

# Introduction to Quantum Computing

## Lecture 16: Variational Quantum Algorithms II

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- 1 Step 2: How to Measure the Energy/Cost
- 2 Step 3: The Ansatz (Family of Quantum States)
- 3 Step 4: Classical Optimisation & VQA Summary

## Step 2: How to Measure the Energy/Cost

## The Mathematical Task

Given a Hermitian matrix  $\mathcal{H}$  (typically called Hamiltonian), compute its smallest eigenvalue (called “ground state energy”)

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## Why is it relevant?

- Can solve it using VQA that is suitable for NISQ devices
- $k$ -local Hamiltonian problem is QMA-complete

# Previously in VQA: the four steps

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Express your desired problem as the ground state of a suitable qubit-Hamiltonian  $\mathcal{H}$

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## Step 2 Energy estimation (the only quantum part)

Given copies of a state  $|\psi\rangle$ , estimate its energy  $\langle\psi|\mathcal{H}|\psi\rangle$

## Step 3 Choice of Ansatz

A family of parametrised quantum states  $|\psi(\vec{\theta})\rangle$  where one of its members approximates best the ground state

## Step 4 Classical optimiser

A classical optimiser that finds the values  $\vec{\theta}^*$  that minimise the cost  $C(\vec{\theta}) := \langle\psi(\vec{\theta})|\mathcal{H}|\psi(\vec{\theta})\rangle$ , ie  $\vec{\theta}^* := \arg \min_{\vec{\theta}} C(\vec{\theta})$

## Step 2: Energy Estimation

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- An efficient, NISQ compatible, description to generate an  $n$ -qubit quantum state  $|\psi\rangle$
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**Task:** Estimate the energy  $E(\psi) := \langle\psi|\mathcal{H}|\psi\rangle$

- 1 Decompose the Hamiltonian to sum of Pauli observables
- 2 Generate multiple copies of  $|\psi\rangle$
- 3 Measure each Pauli suff times to get desired accuracy
- 4 Combine above to get an estimate for the energy  $E(\psi)$

# Decompose Hamiltonian to local Pauli observables

- **Pauli** observables can be measured **locally** and **easily**
- Frequently the Hamiltonian is already as sum of Pauli's

E.g. Ising Hamiltonians

$$\mathcal{H} = - \sum_{(i,j)} J_{ij} Z_i \otimes Z_j - \mu \sum_i h_i Z_i$$

- Other decompositions of the Hamiltonian to simple local observables can and have been considered (not here)

# Decompose Hamiltonian to local Pauli observables

- Any  $n$ -qubit Hermitian operator can be written as sum of products of Pauli matrices  $P_i \in \{I, X, Y, Z\}$   
(is orthonormal basis – Pauli observ:  $\{+1, -1\}$  eigenvalues)

$$\mathcal{H} = \sum c_{i_1, \dots, i_n} P_1^{i_1} \otimes \dots \otimes P_n^{i_n}$$

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- To compute coefficients  $c_{i_1, \dots, i_n}$  use the inner product

$$\langle A, B \rangle := \frac{\text{Tr}(A^\dagger B)}{2^n}$$

$$c_{i_1, \dots, i_n} = \langle P_1^{i_1} \otimes \dots \otimes P_n^{i_n}, \mathcal{H} \rangle$$

# Decompose Hamiltonian to local Pauli observables

## Example:

Decompose  $H = \frac{1}{\sqrt{2}} \begin{pmatrix} 1 & 1 \\ 1 & -1 \end{pmatrix}$  in Pauli's

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As expected

$$H = \frac{1}{\sqrt{2}} (X + Z) = \frac{1}{\sqrt{2}} \left( \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix} + \begin{pmatrix} 1 & 0 \\ 0 & -1 \end{pmatrix} \right)$$

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- Other example (check)

$$\wedge Z = \frac{1}{2} (I \otimes I + I \otimes Z + Z \otimes I - Z \otimes Z)$$

# Estimating the value of Pauli observables $\langle \psi | P | \psi \rangle$

- Prepare-and-measure the state  $N$  times
- Each outcome gives a value  $O_i \in \{+1, -1\}$
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- By Hoeffding (and Chernoff) inequalities we know:

$$Pr(|\bar{O} - \langle O \rangle| \geq \epsilon) \leq e^{-N\epsilon^2}$$

Probability that the true expectation differs by  $\epsilon$  or more from the measured one

- If we require that this probability is also bounded by  $\delta = e^{-N\epsilon^2}$  our confidence, we get above expression

# Implications of the accuracy

- Note that the **resources** required **depend on the problem**
- NP-**complete** problems (or even worse QMA-complete) **cannot be solved in poly-time** with a quantum computer
- Problems outside BQP have negligible “energy gap”, i.e. the ground state differs from the next eigenvalue (1st excited state) by a very small amount
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- To achieve **accuracy that distinguishes** between the two, one needs **super-polynomial repetitions**
- **High accuracy** is also required:
  - To move in a hyper-parametrised space  
(where gradients are negligible)
  - To overcome the effects of noise and determine truly the direction in the parameter space to move.

# Computing the Energy $E(\psi)$

- Using the Pauli decomposition we have:

$$E(\psi) = \langle \psi | \mathcal{H} | \psi \rangle = \sum c_{i_1, \dots, i_n} \langle \psi | P_1^{i_1} \otimes \dots \otimes P_n^{i_n} | \psi \rangle$$

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- The accuracy of the energy estimate depends on the accuracy of individual terms, and the number of terms in the sum
- We will use this value as the “cost” of the state  $|\psi\rangle$
- For the earlier example:  $H = \frac{1}{\sqrt{2}}(X + Z)$  we need to estimate two observables:

$$\langle \psi | X | \psi \rangle = O_1 ; \langle \psi | Z | \psi \rangle = O_2$$

Resulting to  $E(\psi) = \frac{1}{\sqrt{2}}(O_1 + O_2)$

## Step 3: The Ansatz (Family of Quantum States)

# Ansatz: The space we optimise

- To solve the Hamiltonian problem we need to find the quantum state that has **minimum energy** from *all* the states of the **Hilbert space**
- This is **infeasible**. Instead, we select a family of (parametrised) quantum states, and we hope that one **member of the family approx. well the ground state.**

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- Can view the ansatz as a **family of parametrised quantum circuits**:  $|\psi(\vec{\theta})\rangle = U(\vec{\theta})|0\rangle$
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- Two approaches: (i) Hardware Efficient, (ii) Problem Specific

## (i) Hardware Efficient Ansatz

- Generate a family of states that spans evenly the Hilbert space
- Needs to be able to produce high entanglement
- Should choose circuits that are **easy to implement with a given NISQ device**
- Generic Ansatz that **can be used for *any* Hamiltonian problem**

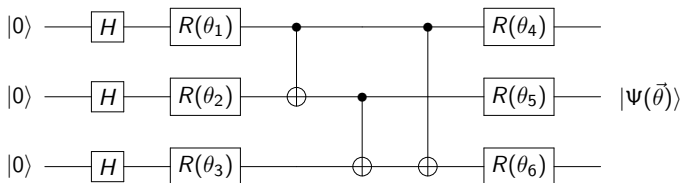
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- **Typical Ansatz:**
  - 1 A number of layers repeating the same circuit with different parameters
  - 2 Single-qubit rotations parametrised by the rotation angle
  - 3 Entangling gates (non-commuting with the rotations)  
Hardware architecture determines entanglement topology (all-to-all Vs nearest-neighbour)



## An Example: 3-qubit, 1-layer, all-to-all entanglement

$$\vec{\theta} := (\theta_1, \theta_2, \theta_3, \theta_4, \theta_5, \theta_6)$$



## (ii) Problem Specific Ansatz

- Generate a family of states that **uses the problem's Hamiltonian**
- Does not span evenly the Hilbert space – hopefully is **more dense around** the region we expect to have **the ground state**
- Not designed with the hardware in mind

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- Many families exist (e.g. unitary coupled cluster, adiabatic, etc).
- We give an important type suitable for optimisation problems:

**Quantum Approximate Optimisation Algorithm (QAOA)**

# Quantum Approximate Optimisation Algorithm

- Let  $H_C$  be the **problem's Hamiltonian** of Ising type (only Pauli-Z, up to quadratic terms)

Recall Max-Cut:  $H_C = \sum_{(i,j) \in E} Z_i \otimes Z_j$

- Let  $H_B = \sum_i X_i$  be the **“mixer” Hamiltonian**

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- Let  $H_B = \sum_i X_i$  be the “**mixer**” **Hamiltonian**
- A 1-layer QAOA ansatz is given by

$$e^{-i\beta H_B} e^{-i\gamma H_C} H^{\otimes n} |0\rangle^{\otimes n}$$

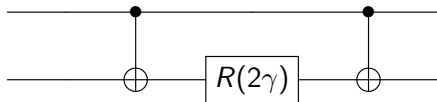
- A single layer has only **two parameters**  $(\beta, \gamma)$ , irrespective of the number of qubits/width of computation

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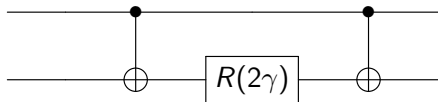
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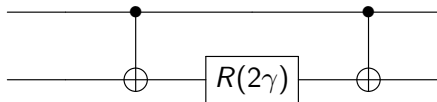
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- **More layers repeat the above** with fresh parameters  $(\beta_2, \gamma_2)$

## Step 4: Classical Optimisation & VQA Summary

# Moving Through Parameter Space

## Given:

- Ansatz: set of q-states parametrised by classical parameters  $\vec{\theta}$
- How to compute the “cost-function”  $E(\vec{\theta}) = \langle \Psi(\vec{\theta}) | \mathcal{H} | \Psi(\vec{\theta}) \rangle$

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## Examples of Techniques Used:

- Gradient Descent
- Monte Carlo - based
- Nelder-Mead method
- COBYLA (Constr. Optimiz. by Linear Approximation)
- Any deterministic or stochastic global optimisation method

# Gradient Descent

- Trick to compute the gradient (generalises).  
Parametrised gates:  $U_P = e^{-i\theta P}$  with  $P$  a Pauli  
(recall  $P^2 = I$ )

$$U_P(\theta) = I \cos(\theta) - iP \sin(\theta) ; \frac{\partial}{\partial \theta} U_P(\theta) = -iPe^{-i\theta P}$$

leading to

$$\frac{\partial}{\partial \theta} E(\theta) = E(\theta + \frac{\pi}{4}) - E(\theta - \frac{\pi}{4})$$

Note: This difference is NOT infinitesimal!

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- Move towards negative direction of the gradient:  
 $\vec{\theta}_{i+1} = \vec{\theta}_i - \gamma \nabla E(\vec{\theta}_i) ; \nabla E(\vec{\theta}_i) = \left( \partial_{\theta_1} E(\vec{\theta}), \partial_{\theta_2} E(\vec{\theta}), \partial_{\theta_3} E(\vec{\theta}) \right)$
- Can find **local minima** (not global)



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- Initial Guess  $\vec{\theta}_0$  and compute  $E(\vec{\theta}_0)$
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- Else, with probability  $p_r = 1 - e^{-\beta(E(\vec{\theta}') - E(\vec{\theta}_0))}$ , **reject guess** (make a fresh guess starting again from  $\vec{\theta}_0$ )
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## Observations

- May keep moving with greater new Energy (esc local min)
- Keeping probability reduces with Energy difference
- $\beta$  is “inverse temperature”. Can increase value with iteration steps  $\beta(i)$  (so it stabilises in time)

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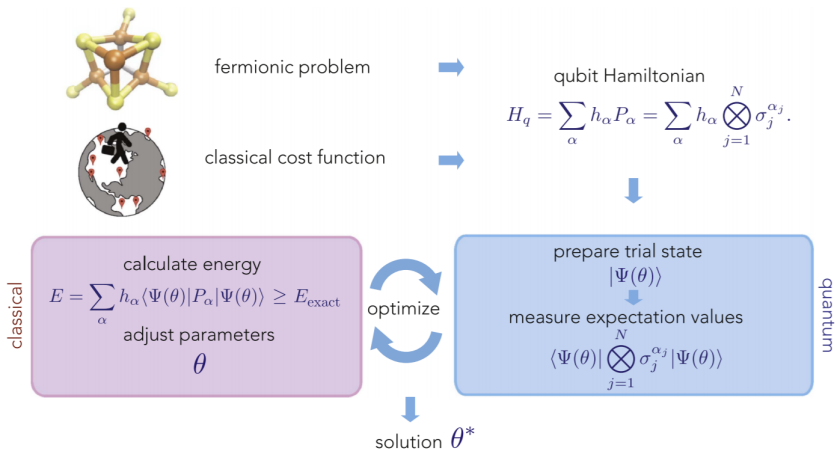
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- **Overall limitations (heuristics):**

- Not exact complexity
- No guarantee for obtaining solution



# VQA Pictorially



Taken from: Nikolaj Moll et al 2018 Quantum Sci. Technol. **3** 030503

# VQA: Summary

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- **Quantum Part** needs small coherence time, works without QECC and can be optimised for given hardware:

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## **Best candidate for quantum speed-up at NISQ devices**

- 1 Map problem to ground state energy of some Hamiltonian  $\mathcal{H}$  (classical)
- 2 Can generate (efficiently) a family of states  $|\Psi(\vec{\theta})\rangle$  (quantum)
- 3 Can compute expectation value  $E_{\vec{\theta}} = \langle \Psi(\vec{\theta}) | \mathcal{H} | \Psi(\vec{\theta}) \rangle$  with local Pauli measurement for any guess  $\vec{\theta}$  (quantum)

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- 4 Comparing  $E(\vec{\theta})$  of existing points, evaluate a new guess  $\vec{\theta}'$  using standard techniques.  
Feedback to step 2 to evaluate  $E(\vec{\theta}')$  (classical)

## Variational Quantum Algorithms Reviews

- 1 *Variational quantum algorithms*, Cerezo, Marco, et al. Nature Reviews Physics (2021): 1-20.
- 2 *Noisy intermediate-scale quantum (NISQ) algorithms*, Bharti, Kishor, et al. arXiv preprint arXiv:2101.08448 (2021).
- 3 *Quantum optimization using variational algorithms on near-term quantum devices*, Moll, Nikolaj, et al. Quantum Science and Technology 3.3 (2018): 030503.