

Hybrid classical/quantum algorithms

Veiko Palge, Ville Kotovirta and Franz Fuchs

Variational Quantum Eigensolver (VQE)

- VQE was the first variational algorithm

Problem:

Find the ground energy of a
Hamiltonian that describes a molecule
or solid state system

- Solution: use variational principle on QC

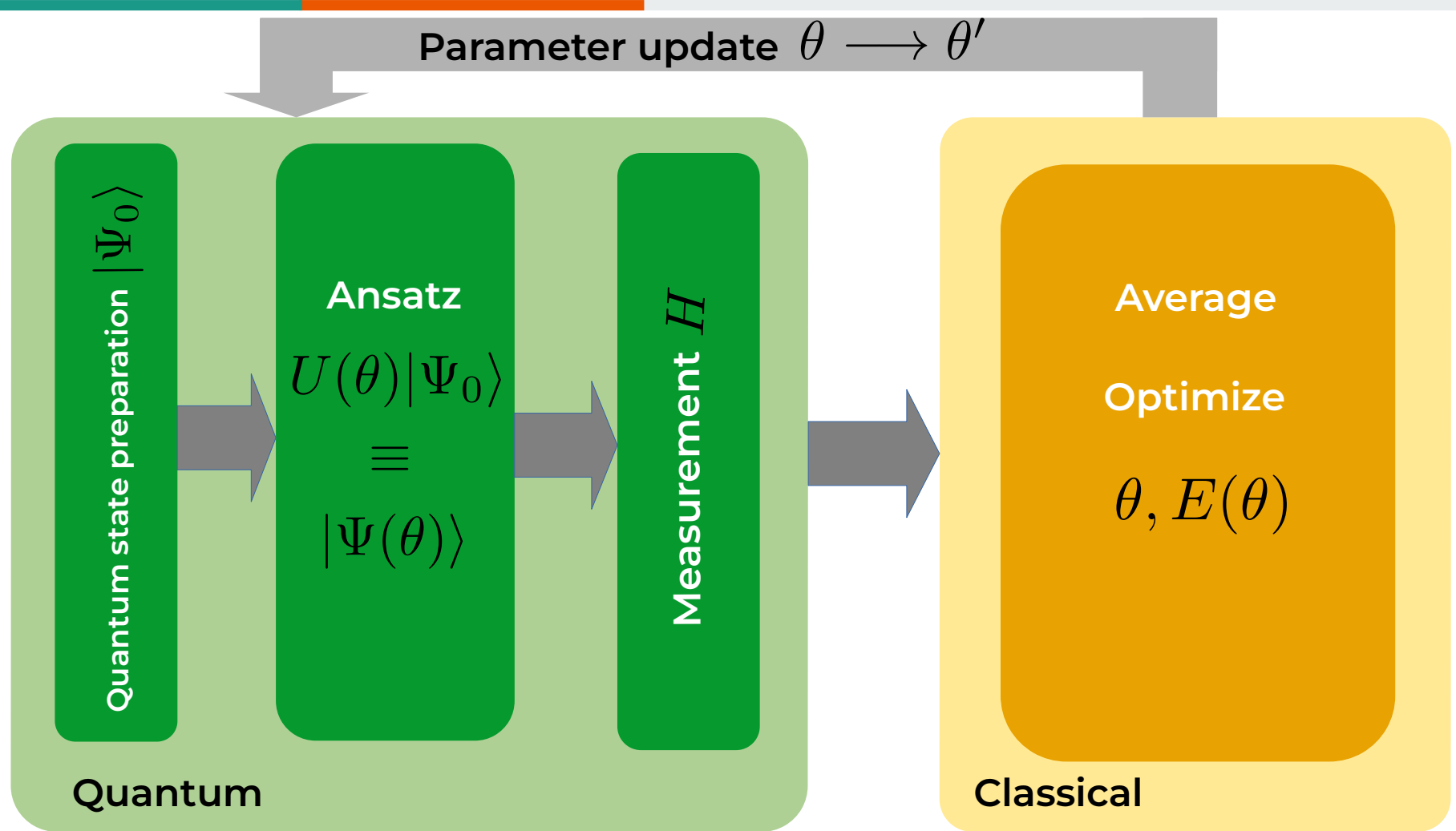
Variational principle

- Hamiltonian H describes the electronic structure of a molecule or solid state system
- Find a trial state $|\Psi(\theta)\rangle$ with parameters θ and compute $\langle H \rangle$
- Vary parameters θ to find the lowest value of $\langle H \rangle$

$$E_0 \leq \langle \Psi(\theta) | H | \Psi(\theta) \rangle \equiv \langle H \rangle$$

Variational principle

- It is extremely powerful and easy to use
- Even if $|\Psi(\theta)\rangle$ has no relation to the actual ground state, one often gets accurate values for the ground state E_0
- Disadvantage: we never know how close we are to the target value — only know that we have upper bound



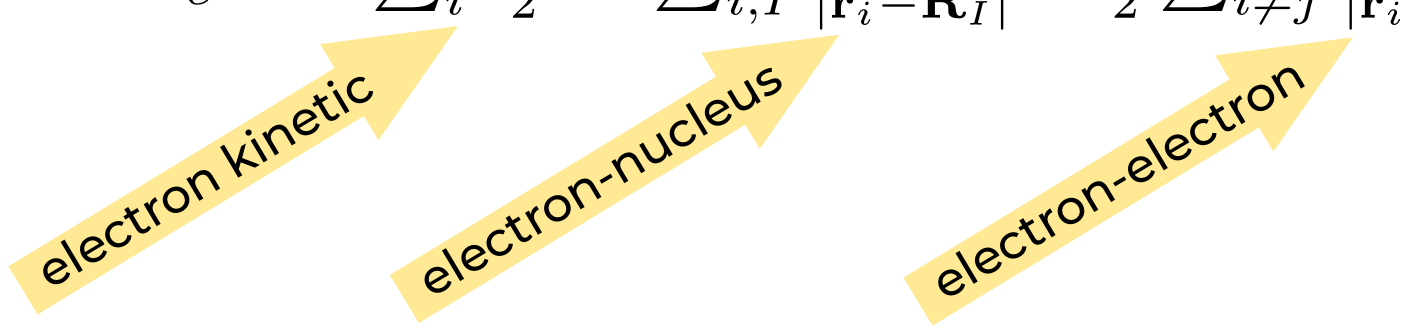
Electronic structure

- Molecule characterized by Hamiltonian

$$H_{\text{mol}} = H_{\text{nucl}} + H_e$$

- Only interested in the electronic structure

$$H_e = - \sum_i \frac{\nabla_i^2}{2} - \sum_{i,I} \frac{Z_I}{|\mathbf{r}_i - \mathbf{R}_I|} + \frac{1}{2} \sum_{i \neq j} \frac{1}{|\mathbf{r}_i - \mathbf{r}_j|}$$



Electronic structure

- Express in second quantization

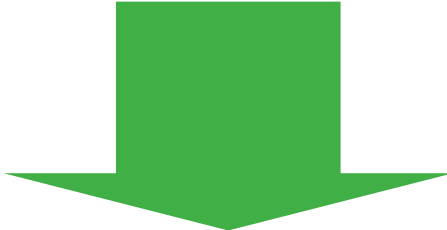
$$H = \sum_{p,q} h_{p,q} a_p^\dagger a_q + \frac{1}{2} \sum_{p,q,r,s} a_p^\dagger a_q^\dagger a_r a_s$$

- $h_{p,q}$ kinetic energy of electrons and Coulomb interaction with nuclei
- $h_{p,q,r,s}$ electron-electron Coulomb repulsion

Map Hamiltonian to quantum computer

- Map **fermionic** Hamiltonian to **qubit** Hamiltonian

$$H = \sum_{p,q} h_{p,q} a_p^\dagger a_q + \frac{1}{2} \sum_{p,q,r,s} a_p^\dagger a_q^\dagger a_r a_s$$

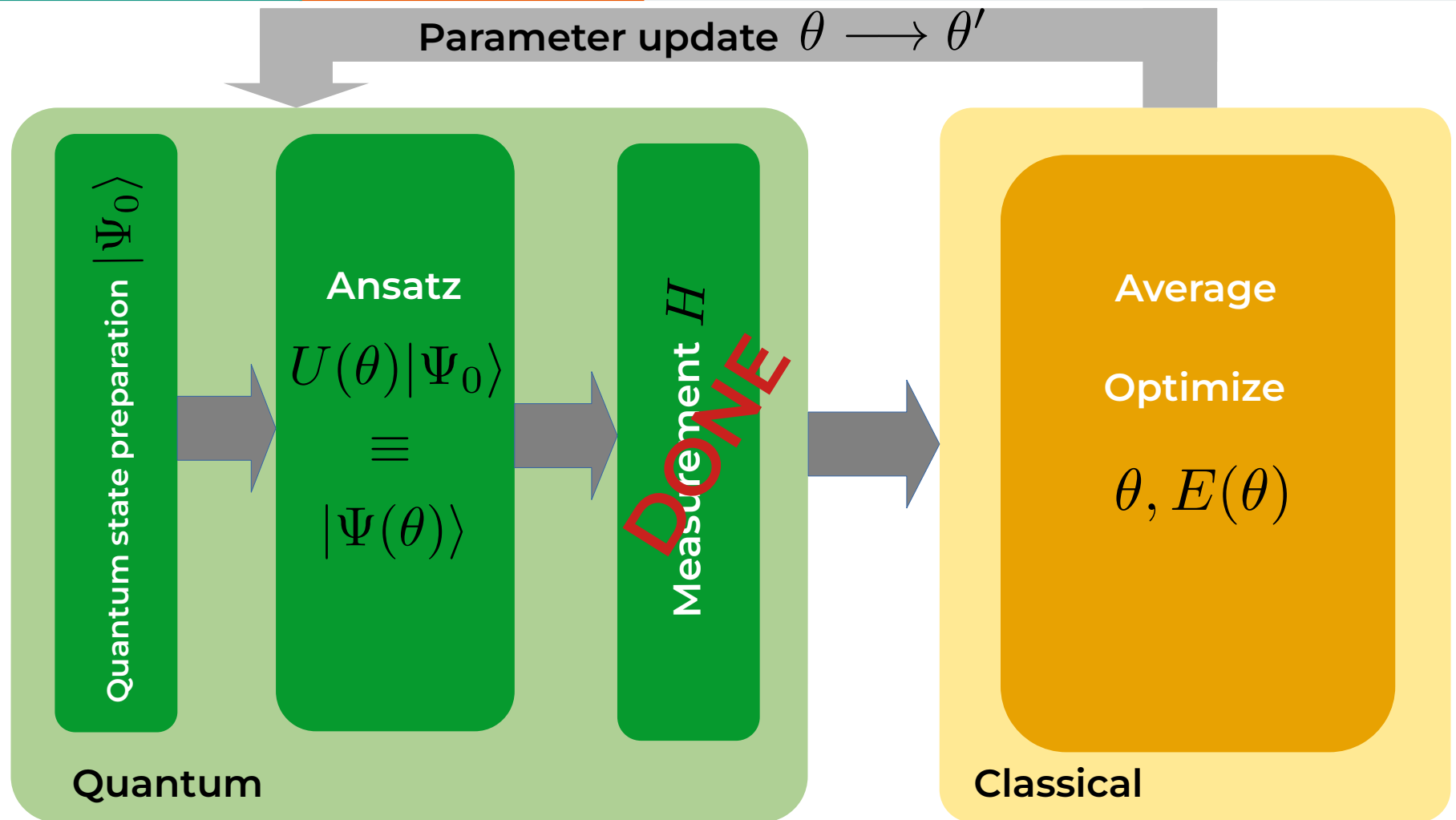


$$H = \sum_j h_j \Pi \sigma_i^j$$

Map Hamiltonian to quantum computer

- Qubit Hamiltonian: sum of Pauli strings
- Example: 4 qubit Hamiltonian for H_2

$$\begin{aligned} H = & h_0I + h_1Z_0 + h_2Z_1 + h_3Z_2 + h_4Z_3 \\ & + h_5Z_0Z_1 + h_6Z_0Z_2 + h_7Z_1Z_2 + h_8Z_0Z_3 + h_9Z_1Z_3 \\ & + h_{10}Z_2Z_3 + h_{11}Y_0Y_1X_2X_3 + h_{12}X_0Y_1Y_2X_3 \\ & + h_{13}Y_0X_1X_2Y_3 + h_{14}X_0X_1Y_2Y_3 \end{aligned}$$



Ansatz circuit

- Inspiration from computational chemistry:
Coupled Cluster (CC) method

$$|\Psi_{CC}\rangle = e^T |\Psi_0\rangle$$

$$T = \sum_i T_i \quad T_1 = \sum_{i,\alpha} t_\alpha^i a_\alpha^\dagger a_i \quad T_2 = \sum_{i,j,\alpha,\beta} t_{\alpha\beta}^{ij} a_\alpha^\dagger a_\beta^\dagger a_i a_j$$

- Unitary Couple Cluster (UCC) ansatz on quantum computers

Ansatz circuit

- UCC is the unitary version of CC

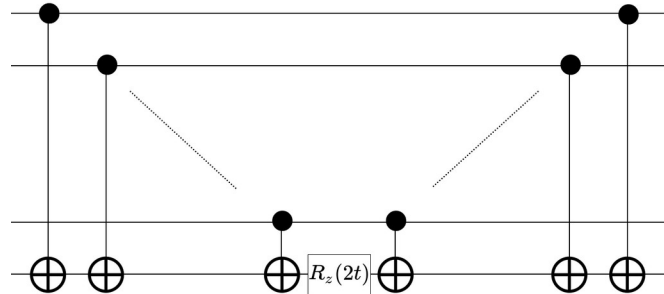
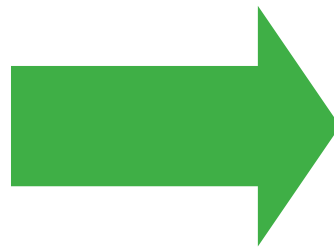
$$|\Psi_{UCC}\rangle = e^{T-T^\dagger} |\Psi_0\rangle$$

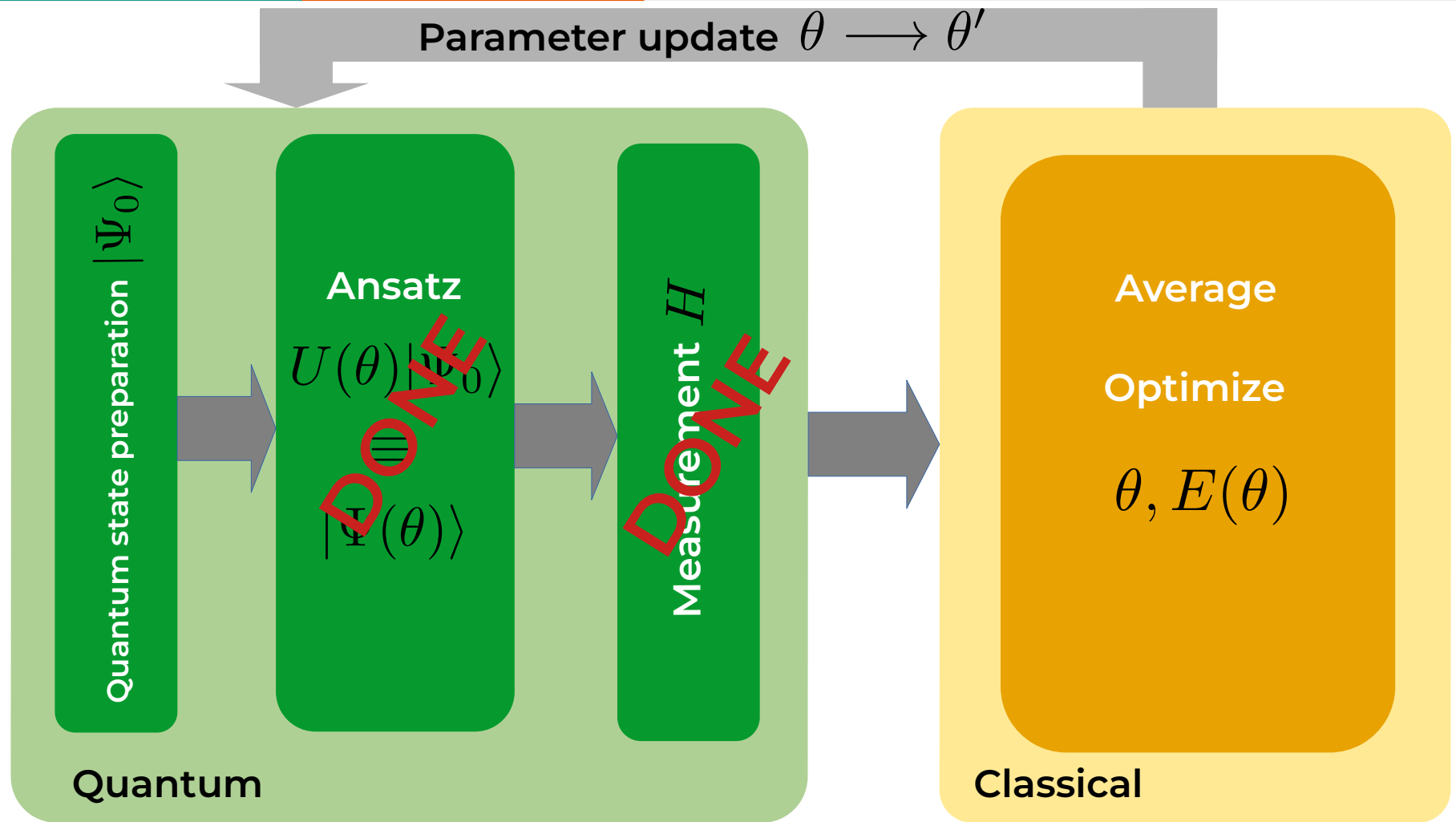
- Unitary e^{T-T^\dagger} can be implemented on a quantum computer
- Usually truncated to Single and Double excitations: UCCSD

Ansatz circuit

- Implement UCC using Trotter-Suzuki decomposition of cluster operator e^{T-T^\dagger}
- Map exponents to qubits
- Qubit exponents realized as circuits

$$e^{-i\theta Z_1 X_2 \dots Y_n}$$



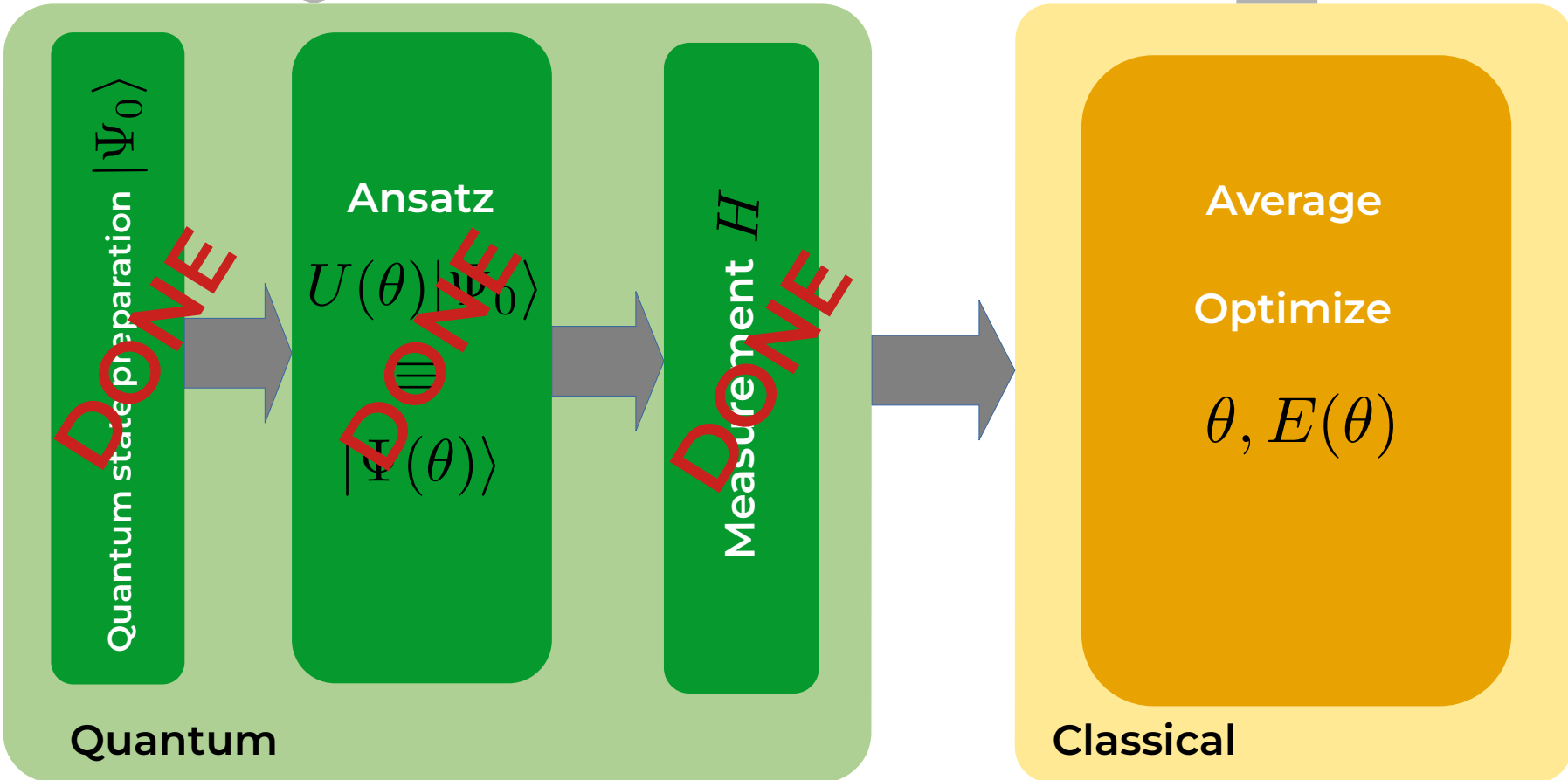


Initial state

- Classically compute the Hartree-Fock (HF) state
- Map to qubit state using Jordan-Wigner mapping, and prepare on the quantum computer
- Example: H₂ with 4 qubits

$$|\Psi_0^{\text{HF}}\rangle = |0011\rangle$$

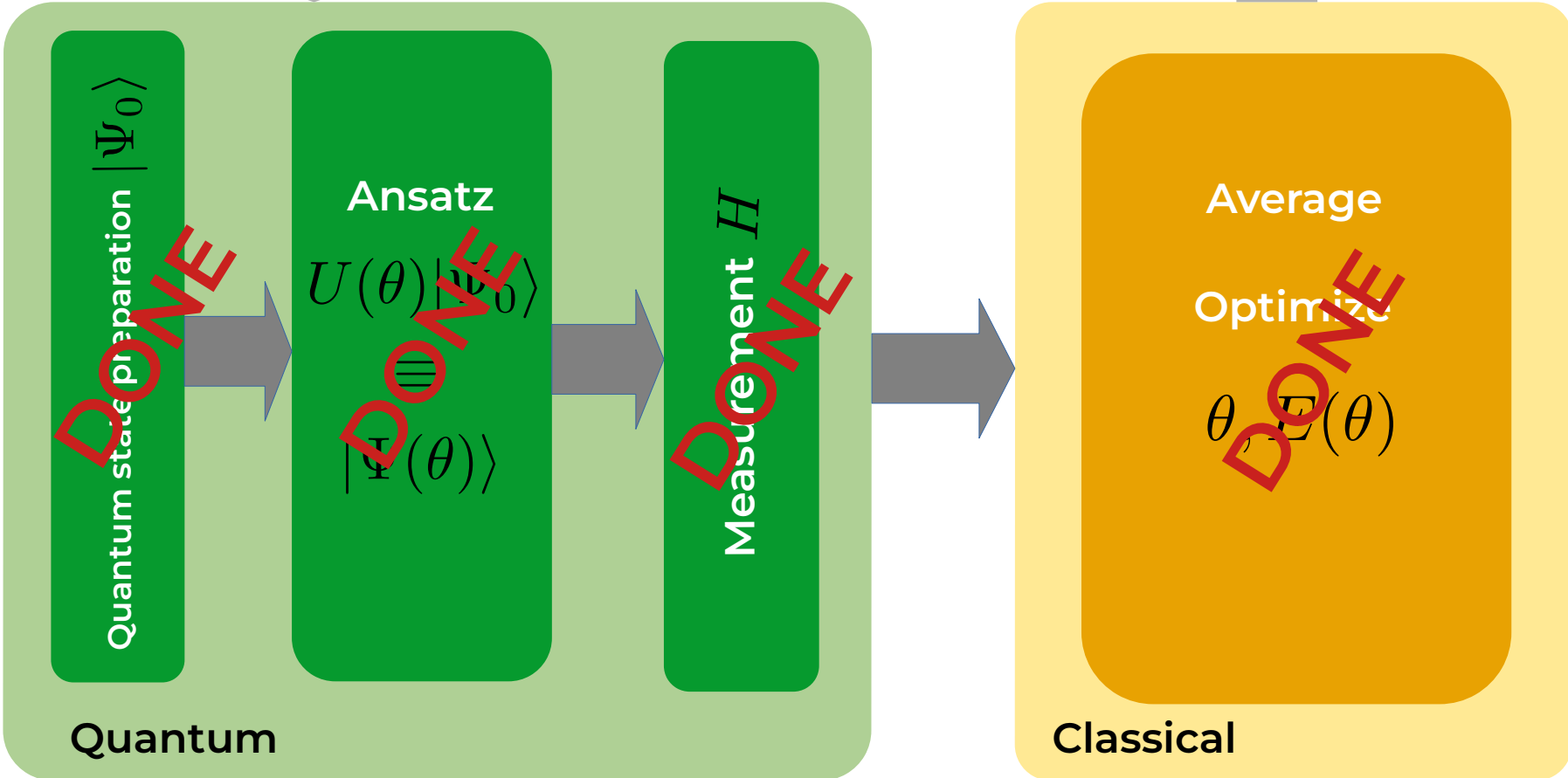
Parameter update $\theta \rightarrow \theta'$



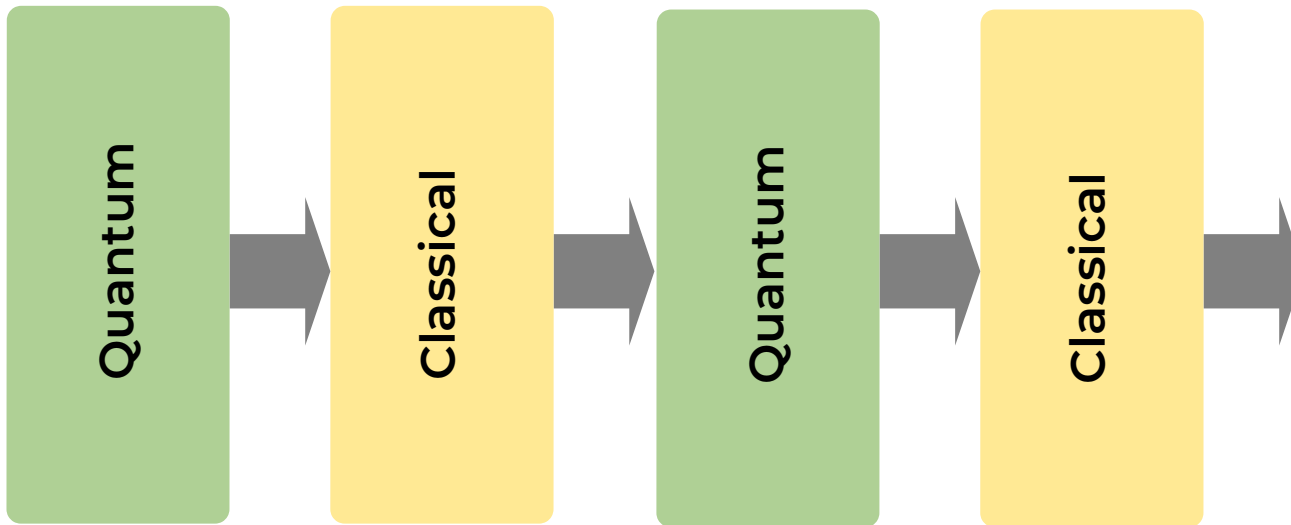
Optimization

- Optimization algorithm (Nelder-Mead, COBYLA, TNC, etc)
- Gradient based algorithms
- Combinations of different approaches

Parameter update $\theta \rightarrow \theta'$



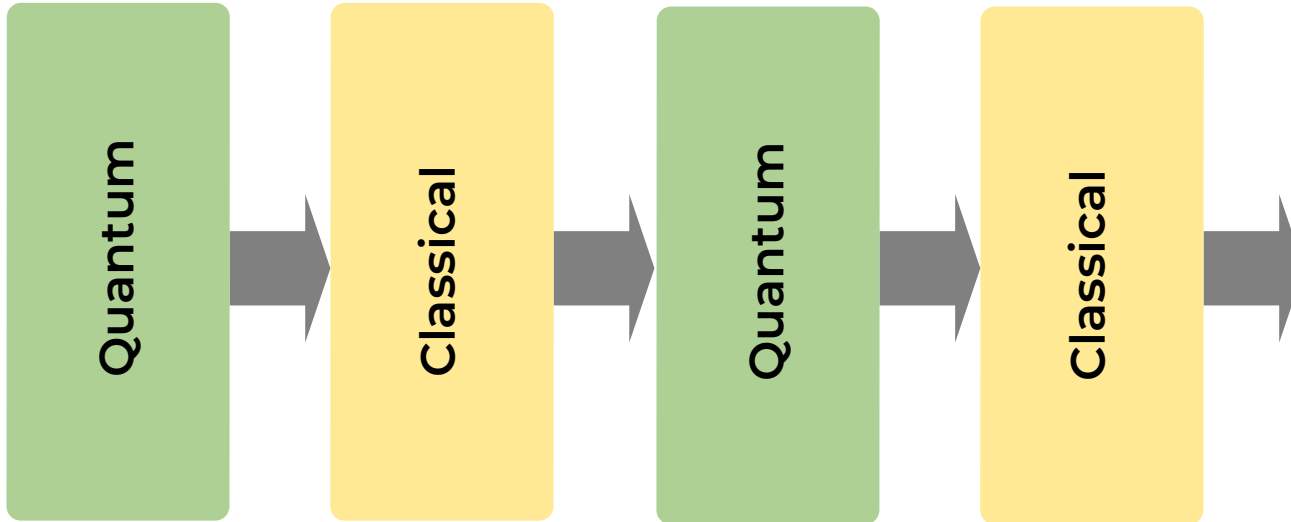
Running VQE



VQE advantage:

after each run the quantum circuit is reset, so
"short" coherence time enough

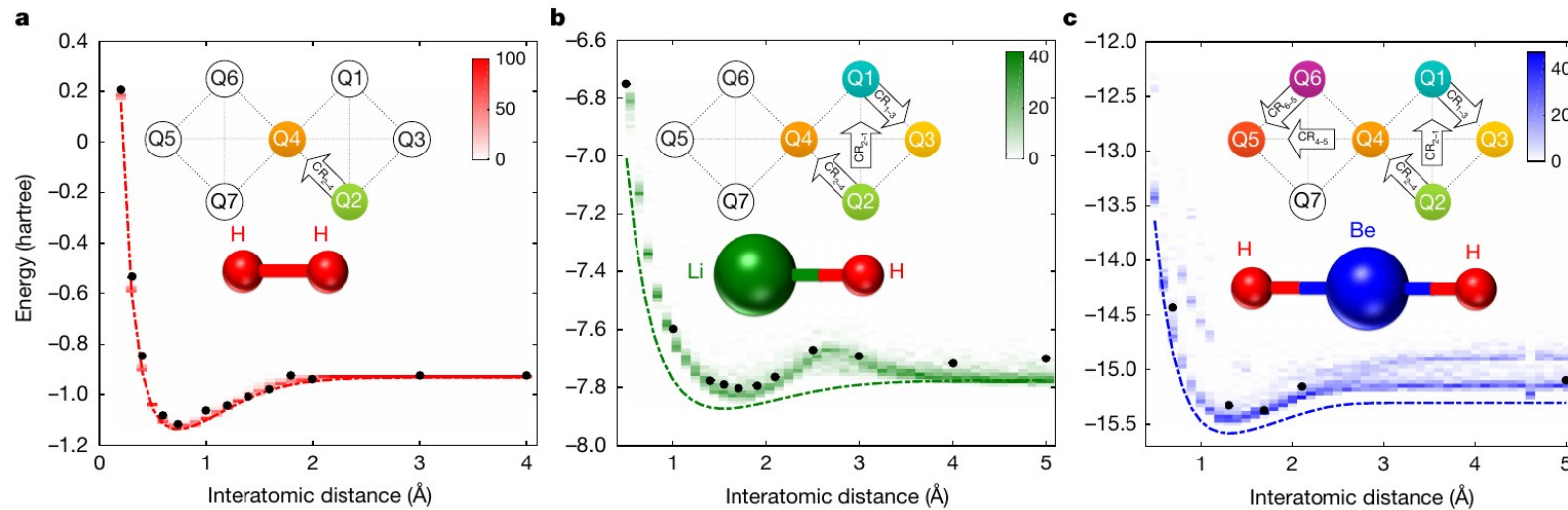
Running VQE



VQE challenge:
is quantum coherence time long enough?

Results: small molecules

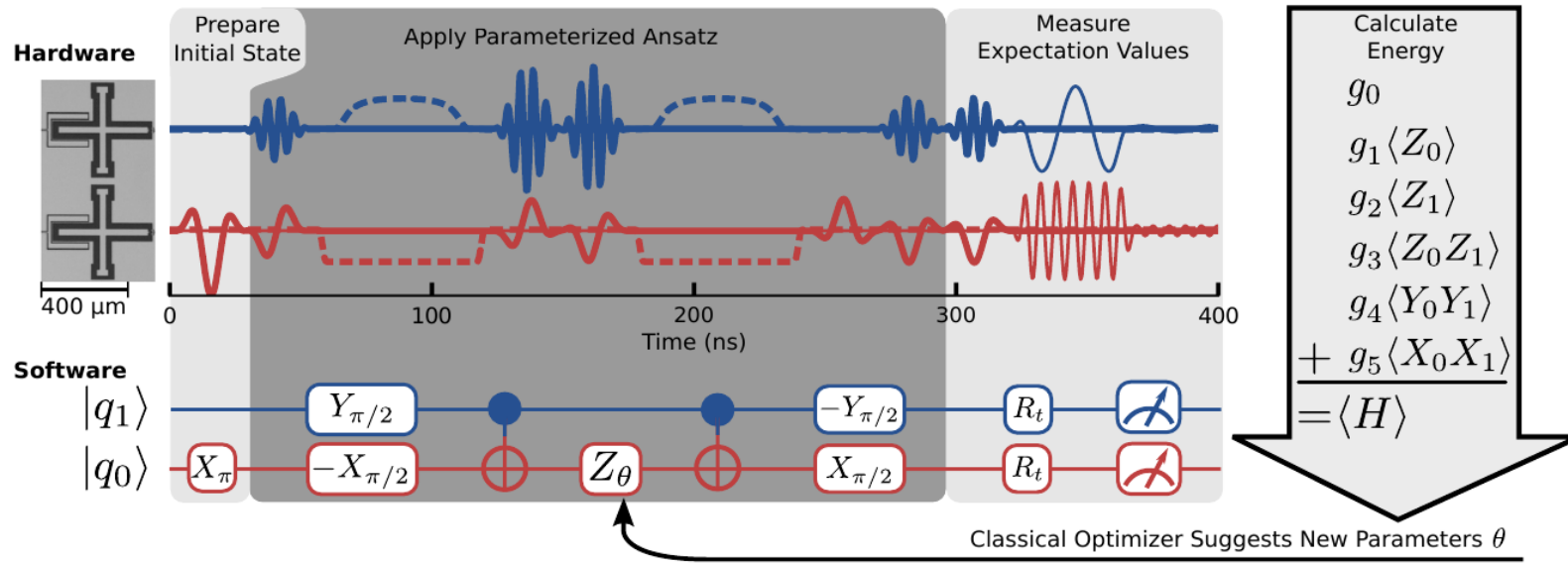
- H₂, LiH, BeH₂ implemented on quantum hardware



Kandala et al 2017, Nature 549, 242-246

H2 implementation

- H2 implemented on quantum hardware



O'Malley et al 2015, 'Scalable Quantum Simulation of Molecular Energies',
arXiv:1512.06860v2 [quant-ph]