteration used control sequences that not only refo-cussed the internal Hamiltonian but also compensated for inhomogeneity in the rf field itself.\(^\text{10}\) Thus, the second set of control sequences was expected to produce much smaller incoherent errors than the first, while the coherent and decoherent errors were expected to be roughly the same for both. The improvement obtained with the rf-compensated set is illustrated in Fig. 6, which plots the correlations and attenuated correlations obtained for each of the 64 product operator basis states used as inputs for the two iterations against one another. Further evidence for substantial improvements with the rf-compensated control sequences may be obtained from the Kraus operator plots and statistics in Fig. 7 (see below for their interpretation). Having demonstrated this clear-cut improvement, all subsequent analyses will be given only for the results obtained with the compensated control pulses.

The correlation provides an estimate of the accuracy of the experimental QFT implementation, but without considering the loss of magnetization due to decoherence or incoherence. The average of the input state correlations over all 64 provide any details regarding the physical relaxation processes operative in alanine, save for an average overall rate of attenuation. Of course, QPT could be performed on a simpler gate, as in Ref. 15 where it was used to derive a relaxation superoperator for free precession, but the goal here is to use QPT to learn about the coherent and incoherent errors...
basis states was 0.96, while the average correlation following application of the QFT was 0.82. A minimization search, however, found pure states which, after operating on them with the experimental superoperator, had a correlation with the same pure state following the theoretical QFT as low as 0.45. The average attenuated correlation (or gate fidelity\(^3\)), which also takes into account the loss of magnetization, was 0.64, indicating that about 22% of the magnetization was lost over the ca. 30 ms needed to implement the QFT (see Sec. IV). These numbers are in-line with expectations based on other recent applications of the rf-compensated control sequences.\(^4\) About half of this magnetization loss was expected due to intrinsic decoherence, and since approximately the same amount of decoherence occurred during the input states’ readout as during the output states’, the remainder is probably due to residual uncompensated incoherence and/or imperfect decoupling of the protons from the carbons used as qubits. These issues will be discussed in more detail in later sections. A complete list of all the output state (attenuated) correlations may be found in Fig. 8 below.

Rather than looking at the action of the QFT on states, one can also look directly at the Kraus operator sum computed from the completely positive part of the experimental supermatrix \(M_{\text{obs}}\) (Sec. III). Each Kraus operator \(A_k\) has an associated amplitude, \(a_k = |A_k|/\sqrt{\lambda_k}\) (where \(\lambda_k\) is the \(k\)th eigenvalue of the associated Choi supermatrix), and since in a perfect implementation the desired unitary operator \(U_{\text{QFT}}\) would be the only Kraus operator with nonzero amplitude, one expects the Kraus operator with the largest amplitude to be at least fairly similar to \(U_{\text{QFT}}\). This is confirmed by the plots of the real part of the largest Kraus operator, shown in Fig. 7, which had an amplitude \(a_1 = 0.86\) and a correlation with the real part \(U_{\text{QFT}}\) of 0.95, implying a net coherent error of roughly 5%. The second largest Kraus operator, also shown, had an amplitude of \(a_2 = 0.34\), and was also rather close to unitary although uncorrelated with \(U_{\text{QFT}}\).

We expect it to be a rough approximation to the largest unitary operator in any sum of unitary transformations making up the incoherent error. Finally, there are another 32 essentially nonunitary Kraus operators in the completely positive part of the experimental QFT supermatrix with smaller amplitudes (Fig. 7).

The non–completely positive part of the QFT supermatrix, which is obtained from the eigenvectors associated with the negative eigenvalues of its Choi supermatrix, will also be of interest in what follows. The positivity of the QFT supermatrix, as defined at the end of Sec. III, was only 0.60, but the ratio of the smallest to the largest Choi supermatrix eigenvalues was \(\approx 0.075\), indicating that the negative eigenvalues were rather small in magnitude in comparison to the positive ones. The ratio with the second largest was \(\approx 0.48\), and the third had almost the same magnitude as the smallest.

These observations, together with the well-known sensitivity of the eigenvectors of nearly degenerate eigenvalues to small perturbations in the elements of the matrix from which they come, imply that rather little information about the errors made in the QFT implementation can be gleaned from the individual Kraus operators after the first two. Only the subspaces spanned by the eigenvectors associated with all of the smaller positive, or perhaps negative, eigenvalues are likely to be statistically significant.

**VII. COMPARISON WITH THE MODEL**

To assess the precision and completeness of our system-plus-apparatus model, the complete set of experiments involved in QPT on the QFT was simulated using the mathematical model of the system-plus-apparatus described in Sec. V. The compatibility of the simulated results with the experimental provides a rigorous test for the accuracy of the model, which takes all the known significant imperfections of the experimental apparatus into account. All of the unitary operations performed during these simulations were implemented using exactly the same strongly modulating control sequences that were used for the experiments, and complete state tomography was performed for each input and output state. It should be noted, however, that these states were reconstructed directly from the observables in the simulated density matrices, without further simulating the spectra and fitting them as required in experimental state tomography (which of course gives rise to additional errors in the actual experiments). Figure 8 shows the correlation and attenuated correlation of the initial and final states obtained from these simulations.

Figure 9 plots the sorted Kraus operator amplitudes obtained from the simulated supermatrix, along with those from the experimental supermatrix for comparison (cf. Fig. 7). The negative values plotted are actually the negative square roots of the corresponding Choi matrix eigenvalues, and are shown to illustrate that the experimental supermatrix was significantly farther from being completely positive than was the simulated (the positivity of the simulated supermatrix was 0.86, as opposed to 0.60 for the experimental). The most likely reason for this is the absence in the simulations of the additional errors expected from fitting the spectra to extract the product operator amplitudes for state tomography. Also shown once again is the real part of the theoretical QFT unitary matrix (cf. Fig. 7), along with the real parts of the matrices of the largest Kraus operators from the experimental and simulated supermatrices. Finally, the corresponding best unitary approximations to the largest Kraus operators, obtained by setting their singular values to unity, have their real parts shown in the bottom right-hand side of Fig. 9. It may be seen that there is a good correspondence between the largest Kraus operators as well as between their best unitary approximations, with a correlation between the simulation and the experiment of 0.90 in both cases. As noted previously, the smaller Kraus operators cannot be expected to correspond significantly to one another.

Since the Fourier transformation converts between the position and momentum bases, it has a very simple interpretation in phase space. Classically, a Fourier transform rotates phase space by 90°. In quantum phase space, the QFT superoperator has only one nonzero element equal to unity in each row and column, and so constitutes a permutation matrix.\(^{33}\) This is shown in Fig. 10, along with the corresponding plots for the simulated and experimental supermatrices. One can see immediately from these plots that there are errors in the QFT implementation. The phase space basis,
however, consists of operators that are neither Hermitian nor products of any underlying Hilbert space basis, and which are not related in any simple way to the physical operators used to implement the QFT. Thus, although the action of the QFT is very easy to understand in the phase space basis, it is very hard to interpret the discrepancies between the theoretical superoperator and observed supermatrix in terms of implementation errors in this basis. It is worth noting, nonetheless, that in the phase space basis the errors appear more white than Gaussian especially in the simulation.

Other bases, in which the discrepancies between the simulated and experimental supermatrices are also clearly manifested, include the computational basis or Zeeman basis, see Fig. 11 and the product operator basis (shown in Fig. 12), where the latter consists of all possible products of the Pauli matrices $\sigma_x$, $\sigma_y$, and $\sigma_z$ of different spins. (see Sec.

FIG. 8. (Color) The differences between the (attenuated) correlations of the output states and their mean values are displayed for each input state, which are indicated by the symbols on the left vertical axis with, e.g., $1X \rightarrow \sigma_x^1$. The simulated vs theoretical state correlations are in blue, and the experimental vs theoretical state correlations are in yellow, while the corresponding attenuated correlations are in cyan and brown, respectively. The corresponding mean values of the (attenuated) correlations were 0.89, 0.82, 0.83, and 0.63, respectively. The pair of dashed vertical lines about the center marks one standard deviation for the input state correlations (not shown), which are expected to be determined primarily by measurement errors so that any values within these bounds are certainly almost entirely due to noise. The larger deviations in the correlations are due to the propagation of coherent error (see text), while the yet larger deviations in the attenuated correlations include the effects of decoherence (modeled as a uniform attenuation in the simulations).
It may be observed that the theoretical superoperator contains several fixed points in the product operator basis, which provide us with another interesting metric for the precision of our implementation. Two of these fixed points are $s_x^1 s_z^3$ and $\frac{s_x^1 + s_z^3}{2}$. The correlations between these fixed points, before and after applying the experimental and simulated superoperators, are 0.91 and 0.93, respectively. Further information can be obtained from plots of the individual rows, which depict how much each input state contributes to a given output state, as shown in the relatively simple case of the $s_x^1 s_z^3$ fixed point in Fig. 13.

The greatest part of the deviation between the simulated and experimental results is expected to be due to the propagation of small coherent errors. Thus it is interesting to compare the simulated and experimental supermatrices after correcting these errors as completely as possible. This is most simply done by left-multiplying by the inverse of the superoperator obtained from the best unitary approximation to the largest Kraus operator, $D_1^\dagger \otimes U_1^\dagger$, followed by the theoretical unitary superoperator $\bar{U}_{\text{QFT}} \otimes U_{\text{QFT}}$, i.e.,

$$\mathcal{M}_{\text{cor,obs|sim}} = (\bar{U}_{\text{QFT}} \otimes U_{\text{QFT}})(\bar{U}_{1,\text{obs|sim}} \otimes U_{1,\text{obs|sim}})^\dagger \mathcal{M}_{\text{obs|sim}}.$$  \hspace{1cm} (29)

The resulting supermatrices in the phase space basis are displayed in Fig. 14, along with the exact QFT supermatrix in the same basis for comparison. It may immediately be seen that there has been a considerable improvement in the similarity of these matrices, as is further confirmed by a correlation between the corrected simulated and corrected experimental supermatrices of 0.94, between the corrected simulated and theoretical of 0.99, and between the corrected experimental and theoretical of 0.95. The correlations...

---

**FIG. 9.** The left-hand plot shows the real amplitudes of experimental QFT Kraus operators (dark gray) and those obtained from the QPT simulation based on the system-plus-apparatus model (light gray), where the negative values plotted are actually the negative square roots of one-eighth the absolute values of the corresponding Choi matrix eigenvalues. It may be seen that the experimental supermatrix deviated significantly more from being completely positive than the simulated, most likely to the absence of errors from fitting the spectra in the latter (see text). The dominant Kraus operator has an amplitude 0.86 for the experiment and 0.87 for the model. The right side of the figure compares the real part of the theoretical unitary QFT operator (top-middle) to the largest Kraus operator from the experiment (middle-left) and simulated (middle-right) supermatrices. Also shown are the real parts of the corresponding best unitary approximations to these largest Kraus operators (bottom line). The correlation between the two largest Kraus operators, and between the corresponding best unitary approximations, was 0.90 in both cases.

**FIG. 10.** (Color) The theoretical (left), simulated (middle), and experimental (right) supermatrices of the QFT in the phase space basis, wherein all their elements are necessarily real. Since the theoretical is simply a permutation matrix in this basis, the differences between it and the other two are easily seen, but the physical meanings of these differences are obscure. For example, the level of "background noise" is noticeably larger in the simulation than in the experiment, although both have the maximum entry of each row (column) in the same place as the theoretical.
between the simulated and experimental correction factors, i.e., the products of the unitary superoperators in the above equation, are, however, only 0.90, indicating that the cumulative effects of coherent errors that were not taken into account by the simulations over the course of the experiments were roughly 10%. It should further be noted that whereas the experimental supermatrix is significantly more strongly correlated with the theoretical than it is with the simulated, after they have been corrected their correlations and attenuated correlations with the theoretical are very nearly the same.

VIII. DISCUSSION

We have shown that our model of the system-plus-apparatus is able to predict many details of the experimental results (see Table II for all the correlations among the theoretical, simulated, and experimental supermatrices, together with the attenuated correlations to the theoretical). Specifically, simulations based on the Hadamard relaxation operator shown in Fig. 5 and Table I have allowed us to establish that the results contain no specific information on the decoherence rates and processes operative in our system, since these are averaged by the complex sequence of transformations that make up the QFT. Their net effect can therefore be modeled as a simple uniform attenuation of all the product operators in the density matrix other than the identity. This is explicitly demonstrated by the close correspondence between the supermatrix eigenvalues shown in Fig. 15, both from simulations using the Hadamard relaxation operator (red ‘‘*’’) and from simulations taking no account of relaxation save by scaling down the nonidentity components of the density operators by a factor of 0.82 (blue ‘‘○’’). In general, of course, relaxation cannot be accounted for by a single attenuation factor, and in fact methods similar to those described here have been used to determine the complete NMR relaxation superoperator of a two-spin system.15

Figure 15 also shows the eigenvalues of the supermatrix obtained from simulations which included incoherent errors from rf field inhomogeneity, first without taking relaxation or readout errors into account (cyan ‘‘●’’), and second from simulations which included input and output state readout and took relaxation into account by scaling the nonidentity components of the density matrices readout by 0.82 (green ‘‘×’’). These plots show rather clearly that most of the additional dispersion seen in the simulations including incoherent errors stems directly from those errors, and was not an unin-
FIG. 13. As may be seen from row 14 of the supermatrices shown in Fig. 12, the operator $\sigma_x^1\sigma_x^2$ is a fixed point of the QFT. The upper bar graph displays the corresponding row of the theoretical (dark gray), simulated (medium gray), and simulated with coherent errors corrected (light gray) supermatrix elements vs the product operator basis, whereas the lower bar graph displays these same statistics for the experimental supermatrix. In contrast to the phase space and Zeeman basis (not shown), there is a significant correlation between the experimental and simulated values; specifically, the simulated-to-experimental correlation coefficients were 0.80 and 0.94 before and after correction, respectively. In the present case these correlations are due almost entirely to the single large diagonal value, but similar correlations were obtained for all other pairs of rows; the average correlations were 0.78 and 0.94 before and after correction, respectively.

Another, probably more important, reason for these apparent discrepancies between the model and the experiments lies in the propagation of many small coherent errors over the course of the QFT implementation.

Although the simulated single-qubit gate fidelities were all better than 0.99, they were also less than unity. The main thing that limited these fidelities was that the number of parameters, and hence the number of time intervals within which the rf amplitude and phase was held constant, had to be kept as low as possible during the optimizations by which the strongly modulating pulses were designed. Although the cumulative effects of such small errors could become significant, since they are included in the simulations they are not likely to be the source of the additional eigenvalue dispersion that is seen in the experimental results. A more important source of coherent errors is expected to be due to the fact that the protons were not included in any of the simulations, and their couplings to the carbons used as qubits will show up as phase rotations. In addition, the model ignores such fine details of the apparatus as the limits on the rise-and-fall times of the transmitter used to generate the rf control sequences, and more generally its frequency response characteristics. Although the simulation of reactive circuits is more complex than the simulation of resistive circuits, in due course we will also include such effects in the model and continue to refine it until the only remaining discrepancies between the simulated and experimental results lie in the intrinsic measurement errors.

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

**FIG. 14.** (Color) Theoretical (left), simulated (middle), and experimental (right) supermatrices in the phase space basis after correcting the latter two by a unitary supermatrix designed to bring them as close as possible to the theoretical (as described in the text). The levels of background noise are a great deal less than in the corresponding uncorrected supermatrices (Fig. 10).

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

**TABLE II.** The correlation coefficients between all pairs of supermatrices, labeled as described in the main text, together with the attenuated correlations (as defined in Sec. II) between each and the theoretically exact supermatrix of the quantum Fourier transform.

<table>
<thead>
<tr>
<th>Supermatrix</th>
<th>Correlations among all pairs of supermatrices</th>
<th>Attenuated correlation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Theoretical</td>
<td>1.00 0.89 0.99 0.82 0.95 1.00</td>
<td></td>
</tr>
<tr>
<td>Simulated</td>
<td>⋮ 1.00 0.90 0.79 0.85 0.68</td>
<td></td>
</tr>
<tr>
<td>Corrected Sim.</td>
<td>⋮ 1.00 0.81 0.94 0.76</td>
<td></td>
</tr>
<tr>
<td>Experimental</td>
<td>⋮ ⋮ 1.00 0.87 0.64</td>
<td></td>
</tr>
<tr>
<td>Corrected Exp.</td>
<td>⋮ ⋮ ⋮ 1.00 0.74</td>
<td></td>
</tr>
</tbody>
</table>

*See text for complete descriptions of the terms in the first column.*
individual spins, using traditional NMR methods. Errors in $U$ primarily single-spin rotations. To test this hypothesis, the errors that were not taken into account by the simulations will be as a consequence, we expect that the residual coherent errors timing of the pulses, which can be controlled very accurately.

the complex plane, specifically, the results of simulations with only coherent errors in the blue "O"), the results including incoherent errors but no relaxation (cyan "●"), and the same following simulation of the full QFT procedure (green "×"), wherein decoherence was mimicked simply by scaling down the simulated QFT superoperators following the state readout pulses by a factor of 0.82 (save for the component with eigenvalue 1, which is needed for trace preservation). To aid the comparison, the eigenvalues of the QFT superoperator including only the coherent errors (blue "O") were scaled to have the same RMS value as those obtained with the simulated relaxation, thereby showing that the net effect of relaxation on the results of the QFT is simply to lower the amplitude of the eigenvalues without otherwise dispersing them in the complex plane. It may also be seen that the effect of incoherent errors on the eigenvalues is both to increase their angular spread and to scale them down proportionately, so that they tend to move along arcs inside of and tangent to the unit circle, and that the additional errors introduced by the tomography procedure do not alter them greatly.

Unwanted bilinear interactions are refocussed during the QFT implementation by the application of π rotations to the individual spins, using traditional NMR methods. Errors in these rotations will contribute to the bilinear errors only to second order, so these bilinear errors depend mainly upon the timing of the pulses, which can be controlled very accurately. As a consequence, we expect that the residual coherent errors that were not taken into account by the simulations will be primarily single-spin rotations. To test this hypothesis, the error operator $U_\Delta = U_{1,exp}^\dagger U_{1,sim}$ was taken, where $U_{1,exp}$ is the best unitary approximation to the largest Kraus operator $A_1$ of the experimental supermatrix (cf. Fig. 9), and $U_{1,sim}$ is similarly the best unitary approximation to the largest Kraus operator of the simulated supermatrix. Using a numerical search, the product of three single-spin rotations was found that fitted $U_\Delta$ best in the least-squares sense. The resulting unitary $U_1^1 \otimes U_2^3 \otimes U_3^3$ had a correlation coefficient with $U_\Delta$ of 0.96, in support of the hypothesis.

The angles and directional cosines of the axes of these three rotations are shown in Table III, while the eigenvalues of the resulting corrected superoperators are shown in Fig. 16. It should be noted that, although the correlation between $U_{1,exp}$ and $U_{1,sim}$ increased from 0.90 to 0.96 on left-multiplying $U_1$ by the Hermitian conjugate of this product of single-spin rotation operators, these rotations are the cumulative result of many small rotation errors and are not simply traced back to any single shortcoming in the experiments. For completeness, the axes and angles of the single-spin ro-

![FIG. 15. (Color) Plot of the eigenvalues of simulated QFT superoperators in the complex plane. Specifically, the results of simulations with only coherent errors (blue "O"), the results including relaxation using the relaxation superoperator shown in Fig. 5 and Table I (red "+"), the results including incoherent errors but no relaxation (cyan "●"), and the same following simulation of the full QFT procedure (green "×"), wherein decoherence was mimicked simply by scaling down the simulated QFT superoperators following the state readout pulses by a factor of 0.82 (save for the component with eigenvalue 1, which is needed for trace preservation). To aid the comparison, the eigenvalues of the QFT superoperator including only the coherent errors (blue "O") were scaled to have the same RMS value as those obtained with the simulated relaxation, thereby showing that the net effect of relaxation on the results of the QFT is simply to lower the amplitude of the eigenvalues without otherwise dispersing them in the complex plane. It may also be seen that the effect of incoherent errors on the eigenvalues is both to increase their angular spread and to scale them down proportionately, so that they tend to move along arcs inside of and tangent to the unit circle, and that the additional errors introduced by the tomography procedure do not alter them greatly.

![FIG. 16. (Color) The blue squares (□) are the eigenvalues of the simulated superoperator $M_{sim}$, including coherent, incoherent, and readout errors, while the magenta triangles (△) are those after correcting $M_{sim}$ by postmultiplication with the product of single-spin rotations $U_1^1 \otimes U_2^3 \otimes U_3^3$ and the orange diamonds after premultiplication with a product of different single-spin rotations (see Table III). Although we cannot quite unambiguously match up pairs of eigenvalues, it is clear that they have been made very similar by these corrections.

<table>
<thead>
<tr>
<th>Side</th>
<th>Spin</th>
<th>$x$, $y$, and $z$-directional cosines of rotation axis</th>
<th>Rotation angle (deg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Left</td>
<td>1</td>
<td>-0.992 0.007 0.123</td>
<td>36.7</td>
</tr>
<tr>
<td>Left</td>
<td>2</td>
<td>-0.386 -0.243 0.890</td>
<td>9.0</td>
</tr>
<tr>
<td>Left</td>
<td>3</td>
<td>0.703 0.701 -0.123</td>
<td>16.9</td>
</tr>
<tr>
<td>Right</td>
<td>1</td>
<td>0.059 -0.031 0.998</td>
<td>14.2</td>
</tr>
<tr>
<td>Right</td>
<td>2</td>
<td>0.227 -0.512 0.829</td>
<td>10.2</td>
</tr>
<tr>
<td>Right</td>
<td>3</td>
<td>-0.092 -0.292 -0.952</td>
<td>38.3</td>
</tr>
</tbody>
</table>
those of the experimental supermatrix $M_{\text{obs}}$, as well as the best unitary approximation to its largest Kraus operator $U_{1,\text{CPTP}} \otimes U_{1,\text{CPTP}}$ and $M_{\text{obs}}$ itself for comparison.

<table>
<thead>
<tr>
<th>Supermatrix</th>
<th>Correlation with $\tilde{U}<em>{1,\text{CPTP}} \otimes U</em>{1,\text{CPTP}}$</th>
<th>Correlation with $M_{\text{CPTP}}$</th>
<th>Correlation with $M_{\text{obs}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Theoretical</td>
<td>0.86</td>
<td>0.89</td>
<td>0.82</td>
</tr>
<tr>
<td>Simulated</td>
<td>0.82</td>
<td>0.82</td>
<td>0.79</td>
</tr>
<tr>
<td>Experimental</td>
<td>0.95</td>
<td>0.97</td>
<td>1.00</td>
</tr>
</tbody>
</table>

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TABLE IV. The correlation coefficients between theoretical, simulated, and experimental supermatrices (see text), and the optimum completely positive and trace-preserving approximation $M_{\text{CPTP}}$, to the experimentally determined supermatrix $M_{\text{obs}}$, as well as the best unitary approximation to its largest Kraus operator $U_{1,\text{CPTP}} \otimes U_{1,\text{CPTP}}$ and $M_{\text{obs}}$ itself for comparison.

...tations that best fit the error operator $U_{\Delta} = U_{1,\text{sim}} \otimes U_{1,\text{exp}}$ are also shown in Table III, together with the eigenvalues of the corresponding corrected superoperators in Fig. 16. The product of these rotations similarly has a 0.97 correlation with $U_{\Delta}$, but in this case one must right-multiply $U_{1,\text{exp}}$ by the Hermitian conjugate of the product to correct it. The close coincidence between the angles of rotation about the $x$ axis on spin 1 in the first case and about the $z$ axis on spin 3 in the latter case is expected, since $U_{QFT} \sigma_x^1 U_{QFT}^\dagger = \sigma_z^3$ (recall $\sigma_x^1 \sigma_z^3$ is a fixed point of the QFT).

Finally, it is of interest to demonstrate that despite a substantial number of negative eigenvalues in the Choi matrix of the experimental supermatrix, it is not necessary to change it much in order to obtain a supermatrix which represents a completely positive and trace-preserving superoperator. For this reason the supermatrix $M_{\text{CPTP}}$ which best fitted the Choi matrix of the experimental supermatrix subject to the constraint that it was both positive semidefinite and satisfied the trace-preservation conditions was computed as described in Sec. III. Although this procedure made essentially no change in the largest Kraus operator (as expected), it did have a significant effect on the experimental supermatrix as a whole. The correlations between this CPTP fit and the other supermatrices that we have dealt with up to now are given in Table IV, along with those of the original experimental supermatrix for comparison. This shows that even though imposing the complete positivity constraint on the experimental observations did not change the supermatrix very much, the change was distinctly in the right direction since it improved the correlation with both the simulated and theoretical supermatrices. This is further confirmed by Fig. 17, which shows the eigenvalues of $M_{\text{CPTP}}$, along with those of the superoperator $\tilde{U}_{1,\text{CPTP}} \otimes U_{1,\text{CPTP}}$ obtained from the best unitary approximation to its largest Kraus operator (scaled down so as to have the same trace as $M_{\text{CPTP}}$) and those of the experimental supermatrix $M_{\text{obs}}$ for comparison. From this we see that the eigenvalues of $M_{\text{CPTP}}$ are closer to being cocircular, indicating that it is closer to an attenuated unitary than was $M_{\text{obs}}$, but that only after taking the unitary part did they become perfectly cocircular.

IX. CONCLUSIONS

In conclusion, we have implemented quantum process tomography of the quantum Fourier transform on a three-qubit NMR quantum information processor. The overall gate fidelity (attenuated correlation between superoperator matrices) was 0.64, whereas the (unattenuated) superoperator correlation was 0.82 (see Table II). Judging by the fact that making the unitary part of the largest Kraus operator correspond as closely as possible to the theoretical QFT gave a correlation of 0.95, we conclude that the loss of fidelity due to incoherence and/or measurement errors during state tomography was of order 5%. The loss of magnetization due to incoherence and decoherence, on the other hand, was 0.64/0.82 = 0.78, i.e., about 22%. This implies that the cumulative effects of coherent errors reduced the fidelity by about 0.82/0.95 = 0.86, or 14%, consistent with the fact that the correlation between the largest Kraus operator and the theoretical QFT unitary was 0.925 = \sqrt{0.86}.

More importantly, QPT of the QFT has enabled us to validate the essential correctness of our model of the system-plus-apparatus used for the experiments in great detail and to isolate its remaining shortcomings. It has further prompted us to develop a range of data analysis and visualization techniques for quantum process tomography, which should be broadly applicable in quantum information processing. The experiments described here demonstrate the precision with which complex quantum dynamics can be controlled, and highlight the significance of liquid-state NMR as a test bed for achieving such control. While full QPT on larger quantum systems will never be practical, the analysis done here should serve as an initial guide as to how information about the errors in quantum information processors can be extracted and, perhaps someday, how to debug a quantum computer.

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