Hybrid Quantum Annealing via Molecular Dynamics

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How to solve combinatorial optimization problems?

• Quantum annealing (QA) is expected to be a powerful solution

Kadowaki-Nishimori (1998)

Rapid progress in QA hardware

but remains to be NISQ device

N-qubit $\rightarrow -\sqrt{N}$ -qubit for fully-connected problem



- 2007: 16 qubit
 2011: 128 qubit
 2013: 512 qubit
 2015: 1152 qubit
 2017: 2048 qubit
 2020: 5640 qubit
- Effective use of classical computer is also crucial in NISQ-era

Quantum-classical hybrid solver

iterative use of QA solver from outer classical solver

find persistent vars by multiple use of classical solver

Chancellor ('17), Okada et al. ('19)

Karimi-Rosenberg ('17)

New idea awaited!

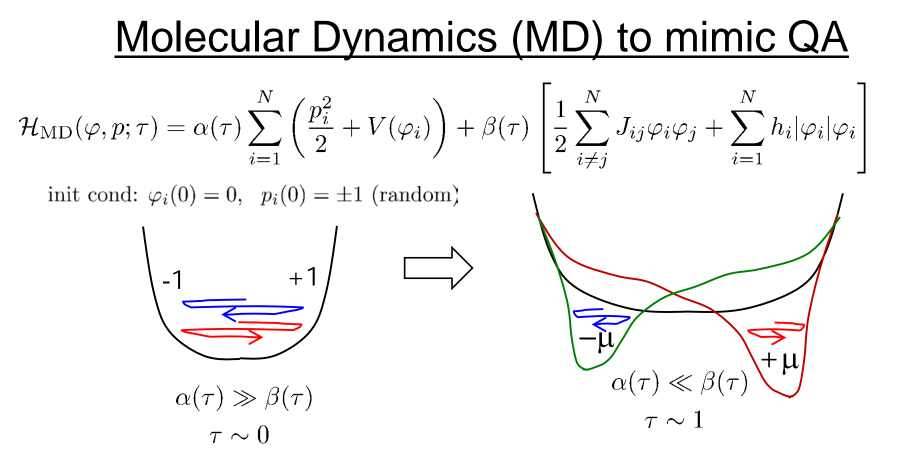
Classical-only solvers also in active development

Coherent Ising machine (optical), CMOS annealing machine (SA/MA), Digital Annealer (SA), ... Simulated bifurcation machine (Molecular-Dynamics (MD)), ...

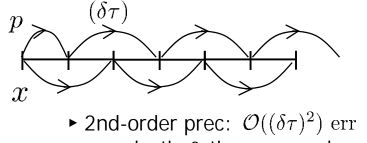
In particular, MD is beneficial in scalability/performance but sys err introduced

Molecular Dynamics (MD) to mimic QA

Quantum annealing (QA)



We solve MD by leap-frog method



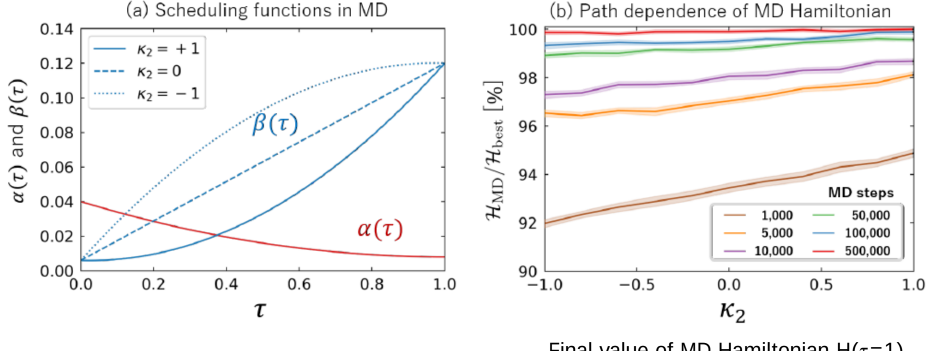
 symplectic & time-reversal (if Hamiltonian does not have explicit τ-dep) Our MD could be better than other MD solvers w/ artificial +/- barrier

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(It can be improved by 
higher-prec solver)
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Adiabaticity in MD

In the case of QA (AQC), it is guaranteed that the system is the ground state thanks to the quantum adiabatic theorem

In the case of MD, there is no such guarantee, so we make numerical check

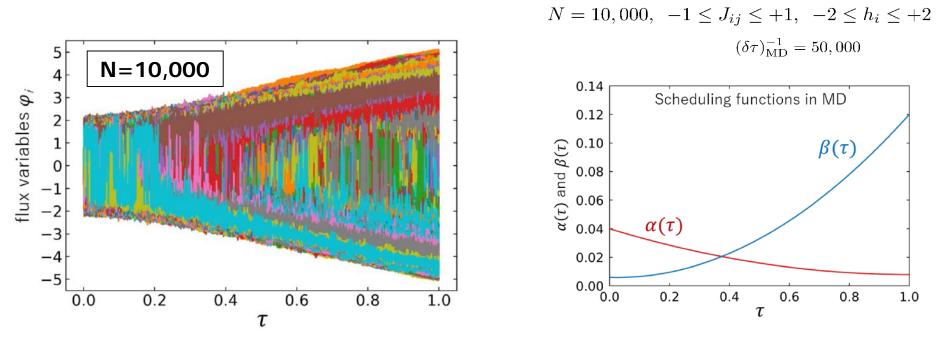


 $\beta(\tau) = \beta_f \left(\tau + \kappa_1(1-\tau) + \kappa_2\tau(\tau-1)\right)$

Final value of MD Hamiltonian $H(\tau=1)$ is indep of schedule function, in particular for finer MD step

→ Good Adiabaticity

Test of MD for Ising spin grass problem



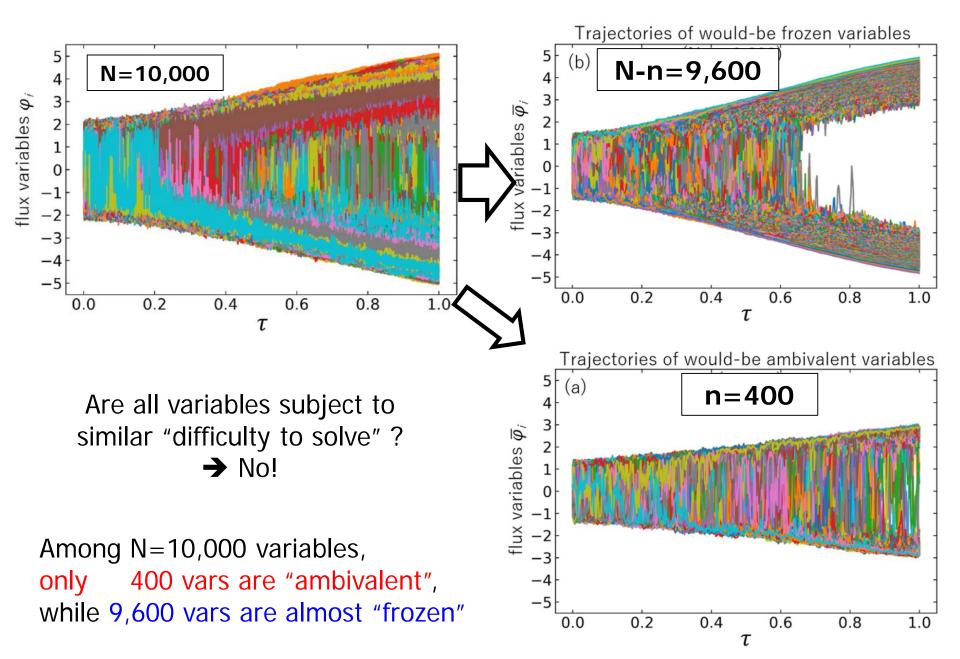
We observe a tendency that variables fall into two (+ or -) categories → MD works rather well

However, it takes additional long-time for "all" variables to be settled down

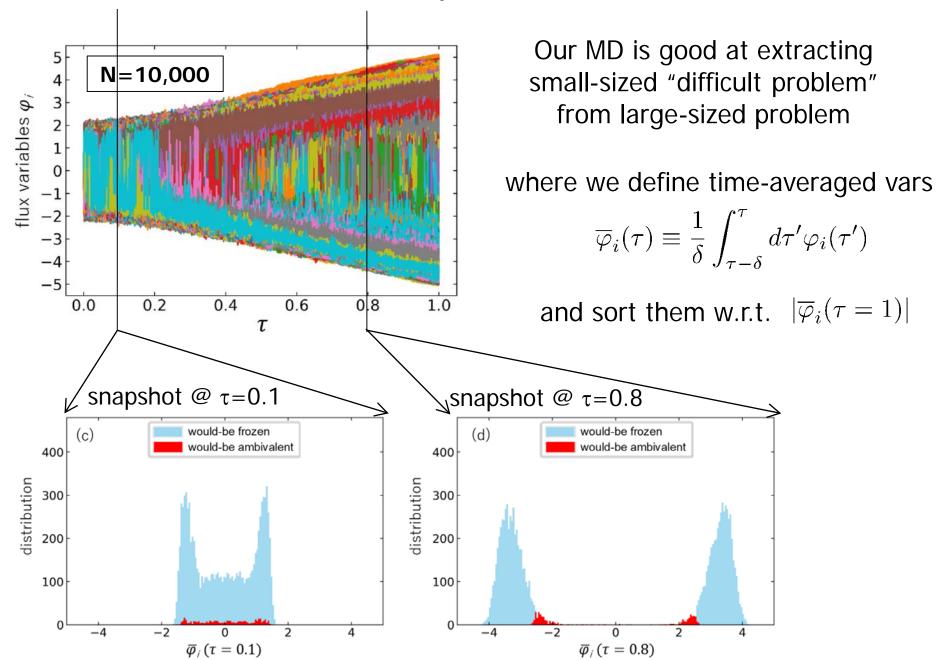
Correspondence between QA and MD $(BJ_{ij} \leftrightarrow \beta J_{ij} | \varphi_i \varphi_j |, Bh_i \leftrightarrow \beta h_i | \varphi_i \varphi_i |)$ can be recovered only when $|\varphi_i| \rightarrow \mu$ for all *i*

→ systematic error from the non-zero distribution of variables

"Hierarchy" in variables

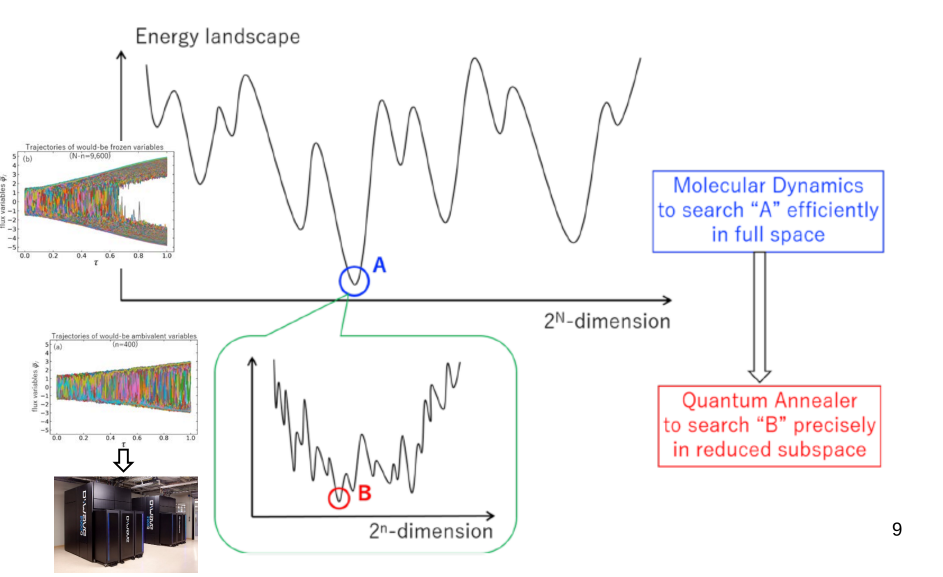


"Hierarchy" in variables

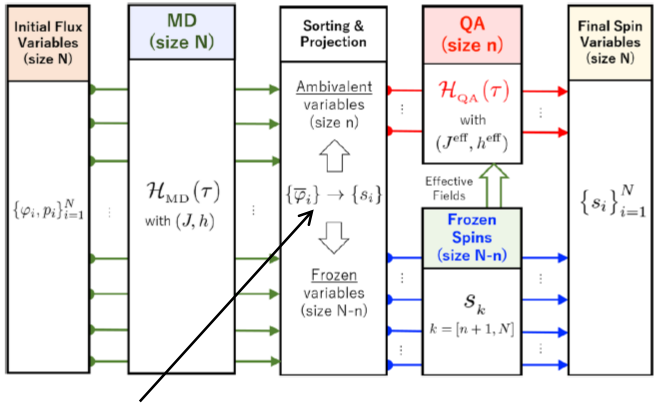


Hybrid Quantum Annealing (HQA) via MD

New Idea: use classical MD as preconditioner for quantum annealing



Flowchart of Hybrid Quantum Annealing (HQA)



We sort w.r.t. continuous time-averaged variables at $\tau = 1$,

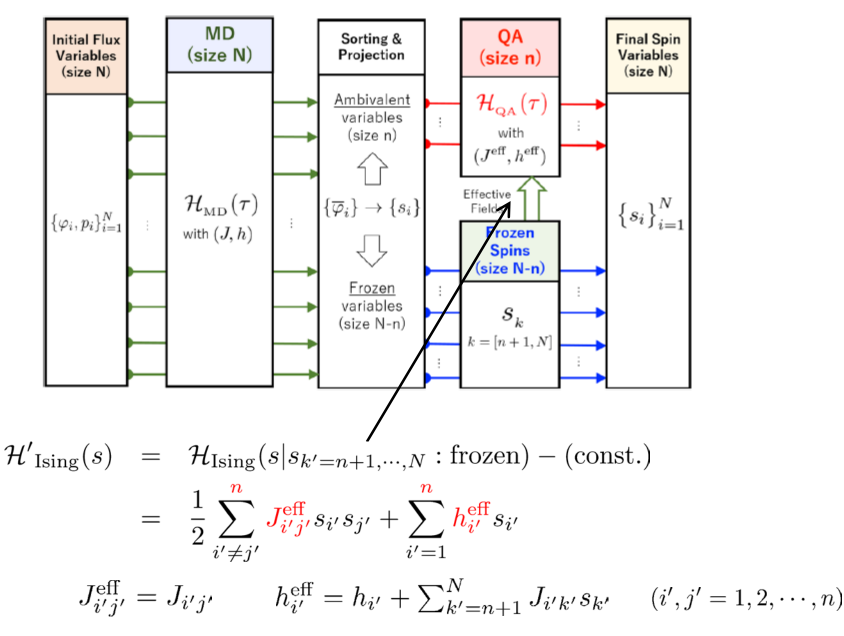
$$\left|\overline{\varphi}_{1'}(\tau=1)\right| \leq \left|\overline{\varphi}_{2'}(\tau=1)\right| \leq \cdots \leq \left|\overline{\varphi}_{n'}(\tau=1)\right| \leq \cdots \leq \left|\overline{\varphi}_{N'}(\tau=1)\right|$$
$$i'=1,2,\cdots,n \text{ (ambivalent)} \quad k'=n+1,n+2,\cdots,N \text{ (frozen)}$$

and then project frozen variables to discrete spin variables

$$s_{k'} = \operatorname{sgn}\left(\overline{\varphi}_{k'}(\tau=1)\right)$$

 \rightarrow we can avoid "continuous-vars" sys err (if frozen spins are correct)

Flowchart of Hybrid Quantum Annealing (HQA)

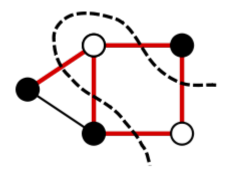


Numerical Results of Hybrid Quantum Annealing (HQA)

MAX-CUT problem

Consider an undirected graph

 $\mathcal{G} = (\mathcal{V}, \mathcal{E})$ $\mathcal{V}: \text{ vertices}$ $\mathcal{E}: \text{ edges w/ weight } \{w_{ij}\}_{(ij)\in\mathcal{E}}$



(Fig from wiki)

Problem:

Find a partition of vertices into 2 sets, $\mathcal{V} = \mathcal{V}_+ \cap \mathcal{V}_-$

which maximizes the sum of weights w_{ij} connecting 2 sets

$$C \equiv \sum_{i \in \mathcal{V}_+, j \in \mathcal{V}_-} w_{ij}$$

Equivalent to minimizing the energy in Ising spin-glass model

$$C(s) = \frac{1}{2} \sum w_{ij} (1 - s_i s_j) = -\frac{1}{2} H_{\text{Ising}}(s) + C_0 \qquad (s_i = \pm 1)$$
$$H_{\text{Ising}}(s) \Big|_{J_{ij} = w_{ij}, h_i = 0} = \frac{1}{2} \sum_{i \neq j} J_{ij} s_i s_j$$

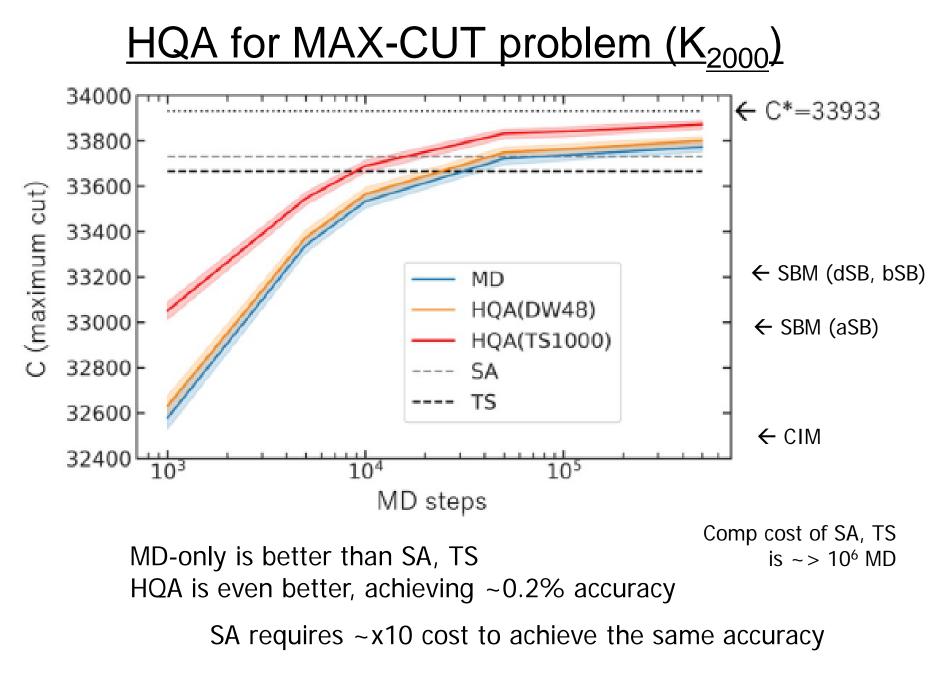
MAX-CUT problem

We consider 2000-node complete graph (K_{2000}) w/ random weights $J_{ij} = \pm 1$ & ensemble (instance) average (we use 100 instances) Classical solver to be benchmarked SA (simulated annealing) Parameters are optimized TS (tabu search) w/ comp cost of SA, TS >~ HQA MD-only (Others) SBM: simulated bifurcation machine (Cost could be different) CIM: coherent ising machine

<u>HQA</u>

HQA (DW48) : HQA w/ n= 48 subsystem solved by D-wave HQA (TS1000) : HQA w/ n=1000 subsystem solved by TS

Dominant cost is MD-part



HQA exhibits significant improvement

HQA for Ising spin-glass problem

We consider N=1,000, 2,000 and 10,000 systems

w/ random parameters & ensemble (instance) average

 $-1 \leq J_{ij} \leq +1, \quad -2 \leq h_i \leq +2 \quad$ w/ uniform distribution

(we use 100 instances)

Classical solver to be benchmarked

SA (simulated annealing) TS (tabu search) MD-only Page 201 Control of the search o

Comp cost of SA, TS is $\sim > 10^6 - 10^7$ MD and thus $> \sim$ HQA

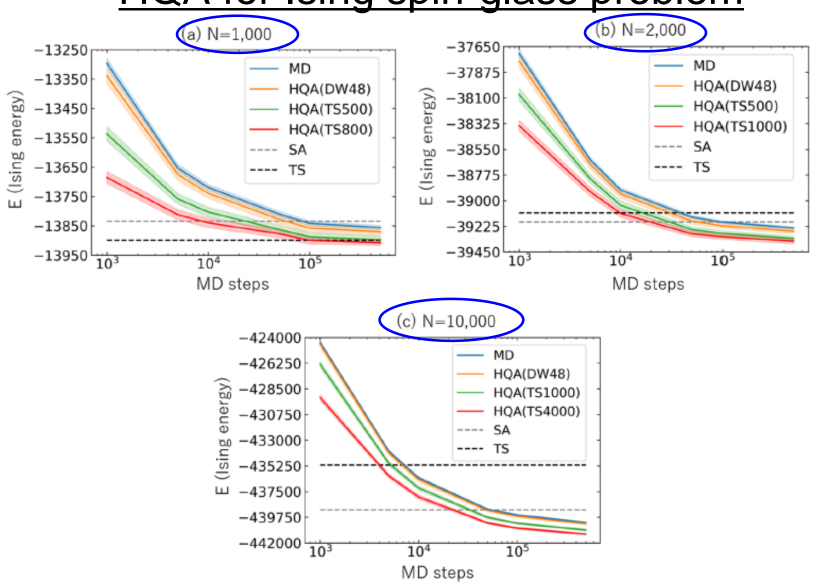
Parameters are optimized

<u>HQA</u>

HQA (DW48) : HQA w/ n= 48 subsystem solved by D-wave HQA (TS-XXX) : HQA w/ n=XXX subsystem solved by TS

Dominant cost is MD-part

HQA for Ising spin-glass problem



HQA exhibits significant improvement

SA requires ~ x100 cost to achieve MD-only accuracy (N=10,000)

Summary

- Hybrid Quantum Annealing (HQA) via Molecular Dynamics
 - New quantum-classical hybrid scheme to solve combinational optimization problem
 - MD can serve as a powerful preconditioner for QA
 - "frozen" / "ambivalent" variables can be identified
 - → "difficult problem" in reduced subspace extracted from full space
 - (NISQ-era) QA solver to search a fine solution in reduced subspace
 - HQA achieves better performance/accuracy than classical SA/TS
 - Larger improvement for a larger system
 - The same concept to extract hierarchy in variables can be utilized w/ any other solver combinations
- Future
 - Extension to binary variables, multi-valued variables
 - More theoretical clarification for classical MD