Experimental study of Forrelation in nuclear spins

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A B S T R A C T

Correlation functions are often employed to quantify the relationships among interdependent variables or sets of data. Recently, a new class of correlation functions, called Forrelation, has been introduced by Aaronson and Ambainis for studying the query complexity of quantum devices. It was found that there exists a quantum query algorithm solving 2-fold Forrelation problems with an exponential quantum speedup over all possible classical means, which represents essentially the largest possible separation between quantum and classical query complexities. Here we report an experimental study probing the 2-fold and 3-fold Forrelations encoded in nuclear spins. The major experimental challenge is to control the spin fluctuation to within a threshold value, which is achieved by developing a set of optimized GRAPE pulse sequences. Overall, our small-scale implementation indicates that the quantum query algorithm is capable of determining the values of Forrelations within an acceptable accuracy required for demonstrating quantum supremacy, given the current technology and in the presence of experimental noise.

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1. Introduction

With the ability of creating exponential number of superposition of states, quantum computation provides an unprecedented computational power over classical computation. For example, Shor’s factoring algorithm [1], the Harrow-Hassidim-Lloyd (HHL) algorithm [2], and other progresses in quantum simulation [3–5] provide strong evidences that quantum computation can gain exponential speed-up in practical problems. Apart from computational decision problems, quantum devices can be exploited for other classically-intractable computational tasks, including sampling distributions of some quantum systems [6–10]. As a result, one may expect to gain “quantum supremacy” [11] in relatively-simple quantum devices in the near future.

Although these results are promising, complete and rigorous proofs supporting claims of gaining quantum supremacy are still unavailable. Recalling that for the case of Shor’s algorithm, we have not excluded the possibility of the existence of a polynomial-time classical algorithm for the factoring problem. For the HHL algorithm, which is BQP-complete, it remains to be determined if quantum computation is indeed more powerful than classical computation, or technically, if it is true that BQP ⊃ BPP. Here BPP (bounded-error probabilistic polynomial time) is the class of decision problems solvable by a probabilistic Turing machine in polynomial time with an error probability of at most 1/3 for all instances and BQP (bounded-error quantum polynomial time) is the quantum analogue of the complexity of BPP in computational complexity theory. Furthermore, the success of the sampling algorithms is founded on several conjectures in the theory of classical computational complexity. Even though boson-sampling devices are capable of creating an exponentially large superposition of quantum states, the transition amplitudes can still be estimated by classical devices within additive errors [12].

On the other hand, query complexity, which counts the number of queries of black-box functions (i.e., without knowledge of the internal structure), provides further evidence supporting quantum speed-up over the classical counterparts. For example, Grover’s search algorithm [13], the Deutsch-Jozsa algorithm [14] and Simon’s algorithm [15] are all characterized in the context of query complexity.

Recently, Aaronson and Ambainis [16] introduced a new concept in query complexity, called Forrelation, which characterizes the multi-fold correlations among different boolean functions. It was found that a quantum computer is capable of solving 2-fold Forrelation problems within a constant O(1) number of queries.
However, classical computers require an exponential number of queries. The difference in the query complexity between quantum and classical methods is shown to be a maximally-achievable separation with quantum methods (see also Refs. [17–19]). Furthermore, multiple-fold FORRELATION problems are as hard as quantum computation [16], i.e., BQP-complete.

Here we report the first experimental study of the 2-fold and 3-fold FORRELATION in a system of nuclear spins, where the NMR quantum circuit for 2-fold FORRELATION involves only 2 queries of the black box functions, but classically, it takes a total of 8 queries for an exact result. Similarly, 3 queries are needed for the NMR implementation of 3-fold FORRELATION, while 12 queries are needed classically if memory is given for the black-box functions; otherwise, it can go up to 192 classical queries.

However, we note that the measurement results come directly from the NMR signals, but a standard implementation of the quantum circuit involves probabilistic measurement outcomes. Furthermore, similar to other experimental demonstrations of Deutsch-Jozsa algorithms [20,21], the applied NMR pulse sequence depends on the knowledge of the functions, which are not strictly “black boxes”. Therefore, the current experimental results cannot be taken as a direct proof for demonstrating quantum supremacy, which is relevant only in the large-N limit.

The purpose of the experiment is to investigate whether a small-size prototype experiment can produce FORRELATION within the accuracy required for demonstrating the quantum advantages (above the threshold 3/5 or below the threshold 1/100), given the current technology and in the presence of experimental noise. In particular, our experimental fluctuation for the spin measurement has to be controlled within 1%. These experimental results allow us to identify the places one can improve for scaling up the size of the experiment in future.

2. Forrelation

Given k Boolean functions, $f_1 = f_1(x_1), \ldots, f_k = f_k(x_k)$, each with n variables, i.e., $x_i \in \{0,1\}^n \rightarrow \{-1,1\}$, the k-fold FORRELATION, $\Phi_k = \Phi_{f_1, f_2, \ldots, f_k}$, of these functions is defined as follows:

$$\Phi_k = \sum_{x_1, x_2, \ldots, x_n} e^{i\langle x_1, x_2, \ldots, x_n \rangle \cdot f_1(x_1) f_2(x_2) \cdots f_k(x_k)}.$$  

where $e^{i\langle x_1, x_2, \ldots, x_n \rangle}$ is the bit-wise inner product between the n-dimensional binary vectors $x$ and $y$. The total number of possible assignment is $N = 2^n$. Essentially, 2-fold FORRELATION is simply the inner product between a boolean function and the Fourier transform of another boolean function, i.e.,

$$\Phi_{f,g} = \frac{1}{2\pi} \sum_{x,y} (-1)^{x \cdot y} f(x) e^{i xy}.$$  

Importantly, an exact determination of 2-fold FORRELATION $\Phi_{f,g}$ is a computationally-hard problem for classical devices, which can be justified by the following challenge [16]: given a pair of Boolean functions $f$ and $g$, suppose it is known that either (1) $\Phi_{f,g} \leq 1/100$ or (2) $\Phi_{f,g} > 3/5$ is true, all classical methods require an exponential number $\Omega(\sqrt{N}/\log N)$ of queries to the black-box functions, but quantum computers can finish the task with a constant number of queries. The separation between the quantum and classical query complexity is (almost) possibly largest one can achieve [16].

Quantum circuits for solving 2-fold and 3-fold FORRELATION problems [16] are shown in Fig. 1. For 2-fold FORRELATION problems, there are 2 query operators $O_{1}$ and $O_{2}$, which map each input basis state $|x\rangle$ to $f_1(x)|x\rangle$ and $f_2(x)|x\rangle$ respectively, i.e., $O_j |x\rangle = f_j(x)|x\rangle$.

3. Experimental background

Nuclear magnetic resonance (NMR) is a reliable technology for studying small-to-medium size quantum information experiments [22,23], and quantum simulation [24–28]. Motivated by the needs of studying quantum information, many sophisticated techniques of controlling nuclear spins have been developed.

Here all the experiments are carried out at room temperature (295 K) on a Bruker Avance III 400 MHz spectrometer and the $^{13}$C labelled Diethyl-fluoromalonate dissolved in d6 acetone is used as a 3-qubit NMR quantum information processor. The structure and Hamiltonian parameters of Diethyl-fluoromalonate are shown in Fig. 2a where $^{13}$C, $^1$H and $^{19}$F nuclear spins respectively act as an ancillary qubit and two work qubits. Moreover, the internal Hamiltonian of the system is given by

$$H_{int} = \sum_{i=1}^{3} \pi_i \sigma_i^z + \sum_{j=1}^{3} \pi_j \sigma_j^2.$$  

The whole experimental procedure consists of three parts: (1) state initialization, (2) realization of the quantum algorithm for solving 2 (or 3)-fold FORRELATION problem, and (3) readout of the expectation value of $\sigma_i^z$ of the ancillary qubit $^{13}$C, which is equal to the FORRELATION, i.e.,

$$\langle \sigma_i^z \rangle = \Phi_i$$  

for any $k \geq 2$. We note that for the NMR quantum computing, the whole system, starting from the thermal equilibrium state, can be converted to the pseudo-pure state (PPS) [29,30] $\rho_{PPS} = (1 - \nu)/8 + \nu|00\rangle \langle 00|$, using the spatial average technique [31]. To check the success of preparing the PPS, a full quantum state tomography (QST) [32] is carried out. The fidelity between the density matrix prepared in experiment ($\rho_{exp}$) and the target one in theory ($\rho_{th}$) is given by the following expression,

$$F(\rho_{exp}, \rho_{th}) = \text{tr}(\sqrt{\rho_{exp}} \rho_{th} \sqrt{\rho_{exp}})/\sqrt{\text{tr}(\rho_{exp}^2) \text{tr}(\rho_{th}^2)}.$$  

A spectrum of the PPS observed on $^{13}$C is shown in Fig. 3a. The real parts of the initial state are shown in the last figure as $\rho_0$. Overall, the initial state can be well prepared in our setup: the fidelity can reach up to 96.9%.

4. Experimental details

To solve the $k$-fold FORRELATION problem, a quantum circuit is designed to obtain FORRELATION $\Phi_k = \Phi_{f_1, \ldots, f_k}$ by measuring the
Fig. 2. (Color online) (a) Molecular structure and Hamiltonian parameters of Diethyl-fluoromalonate. The chemical shifts and scalar coupling constants of the molecule are on and below the diagonal (in Hz) in the table, respectively. (b) An example of pulse sequences for solving the 3-fold Forrelation problem. $^{13}$C acts as the probe qubit while $^{1}$H and $^{19}$F are the work qubits. The circuit is comprised of 52 $\pi/2$ ($\pi$) hard pulses and 31 free evolution periods under spin–spin $J$-couplings in total (refocusing pulses are omitted for clarity).

Fig. 3. (Color online) NMR spectra of the probe qubit $^{13}$C after a readout pulse. (a) Spectrum of the PPS. (b), (c) and (d) Spectra of experimental instances 1, 2 and 3 in 2-fold case, respectively. (d) and (e) Spectra of experimental instances 4 and 5 in 3-fold case, respectively.

Table 1
Ten selected experimental instances for 2-fold and 3-fold Forrelation problems and their respective target Forrelation $\Phi_{\text{th}}$, where $D\{a, b, c, d\}$ indicates a $4 \times 4$ diagonal matrix with diagonal elements $a$, $b$, $c$ and $d$.

| Case | Instance | $Q_1$ | $Q_2$ | $Q_3$ | $\Phi_{\text{th}}$ |
|------|----------|-------|-------|-------|-------------------|
| 2-fold | 1 | $D\{11 - 11\}$ | $D\{1 - 1\}$ | 1 |
|       | 2 | $D\{11 - 11\}$ | $D\{1 - 1\}$ | 0.5 |
|       | 3 | $D\{111 - 1\}$ | $D\{11\}$ | 0 |
|       | 4 | $D\{-111\}$ | $D\{-11\}$ | -0.5 |
|       | 5 | $D\{11 - 1\}$ | $D\{-11\}$ | -1 |
| 3-fold | 1 | $D\{11 - 1\}$ | $D\{1 - 1\}$ | 1 |
|       | 2 | $D\{11 - 1\}$ | $D\{1 - 1\}$ | 0.5 |
|       | 3 | $D\{11 - 1\}$ | $D\{1 - 1\}$ | 0 |
|       | 4 | $D\{-111\}$ | $D\{-11\}$ | -0.5 |
|       | 5 | $D\{-11\}$ | $D\{-11\}$ | -1 |
Here we focus on the experimental results of 2-fold and 3-fold FORRELATION in nuclear spins. There are in total five possible values for the FORRELATION $\Phi_{f_1, ... , f_k}$ in both cases, namely, $(1, 0.5, 0, -0.5, -1)$. For each theoretical value of $\Phi_{f_1, ... , f_k}$, we associate it with a set of functions listed in Table 1. The operators $O_1$, $O_2$ and $O_3$ are $4 \times 4$ diagonal matrices in the computational basis, i.e., $D([a, b, c, d])$, where $a, b, c, d \in \{1, -1\}$.

The quantum algorithm for solving the $k$-fold FORRELATION problem can be decomposed into several elemental $\pi/2$ (and $\pi$) hard pulses and evolutions under spin–spin $J$-couplings of the internal Hamiltonian in experiment [33,34], and the whole pulse sequences can be compiled with phase tracking and numerical optimization of the refocusing scheme. For example, the pulse sequences for the instance 4 in the 3-fold case are shown in Fig. 2b. There are at least $52 \pi/2 (\pi)$ hard pulses (without adding the refocusing pulses) and 31 free evolution periods in total are required. In experiment, we utilized the gradient ascent pulse engineering (GRAPE) method [35] to pack the whole algorithm for each instance into one shaped pulse, with the length of each pulse being 15 ms and the number of segments being 5000. All the shaped pulses are calculated with their fidelities reaching 99.5% and are guaranteed to be robust to the inhomogeneity of radio-frequency pulses.

### Table 1

| $f_1$ | $f_2$ | $f_3$ | $f_4$ | $f_5$ |
|-------|-------|-------|-------|-------|
| 1     | 0     | 0     | 0     | -1    |
| 1     | 0     | 0.5   | 0     | -0.5  |
| 1     | 0     | 0.5   | 0.5   | -0.5  |
| 1     | 0     | 1     | -1    | 0.5   |
| 1     | 0     | 1     | 0     | 0.5   |

Fig. 4. (Color online) The experimental results of Forrelation $\Phi$. The dashed lines are the criterion values for the Forrelation decision problem. The inset is zooming in on the 3rd instance in both plot. Only instances 1 and 3 satisfy $|\Phi| \leq 1/100$ or $\Phi \geq 3/5$ in both 2-fold and 3-fold cases.

Fig. 5. (Color online) Real parts of theoretical and experimental density matrices of $\rho_0$ (PPS), $\rho_{2,1}$, $\rho_{2,2}$, $\rho_{2,3}$, $\rho_{3,4}$ and $\rho_{3,5}$. In each figure, the outer transparent columns stand for the theoretical values, while the inner colored stand for the experimental results.
5. Experimental results

Finally, an observation of the final state on the probe qubit \( \sigma_2 \) is conducted in each run of the experiment to get the probabilities of the probe qubit in the \( |0\rangle \) state, by integrating the whole spectrum. Fig. 3 shows the NMR spectra of the probe qubit \(^{13}\text{C}\), where \( a \) is the spectrum of \(^{13}\text{C}\) after a readout pulse when the system is initialized in PPS taken as a calibration. (b), (c) and (d) are the \(^{13}\text{C}\) spectra of the final state after conducting quantum algorithms of the selected transformation operators instances 1, 2 and 3 in 2-fold case, respectively, while (e) and (f) show the spectra after conducting quantum algorithms of the selected transformation operators instances 4 and 5 in 3-fold case, respectively.

The experimental results of the 5 selected instances in 2-fold (3-fold) case are respectively 0.9867, 0.4509, \(-0.0011, -0.4454\) and \(-0.9516 (0.9791, 0.4659, -0.0068, -0.4871 and -0.9355)\), as shown in Fig. 4. From the results, we can distinguish for the 10 selected instances which interval their values locate. i.e., instance 1 is in the case of \( \Phi < 3/5 \), instance 3 is in the case of \(|\Phi| < 1/100 \) with bounded probability of error in both 2-fold and 3-fold cases; therefore our experimental results indicate that 2-fold and 3-fold FORRELATION problems can be solved by making only 1 quantum query to each of \( f_1, f_2 \) and \( f_3 \).

Furthermore, we performed a full QST on the final states. Indeed, we conducted QST on instances 1, 2, and 3 in 2-fold case, and instances 4 and 5 in 3-fold case. To be noticed, it is not necessary to perform QST on all the final states because of the same radio-frequency pulses generation method and the similar experimental process. To describe the density matrices of the final states, we label them as \( \rho_{kn} \) for instance \( n \) in k-fold case. The real parts of the density matrices for the final states of \( p_{21}, p_{22}, p_{23}, p_{31}, p_{32}, p_{33} \) and \( p_{34} \) are presented in Fig. 5 (the imaginary parts are very close to zero). Since the fidelities of all the shaped pulses generated by GRAPE are almost 99.5\%, the experimental final density matrices are indeed very close to the theoretical ones, as shown in the Fig. 5. The five selected experimental fidelities are 95.27\%, 96.26\%, 94.69\%, 94.73\% and 94.64\%, respectively, indicating a very good implementation of the quantum algorithm in experiment.

6. Conclusions

In summary, we tested a quantum implementation of solving the \( k \)-fold FORRELATION problem [16] in a prototype experiment. The FORRELATION \( \Phi_{k-1, k} \) among a set of “black-box” functions are obtained by the spin polarization \( \langle \sigma_2 \rangle \) of an ancillary qubit. The goal of the experiment is to determine if \( |\Phi_{k-1,k}| \leq 1/100 \) or \( \Phi_{k-1,k} \geq 3/5 \). In our experiments, 5 selected instances of both the 2-fold and 3-fold FORRELATION problems are solved on a three-qubit NMR quantum information processor. The experimental results successfully identify that instances 1 and 3 are in the case of \(|\Phi_{k-1,k}| \leq 1/100 \) or \( \Phi_{k-1,k} \geq 3/5 \) in both 2-fold and 3-fold cases. The quality in the preparation of the PPS and the implementation of the quantum algorithm are benchmarked by a full quantum state tomography for both the initial and the final states. Besides, all the shaped pulses are designed to be robust to the inhomogeneity of the radio-frequency pulses. The main source of errors are caused by the imperfection of GRAPE pulses and the instrumental-related imperfection of the shaped pulse. The total length of each shaped GRAPE pulse is only 15 ms, which is much shorter than the relaxation time of our system. To our knowledge, this is the first implementation of solving FORRELATION problem reported in the literature, and the experimental method can be extended to a more complex version of the multiple-fold FORRELATION in \( 2^n \)-dimensional space, and be implemented in other platforms such as superconducting devices and trapped ions.

Conflict of interest

The authors declare that they have no conflict of interest.

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