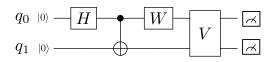
# Quantum phase estimation -

An example from chemistry

January 4, 2021

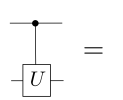
### prequisite: quantum circuits



- ▶ qubit: 2-level quantum system ( $|0\rangle$ ,  $|1\rangle$ ). quantum computer = n qubits.
- ▶ quantum circuit ≃ quantum program = list of quantum instructions/gates.
- ► The circuit above computes:

$$|\psi\rangle = V \cdot (W \otimes I) \cdot CNOT_{0 \to 1} \cdot (H \otimes I)|00\rangle$$

Note: "controlled-gates" means:



$q_0 = 0$ $q_1, 0 : q_1, 1$	$q_0 = 1$ $q_1, 0; q_1, 1$
$ \left(\begin{array}{cc} 1 & 0 \\ 0 & 1 \end{array}\right) $	(0)
(0)	$\overline{U}$

## Quantum simulation: general motivation

- ▶ Efficient numerical simulation quantum phenomena on classical machines seems hard [1].
- ► Therefore quantum machines might be able to perform classically-intractable computation<sup>1</sup>
- And in particular they should be good at simulating quantum phenomena (e.g simulating QCD, solving condensed-matter problems, computing molecular energies etc.)

 $\rightarrow$  But in practice, how would that work ? What **quantum circuits** would I run on my hypothetical quantum computers to solve these problems ?

<sup>&</sup>lt;sup>1</sup>basic motivation behind quantum computing research

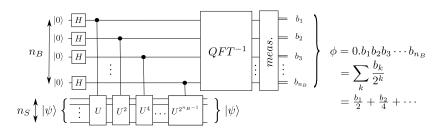
## Quantum Simulation: more precisely

Most of the time in the potential applications I mentioned (particle phycics, condensed matter, chemistry), the computation consists in diagonalizing Hamiltonians and in particular find their ground state energy.

You will see in these practical quantum programming sessions the two main frameworks for doing so on a quantum computer:

- phase estimation (this week) → the "classic guaranteed-speedup quantum algorithmics approach".
- variational quantum algorithms [2] (next week, in a slightly more "combinatorial optimization"-oriented context) → the "more recent, empirical, practical approach".

## This week's answer: quantum phase estimation



With 
$$|\psi\rangle$$
 eigenvector of  $U$  with eigenvalue  $e^{-2i\pi\phi}$ ,  $(U|\psi\rangle=e^{-2i\pi\phi}|\psi\rangle$ ,  $0\leq\phi<1$ )

The circuit above allows to compute the  $n_B$  most significant bits of  $\phi$  in a **binary decomposition**.

**Template** for *all* quantum algorithms with **exponential speedup**, both in quantum simulation and other application contexts.

## iterative quantum phase estimation: what we will use

for k in range  $(n_B, 0, -1)^2$ :

- 1. compute  $\phi_k = \sum_{l=k+1}^{n_B} \frac{b_l}{2^{l-k+1}}$
- 2. execute the following circuit:

$$|0\rangle - H - RZ(\phi_k) - H - meas = b_k$$

$$n_S \downarrow |\psi\rangle \left\{ - U^{2^{k-1}} \right\} |\psi\rangle$$

which yields  $b_{k}$ .

With 
$$H=rac{1}{\sqrt{2}}egin{pmatrix}1&1\\1&-1\end{pmatrix}$$
 and  $RZ(\phi_k)=egin{pmatrix}1&0\\0&e^{i\phi_k}\end{pmatrix}$ 

<sup>&</sup>lt;sup>2</sup>Python style

Recall that  $\phi=0.b_1\cdots b_{n_B}$  in binary decomposition,  $\phi_k=\sum_{l=k+1}^{n_B}\frac{b_l}{2^{l-k+1}}=0.0b_{k+1}\cdots b_{n_B}$  and  $U|\psi\rangle=e^{-2i\pi\phi}|\psi\rangle$ .

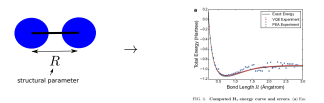
### **Evolution of the system:**

$$\begin{split} &|0\rangle|\psi\rangle \underset{H}{\rightarrow} \left(\frac{|0\rangle+|1\rangle}{\sqrt{2}}\right)|\psi\rangle = \left(\frac{|0\rangle|\psi\rangle+|1\rangle|\psi\rangle}{\sqrt{2}}\right) \\ &\underset{\bullet - U}{\rightarrow} \left(\frac{|0\rangle|\psi\rangle+e^{-2i\pi\cdot 2^{k-1}\phi}|1\rangle|\psi\rangle}{\sqrt{2}}\right) = \left(\frac{|0\rangle+e^{-2i\pi\cdot 0.b_k\cdots b_{n_B}}|1\rangle}{\sqrt{2}}\right)|\psi\rangle \\ &\underset{RZ(\phi_k)}{\rightarrow} \left(\frac{|0\rangle+e^{-2i\pi\cdot 0.b_k}|1\rangle}{\sqrt{2}}\right)|\psi\rangle = \left(\frac{|0\rangle+(-1)^{b_k}|1\rangle}{\sqrt{2}}\right)|\psi\rangle \underset{H}{\rightarrow} |b_k\rangle|\psi\rangle \end{split}$$

## Our example: molecular ground state energy calculation

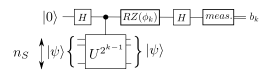
**General problem:** Given  $\mathcal{H}$ , electronic structure Hamiltonian of a molecule, we want to compute its ground state energy.

Actually, usually  ${\cal H}$  depends on some structural/geometric parameters. The purpose is then to compute a *ground state energy landscape* as a function of these parameters. May provide insight on reaction mechanisms  $^3$ .



**Our instance:**  $H_2$  and its dissociation curve. We will use data from [4], where they solve it on an actual small quantum chip.

<sup>&</sup>lt;sup>3</sup>Potential futuristic applications:  $CO_2$  capture [3],  $H_2O$  cracking...



We will therefore work with:

- $U = e^{-i\mathcal{H}dt}$ , for some value of dt to be chosen.
- $\blacktriangleright$   $|\psi\rangle$  the ground state of H, whose energy E we want to compute.
- we then have  $U|\psi\rangle=e^{-iEdt}|\psi\rangle=e^{-2i\pi\frac{Edt}{2\pi}}$

**Problem 1:** the algorithm only allows us to compute  $\phi = \frac{Edt}{2\pi} \mod [1] \ (\in [0,1])$ 

Following [5], we will use an interval guess  $[E_{min}, E_{max}]$  for E. Then  $\mathcal{H} \leftarrow \mathcal{H} - E_{min} \cdot I$  ensures E > 0 and choosing  $dt = \frac{2\pi}{E_{max}}$  ensures  $\frac{Edt}{2\pi} < 1$ 

## Other problems

**Problem 2** In general we do not have access to the ground state.

ightarrowuse an **ansatz**<sup>4</sup> with theoretical good overlap with ground state. As suggested in [4], we will use here the Hartree-Fock approxiamtion to our problem, namely  $|01\rangle = I \otimes X |00\rangle$ 

**Problem 3** How do we implement  $U^{2^k} = e^{-i\mathcal{H}2^k dt}$ ?

 $\rightarrow$  usually  $\mathcal{H}$  comes as a sum of simpler terms  $\sum_{l} \mathcal{H}_{l}$  (e.g Pauli products) for which  $e^{-i\mathcal{H}_{l}dt}$  is simple to implement.

→ and use **Trotter's formula** [5]

$$e^{-i\left(\sum_{l}\mathcal{H}_{l}\right)dt}\simeq\left(\prod_{l}e^{-rac{i\mathcal{H}_{l}dt}{p}}
ight)^{p}$$

with p integer, as large as possible.

<sup>&</sup>lt;sup>4</sup>easy-to-prepare approximation

- ▶ We will reproduce the curves from [4].
- ▶ In this example:
  - $ightharpoonup n_S = 2$
  - $H(R) = g_0I + g_1ZI + g_2IZ + g_3ZZ + g_4YY + g_5XX^5$
  - ► The coefficients are stored in hamiltonian\_data.json
- ► The purpose is to reproduce

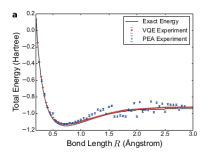


FIG. 3. Computed H<sub>2</sub> energy curve and errors. (a) Ene

<sup>&</sup>lt;sup>5</sup>This Hamiltonian is the output of (1) a conversion from an *electronic* structure Hamiltonian to a spin Hamiltonian expressed with Pauli matrices and (2) simplifications leveraging theoretical chemistry knowledge

## The programming framework we will use: myqlm

#### What you need to know:

- quantum circuits are first written as Program and then exported as circuit.
- ► A **job** is created from a circuit and sent to a **qpu** (a simulator)
- ► **AbstractGates** allow to define custom parameterized gates. They will come handy in programming Hamiltonian simulation.
- QRoutines allow to program arbitrary parametrized sub-circuits, which are then pluggable anywhere into a larger circuit.
- see the minimal notebook and https://myqlm.github.io/ for more details

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