Good dynamics versus bad kinematics. Is entanglement needed for quantum computation?

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We consider quantum computing with pseudo-pure states. This framework arises in certain implementations of quantum computing using NMR. We analyze quantum computational protocols which aim to solve exponential classical problems with polynomial resources and ask whether or not entanglement of the pseudo-pure states is needed to achieve this aim. We show that for a large class of such protocols, including Shor’s factorization, entanglement is necessary. We also show that achieving entanglement is not sufficient: if the noise in the state is sufficiently large, exponential resources are needed even if entanglement is present.

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In a beautiful example of how technology can stimulate fundamental physics, the proposals for implementing quantum computing via liquid state NMR have sparked a debate recently on the very nature of quantum computing. More precisely, doubts have been raised as to whether entanglement is a necessary requirement for a quantum computer to be able to speed-up a computation (exponentially) relative to a classical computer.

The proposal to use liquid state NMR with pseudo-pure states for quantum computing has two important aspects.

- **Bad kinematics.** On the one hand liquid state NMR quantum computing has a great disadvantage: One cannot prepare pure states. This situation is different from the original quantum computational protocols which considered the quantum computer in a pure state $|\Psi\rangle$. Instead, in the NMR protocol one prepares “pseudo-pure” states, i.e. mixed states of the form

$$\rho = (1 - \epsilon)M + \epsilon|\Psi\rangle\langle\Psi|$$

(1)

where $M$ is the maximally mixed state (i.e. the identity density matrix normalized to have trace 1). In other words, in NMR the pure state $|\Psi\rangle$ is contaminated by noise.

Furthermore it has been shown that in all experiments to date the noise is so large that the “pseudo-pure” state $\rho$ is non-entangled even if the pure-state component $|\Psi\rangle$ is entangled. Since entanglement is widely considered to be the main ingredient in quantum computing, these results lead to the question as to whether the NMR scheme is a “true” quantum computation.

- **Good dynamics.** On the other hand, the NMR experiments have a great advantage: one can produce correct dynamics, that is, the interactions between the spins are exactly as required in the theoretical quantum computational protocols. Thus if the initial state of the spins would be pure instead of “pseudo-pure”, NMR experiments would completely implement the original quantum computation protocols.

Furthermore, the noise in the “pseudo-pure” state looks quite benign - it averages to zero (without loss of generality we can consider our observables to be traceless). Thus the expectation value of any operator $A$ when the quantum system is in a given pure state $\Psi$ is the same, up to normalization, as the average in the corresponding pseudo-pure state $\rho = (1 - \epsilon)M + \epsilon|\Psi\rangle\langle\Psi|$, i.e.

$$\text{Tr}(A \rho) = \epsilon \langle \Psi | A | \Psi \rangle$$

(2)

Given the good dynamics some authors have suggested that in fact liquid-state NMR computing is nonetheless a “true” quantum computation, capable of speeding-up computations relative to classical computers. As a corollary, it was suggested that perhaps entanglement is not a sine-qua-non requirement for quantum computing.

Whether or not entanglement is a necessary condition for quantum computation is a question of fundamental importance, which obviously goes far beyond the NMR computing context in which it arose. In the present Letter we study this question for the pseudo-pure state quantum computing. Specifically we analyze quantum computational protocols which aim to solve exponential classical problems with polynomial resources and ask whether or not entanglement of the pseudo-pure states is needed to achieve this aim. We show that for a large class of such protocols, including Shor’s factorization, entanglement of the pseudo-pure states is necessary: unless the pseudo-pure state of the quantum computer becomes entangled during the computation, the aim of transforming exponential problems to polynomial ones cannot be achieved.

We will first consider the general effect of noise on the computation. Then the relationship between separability and noise.

\[
\text{Tr}(A \rho) = \epsilon \langle \Psi | A | \Psi \rangle
\]

(2)
Consider then a pure state computational protocol in which the computer starts in the state $|\Psi_0\rangle$ and ends in the state $|\Psi_f\rangle = U|\Psi_0\rangle$ where $U$ is the unitary time evolution operator which describes the computation. The corresponding computational starting with pseudo-pure state

$$\rho = (1 - \epsilon)M + \epsilon|\Psi_0\rangle\langle\Psi_0|$$

ends up in the state

$$\rho = (1 - \epsilon)M + \epsilon|\Psi_f\rangle\langle\Psi_f|.$$  \hspace{1cm} (3)

Upon reaching the final state, a measurement is carried out and the result of the computation is inferred from the result of the measurement.

We will assume the most favorable case that the pure state protocol gives the correct answer with certainty with a single repetition of the protocol and that if the result of the computation is found, one can check it by using the computational problem with certainty with one repetition. Thus if $\epsilon$ becomes exponentially small with $N$, the number governing the scaling of the classical problem, (in other words the noise becomes exponentially large with $N$), the protocol requires an exponential number of repetitions to get the correct answer. So for this amount of noise, the quantum protocol with a pseudo-pure state cannot transform an exponential problem into a polynomial one: even in the best possible case that the pure-state protocol takes one computational step, the protocol with noise takes exponentially many steps. We emphasize that this conclusion applies quite generally to pseudo-pure state quantum computing and is independent of the discussion of separability which follows later.

In the state $|\Psi_0\rangle$ there is a probability $\epsilon$ of finding the computer in the “correct” final state $|\Psi_f\rangle$ arising from the term

$$\epsilon|\Psi_f\rangle\langle\Psi_f|$$

in $|\Psi_0\rangle$. As stated above, will assume here the most favorable case, that if the state is $|\Psi_f\rangle$ then from the outcome of the final measurement, one can infer the solution to the computational problem with certainty with one repetition. We note that general protocols, such as Shor’s algorithm for example, a single repetition of the protocol is not sufficient to find the correct answer.

There is also the probability $(1 - \epsilon)$ of finding the computer in the maximally mixed state $M$. In this case there is a possibility that the correct answer will be found, since the noise term contains all possible outcomes with some probability. However, the probability of finding the correct answer from the noise term must be at least exponentially small with $N$. Otherwise there would be no need to prepare the computer at all: one could find the correct answer from the noise term simply by repeating the computation a polynomial number of times. In fact, if the probability of finding the correct answer from the noise term did not become exponentially small with $N$ we could dispense with the computer altogether. For using a classical probabilistic protocol which selected from all the possibilities at random, we would get the correct answer with probability of the order of one with only a polynomial number of trials.

Thus we may say that the probability of finding the correct answer from the state $|\Psi_0\rangle$ is essentially $\epsilon$ and so the computation must be repeated $\frac{1}{\epsilon}$ times on average to find the correct answer with probability of order one.

We now consider whether reaching entangled states during the computation is a necessary condition for exponential speed-up. We address this by investigating what can be achieved with separable states. Specifically we impose the condition that the pseudo-pure state remains separable during the entire computation. For an important class of computational protocols we show that this condition implies an exponential amount of noise.

The protocols which we consider use $n = n_1 + n_2$ qubits of which $n_1$ are considered to be the input registers and the remaining $n_2$, the output registers. We assume that $n_1$ and $n_2$ are polynomial in the number $N$ which describes how the classical problem scales. As stated earlier we consider problems in which the quantum protocol gives and exponential speed-up over the classical protocol, specifically the classical protocol is exponential in $N$ whereas the quantum protocol is polynomial in $N$. (For example in the factorization problem, the aim is to factor a number of the order of $2^N$. The classical protocol is exponential in $N$ and in Shor’s algorithm, $n_1$ and $n_2$ are linear in $N$.)

We first describe the protocols as applied to pure states. The first steps are

1. Prepare system in the ground state

$$|\Psi_0\rangle = |000...0\rangle \otimes |000.0\rangle$$

2. Perform a Hadamard transform on the input register, so that the state becomes

$$|\Psi_1\rangle = \frac{1}{2^{n_1/2}} \sum_{x=0}^{2^{n_1}-1} |x\rangle \otimes |000..0\rangle$$

3. Evaluate the function $f : \{0,1\}^{n_1} \rightarrow \{0,1\}^{n_2}$. The state becomes

$$|\Psi_2\rangle = \frac{1}{2^{n_1/2}} \sum_{x=0}^{2^{n_1}-1} |x\rangle \otimes |f(x)\rangle$$
\[ \rho_0 = (1 - \epsilon)M_{2^n} + \epsilon |\Psi_0\rangle \langle \Psi_0| \tag{9} \]

where \(|\Psi_0\rangle\) is given in (4), and \(M_{2^n}\) is the maximally mixed state in the \(2^n\) dimensional Hilbert space. After the second computational step the state is

\[ \rho_0 = (1 - \epsilon)M_{2^n} + \epsilon |\Psi_2\rangle \langle \Psi_2|. \tag{10} \]

Consider now protocols in which the function \(f(x)\) is not constant. Let \(x_1\) and \(x_2\) be values of \(x\) such that \(f(x_1) \neq f(x_2)\). Thus we may write the state \(|\Psi_2\rangle\) as

\[ |\Psi_2\rangle = \frac{1}{2^{n_1/2}} (|x_1\rangle |f(x_1)| + |x_2\rangle |f(x_2)| + |\Psi_r\rangle) \tag{11} \]

where \(|\Psi_r\rangle\) has no components in the subspace spanned by \(|x_1\rangle |f(x_1)|, |x_1\rangle |f(x_2)|, |x_2\rangle |f(x_1)|, |x_2\rangle |f(x_2)|\). It is convenient to relabel these states and write

\[ |\Psi_2\rangle = \frac{1}{2^{n_1/2}} ((1)|1\rangle + |2\rangle |2\rangle + |\Psi_r\rangle) \tag{12} \]

where \(|\Psi_r\rangle\) has no components in the subspace spanned by \(|1\rangle|1\rangle, |1\rangle|2\rangle, |2\rangle|1\rangle, |2\rangle|2\rangle\).

We now derive a necessary condition on \(\epsilon\) for the state of the system to be separable throughout the computation. For consider projecting each particle onto the subspace spanned by \(|1\rangle\) and \(|2\rangle\). The state after projection is

\[
\rho'_{2} = \frac{1}{A} \left[ \frac{4(1 - \epsilon)}{2^{n_1+n_2}} M_4 \\
+ \frac{2\epsilon}{2^{n_1}} \left( \frac{|1\rangle|1\rangle + |2\rangle|2\rangle}{\sqrt{2}} \right) \left( \frac{|1\rangle|1\rangle + |2\rangle|2\rangle}{\sqrt{2}} \right) \right]
= (1 - \epsilon')M_4 \\
+ \epsilon' \left( \frac{|1\rangle|1\rangle + |2\rangle|2\rangle}{\sqrt{2}} \right) \left( \frac{|1\rangle|1\rangle + |2\rangle|2\rangle}{\sqrt{2}} \right), \tag{13}
\]

where

\[ A = \left( \frac{4(1 - \epsilon)}{2^{n_1+n_2}} + \frac{2\epsilon}{2^{n_1}} \right) \tag{14}\]

is the normalization factor, \(M_4\) is the maximally mixed state in the 4-dimensional Hilbert space spanned by \(|1\rangle|1\rangle, |1\rangle|2\rangle, |2\rangle|1\rangle, |2\rangle|2\rangle\) and

\[ \epsilon' = \frac{2\epsilon}{2^{n_2} A} = \frac{\epsilon}{(1 - \epsilon)2^{-n_2+1} + \epsilon}. \tag{15}\]

Now a 2 qubit state of the form

\[ (1 - \delta)M_4 + \delta \left( \frac{|1\rangle|1\rangle + |2\rangle|2\rangle}{\sqrt{2}} \right) \left( \frac{|1\rangle|1\rangle + |2\rangle|2\rangle}{\sqrt{2}} \right) \tag{16}\]

is entangled for \(\delta > 1/3\). Therefore the original state (11) must have been entangled unless

\[ \epsilon' \leq 1/3 \implies \epsilon \leq \frac{1}{1 + 2^{n_2}}. \tag{17}\]

since local projections cannot create entangled states from unentangled ones.

Therefore we conclude that if we have a computational protocol (for non-constant \(f\)) starting with a mixed state of the form (1) and if we require that the state remains separable throughout the protocol, then we certainly need

\[ \epsilon \leq \frac{1}{1 + 2^{n_2}}. \tag{18}\]

However we have shown earlier that, even in favorable circumstances, a computation with noise \(\epsilon\) takes of the order of \(1/\epsilon\) repetitions to get the correct answer with probability of the order of one.

Thus we reach our main result that computational protocols of the sort we have considered require exponentially many repetitions. So no matter how efficient the original pure state protocol is, the mixed state protocol which is sufficiently noisy that it remains separable for all \(N\), will \textit{not} transform an exponential classical problem into a polynomial one.

We note that while we have considered protocols of a specific form, many of the details are unimportant. As long as the number of qubits \(n_1\) and \(n_2\) in the output register is polynomial in the number \(N\) which governs the classical problem, and the pure state protocol goes through a state which has a non-negligible amount of entanglement, similar conclusions can be drawn.

We repeat here that our conclusions only apply to separable states of pseudo-pure state form (4). We have nothing to say at this stage about separable states of other forms. We have also only considered exponential speed up so that our results do not apply to Grover's algorithm for example (10). Furthermore, we cannot rule out the possibility of the future discovery of more efficient algorithms of Shor type to which our results do not apply.

We have shown earlier that having entanglement is a necessary condition: is it sufficient? Our earlier results show that it is not. As long as the noise is exponential (i.e. \(\epsilon\) decreases exponentially with \(n\)), the computation has to be repeated an exponential number of times, even if entangled states are reached during the computation (we note that it is known that there are entangled states with an exponential amount of noise (5)).

Finally let us return to NMR quantum computation which gave rise to the issues we have been discussing. In our previous discussion we had in mind that one has a single quantum computer and \(1/\epsilon\) then gives the number of times the computation has to be repeated. In NMR, each molecule in the sample is considered to be a quantum computer. Here, rather than repeating the computation on the same computer, one treats a large number of computers in parallel. Thus, if each molecule could be accessed individually, \(1/\epsilon\) would the number of individual molecular computers one would need in the sample.
In fact, rather than being able to address each molecule individually, one can only measure bulk properties of the sample - a far less favorable situation - so that $1/\epsilon$ is a lower bound on the sample scaling required.

The usual construction of pseudo-pure states in NMR has $\epsilon \sim \frac{2}{n^2}$. In the light of the discussion in the previous paragraph the sample size would therefore have to grow exponentially with $n$. Thus this framework does not allow one to convert exponential classical problems into polynomial ones via Shor-type protocols.

Of course, other ways of using liquid state NMR as a quantum computer might be found with more favorable scaling than current techniques. Whether or not this is possible is obviously beyond the scope of this paper.

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