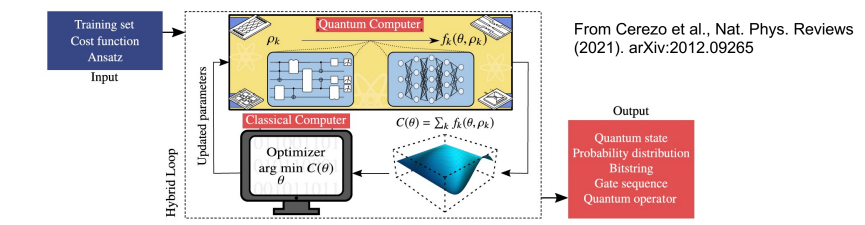
IOWA STATE UNIVERSITY

Department of Physics and Astronomy

Variational quantum algorithms 1

Peter P. Orth (Iowa State University & Ames Laboratory)

SQMS/GGI Summer School, Florence, Italy, 26 July 2022

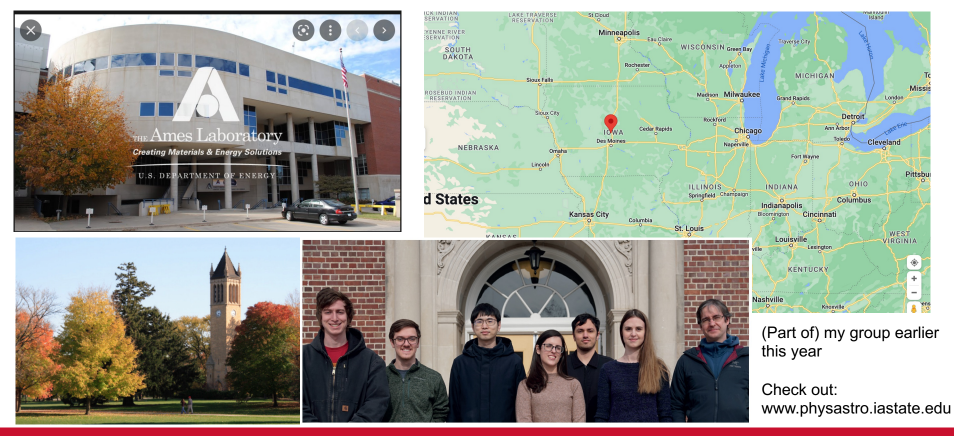




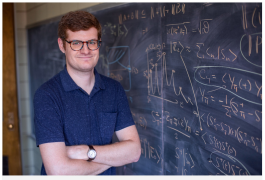




Ames National Lab & Iowa State University



Quantum Computing at Ames National Laboratory & Iowa State



Thomas ladecola studies how states of matter emerge from collections of atoms and subatomic particles and recently contributed theoretical work and data analysis to a paper published by the journal Nature. Larger photo. Photo by Christopher Gannon.

Srimoyee Sen

Tom ladecola (Nonequilibrium dynamics, topological field theory, quantum computing)

Srimoyee Sen (QCD, quantum field theory)

- > Ames Lab participates in two national quantum initiative centers of the Department of Energy (SQMS & C2QA)
- Strong collaborations across fields and between theory and experiment at Ames National Lab

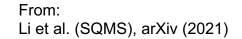


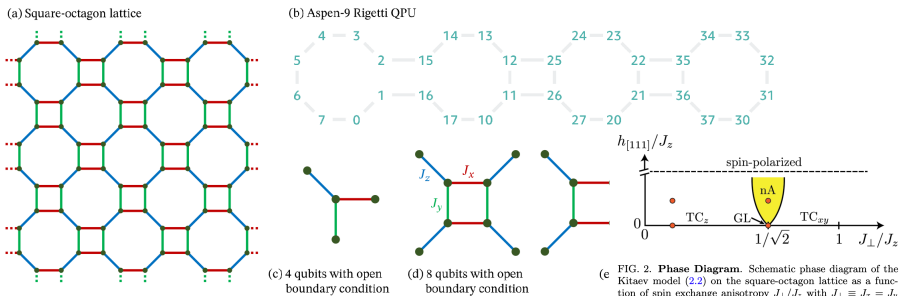
Yongxin Yao (materials simulations, quantum computing)

Joint group meeting of my group and the groups of Prof. Flint and Prof. ladecola



VQE for Kitaev spin model





Kitaev model on square-octagon lattice matches Rigetti's QPU geometry. No SWAP gates needed as connectivities match.

FIG. 2. Phase Diagram. Schematic phase diagram of the Kitaev model (2.2) on the square-octagon lattice as a function of spin exchange anisotropy J_{\perp}/J_z with $J_{\perp} \equiv J_x = J_y$ and magnetic field in [111] direction $h_{[111]}$. It includes gapped toric code phases (TC_z , TC_{xy}) that are stable with respect to small fields, the gapless line (GL) at $J_{\perp}/J_z = 1/\sqrt{2}$ and a phase with non-Abelian (nA) Majorana excitations that emerges in field above the gapless line. At large magnetic fields the system enters a spin-polarized paramagnetic phase. The red circles denote the different, representative model parameter points that are studied in our benchmark simulations.

VQE ansatz

From: Li et al. (SQMS), arXiv (2021)

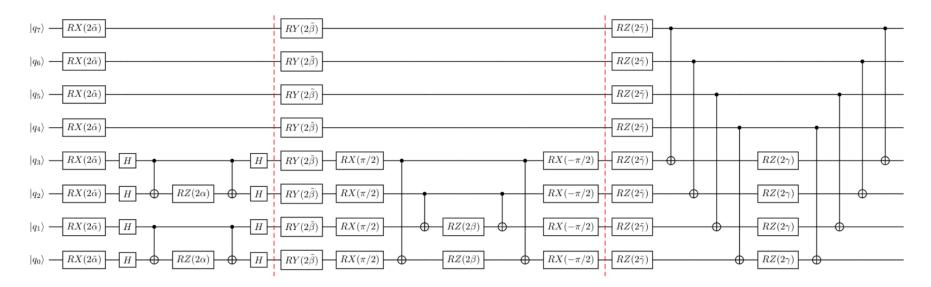
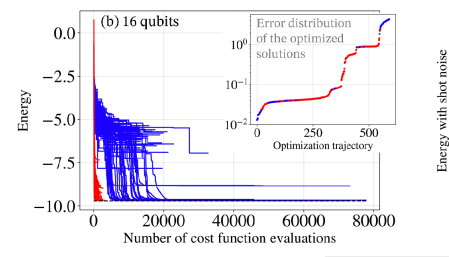


FIG. 3. **HVA with one layer on eight qubits**. The Hamiltonian Variational Ansatz (HVA) with one layer on eight qubits, split into commuting blocks. The first block corresponds to the operation $e^{-i\tilde{\alpha}\sum_{q}X_{q}} e^{-i\alpha\sum_{(i,j)\in X-\text{links}}X_{i}X_{j}}$, the second to $e^{-i\tilde{\beta}\sum_{q}Y_{q}} e^{-i\beta\sum_{(i,j)\in Y-\text{links}}Y_{i}Y_{j}}$, and the third to $e^{-i\tilde{\gamma}\sum_{q}Z_{q}} e^{-i\gamma\sum_{(i,j)\in Z-\text{links}}Z_{i}Z_{j}}$. For the circuit shown here, we used $X-\text{links} = \{(q_{0}, q_{1}), (q_{2}, q_{3})\}, Y-\text{links} = \{(q_{0}, q_{3}), (q_{1}, q_{2})\}$, and $Z-\text{links} = \{(q_{0}, q_{4}), (q_{1}, q_{5}), (q_{2}, q_{6}), (q_{3}, q_{7})\}$.

Noiseless and noisy (shots) VQE optimization

From: Li et al. (SQMS), arXiv (2021)

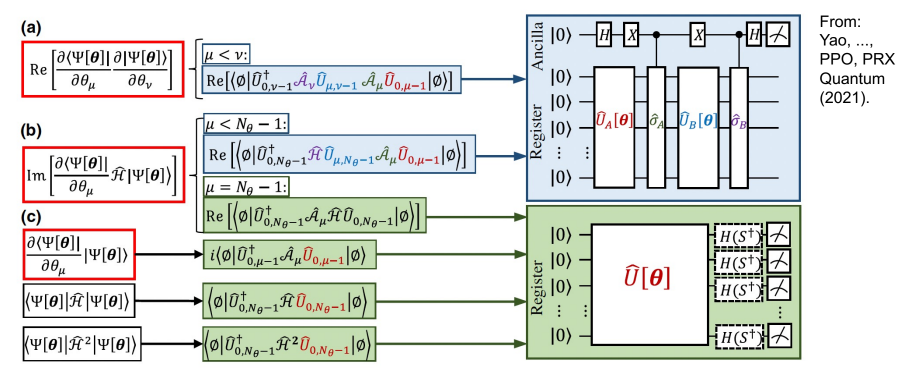


0	BOBYQA-noisy -2 Optimized energy i CMA-ES -2 Distribution (noiseless)
-2	-4 0 250 500 Optimization trajectory
-4	
(0 10000 20000 30000 40000 50000 Number of cost function evaluations

Optimizer	Error (noiseless)	Measured deviation	Cost function evaluations
BFGS, 501 initial values	0.45069	0.42052	mean: 747, max: 1994
BOBYQA, 501 initial values	0.27485	0.21843	mean: 471, max: 610
BOBYQA-noisy, 501 initial values	0.07989	-0.00453	mean: 3532, max: 4004
CMA-ES	0.02416	-0.06462	37570
CMA-ES, 80 initial values	0.01610	-0.07125	mean: 21042, max: 52000
Dual annealing	0.04534	-0.01631	60101
SPSA	0.00612	0.00879	100000 (cutoff)

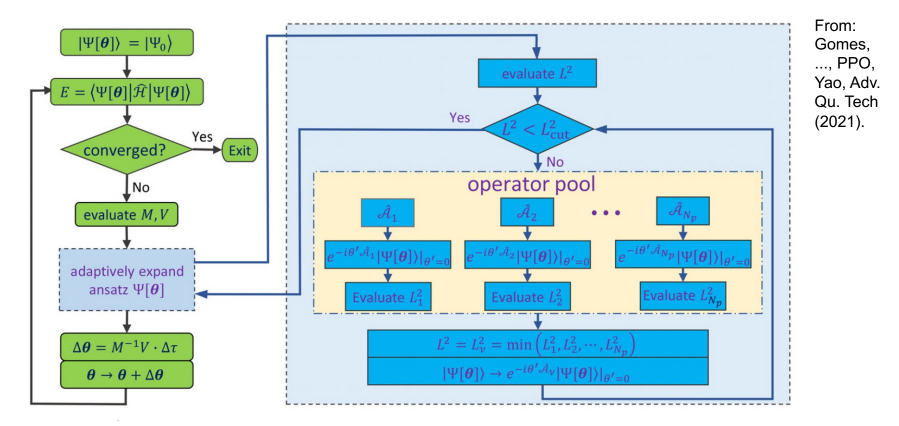
Noiseless: 16 qubits Noisy: 8 qubits (8000 shots)

Quantum circuits for measurement of Re[G] and energy gradient



These circuits are for real-time evolution. For imaginary time evolution one applies an additional S^\dag gate after the first Hadamard gate (and removes the X gates) in order to measure the imaginary part in the Hadamard test.

Adaptive variational quantum imaginary time evolution



Adaptive variational quantum imaginary time evolution

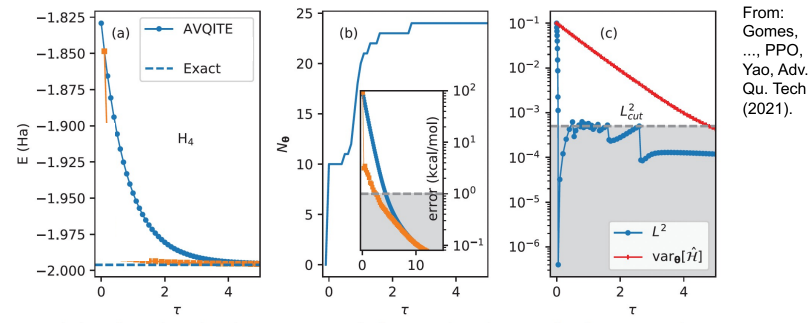
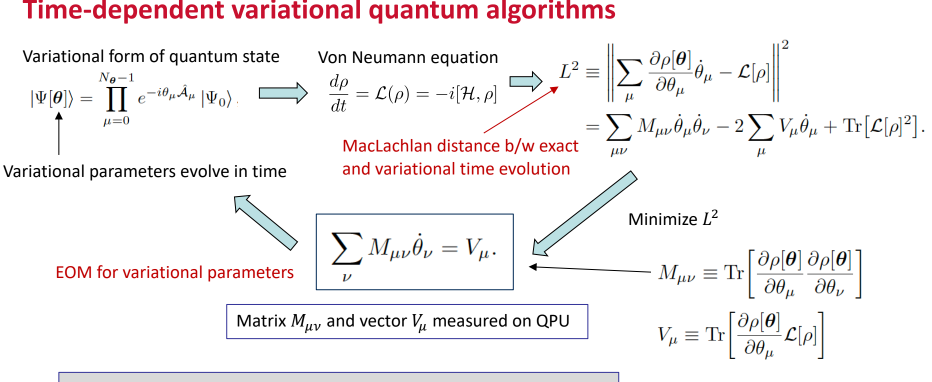


Figure 2. AVQITE calculation for H₄ chain. Along the imaginary time path of $\tau = N\Delta\tau$, the total energies of H₄ from AVQITE and quantum Lanczos calculations are shown in panel (a), the number of variational parameters N_{θ} in (b), and the McLachlan distance L^2 and energy variance $var_{\theta}[\hat{H}]$ in (c). The exact full configuration interaction result is shown as the dashed line in (a) for reference. The total energy errors $E - E_{Exact}$ are plotted in the inset of (b), with the shaded area denoting chemical accuracy. The chosen threshold $L_{cut}^2 = 5 \times 10^{-4}$ is indicated by a dashed line in (c).

Time-dependent variational quantum algorithms

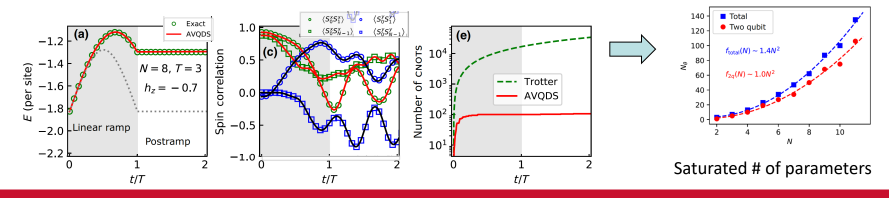


Only so good as the ansatz can follow the dynamics How to select an efficient yet flexible variational ansatz?

[1] Li, Benjamin, Endo, Yuan (2019).

Application I: continuous quench in integrable spin chain

- > Linear quench of anisotropic XY chain in transverse magnetic field $\hat{\mathcal{H}} = -J \sum_{i=0}^{N-2} \left[(1+\gamma) \hat{X}_i \hat{X}_{i+1} + (1-\gamma) \hat{Y}_i \hat{Y}_{i+1} \right] + h_z \sum_{i=0}^{N-1} \hat{Z}_i \text{ with } \gamma(t) = 1 - \frac{2t}{T}$
- > AVQDS follows exact solution during and after quench, shown for N = 8
- > Circuit depth saturates at 100 CNOTs << Trotter circuit depth 10^4 CNOTs
- > Can simulate system with gate depth independent of time t recan simulate to arbitrary times!



8 (-2, 0) γ (-0.7, 1) (-0.7, 1) (-0.7, -1)(-0.7, -1)

Application II: sudden quench in nonintegrable spin chain

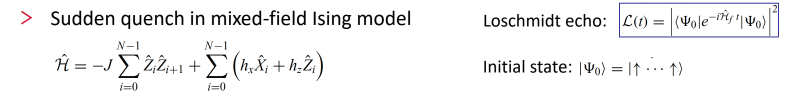
> Sudden quench in mixed-field Ising model $\hat{\mathcal{H}} = -J \sum_{i=0}^{N-1} \hat{Z}_i \hat{Z}_{i+1} + \sum_{i=0}^{N-1} \left(h_x \hat{X}_i + h_z \hat{Z}_i \right)$

Loschmidt echo:
$$\mathcal{L}(t) = \left| \langle \Psi_0 | e^{-i\hat{\mathcal{H}}_f t} | \Psi_0 \rangle \right|^2$$

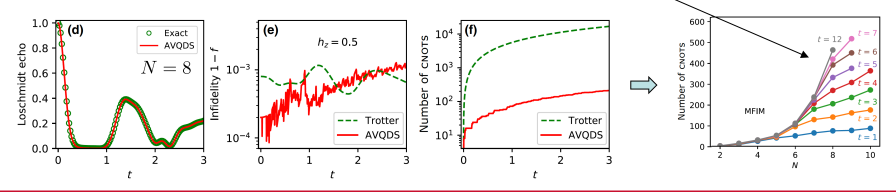
Initial state: $|\Psi_0\rangle = |\uparrow \cdots \uparrow\rangle$

- > AVQDS follows exact solution, shown for $N \leq 10$
- > Circuit depth two orders of magnitude smaller than Trotter circuit depth
- Saturation timescale AVQDS circuit depth saturates at system size N dependent time scale 1.0 600 (a) (d) (e) 104 Exact Number of CNOTS 0.8 Poschmidt echo 0.4 0.4 0.2 $h_7 = 0.5$ AVQDS SLOND 400 N = 8nfidelity 1 – f N = 810³ 10-3 Ъ 300 Number N = 710² 200 Trotter Trotter N = 6100 AVQDS 10¹ AVQDS N = 5 10^{-4} 0.0 10 2 1 2

Application II: sudden quench in nonintegrable spin chain



- > Circuit depth two orders of magnitude smaller than Trotter circuit depth
- Saturated AVQDS circuit depth scales exponentially with system size N
- > # measurements is main bottleneck of algorithm



VQE-X: variational quantum eigensolver for highly excited states

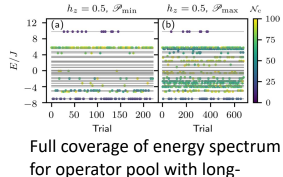


Adaptive variational quantum eigensolver for excited states

> Long-time dynamics of observable determined by diagonal ensemble of excited states

$$\langle \mathcal{O}(t) \rangle = \sum_{n,m} c_n c_m^* e^{i(E_m - E_n)t} \langle m | \mathcal{O} | n \rangle \xrightarrow{t \to \infty} \langle \mathcal{O}(t) \rangle \approx \sum_n |c_n|^2 \langle n | \mathcal{O} | n \rangle$$

- > Adaptive variational quantum eigensolver to prepare highly excited states (VQE-X)
 - > Minimize energy variance (instead of energy): $C(|\psi(\theta)\rangle = \langle \psi(\theta)|H^2|\psi(\theta)\rangle \langle \psi(\theta)|H|\psi(\theta)\rangle^2$



range Pauli strings

⁹ Nontrivial pool

dependence

$$\mathscr{P}_{\min} = \{Y_i\}_{i=1}^N \cup \{Y_iZ_{i+1}\}_{i=1}^N$$
$$\mathscr{P}_{\max} = \{Y_i\}_{i=1}^N \cup \{Y_iZ_j\}_{i,j=1}^N \cup \{Y_iX_j\}_{i,j=1}^N$$

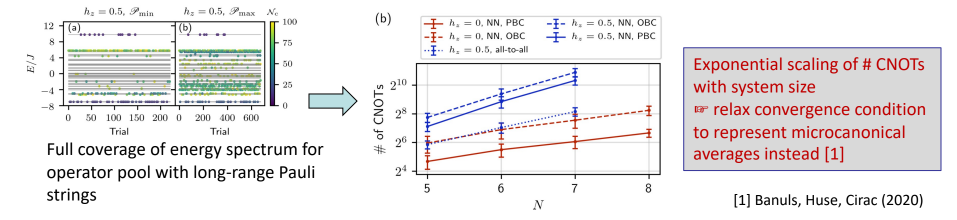
[1] Zhang, Gomes, Yao, PPO, Iadecola, PRB **104**, 075159 (2021). Can investigate properties of volume law highly excited states

Adaptive variational quantum eigensolver for excited states

> Long-time dynamics of observable determined by diagonal ensemble of excited states

$$\langle \mathcal{O}(t) \rangle = \sum_{n,m} c_n c_m^* e^{i(E_m - E_n)t} \langle m | \mathcal{O} | n \rangle \xrightarrow{t \to \infty} \langle \mathcal{O}(t) \rangle \approx \sum_n |c_n|^2 \langle n | \mathcal{O} | n \rangle$$

- > Adaptive variational quantum eigensolver to prepare highly excited states (VQE-X)
 - > Minimize energy variance (instead of energy): $C(|\psi(\theta)\rangle = \langle \psi(\theta)|H^2|\psi(\theta)\rangle \langle \psi(\theta)|H|\psi(\theta)\rangle^2$



Summary and outlook

- > Adaptive variational quantum dynamics simulation (AVQDS) framework
 - > Orders of magnitude shallower circuits than Trotter simulations
 - Linear growth of # CNOTs with time initially, then saturation at e^N number of gates
 - > Allows simulation out to arbitrary times for small systems
- > Adaptive variational quantum eigensolver for highly excited states (adaptive VQE-X)
 - > Minimize energy variance allows preparing arbitrary excited states
 - > Generate volume-law states requires exponentially many parameters
 - > May be alleviated in simulations of microcanonical averages

References:

- N. Gomes et al., arXiv:2102.01544 (2021)
- Y. Yao et al., PRX Quantum 2, 030307 (2021)
- F. Zhang et al., PRB 104, 075159 (2021)

Thank you for your attention!

