Introduction to Quantum Computers and Quantum Algorithms

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August 7, 2021

Abstract

This is a short introduction to Quantum Computing intended for physicists. The basic idea of a quantum computer is introduced. Then we concentrate on Shor’s integer factoring algorithm.

1 What is a Quantum Computer?

In this first lesson I want to introduce the basics necessary to understand quantum algorithms: What a quantum computer is, quantum gates and how to do usual computations on a quantum computer, namely “reversible computation”.

1.1 A Quantum Computer = many 2-level systems (qubits)

A “qubit” (or “quantum bit”) is simply a quantum mechanical 2 level system. In theoretical considerations, for convenience we usually assume that the 2 levels are energy degenerate. So a qubit is any quantum mechanical system with a 2 dimensional Hilbert space, or a system of which we use only a 2 dimensional subspace. Examples are the spin degree of freedom of a spin 1/2 particle, the polarization of a single photon or an atom or ion in the ground state or a particular excited state. Also we can use the ground- and first excited state of a harmonic oscillator or of a mode of the electromagnetic field, thus no photon (vacuum) or 1 photon. In the 2 dimensional Hilbert space we choose 2 orthonormal basis states and denote them by $|0\rangle$ and $|1\rangle$. (For a non energy-degenerate qubit these would usually be the energy eigenstates, like the electronic ground state and some excited state of an atom.)

A quantum computer simply consists of many such qubits. For convenience we imagine that usually they do not interact, thus when we don’t intervene, the time evolution of the quantum computer is trivial (say $H_0 = 0$). Now the
Hilbert space of many qubits is the tensor product of their Hilbert spaces. A natural basis is given by states of the form

\[ |1101001 \ldots \rangle = |1\rangle |1\rangle |0\rangle |1\rangle |0\rangle |1\rangle \ldots \]

Thus the overall Hilbert space is spanned by all binary strings of some length (corresponding to each qubit being either in state |0\rangle or in state |1\rangle). Note that the dimension of this space increases exponentially with the number of qubits, and so the number of probability amplitudes necessary to specify a state increases exponentially.

As with conventional computers, we often imagine that bit strings stand for numbers in binary, so we can label such basis states with numbers. Then a (pure) state of a \( n \) qubit quantum computer (QC) can be written

\[ |\Psi_{QC}\rangle = \sum_{x=0}^{2^n-1} a_x |x\rangle \]

with the normalization \( \sum_x |a_x|^2 = 1 \).

**exercise:** Estimate the maximal number of qubits such that it is still possible to store the amplitudes describing their overall state on some presently available computer. What about a computer using all matter in the visible universe?

### 1.2 Computing: “quantum gates”

Usually (and without loss of generality) we assume that the quantum computer is initially in the “all 0” state, thus the product state where all qubits are in state |0\rangle:

\[ |QC_{ini}\rangle = |0000\ldots \rangle \]

The computation consists of a sequence of unitary operations that are realistically assumed to only act on few qubits at a time. Formally, when we apply e.g. some \( U(8) \) transformation to 3 qubits, the overall transformation is the tensor product of this transformation with the identity on the remaining qubits. Clearly, it is necessary to apply unitary transformations to more than just individual qubits in order to arrive at interesting states, as otherwise we will be stuck with (unentangled) product states. **exercise:** Check that typically all amplitudes describing the state of the quantum computer are changed when a quantum gate (a unitary operation) is applied to, say, one qubit.

It turns out that it is enough if we are able to apply unitary transformations (= “quantum gates”) to any two qubits (thus \( U(4) \)), and of course also to individual qubits (\( U(2) \)). In the following it will also be convenient to assume that we can directly do some 3 qubit quantum gate, although it could really be decomposed into 1- and 2 qubit gates.

Below is a graphical representation of a quantum computation on 5 qubits (which is little for theoreticians, but much for experimentalists...). The single
Figure 1: A sequence of quantum gates on 5 qubits, all initialized in state $|0\rangle$. Time runs from left to right. Only 1-qubit and 2-qubit gates are used, as this is sufficient for universal quantum computation. A complete quantum computation would also include a measurement of each qubit at the end (on the right).

dots represent some $U(2)$ transformations on single qubits while the pairs of dots joined by a vertical line stand for 2-qubit gates, thus some $U(4)$s. Time runs from left to right. Take care not to confuse the qubits (the 5 horizontal lines) with the 32 basis states.

1.2.1 How “quantum gates” are carried out

Quantum gates are done by “switching on” terms in the Hamilton operator for an appropriate time. Physically, this is done by applying “exterior fields” to the qubits, thus fields which can be treated as classical. So called “coherent states” as e.g. produced by a laser or radio transmitter are like this, thus they behave nearly classically. It is remarkable that such states exist which allow us to manipulate a quantum system without getting entangled with it, thus without causing decoherence. Also it is not obvious that these states occur naturally.

1-qubit gates are usually rather easy to carry out. In an ion trap quantum computer, one can shine a laser with the right frequency for some time at an ion. By tuning the frequency to the energy difference between the ground- and excited states (thus resonance), we induce “Rabi oscillations” between $|0\rangle$ and $|1\rangle$. Note that to carry out the gate correctly we also have to control the intensity, time and phase of the laser.

In most hardware proposals, 2-qubit gates are more difficult to carry out. One possibility is to simply bring 2 qubits closely together so that they interact.
1.3 conventional computation on a quantum computer

Most of Shor's algorithm is simply a conventional computation, but applied to a superposition of (exponentially) many “conventional” states. For a computation to work on superpositions it has to be coherent, thus it must consists of (unitary) quantum gates, excluding e.g. measurements. Also “throwing away” qubits would not be allowed, as it would destroy coherence. But for Shor’s algorithm it is necessary that at the end only a certain function is calculated and not also some intermediate results. In the following I will also show how the unwanted intermediate results can be “deleted” in a coherent way.

1.3.1 reversible gates

First of all we want only to use gates which have as many output bits as input bits (unlike e.g. an AND gate which is $2 \rightarrow 1$), as we don’t want to destroy qubits.

It is easy to imagine a classical computation with such gates: The computer is a bit string of constant length. We can now act on individual bits with, say, NOT or RESET (reset to 0). In addition, we can pick any 2 bits in the bit string and e.g. leave the first bit unchanged and set the second one to the AND of the 2 bits, or SWAP the 2 bits. Clearly in this way any computation can be done. (It is well known that any classical computation can be done by using just NAND (= not AND), actually there used to be chips with several NAND gates on them which could then be wired into logic circuits by engineers.)

We want to do classical computation with unitary gates to preserve coherence. The only unitary transformations which map all “computational basis states” (binary strings) to such basis states (and not superpositions), are permutations of these basis states. (Thus in each row and column of the unitary matrix there is one 1.) Such classical gates are called “reversible”, as they are 1 to 1. There is only one non-trivial such 1-qubit gate, the NOT:

$$U_{NOT} : |0\rangle \rightarrow |1\rangle \quad |1\rangle \rightarrow |0\rangle$$

An important 2-qubit gate is the “controlled-NOT” (CNOT or XOR), which doesn’t change the first bit but changes the second one if the first is a 1:

$$U_{CNOT} : |a, b\rangle \rightarrow |a, a \ XOR \ b\rangle = |a, a \oplus b\rangle$$

(XOR means “exclusive OR”, thus either $a$ or $b$ but not both; $\oplus$ is addition modulo 2) [exercise: check that CNOT is a permutation of the 4 possible 2-bit strings, and thus is unitary. Also write down the 4x4 matrix representing the overall transformation when we act on each of 2 qubits with a NOT (thus find the tensor product of the two NOTs).]

It is not difficult to show that the above 2 gates are not enough to do universal computation. We still lack something that allows to compute an AND, but this is not possible with reversible 2-bit gates [exercise: show that if one of the output bits of a 2-bit gate is the AND of the inputs, then the gate can’t be
reversible. For this we use the CCNOT (= "Toffoli gate") which acts as follows on 3 (qu)bits (thus it’s a $U(8)$):

$$U_{CCNOT} : |a, b, c⟩ \rightarrow |a, b, c \ XOR (a \ AND \ b)⟩ = |a, b, c \oplus (a \cdot b)⟩$$

Thus the last bit is flipped only if $a$ and $b$ are both 1.

1.4 "garbage uncomputing"

It is now clear that starting from some input $|x⟩$ (thus the binary string representing $x$) we can compute any function $f(x)$. But we will probably also produce some unwanted output $g(x)$. E.g. if we compute the AND of two bits with the CCNOT, the two original bits will still be around. Of course besides $|x⟩$, we will also need qubits initialized to $|0⟩$. In summary, what we can do is:

$$|x, 0⟩ → |f(x), g(x)⟩$$

where the 0 stands for several qubits. In the following, I will not always indicate when in the process of computing we will add such “fresh” qubits in state $|0⟩$.

As they are unitary, all the gates we use are reversible; thus, we can also undo the whole computation by applying the inverse gates in reversed order. The trick to get rid of the unwanted “garbage” $g(x)$ now is to first copy the wanted $f(x)$ into a “save place” and then undo the original computation (which doesn’t touch the “save place”). Copying into a “save place” can simply be done by “XORing” each bit of $f(x)$ into a fresh bit in state $0$. Thus overall:

$$|x, 0, 0⟩ \overset{U_f}{\rightarrow} |f(x), g(x), 0⟩ \overset{\text{copy } f(x)}{\rightarrow} |f(x), g(x), f(x)⟩ \overset{U_f^{-1}}{\rightarrow} |x, 0, f(x)⟩$$

Thus we will still have $x$ in the final state, but this is unavoidable if $f(x)$ is not bijective (1 to 1), as otherwise the overall computation would not be unitary. The “work qubits”, which at the end of the computation are again in state $|0⟩$, pose no problem, as they are now again unentangled with the other qubits and could thus safely be thrown away.

**exercise:** If $f(x)$ is 1 to 1 and if we know an (efficient) algorithm for its inverse, we can do $|x⟩ \rightarrow |f(x)⟩$ by using similar tricks as above. Hint: use the reverse of the algorithm which computes $f^{-1}$, thus $(U_f^{-1})^{-1}$.

1.5 measurement of the quantum computer

At the end of a computation, the quantum computer is observed and thus its state “collapses” to some binary string that will be the (classical) output of the computation. Thus we assume that we measure an observable whose (non-degenerate) eigenstates are the computational basis states. This corresponds to measuring each qubit separately such that it gets projected to either $|0⟩$ or $|1⟩$. It would be unrealistic to assume that we can simply measure any observable
on the whole quantum computer. On the other hand, if we would like to make a measurement corresponding to another basis, this is equivalent to first doing some unitary transformation (a sequence of quantum gates) and then doing the usual measurement. In this way, one can e.g. effectively measure two qubits in the (maximally entangled) “Bell basis”.

1.6 summary
We have shown that, given a conventional algorithm to compute \( f(x) \), we can construct a sequence of quantum gates which act on basis states as \( |x\rangle \rightarrow |x, f(x)\rangle \). Because time evolution in quantum theory is linear (unitary maps are linear), we can apply this to superpositions and will get a superposition of the outputs:

\[
\sum_x a_x |x\rangle \rightarrow \sum_x a_x |x, f(x)\rangle
\]

Thus in a way we can compute the function \( f(x) \) in one go in “quantum parallelism” for many inputs \( x \). Of course simply observing the final superposition will simply give us a basis state at random (with probability \( |a_x|^2 \)), thus in Shor’s algorithm we first do something else before measuring.

2 Shor’s quantum factoring algorithm

In 1994 Peter Shor found his famous algorithm which created much interest in quantum computing. Probably this algorithm is still the most important quantum algorithm. (In other words, there has not been as much progress as many people had hoped for.)

2.1 The problem: finding prim factors
The problem is to write a given positive integer as a product of prim numbers, e.g. 12827 = 101 × 127. It is not so difficult to find small prime factors, but if there are several large prime factors, no fast classical algorithm for factoring is known. Much better algorithms are known than just trying to divide a number by all possible prim factors (up to the square root). But the presently best algorithm still takes time which is roughly exponential in the third root of the number of digits: the number of elementary operations to factor a number \( N \) is roughly \( 10^{\Omega(\log N)^{1/3}} \). The largest numbers that can presently be factored (with much computer power and time) have about 150 decimal digits.

2.1.1 public key cryptography: RSA
Factoring is important because it would allow to break the important “public key” cryptosystem RSA. With this system, it is possible to make public how to encrypt messages, but this will not allow the public to decrypt messages. (In
principle of course, decryption is possible, but it is thought that it would take too long.) Simplifying things a bit, in RSA the public key needed to encrypt, is the product $p \times q$ of two very large prim numbers, while the “private key” e.g. allowing a military headquarters to decrypt messages coming by radio from the field, are the two large prim numbers $p$ and $q$ separately. So factoring $pq$ would allow breaking the code.

It is possible to calculate rather quickly whether a given number is a prim number or not, so a computer can easily generate some large prim numbers $p$ and $q$. (So it is also easy to find that a number is composite (not prim), but this is not the same as actually finding the factors.)

2.2 Fermat’s little theorem and its generalization

Shor’s algorithm relies on the fact that factoring can be reduced to finding the period $r$ of some periodic function from the integers to the positive integers: $f: \mathbb{Z} \rightarrow \mathbb{N}$, with $f(x + r) = f(x)$ for all $x$.

First consider Fermat’s little theorem which says that $a^{p-1} \mod p = 1$, for a prim $p$ and any non-zero integer $a \in \mathbb{Z}_p$ (thus $0 < a < p$). The reason is simply that those $p-1$ numbers form the multiplicative group mod $p$. However, in any finite group an element to the power order of the group (number of elements) is the neutral element: $g^{[G]} = e$.

The same can be done for the ring mod $pq$. In this ring all numbers $< pq$ which are coprim to $pq$ (thus their greatest common divisor (gcd) is 1) have a multiplicative inverse. There are $(p-1)(q-1)$ such numbers, so this is the order of the multiplicative group.

[exercise: Show that mod $n$ a number $m$ which is coprim to $n$ has a multiplicative inverse, thus there is an integer $m'$ such that $m'm = kn + 1$ for some integer $k$. Hint: Euclid’s (non-quantum...) algorithm to find gcd(m,n) will also give $k$ and $m'$.

Thus we have for any $a$ coprim to $pq$ that $a^{(p-1)(q-1)} \mod pq = 1$. Now consider the function $f_a(x) = a^x \mod pq$. It is periodic with period $r$ where $r$ is the smallest number such that $a^r \mod pq = 1$. In addition, within a period it is 1 to 1, thus

$$f(x) = f(y) \iff x - y = kr \quad (r = \text{period}, k \text{ integer})$$

In Shor’s algorithm the period $r$ of $f(x)$ is determined. It is clear that $r$ has to be a fraction of $(p-1)(q-1)$, and usually it is not a very small fraction. If we knew $(p-1)(q-1)$ (and of course $pq$) we could easily find $p$ and $q$ by solving a quadratic equation. I hope it is therefore plausible that finding $r$ also essentially solves the problem. I will not show the actual way Shor proposed to get the prim factors after finding $r$. Also for simplicity let us imagine that, as in RSA, we know that we have a product of just 2 primes, although Shor’s algorithm works for factoring any number.
2.3 Shor’s algorithm

Shor’s algorithm starts by preparing the “uniform amplitude superposition” (or “equally weighted superposition”) for some number \( n \) of qubits:

\[
\frac{1}{\sqrt{2^n}} \sum_{x=0}^{2^n-1} |x\rangle
\]

Note that in this state every qubit is in the state \( \frac{1}{\sqrt{2}} (|0\rangle + |1\rangle) \), thus this is an unentangled state which can easily be obtained.

In the main part of the algorithm we carry out the (really classical) algorithm to compute the function \( f(x) \) (thereby also adding some “fresh” qubits, and doing the “garbage uncomputing” as described above):

\[
\sum_{x} |x\rangle \rightarrow \sum_{x} |x, a^x \mod pq\rangle
\]

Above and in the following I will often leave away the state normalizations.

[exercise: Show how to quickly compute “modular exponentiation” even with big numbers. Compute (it’s possible by hand) \( 8^{65} \mod 37 \). Clearly first computing \( 8^{65} \) and only then doing \( \mod 37 \) is not the best. With what power of the number of digits of the three numbers involved (say all have \( n \) digits) does the computation time grow?]

Note: for Shor’s algorithm it is also helpful that the necessary “garbage uncomputing” can be done periodically and not only at the end, as this would use a lot of work space (qubits).

2.3.1 period finding

Let’s now imagine that we measure the “quantum register” in which \( f(x) \) is (this is actually not necessary, but makes the presentation easier):

\[
\sum_{x} |x, f(x)\rangle \rightarrow \sum_{k} |x_0 + kr, f(x_0)\rangle
\]

Where we assume that we chose the range \( 0 \ldots 2^n - 1 \) of the input \( x \) to cover many periods \( r \) (see later). Thus we will observe at random some value \( f(x_0) \) and collapse the state of the QC to the corresponding subspace. Because of the periodicity of \( f(x) \) many input values \( x \) will give the same value, thus with the collapse we will still have a superposition in the input register. As it is now unentangled with the “output register” (which is in a fixed basis state), we can only look at the input register:

\[
\sum_{k} |x_0 + kr\rangle
\]

where \( k \) runs from 0 to about \( 2^n/r \). If we plot the amplitudes as a function of the number of the basis state, we get equally spaced peaks, beginning not at 0 but with an offset \( x_0 \) (see figure 2). Observing this superposition will not help
in finding the period \( r \), as in each run of the algorithm we will get a different random offset.

### 2.3.2 The Quantum (Fast) Fourier Transform

The Quantum (Fast) Fourier Transform applies the discrete Fourier Transform to the amplitudes of a quantum register, thus:

\[
QFFT : \sum_{x=0}^{2^n-1} a_x |x\rangle \rightarrow \sum_{x=0}^{2^n-1} \tilde{a}_x |x\rangle \quad \text{with} \quad \tilde{a}_x = 2^{-n/2} \sum_{y=0}^{2^n-1} e^{2\pi ixy/2^n} a_y
\]

Note that this is a unitary transformation. It can be carried out very quickly with a quantum version of the well-known Fast Fourier Transform algorithm (FFT). Actually applying the QFFT to a \( n \) qubit register takes only \( O(n^2) \) quantum gates, or even less. Of course, these have to be “non-classical” gates (transforming “computational basis states” into superpositions). Without going into details, just note that two types of gates are used: the 1-qubit “Hadamard transform”

\[
H : |0\rangle \rightarrow \frac{1}{\sqrt{2}} (|0\rangle + |1\rangle) \quad |1\rangle \rightarrow \frac{1}{\sqrt{2}} (|0\rangle - |1\rangle)
\]

and 2-qubit “controlled phase shift” gates which do \( |11\rangle \rightarrow e^{i\phi} |11\rangle \), and leave the other 3 basis states unchanged.

Fourier transforming a periodic function will result in equally spaced peaks with no offset (thus, the first peak is at 0). In particular when transforming our equally spaced peaks we again get peaks and the offset will only show up in the phases of the resulting amplitudes. (A minor complication is that the peaks are now a bit “smeared out”. This is because in general the size \( 2^n \) of the Fourier transformation is not a multiple of the period \( r \).)

Below are pictures of the probability (amplitude absolute value squared) for the 64 basis states of a 6 qubit register before and after the QFFT. (For the first picture, we could also say that it shows the amplitudes themselves, as they are anyway real and positive.)
exercise: (maybe a bit cumbersome) Check that the discrete Fourier transform qualitatively really does what’s indicated in the picture above: The resulting peaks start at 0 and are spaced by $2^n/r$. Check that the width of the resulting peaks is of order $O(1)$ for $2^n \approx r^2$ or larger. (In Shor’s algorithm $n$ is chosen such that $2^n \approx (pq)^2$ or larger.)

(Note: the QFFT Fourier transforms $2^n$ amplitudes in only some $O(n^2)$ steps, while a conventional FFT would take $O(2^n \log 2^n)$ steps to transform $2^n$ numbers. Still, we cannot say that a quantum computer can Fourier transform much faster than a classical computer, as the operations are not comparable. Fourier transforming the amplitudes of a register has no classical analogue and cannot be used to do the job of a classical FFT.)

After the QFFT, we measure the register. Thus with high probability we will measure an integer close to $k \cdot 2^n / r$ for some random integer $k$. By dividing the observed integer by $2^n$ we get a fraction close to $k/r$. Then we can use the continued fraction expansion of this fraction, which gives a sequence of best rational approximations. With luck a single run will be enough to find $k$ (which doesn’t interest us) and $r$. Otherwise combining the output of several runs of the algorithm (different $k$’s) will allow to find $r$ with high probability.

As mentioned above, measuring the output register (which holds $f(x)$) was not necessary. After all we didn’t use the measured value $f(x_0)$.

What would have happened if we had not uncomputed the “garbage”? Then the computed function would have been the pair $f(x), g(x)$ and it would not have been periodical, thus the algorithm wouldn’t have worked.

3 Computer science aspects and further remarks

3.1 On the “power” of quantum computation

In what respect is a quantum computer thought to be more powerful than a conventional computer? In computer science, people typically ask about the
scaling of the computation time (number of elementary operations) with the size of a problem (number of bits of the input). So multiplying two \( n \)-digit numbers normally takes \( O(n^2) \) gates (it can also be done a bit faster), while in classical factoring algorithms the computation time grows faster than any power of the length (number of digits) of the number to be factored.

In “computational complexity theory” people are interested in how the best possible algorithm for a problem scales. The main distinction is between problems that can be solved in “polynomial time” (time grows at most like some power of the input size) and those which take longer, in particular exponential time. It is believed that many interesting problems like factoring cannot be solved in polynomial time, although so far it hasn’t been proven for any natural problem. Shor’s algorithm factors in polynomial time (actually \( O(n^3) \) because of modular exponentiation), so if classically factoring really is hard (can’t be done in polynomial time), it follows that quantum computers are more “powerful” than conventional ones.

### 3.2 Trying to solve NP-complete problems

Problems of the type NP (NP stands for “nondeterministic polynomial”) are roughly those where, once one has an answer, one can quickly check whether it really is correct. More formally, in NP problems, the answer is just “yes/no”, and if it is “yes”, there exists a short (polynomial time) proof that it really is “yes”. Most natural such problems are related, in that any one can be reduced to any other one. These problems are called “NP-complete”. Thus if one finds an
algorithm that can solve such a problem (in polynomial time), one can solve all of them (actually all NP problems). The question whether there is a conventional, efficient (polynomial time) such algorithm is the famous “NP=P(?)” question. (It is widely believed that NP≠P, but it’s still unproven...)

Factoring is not NP-complete, but it is NP (checking whether a given number really is a factor is easy). It would be a big breakthrough if somebody would manage to find an efficient (polynomial time) quantum algorithm for an NP-complete problem. But it seems not to be easy and may well not be possible at all.

3.3 Subtle questions about scaling of precision; fault tolerance

When talking about what kind of computers are possible, questions about the precision of their parts and whether this precision has to increase for longer computations should not be forgotten. For classical computers, digital (electronic) hardware seems not to have much of a scaling problem.

However, quantum computers seem to have a problem similar to that of classical analogue computers: There is a continuum of states and gates. If we shine a laser a bit too long at an ion, there is no automatic resetting like in digital electronics. Rather, the errors in the amplitudes will grow larger and larger as we do more gates. Also the very non-classical state of the quantum computer is sensitive to decoherence. Thus, it seems that the longer a quantum computation is, the more precise the gates would have to be.

Fortunately, it has been shown that clever error-correcting techniques are possible which make quantum computations “fault tolerant”. In these techniques, only a subspace of the whole QC is used. Qubits are encoded in complicated entangled subspaces of several physical qubits, and gates are done by really doing a whole sequence of gates on the physical qubits.

Fault tolerant quantum computing is maybe the second most important development after Shor’s algorithm in this field. Without it, running interesting quantum algorithms would be only a very distant possibility.

3.3.1 Is entanglement necessary for quantum computation?

One could argue that a quantum computation could be carried out with any high dimensional quantum system. E.g., instead of using many qubits one could use the energy eigenstates of a hydrogen atom. Because this would be a single system, there would be no entanglement. The problem is that in such systems the precision problem scales much worse and effectively destroys the (possible) advantage of quantum computing.

Therefore, the reason why entanglement is necessary for “real” quantum computing is because when we talk about the power of a computing model we talk about scaling issues, and precision is an important aspect of this.

So what went wrong when computer scientists thought that their formal classical computing models (in particular the Turing machine) captured the full
computational power of nature? One can argue that the reason is, that they didn’t think of the possibility of entangled states, thus that the state of a system cannot be described by describing the state of each part separately.

Maybe this is not the full truth: A different aspect of the question whether entanglement is necessary has arisen from thinking about quantum computation with highly mixed states, as in NMR (nuclear magnetic resonance) quantum computing experiments. If the state is close enough to the maximally mixed state, it necessarily is separable, thus it can be viewed as a probabilistic mixture of product states. Still it is not clear whether such an “unentangled quantum computation” can efficiently be simulated with probabilistic classical computation. The problem is how to efficiently arrive from the statistical ensemble before a quantum gate to the one after the quantum gate. Although none of the major quantum algorithms (Shor’s, Grover’s...) seem to work on such a quantum computer, it still may outperform conventional computers on some other problems.

3.4 Other quantum algorithms

There are a number of other quantum algorithms besides Shor’s, but many of them solve rather “academic” and somewhat unnatural problems which are primarily of interest to computer scientists. Often something has to be found out about “black box” functions, thus we can only evaluate the function, but we do not see how it is calculated. Grover’s, Simon’s and the Deutsch-Jozsa algorithm are all of this kind.

3.4.1 Grover’s “quantum searching” algorithm

The problem that Grover’s algorithm solves can be described like this: Say we have a classical algorithm which searches through a big number $N$ of cases (e.g. numbers) to find the single one which fulfills some criteria. Then Grover’s algorithm can find such an object in only about $\sqrt{N}$ instead of $N$ search steps. One could e.g. search through all possible next 10 moves of a chess player. However, usually there are better ways to find solutions than to simply search through all possibilities, so Grover’s algorithm may not be so useful.

Also one can solve any NP problem by searching through exponentially many cases, but then Grover’s algorithm would still take exponentially many steps, so it doesn’t provide a fast solution. Also it has been shown that such “simple” unstructured searches can’t be done faster than by Grover’s algorithm. Thus this “black box” approach to efficiently solve NP problems does not work.

3.4.2 communication complexity

Also many algorithms solve “communication complexity” problems, where e.g. the input is split between two distant places and the problem should be solved with as little communication as possible. There are such algorithms where
exponentially fewer qubits have to be sent that the number of bits that would have to be sent classically.

3.5 Are even more powerful computers possible?

First: can quantum computers even work? In a way, a working quantum computer would test quantum theory in a way that has not been done so far. In particular the “existence” of the large Hilbert spaces of systems with many parts would be tested. A quantum computer could more or less at random produce states in this space and check whether they behave as expected. Thus, the question is whether quantum theory really is correct or whether it is only an “effective” theory that works for not too high dimensional Hilbert spaces.

Probably quantum theory is correct. Very few physicists would doubt that at least the basic framework (superposition principle, etc.) is correct. Still, a working QC would provide a very strong test. Also theoretical quantum computing could be of interest to quantum physics. In a way, the complicated quantum formalism with the huge information content of states is only a tool to arrive at relatively simple results. If it could be shown that quantum computers are more powerful than classical ones, it would be clear that this formalism could not be simplified a lot while still producing the same results.

Could there be computers that are more powerful? So far, there is no indication of this in physics, but it is at least imaginable. E.g., it is known that more powerful quantum algorithms would exist if quantum theoretic time evolution were not exactly linear. This would allow to efficiently solve NP-complete problems and even more, but it seems improbable that such nonlinearities exist.

recommended literature:
Shor’s original 26 page article [1] is a good entry point.

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