TROTTER CIRCUIT OPTIMIZATION THROUGH ADIABATIC COMPUTATION

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August 31, 2023

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Part I

QUBO PROBLEMS IN CIRCUIT OPTIMIZATION

LIE-TROTTER FORMULA AND CIRCUIT

TROTTERIZATION

To simulate time-evolving process such as adiabatic quantum process, we approximate continuous process with discrete steps.

We call the discretized approximation as **Trotter** formula.

$$\exp(-i\mathcal{H}t) \approx \Pi_i^n \exp\left(-i\mathcal{H}_i \frac{t}{n}\right)$$
 (1)

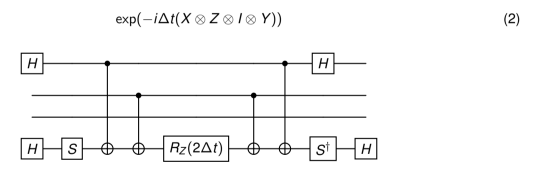
where, *n* is a trotter steps.

As we increase the step number n, we get more precise unitary trnasformation.

LIE-TROTTER FORMULA AND CIRCUIT

TROTTERIZATION

Practically, each terms of Hamiltonian are described with **Pauli string**. A single Pauli string, for example *XZIY*, Hamiltonian has a well known corresponding circuit.



Optimization of evolution circuit is a combination of two parts.

- ► Mutually Commuting Partition
- ► Pauli-Frame

MUTUALLY COMMUTING PARTITION

Pauli strings are always anti-commute or commute each other. For given two Pauli strings, P_i , P_j ,

either
$$[P_i, P_j] = 0$$
 or $\{P_i, P_j\} = 0$ (3)

where, [] is a commutator, and $\{\ \}$ is an anti-commutator.

If all Pauli-terms of Hamiltonain are mutually commute each other, Eq(1) becomes an unitary operator of total Hamiltonain evolution of time t.

$$\exp(-i\mathcal{H}t) = \prod_{i=1}^{n} \exp(-i\mathcal{H}_{i}t)$$
(4)

MUTUALLY COMMUTING PARTITION

- 1. We must know all commuting relation of the given Pauli-stirng set.
- 2. How to make a mutually partitions of the given set?

MUTUALLY COMMUTING PARTITION

To make a mutually commuting partition, we have to know all commuting relationships of the given Pauli-terms of Hamiltonian. We can check the commutation with General commutativity(GC), see Gokhale et al., 2020.

If a system is n qubits system and there are m number of Pauli-terms, total operation would be, roughly,

$$\binom{m}{2} * n = O(m^2 n) \tag{5}$$

Unfortunately, $max(m) = 2^n$ for *n*-qubit system Hamiltonian, it could be expoentially growth.

MUTUALLY COMMUTING PARTITION

Chapuis et al., 2018 suggested acceleration of commuting term determination. They decompose single Pauli-string into X and Z families.

- ► X-family: IIIX, XIXI, IIXI, XXII, IXXX, . . .
- ► Z-family: IIIZ, ZIZI, IIZI, ZZII, IZZZ, . . .

$$YZIX = XIIX \cdot ZZII = x_i \cdot z_i \tag{6}$$

$$[P_i, P_j] = [x_k z_l, x_m, z_n] = \begin{cases} 0 & \text{if}[z_l, x_m] = [x_k, z_n] \\ -2P_i P_j & \text{otherwise} \end{cases}$$
(7)

MUTUALLY COMMUTING PARTITION

Now, if we have compatible grpah of Pauli-set, we can extract mutually commuting partition by solving a sequential Max-Clique problem of the commute graph.

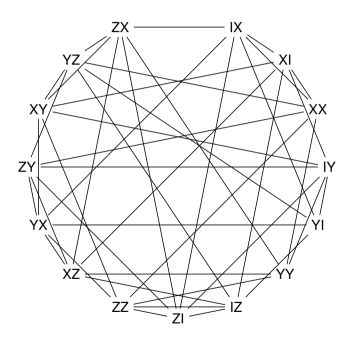
It is well known NP-complete problem, from 21-complete problems. See Karp, 1972.

Kurita et al., 2023 suggested Ising formulation for finding Max-clique finding problem of compatible graph.

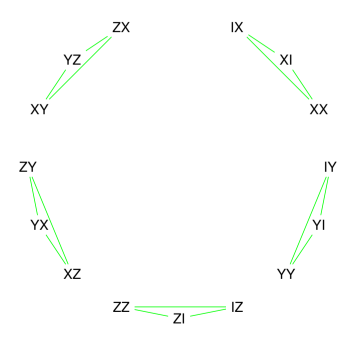
$$\mathcal{H} = -\mu_0 \sum Z_i + \mu_1 \sum h_{ij} Z_i Z_j \tag{8}$$

where, $h_{ij}=0$ if Z_i-Z_j edge weight is 0 otherwise 1, $\mu_0=1, \mu_1=2$ in Kurita et al..

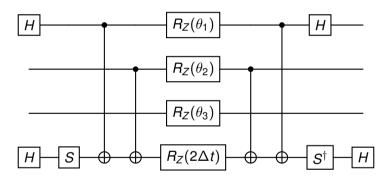
MUTUALLY COMMUTING PARTITION



MUTUALLY COMMUTING PARTITION

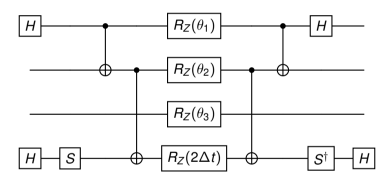


PAULI FRAME



 $\mathcal{H} = tXZIY + \theta_1XIII + \theta_2IZII + \theta_3IIII$

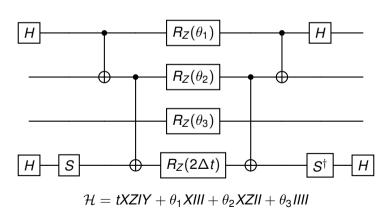
PAULI FRAME



 $\mathcal{H} = tXZIY + \theta_1XIII + \theta_2XZII + \theta_3IIII$

PAULI FRAME

Schmitz et al., 2023 analyzed and Pauli-Frame method and optimized circuit with minimum cost of *CNOT*, *H*, *S* operations to



If there are two max clique on graph, sharing same number of nodes, the next Hamiltonian pick one of them randomly.

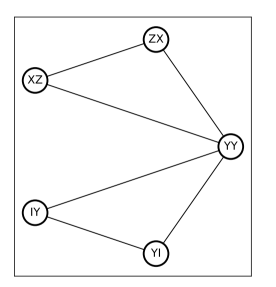
$$\mathcal{H} = -\mu_0 \sum Z_i + \mu_1 \sum h_{ij} Z_i Z_j \tag{10}$$

Eventhough, they are same in commutation graph, frame change cost can be different. In this project, we only consider H, S costs. The weight of each Pauli-terms would be calculated with function w(,), such that

- w(,) = 0: (X, X), (Y, Y), (Z, Z), (Z, I)
- w(,) = 1: (X, Z), (X, Y), (X, I)
- $\mathbf{v}(,) = 2: (Y, I), (Y, Z)$

For *N*-qubit system, extended weight function W(,) is defined as,

$$W(S_i, S_j) := \frac{1}{N} \sum_{k=1}^{N} w((S_i)_k, (S_j)_k)$$
 (11)



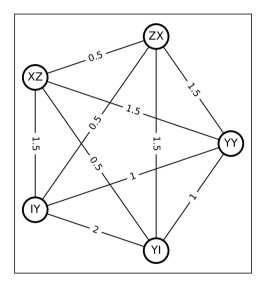


Figure. Compatible and basis transform weight graph example. Left graph is a compatible graph of 5 Pauli basis of 2 qubits system and edges are indicating commutation relationship. Right graph is a basis transform weight graph of the same Pauli-basis set of the left.

We can redefine a Hamiltonian for optimization,

$$\mathcal{H} = -\mu_0 \sum_{i < j} Z_i + \mu_1 \sum_{i < j} h_{ij} Z_i Z_j + \mu_2 \sum_{i < j} w_{ij} Z_i Z_j$$
 (12)

To avoid the degeneration of energy and to conserve max and commuting condition, the coefficients, μ_0, μ_1, μ_2 have next relationship. For N gubits system,

$$\|\mu_1\| > N\|\mu_0\| \|\mu_0\| > \frac{1}{2}N(N-1)\|\mu_2\|$$
 (13)

Full procedure of algorithm.

- 1. Find a compatible graph of the given Hamiltonian
- 2. Calculate weight between Pauli-strings with Eq(11)
- 3. Find a min-number of mutually commuting partition, p_1, p_2, \ldots , using **adiabatic computer**.
- 4. Find a shortest hamilton path of each local partition p_i , <- reduced problem, you can use classic algorithm.
- 5. Connecting p_i in order to following 4 step result.

HEH+ MOLCULAR HAMILTONIAN

Pennylane HeH+ molcule Hamiltonian:

4 qubits are required and consist of 25 Pauli-terms.

'ZXZX','IYIY','ZYZY','IZIZ','XZXZ','XIXI','YZYZ','YIYI','ZIZI', ,'IIIZ', 'ZZII', 'IZZI', 'ZIIZ', 'IZII', 'IIZI', 'ZIII', 'IIZZ', ,'XZXI', 'YZYI', 'XXYY', 'YXXX', 'XYYX', 'IYZY', 'IXZX'

HEH+ MOLCULAR HAMILTONIAN

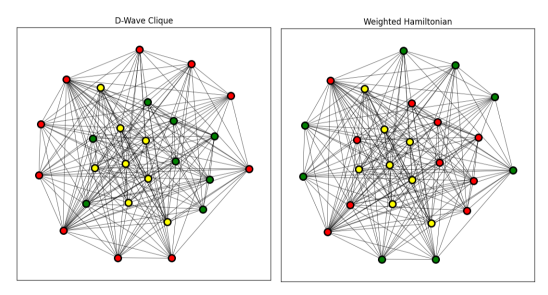


Figure. Commuting Partition HeH+ Hamiltonian Pauli-terms. Left: Ising formula solution of D-Wave. Right: Basis cost term weight added optimization.

HEH+ MOLCULAR HAMILTONIAN

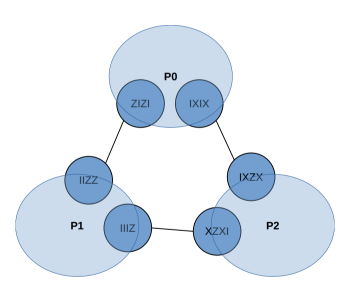
The optimization result is 3 number of partition.

 p_0 ['ZXZX','IYIY','ZYZY','IZIZ','XZXZ','XIXI','YZYZ','YIYI','ZIZI']

 p_1 ['IIIZ', 'ZZII', 'IZZI', 'ZIIZ', 'IZII', 'IIZI', 'ZIII', 'IIZZ']

p₂ ['XZXI', 'YZYI', 'XXYY', 'YXXY', 'YYXX', 'XYYX', 'IYZY', 'IXZX']

HEH+ MOLCULAR HAMILTONIAN



HEH+ MOLCULAR HAMILTONIAN

Compare to Pennylane ApproxTimeEvovle() circuit

```
gates: 270
depth: 169
shots: Shots(total=None)
gate_types:
{'RZ': 106, 'CNOT': 84, 'RX': 80}
gate_sizes:
{1: 186, 2: 84}
gates: 137
depth: 107
shots: Shots(total=None)
gate_types:
{'Hadamard': 16, 'CNOT': 74, 'RZ': 25, 'S': 11, 'Adjoint(S)': 11}
gate_sizes:
{1: 63, 2: 74}
```

Figure. Left: Pennylane ApproxTimeEvolve() trotter number =1 circuit. Right: Optimized evolve circuit.

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