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Quantum simulation: From basic principles to applications

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Envisioned by Richard Feynman in the early 1980s, quantum simulation has received dramatic impetus thanks to the development of a variety of platforms able to emulate a wide class of quantum Hamiltonians. During the past decade, most of the quantum simulators have implemented rather well-known models, hence permitting a direct comparison with theoretical calculations and a precise benchmarking of their reliability. The field has now reached a maturity such that one can address difficult problems, which cannot be solved efficiently using classical algorithms. These advances provide unprecedented opportunities to explore previously unreachable fields, test theoretical predictions, and inspire novel approaches.

This contribution is an elementary introduction to quantum simulation. We discuss the challenges, define both digital and analog quantum simulators, and list the demanding conditions they require. We also provide a brief account of the contributions gathered in the dossier on Quantum Simulation of the Comptes-Rendus de Physique of the French Academy of Sciences \cite{1,2,3,4,5,6}. The latter completes excellent reviews that appeared previously, see for instance Refs. \cite{7,8,9,10,11,12,13,14}.

Universal models and the role of simulations in many-body physics

Understanding the behavior of macroscopic quantum systems is a major challenge of modern physics. The basic laws of low-energy physics are by now quite well known at the microscopic, say atomic, scale. Conversely, many fundamental questions remain open, and even debated, about the collective dynamics at the macroscopic scale. By "macroscopic scale", here we mean systems made up of a huge number of constituents, or degrees of freedom, say $10^6$, $10^{23}$, or even more, as relevant in condensed matter physics or in astrophysics, for instance. Such huge systems cannot be treated exactly, be it at the classical level and, even worse, at the quantum level. Yet, it is the main outcome of the thermodynamic approach that the collective behavior of a macroscopic system can drastically differ from that of its elementary constituents. For instance, the elementary interactions between the \text{H}_2\text{O} molecules are fundamentally unchanged when a water bucket turns from the liquid phase to the solid phase at zero degree Celsius. Similarly, the interactions between the microscopic magnetic moments do not show any brutal change when a magnetic material gets magnetized underneath the Curie temperature. Hence, the dramatic effects observed in macroscopic systems are governed by large-scale instabilities, without obvious counterparts at the microscopic level. This observation takes a universal character, summarized in the celebrated motto "More is Different" \cite{15}. Such so-called emerging phenomena also appear in quantum systems, where new effects that are impossible in the classical world show up below some critical temperature or at zero temperature when some interaction parameter passes through a critical value. Celebrated examples include the superfluid $\Lambda$ transition in helium or super-conducting transitions and other metal-insulator transitions in electronic systems.

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Strikingly enough at first sight, while emerging phenomena only appear in very large scale macroscopic systems, their germs are contained in the mutual interactions of their elementary constituents. Hence, local two-body interactions are sufficient to explain a huge class of phase transitions, such as the liquid-solid and magnetic transitions mentioned above. The Boltzmann statistical approach proved particularly successful in describing this connection between the microscopic and macroscopic scales. To understand emerging phenomena on grounds as fundamental as possible, the programme is now well established: One tries to identify the basic microscopic terms that seem to be relevant and disregards all the other microscopic details. One then elaborates a model, as generic as possible, likely to reproduce the main experimental observations. The most usual examples are the Ising and Heisenberg models for magnetic transitions, or the Hubbard model for metal-insulator transitions [16,17,18]. Then all is left to do is to check that the phenomenon of interest indeed emerges from the dynamics of the simplified model. The realization of this programme is nothing but a simulation. It consists in building up a simplified system that mimics the main properties of a real system.

The difficulty of simulating generic quantum systems using classical computers
Unfortunately, severe problems usually appear at the last stage because limiting ourselves to a simple model does not guarantee that the solution is straightforward. In a few cases, exact solutions are known, a significant class of which is Bethe-ansatz integrable models. In some other cases, it is possible to make relevant approximations and build up tractable theories that yield accurate predictions. This is the case of mean field approximations, which work well to explain some superfluid transitions for instance. In most cases, however, exact or quasi-exact solutions are not known. One then traditionally resorts to numerical simulations. In this respect, the development of advanced approaches, such as Monte Carlo techniques, density functional theory, molecular dynamics, tensor-network approaches, and dynamical mean field theory, to name a few, have dramatically contributed to enlarging the class of models whose solution is known. However, even the most advanced numerical approaches have their own limitations and use approximations or representations that do not hold in all cases. Then, it is necessary to turn to exact computations. While it may be possible for some reasonably large classical systems, it is practically impossible for a quantum system.

Consider the most simple example put forward by Richard Feynman in Ref. [19] of a system made of $N$ spins 1/2. Since each spin can be either in the spin-up or spin-down state, there are $2^N$ possible configurations. A pure classical state is parametrized by $N$ binary numbers, the values of which, 0 or 1, represent the state of each spin. A computer with a memory of 100 Go $\sim 10^{12}$ bits can thus efficiently simulate about $10^{12}$ spins 1/2. Conversely, a generic pure quantum state is the coherent superposition of all the possible configurations. This requires $2^N$ C-coefficients and the same memory of 100 Go can only store the spin state of only $\log(10^{12})/\log(2) \sim 40$ particles. It is thus impossible to simulate the exact quantum state of more than a few tens of spins. More generally, it is practically impossible to store and manipulate the state of a macroscopic quantum system, owing to the exponential growth of its Hilbert space in the system’s physical size.

A careful reader might note that a similar issue occurs if one wants to simulate the full statistical distribution of the classical counterpart of the same spin systems. In this case, one would need to store the probabilities of each of the $2^N$ configurations, the number of which also grows exponentially with the number of constituents, $N$. However, this issue can be easily solved by using a stochastic algorithm, which amounts to introduce random jumps between the configurations. This is what Monte Carlo algorithms do for instance. Then, relevant quantities may be found by running the simulation a large, but not exponentially large, number of times, and averaging the results. As noticed by Feynman, this just simulates what Nature indeed does when we acquire experimental data [19].

The issue is much more serious in the quantum world, because even pure states cannot be simulated efficiently on truly large scales. Then, Feynman pointed out that the only reasonable thing to do is to make the computer quantum itself. Then, the register would naturally be exponentially large in the number of bits and one would be able to store an exponentially large number of coefficients [19]. Feynman also postulated that it should be possible to simulate the time evolution of any quantum system in a reasonable computer time. This statement was proven for local quantum Hamiltonians a few years later [20]. Seith Lloyd showed that the simulation can be performed with arbitrary precision in a computer time that grows at
most polynomially with the physical time, using Trotter-Suzuki’s decomposition of the many-body evolution
operator into sequences of local Hamiltonian evolutions. A machine, exploiting quantum resources to simulate
a given quantum model, is called a quantum simulator.

Up to now, we have focused on the dynamics of many-body systems at thermodynamic equilibrium.
However, similar questions and challenges appear when one considers the unitary time evolution of large
systems. Realizing quantum simulations in this case is actually even more important for several reasons. On
the one hand, we do not know (yet) a universal approach, which would be the out-of-equilibrium counter-
part of what is statistical physics for equilibrium thermodynamics. It follows that many questions are still
completely open, regarding for instance the spreading of information or the thermalization of these systems.
On the other hand, out-of-equilibrium quantum systems are likely to develop an entanglement entropy that
grows unbounded during the time evolution. It results that simulations using classical machines rapidly
reach their limits.

What is a quantum simulator?
Building up a quantum simulator is still a tour de force. Of course, the aim here is not to reproduce exactly
the initial system, for instance a complete material with all its microscopic details. Should we do this,
we would loop back to the initial problem and we would gain no information about the relevance of the
theoretical model. In the best case, we would address the specific initial problem and miss the basic physical
ingredients at the origin of the considered phenomenon. Instead, the aim here is to build up a clean system,
exactly governed by a basic model Hamiltonian. The latter is proposed in a theoretical context to reproduce
some physical phenomenon observed in a class of real systems. This is the basic idea of quantum simulation.

More precisely, a quantum simulator is a controlled device that allows us to (i) engineer a class of quantum
Hamiltonians exactly, (ii) control its dynamics, and (iii) make sufficiently precise measurements to consider
that the problem is solved. In practice, a quantum simulator is useful provided it is able to solve a hard
problem. In this context, one considers that a problem is "hard" when it cannot be solved, either analytically
or numerically with a classical computer in a time that grows at most as a power of the system’s size. This
feature is often retained as the main criterion for a quantum simulator. However, it cannot be considered
a perennial definition owing to continuous progress in many-body analytical and numerical approaches. In
fact, it is very hard to prove that a given quantum problem cannot be solved with a classical algorithm in
polynomial time. However, it is a common belief that a large amount of entanglement is a good working
criterion. Such a situation appears, for instance, in the vicinity of quantum phase transitions, in gapless
systems at equilibrium or in far-from-equilibrium systems where the entanglement may grow unbounded
during the time evolution. The applications of quantum simulation are potentially unlimited and range from
condensed-matter and high-energy physics to chemistry, biology, and cosmology, for instance [7,8,21].

To be more specific, one distinguishes two classes of quantum simulators:

Digital quantum simulators – A digital quantum simulator is a universal machine, fully reprogrammable to
simulate the thermodynamics or the real-time evolution of any quantum model. This is the kind of simulators
envisioned by Feynman and Lloyd. It was proven that, at least for local Hamiltonians, i.e., Hamiltonians con-
taining only finite-range interactions, such a universal quantum simulator can indeed be implemented [19,20].
Such a machine would exploit a fully reconfigurable register of qubits and a programmable sequence of log-
gical gates to realize the desired simulation. This is actually nothing but a quantum computer [22,23,24,25].
Hence, the distinction between a digital quantum simulator and a quantum computer mainly lies on the use
we make of it. While a "quantum computer" would be used to implement a variety of quantum algorithms,
the most celebrated example of which are the Shor factorization algorithm or the Grover search algorithm,
a "universal quantum simulator" would be dedicated to optimization problems, such as the determination
of the ground state of some Hamiltonian, or to its unitary, real-time, dynamics. In principle, building a
universal quantum simulator would thus be as difficult as building a quantum computer. In particular, it
would be sensitive to the same decoherence issues and require the implementation of error-correction codes.
However, real or imaginary time evolution by a local Hamiltonian usually requires much less resources than
generic quantum algorithms, and, in practice, are significantly more robust.
Analog quantum simulators – An alternative approach consists in building up a physical system from scratch to simulate each specific model. Assume one wants to determine the ground state or the time evolution of some Hamiltonian $\hat{H}$, say a spin-1/2 model. The idea behind analog quantum simulation is to create an ensemble of elements with two well-identified states, e.g., the two polarizations of a photon, the two internal states of an atom, or a quantum dot. These two states are used to represent the two spin states. Then isolate the system from its environment and engineer interactions between these two-state elements according to the Hamiltonian $\hat{H}$. This may be realized by coupling the system with a cavity in the case of photons or via laser fields in the case of atoms, for instance. In order to find the ground state, cool down the system; in order to determine its time evolution, prepare it in a well-defined initial state and let it run. Then, one realizes from scratch a system exactly governed by the desired model Hamiltonian and Nature simply works for us, with all its quantum properties. The last thing to be done is to measure the outcome.

Analog quantum simulation has pros and cons compared to digital quantum simulation. On the one hand, one needs to build up a new analog simulator for each studied model, while a unique digital simulator would be sufficient. On the other hand, the architecture of the analog simulator can be optimized for the considered problem, while that of a digital simulator is the result of a compromise between all possible cases. In practice, an analog quantum simulator is much easier to build than a digital quantum simulator and there are now many examples of successful implementations of the former, see for instance Ref. [21] and the contributions in this special issue [1,2,3,4,5,6].

Requirements and challenges
By analogy with standard numerical simulation, one may say that digital quantum simulation assumes that the hardware is available and focuses on the software part. Conversely, analog quantum simulation works directly on the former and the software part is built on the latter. In both cases, realizing an efficient quantum simulator requires to take up several major challenges that we summarize here (see also Ref. [9]):

(i) Build up. Create a quantum system that can be manipulated almost at will using external fields. It should be isolated from its environment to avoid decoherence issues. Depending on the problem at hand, the system can be made of bosons, fermions, spins, or mixtures of the latter.

(ii) Quantum engineering. Design the desired Hamiltonian with at least one adjustable relevant parameter. One should be able to tune the latter from a regime where the problem can be solved by other means to a regime where it cannot be solved easily. It allows benchmarking of the quantum simulator, similarly as it is done in traditional numerical work.

(iii) Initialization. Prepare the system in a well-defined state. It allows one to target the ground state using cooling techniques or to prepare some initial state to explore a specific trajectory in the system’s Hilbert space, for instance. The initial state is often a pure state, but one can also prepare a mixed state by interaction with a controlled bath. Furthermore, the bath may be used for simulating open quantum systems.

(iv) Detection. Measure relevant, local or non-local, observables that yield sufficient information to "solve" the problem with sufficient fidelity. Depending on the problem at hand, one may use destructive or non-destructive measurement techniques.

Since the goal is to use quantum simulators to solve problems than cannot be solved by other means, a major concern is of course the reliability of the used simulator. There are several ways to address this issue [26]. First, as mentioned above, one can make sufficient benchmarking of the simulator by addressing regimes where other solutions exists. Second, in the case of isolated systems, one can check that quantum coherence is maintained using an adiabaticity property: one back-propagates the system and checks the fidelity of the final state to the initial state. Third, it is crucial to solve a given problem on several simulators based on significantly different platforms. Should the results be consistent, one would consider that the solution is reliable. In this respect, dramatic progress has been realized starting by the early 2000s in a variety of fields, including:

(*) Ultracold quantum gases,
(*) Artificial ion crystals,
(*) Photonic systems, including polaritons in cavities,
(*) Superconducting circuits,
Magnetic insultors, Electronic spins in quantum dots, for instance. During the past decade, a number of quantum simulators have been demonstrated. So far, the results of most of them could be directly compared to theoretical calculations, which allowed extensive benchmarking. Now, some of the most advanced implementations are likely to address really hard problems, which cannot be solved efficiently using classical algorithms. To make this exciting perspective a reality, continued development of a variety of platforms, including those mentioned above and hopefully new ones, will be pivotal. It will allow us to address complementary questions as well as to compare results obtained by different approaches.

Contributions to this dossier
The dossier on Quantum simulation makes a point on recent advances of the field and discusses perspectives via a selection of contributions in various areas from ultracold atoms and quantum optics to statistical physics and condensed matter. Tarruell and myself review progress on the quantum simulation of the celebrated Hubbard model using ultracold Fermi gases. Aidelsburger et al.provide an introduction to novel approaches to engineer artificial gauge fields within a wide class of systems ranging from quantum optics to solid-state systems. Lebreuilly and Carusotto discuss realizations of strongly correlated quantum fluids of light in driven-dissipative photonic devices with applications to the generation of Mott insulator and fractional quantum Hall states of light. Le Hur et al.review advances in the study of real-time dynamics of impurity models and their realizations in quantum devices, including superconducting circuits, quantum electrical circuits, and ultracold-atom architectures. Bell et al.propose a novel platform based on superconducting quantum interference devices (SQUIDs) to emulate quantum phase transitions in one dimension, as well as perspectives to address non-integrable and disordered systems. Finally, Alet and Laflorencie discuss recent advances on many-body localization in isolated quantum systems and current experimental efforts to probing this physics.

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