OptGraphState Tutorials

Version 0.3.1

 ${\bf Github:\ https://github.com/seokhyung-lee/OptGraphState}$

API Reference: https://seokhyung-lee.github.io/OptGraphState

$Seok\text{-}Hyung \ Lee \ \texttt{sh.lee1524@gmail.com}$

November 8, 2023

Contents

1	Prerequisites	3
2	Installation	4
3	Defining a GraphState object	4
4	Executing the strategy	7
5	Obtaining instructions for fusion-based generation	11
6	Success probability of graph state generation	18
7	List of features 7.1 Class variables of GraphState	19 20 20 20 21 21 21
8	More on the graphs and fusion network 8.1 Original graph GraphState.graph	

O.D PHSION NELWORK GLADINGLACE. LUSTON NELWOLK	twork GraphState.fusion_network	. 22
--	---------------------------------	------

OptGraphState is a python package that implements the graph-theoretical strategy to optimize the fusion-based generation of an arbitrary graph state that is proposed in Ref. [1]. The package has the following features:

- Finding a resource-efficient method of generating a given graph state through type-II fusions from multiple three-qubit linear graph states $(\left|G_*^{(3)}\right\rangle)$.
- Computing the corresponding resource overhead, which is quantified by the average number of required $\left|G_*^{(3)}\right\rangle$ states or fusion attempts.
- Visualizing the original graph (of the graph state you want to generate), unraveled graphs, and fusion networks. An unraveled graph is a simplified graph where the corresponding graph state is equivalent to the desired graph state up to fusions and single-qubit Clifford operations. A fusion network is a graph that instructs the fusions between $G_*^{(3)}$ states required to generate the desired graph state.
- Various predefined sample graphs for input.

This tutorial is written in such a way that it is as understandable as possible without having a background in Ref. [1] except for Sec. 8. The API Reference is available in https://seokhyung-lee.github.io/OptGraphState.

1 Prerequisites

OptGraphState has the following prerequisites:

- python == 3.9
- numpy == 1.24.2
- python-igraph == 0.10.4
- networkx == 3.1
- matplotlib == 3.7.1
- parmap == 1.6.0 (Optional, for multiprocessing)
- tqdm == 4.66.1 (Optional, for progress bar)

The specified versions of these requirements are the ones with which we has tested OptGraphState; it does not mean that OptGraphState does not work with other versions.

OptGraphState heavily relies on the *python-igraph* library, which offers a convenient interface for creating, manipulating, analyzing, and visualizing graphs [2]. The majority of graph-theoretical algorithms used in the strategy are implemented using *python-igraph* (which is faster than *NetworkX* since it is written in C), with the exception of the maximum matching problem, which utilizes the *NetworkX* library. Nevertheless, users are not required to have in-depth knowledge of these libraries to use OptGraphState, though it may be helpful for more thorough investigations.

2 Installation

OptGraphState can be installed with pip:

\$ pip install optgraphstate

After starting Python, import OptGraphState with

```
>>> import optgraphstate as ogs
```

3 Defining a GraphState object

Most features of OptGraphState are implemented via the GraphState class. A GraphState object is defined by providing the graph of the graph state that you want to generate. This can be done in the following three ways:

(1) Given by a list of edges. Each edge is given by a 2-tuple of vertex labels in $\{0, 1, 2, \dots, n_V - 1\}$, where n_V is the number of vertices. For example, to define a GraphState object with a four-vertex linear graph:

```
>>> gs = ogs.GraphState(edges=[(0, 1), (1, 2), (2, 3)])
```

Note that the number of vertices is automatically set to be the largest vertex label plus one.

(2) Given directly as an igraph. Graph or a networkx. Graph object. If a networkx. Graph object is given, it is internally converted to an igraph. Graph object. For example, to define a GraphState object with a four-vertex linear graph:

```
>>> import igraph as ig
>>> g = ig.Graph(edges=[(0, 1), (1, 2), (2, 3)])
>>> gs = ogs.GraphState(graph=g)
```

or

```
>>> import networkx as nx
>>> g = nx.Graph()
>>> g.add_nodes_from([0, 1, 2, 3])
>>> g.add_edges_from([(0, 1), (1, 2), (2, 3)])
>>> gs = ogs.GraphState(graph=g)
```

(3) Chosen among predefined graphs. OptGraphState provides a range of predefined families of graphs. A GraphState object can be defined by giving the shape and parameters of the desired graph in the arguments shape (as a str) and prms (as a tuple or int), respectively. For example, to define a GraphState object with a six-vertex cycle graph:

```
>>> gs = ogs.GraphState(shape='cycle', prms=6)
```

To define a GraphState object with a 2D lattice graph that has dimensions of (3,3):

```
>>> gs = ogs.GraphState(shape='lattice', prms=(3, 3))
```

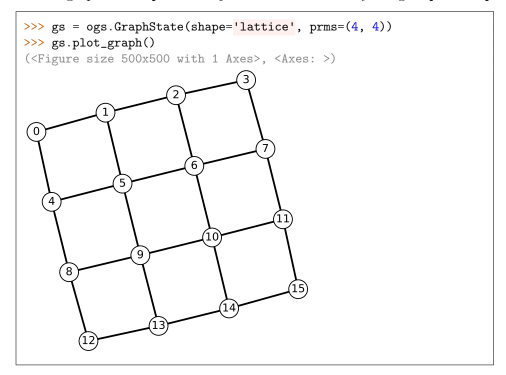
The full list of supported graphs are as follows (see Sec. 1.1, Fig. 2, and Sec. 3.3 of Ref. [1] for their descriptions and visualizations):

- shape='random': Random graph for fixed numbers of vertices and edges, sampled by the Erdős–Rényi model [3].
 - prms[0] <int>: Number of vertices.
 - prms[1] <int>: Number of edges.
 - (Optional) prms[2] <None or int>: Random seed. If None, the current system time is used as the seed. If not given, the random number generator is not initialized.

- shape='complete', 'star', 'linear', or 'cycle': Complete, star, linear, or cycle graph, respectively.
 - prms <int>: Number of vertices.
- shape='lattice': Lattice graph.
 - prms <tuple of int>: Numbers of repeated vertices along the len(prms) axes.
- shape='tree': Tree graph where all branches in each generation have an equal number of children.
 - prms[0] <int>: Degree of the root vertex.
 - prms[i] (i \geq 1) <int>: Number of the children of each ith-generation branch.
- shape='rhg': Raussendorf-Harrington-Goyal lattice with primal boundaries only.
 - prms[0], prms[1], prms[2] <int>: Sizes of the lattice along the three axes in the unit of a cell, respectively.
- shape='repeater': Repeater graph with 4m vertices.
 - prms <int>: Parameter m.
- shape='parity_encoding': (n,m) parity-encoded graph [4, 5], where each vertex of the logical-level graph corresponds to a qubit encoded in the basis of either $\left\{ (|0\rangle^{\otimes m} + |1\rangle^{\otimes m})^{\otimes n} \pm (|0\rangle^{\otimes m} |1\rangle^{\otimes m})^{\otimes n} \right\}$ (Orientation 1) or $\left\{ [(|0\rangle + |1\rangle)^{\otimes m} \pm (|0\rangle |1\rangle)^{\otimes m} \right\}$ (Orientation 2).
 - prms[0] <str or igraph.Graph>: Logical-level graph given by either its name or graph structure. Currently 3-star ("3-star" or "3-linear) and 6-cycle ("6-cycle" or "6-ring") graphs can be given by their names. For the other graphs, it should be given by an igraph.Graph object. It can be generated directly via the python-igraph library or from the function get_graph_from_edges() or get_sample_graph().
 - prms[1], prms[2] <int>: Parameters n and m of the parity encoding.

- (Optional) prms[3] <bool>: Orientation of the parity encoding. If True (default), "Orientation 1" is used. If False, "Orientation 2" is used.
- shape='ptqc': Microcluster for parity-encoding-based topological quantum computing protocol [5].
 - prms[0], prms[1] <int>: Parameter n and m of the parity encoding.
 - prms[2] <bool>: Whether the H-configuration is HIC (True) or HIS (False).
 - prms[3] <bool>: Whether the microcluster is a central (True) or side (False) one.

The graph of a GraphState object can be visualized by using GraphState.plot_graph():



4 Executing the strategy

The strategy is performed by iterating the three steps: unraveling the original graph, building a fusion network, and computing the resource overhead by

determining the fusion order. To briefly explain these steps:

- 1. Unraveling the graph: Find an unraveled graph where the corresponding graph state is equivalent to the original graph state up to several fusions and single-qubit Clifford gates and is easier to generate than the original graph state. It is done by unraveling specific subgraph structures of the original graph, such as bipartitely-complete graphs and cliques.
- 2. Building a fusion network: A fusion network is graph where the nodes indicate individual three-qubit linear graph states $(G_*^{(3)})$ and the links indicate fusions between them that are required to generate the desired graph state. We use the term "node" and "link" instead of "vertex" and "edge" to distinguish them from the vertices and edges in the original and unraveled graphs.
- 3. **Determining the fusion order and computing the resource overhead**: Based on the built fusion network, the order of fusions is determined using the *min-weight-maximum-matching-first* strategy. The resource overhead, which is quantified by the average number of three-qubit linear graph states required to generate the desired graph state, is computed during this process.

See Ref. [1] for more details on the strategy. Note that these three steps are implemented in GraphState.unravel_graph(), GraphState.build_fusion_network(), and GraphState.calculate_overhead(), respectively.

Since the above process contains randomness, a different value is obtained each time the resource overhead is calculated. Therefore, we take an approach to find the optimal one by iterating the entire steps sufficiently.

You have two choices for the iteration: (i) Iterate for a fixed number of times or (ii) use the adaptive iteration method. Here, the adaptive iteration method with the given initial iteration number $n_{\rm init}$ and the multiplicative factor M means the following process: Denoting the process to iterate the three steps n times as R(n), first perform $R(n_{\rm init})$ and obtain $Q_{\rm opt}^{(1)}$, which is the minimal resource overhead obtained so far. Then, perform $R(Mn_{\rm init})$ and obtain $Q_{\rm opt}^{(2)}$ similarly. If $Q_{\rm opt}^{(1)} \leq Q_{\rm opt}^{(2)}$, stop the iteration and return $Q_{\rm opt}^{(1)}$. If otherwise, perform $R(M^2n_{\rm init})$, obtain $Q_{\rm opt}^{(3)}$, and stop the iteration if $Q_{\rm opt}^{(2)} \leq Q_{\rm opt}^{(3)}$, which returns $Q_{\rm opt}^{(2)}$. If $Q_{\rm opt}^{(2)} > Q_{\rm opt}^{(3)}$, perform $R(M^3n_{\rm init})$, obtain $Q_{\rm opt}^{(4)}$, and so on.

(i) With a fixed iteration number: Use GraphState.simulate(). For example, to iterate 1000 times with the fusion success rate of 0.5 for a (5,5)-lattice graph through multiprocessing:

```
>>> gs = ogs.GraphState(shape='lattice', prms=(5, 5))
>>> res = gs.simulate(
       1000,
                # iteration number
. . .
        p_succ=0.5,  # fusion success rate
. . .
                   # whether to use multiprocessing or not
       mp=True,
        n_procs=None # number of simultaneous processes
. . .
                      # (number of CPU cores if None)
. . .
...)
Multiprocessing ON: n_procs = 8
Calculating for n_{iter} = 1000... Done. Best: 100352.00 (2.77 s)
{'best_overhead': 100352.0,
'best_num_fusions': 67262.0,
'best_num_steps': 12,
 'best_seed': 1910731091,
 'n_iter': 1000,
 'unravel_bcs_first': False}
```

(ii) With the adaptive iteration method: Use GraphState.simulate_adaptive(). For example, if the initial iteration number is 200 and the multiplicative factor is 2:

```
>>> gs = ogs.GraphState(shape='lattice', prms=(5, 5))
>>> res = gs.simulate_adaptive(
        200,
                     # initial iteration number
       mul=2,
                    # multiplicative factor
       p_succ=0.5,  # fusion success rate
       mp=True,
                     # whether to use multiprocessing or not
       n_procs=None # number of simultaneous processes
. . .
                      # (number of CPU cores if None)
Multiprocessing (n_procs = 8)
Calculating for n_iter = 200... Done. Best: 112640.00 (1.59 s)
Calculating for n_iter = 400... Done. Best: 100352.00 (1.83 s)
Calculating for n_iter = 800... Done. Best: 100352.00 (2.41 s)
>>> res
{'best_overhead': 100352.0,
 'best_num_fusions': 67262.0,
 'best_num_steps': 12,
 'best_seed': 3704429497,
 'n_iter': 1400,
 'unravel_bcs_first': True}
```

For both methods, the outcome of a simulation is returned as a dictionary with the following keys and values:

- 'best_overhead': Smallest overhead value (i.e., average number of basic resource states) among the samples.
- 'best_num_fusions': Average number of required fusion attempts for the best sample (that gives the smallest overhead).
- 'best_num_steps': Minimal number of steps (i.e., groups of fusions that can be done in parallel) for the best sample.
- 'best_seed': Random seed of the best sample, which can be used to reproduce the result; e.g.,

```
>>> gs.simulate(1, p_succ=0.5, seed=3704429497)
```

- 'n_iter': Total number of iterations.
- 'unravel_bcs_first': Whether bipartitely-complete subgraphs (BCSs) are unraveled before cliques are unraveled, for the best sample.

Only the data of the best sample is obtained by default. If you want to get the data of all samples, set the keyword argument get_all_data=True. Similarly, set get_all_graphs=True to get all the unraveled graphs, and set get_all_fusion_networks=True to get all the fusion networks.

If you want to optimize the average number of fusion attempts instead of the average number of basic resource states, set optimize_num_fusions=True.

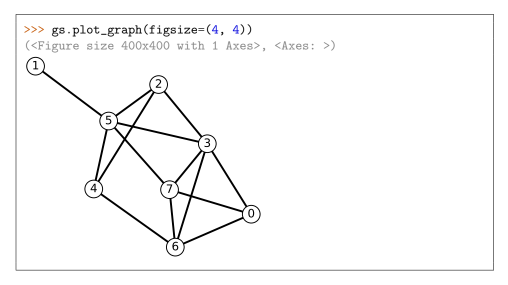
5 Obtaining instructions for fusion-based generation

The outcome of GraphState.simulate() or GraphState.simulate_adaptive() does not provide detailed instructions on how to generate the desired graph state. However, after running the simulation, all information of the best sample is stored in the GraphState object: the original graph in GraphState.graph, the unraveled graph in GraphState.unraveled_graph, and the fusion network in GraphState.fusion_network.

Let us consider the following example:

```
>>> gs = ogs.GraphState(shape='random', prm=(8, 14, 4))
>>> res = gs.simulate(1000)
Multiprocessing OFF.
Calculating for n_iter = 1000...
Done. Best: 384.00 (1.45 s)
>>> res
{'best_overhead': 384.0,
   'best_num_fusions': 262.0,
   'best_num_steps': 6,
   'best_seed': 4248262454,
   'n_iter': 1000,
   'unravel_bcs_first': False}
```

which has the following graph



The number inside each vertex is referred to as its name, which is unique. The easiest way to learn how to generate the desired graph state from initial $\left|G_*^{(3)}\right>$ states is to use GraphState.get_instructions():

```
>>> node_names, fusions, final_qubits = gs.get_instructions()
>>> node_names
['0', '2', '3', '5', '5-1', '6', '7', '9', '10', '11']
>>> fusions
{1: [(('5', 'L', None), ('5-1', 'R', None)),
  (('0', 'L', None), ('7', 'L', None)),
  (('6', 'L', None), ('9', 'L', None)),
  (('2', 'L', None), ('10', 'L', None)),
  (('3', 'L', None), ('11', 'L', None))],
 2: [(('5-1', 'L', None), ('7', 'L', None)),
  (('6', 'L', None), ('10', 'L', None))],
 3: [(('10', 'R', 'RZ'), ('11', 'R', None))],
 4: [(('3', 'L', None), ('9', 'L', None))],
 5: [(('2', 'L', None), ('5', 'L', None))],
 6: [(('0', 'L', 'RZ'), ('9', 'R', None))]}
>>> final_qubits
{'0': ('0', 'R', 'RXd'),
 '1': ('5-1', 'L', None),
 '2': ('2', 'R', 'RXd'),
 '3': ('3', 'R', None),
 '4': ('11', 'L', None),
 '5': ('5', 'R', 'RZ'),
 '6': ('6', 'R', None),
 '7': ('7', 'R', 'RZ')}
```

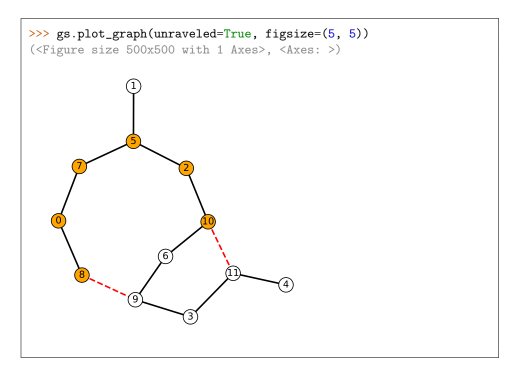
The desired graph state can be generated as follows (assuming that all fusions succeed):

- 1. Prepare len(node_names) copies of $\left|G_*^{(3)}\right\rangle$ states, which are called *nodes* and respectively named by the elements in the list node_names.
- 2. Perform fusions between the nodes according to the list fusions[1] whose elements indicate fusions that can be done in parallel. Each element has the form of ((n1, q1, c11), (n2, q2, c12)), which means:
 - n1, n2: Names of the nodes involved in the fusion.
 - q1, q2: Qubits (in the two nodes) that participate in the fusion. Each of them is either 'R' or 'L', which respectively means that the qubit is the root qubit of the node (central qubit of the $|G_*^{(3)}\rangle$ state) or any one of its leaf qubits (non-central qubits).

- c11, c12: Clifford gates that should be applied to the qubits before the fusion is performed. 'RXd' and 'RZ' respectively means $R_X(\pi/2) := \exp\left[-i(\pi/4)\hat{X}\right]$ and $R_Z(-\pi/2) := \exp\left[i(\pi/4)\hat{Z}\right]$. Multiple $\pi/2$ rotations are expressed as 'RZ-RXd', which means $R_X(\pi/2)R_Z(-\pi/2)$.
- 3. Similarly, perform fusions according to the lists fusion[2], fusions[3], and so on, in order.
- 4. The dictionary final_qubits shows the correspondence between the vertices in the original graph and the final remaining qubits after all the fusions are performed. For each vertex (with a name vname), final_qubits[vname] has the form of (node, qubit, cl), where
 - node: Node containing the qubit.
 - qubit: Either 'R' or 'L' that indicates the qubit in the node.
 - cl: Clifford gate that should be applied to the qubit.

In real implementation, fusions may fail due to theoretical limitations or environmental noises. If such failures are considered, whenever a fusion (say, on qubits q_1 and q_2) fails, you need to discard qubits that were entangled with q_1 or q_2 and reprepare the state on the qubits. Our software does not provide detailed instructions to accomplish such processes.

For more advanced analysis, you may need to visualize the obtained unraveled graph and fusion network. First, use GraphState.plot_graph() to visualize the unraveled graph:



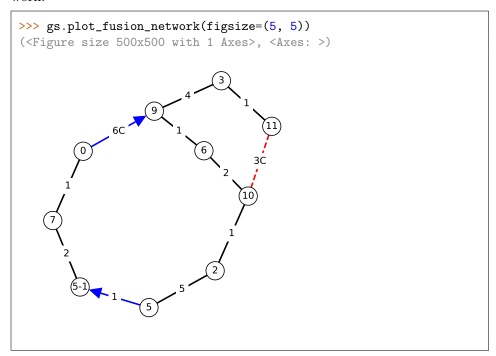
It can be interpreted as follows:

- Each vertex is a qubit, each black solid line is an edge of the unraveled graph, and each red dashed line indicates a fusion that should be done to recover the original graph state.
- The numbers inside the vertices are their unique names. The vertex names in the original graph are here as well, which mean that they are the same qubit. Other vertices in the unraveled graph ('8'-'11') are for new qubits.
- The orange vertices mean that non-trivial Clifford gates should be applied to the qubits to recover the original graph state. Such gates can be identified by using GraphState.get_vertex_clifford():

```
>>> gs.get_vertex_clifford(0)
'RXd'
>>> gs.get_vertex_clifford(10)
'RZ'
```

Namely, the vertices '0' and '10' should be subjected to $R_X(\pi/2)$ and $R_Z(-\pi/2)$, respectively.

Next, use GraphState.plot_fusion_network() to visualize the fusion network:



It can be interpreted as follows:

- Each node is a $|G_*^{(3)}\rangle$ state and each link indicates a fusion required between the $|G_*^{(3)}\rangle$ states. We refer to the central qubit of a $|G_*^{(3)}\rangle$ state as its *root* qubit and the other qubits as its *leaf* qubits. The numbers inside the nodes are their unique names.
- A number placed on each link means the order of the fusion. The fusions should be performed in the order in which these numbers increase (from 1 to 6) and those with the same number can be done simultaneously.
- The line style and color of each link indicates which qubits in the $\left|G_*^{(3)}\right\rangle$ states are involved in the fusion:
 - Black solid line: Two leaf qubits are fused.

- Red dashed line: Two root qubits are fused.
- Blue arrow: One leaf qubit and one root qubit are fused. The arrow points in the direction from the leaf qubit to the root qubit.
 (For example, the leaf qubit of the node '5' and the root qubit of the node '5-1' are fused.)
- If 'C' is written on a link (e.g., '3C' on the link connecting the nodes '10' and '11'), it means that the fusion of the link is accompanied by non-trivial Clifford gates applied to one or both of the qubits before the fusion is performed. To identify such Clifford gates, use GraphState.get_link_clifford():

```
>>> gs.get_link_clifford(10, 11)
('RZ', None)
```

Namely, for the link connecting the nodes '10' and '11', the (leaf) qubit of the node '10' involved in the fusion should be subjected to $R_Z(-\pi/2)$ beforehand.

- We need to know how the final remaining qubits after performing all the fusions correspond to the vertices in the original graph. For a vertex in the original graph, it is as follows:
 - If the vertex with the same name in the unraveled graph is connected to two or more vertices (with solid lines), it corresponds to the root qubit of the node with the same name. (For example, the vertex '0' corresponds to the root qubit of the node '0'.)
 - If the vertex with the same name in the unraveled graph is connected to only one vertex u, it corresponds to any one of the final remaining leaf qubits of the nodes that have names starting with the name of u. (For example, the vertices '1' and '4' correspond to the final remaining leaf qubits of the nodes '5-1' and '11', respectively.)
- Note that, after all the fusions are performed, all the required Clifford gates should be applied to final remaining qubits according to the unraveled graph. These Clifford gates are not visualized in the fusion network, which is one of the reasons that you should draw the unraveled graph.

6 Success probability of graph state generation

OptGraphState provides a function to calculate the probability of successfully generating a graph state when the number of provided basic resource states is limited. See Sec. 3.4 of Ref [1] for theoretical details.

Before the calculation, you first need to run the simulation by GraphState.simulate() or GraphState.simulate_adaptive(). For example,

```
>>> gs = ogs.GraphState(shape='parity_encoding',
>>> prms=('6-cycle', 4, 4))
>>> res = gs.simulate_adaptive(1200, mp=True)
Multiprocessing (n_procs = 8)
Calculating for n_iter = 1200... Done. Best: 21248.00 (5.66 s)
Calculating for n_iter = 2400... Done. Best: 20864.00 (8.66 s)
Calculating for n_iter = 4800... Done. Best: 20480.00 (14.83 s)
Calculating for n_iter = 9600... Done. Best: 20480.00 (30.79 s)
```

After that, execute GraphState.calculate_succ_probs():

```
>>> c_start, probs, overhead = gs.calculate_succ_probs(cutoff=0.99)
prec = 30: Error
prec = 35: 2677.0180012995289751721232292276553
prec = 40: 14321.06300108698315566255058239658324074
prec = 45: 20132.9839359893134477038277196303371420926776
prec = 50: 20133.056172721261728678099981064827212204339410780
>>> c_start
96
>>> overhead
20133.056172721261728678099981064827212204339410780
>>> for i, p in enumerate(probs[:10]):
       print(f"c={c_start+i}: {float(p):.2e}")
c=96: 1.26e-29
c=97: 1.26e-29
c=98: 2.90e-28
c=99: 3.16e-28
c=100: 3.46e-27
c=101: 4.03e-27
c=102: 2.86e-26
c=103: 3.52e-26
c=104: 1.83e-25
c=105: 2.36e-25
```

The method computes the success probability while increasing the resource

count until the calculated probability reaches the cutoff threshold (given by the cutoff parameter) that is smaller than one. It returns three values:

- 1. c_start: First resource count with nonzero success probability.
- 2. probs: Array of decimal.Decimal objects where probs[i] is the success probability for the resource count of c_start + i.
- 3. overhead: Estimated resource overhead. Since the cutoff threshold is smaller than one, overhead is smaller than the true value obtained from the simulation.

The execution of the method involves the summation of many extremely small numbers, thus low precision of numbers may lead to wrong results. Therefore, by default, it executes the computation multiple times while increasing the precision (i.e., number of significant figures for the decimal package) by 5 starting from 30. When the estimated overhead does not change by more than 1% in two consecutive trials, the iteration is terminated and the results of the final trial are returned. If you do not want this iteration, set auto_increasing_prec=False.

7 List of features

7.1 Class variables of GraphState

- GraphState.data <dict>: Any data obtained during unraveling the graph, building the fusion network, and calculating the resource overhead.
- GraphState.graph <igraph.Graph>: Graph of the given desired graph state
- GraphState.unraveled_graph <None or igraph.Graph>: Graph of the generated unraveled graph state.
- GraphState.fusion_network <None or igraph.Graph>: Generated fusion network.

See Sec. 8 for the list of vertex and edge attributes of these igraph. Graph class variables.

7.2 Execution of the strategy

- GraphState.unravel_bcss(): Unravel bipartitely-complete subgraphs of the original graph.
- GraphState.unravel_cliques(): Unravel cliques of the original graph.
- GraphState.unravel_graph(): Unravel bipartitely-complete subgraphs and cliques of the original graph in a random order (by default).
- GraphState.build_fusion_network(): Build a fusion network from the original or unraveled graph.
- GraphState.calculate_overhead(): Determine the fusion order and calculate the resource overhead.
- GraphState.initialize(): Initialize the generated unraveled graph, fusion network, and calculated data.
- GraphState.simulate(): Iterate the strategy for a fixed number of times.
- GraphState.simulate_adaptive(): Iterate the strategy with the adaptive iteration method.

7.3 Extracting information from the graphs and fusion network

- GraphState.get_instructions(): Get the instructions for the fusion-based generation of the desired graph state.
- GraphState.get_vertex_clifford(): Get the Clifford gate applied to a given vertex in the unraveled or original graph.
- GraphState.get_link_clifford(): Get the Clifford gates that need to be applied to two qubits involved in the fusion of a given link in the fusion network.

7.4 Calculation of success probability of graph state generation

• GraphState.calculate_succ_probs(): Calculate the success probability of the graph state generation for each number of provided basic resource states.

- get_ftpdf_contracted: Compute the fourier-transformed probability distribution function (FTPDF) for a new node created by contracting a link of a fusion network from the FTPDFs for the two endpoints of the link. See Sec. 3.4 of Ref. [1] for more details.
- recover_probs_from_ftpdf: Recover the cumulative mass function (CMF) of the resource count for a node in a fusion network from its FTPDF. See Sec. 3.4 of Ref. [1] for more details.

7.5 Visualization

- plot_graph(): Plot a given graph.
- plot_fusion_network(): Plot a given fusion network.
- GraphState.plot_graph(): Plot the original or unraveled graph. It is equal to plot_graph(GraphState.graph) or plot_graph(GraphState.unraveled_graph).
- GraphState.plot_fusion_network(): Plot the fusion network. It is equal to plot_fusion_network(GraphState.fusion_network)

7.6 Graph tools

- get_graph_from_edges(): Generate a graph (igraph.Graph) from a given list of edges.
- get_sample_graph(): Generate a predefined graph (igraph.Graph) from a given shape and parameters.
- find_nonoverlapping_bcss(): Find a maximum set of bipartitely-complete subgraphs that do not share any vertices.
- find_nonoverlapping_cliques(): Find a maximum set of cliques that do not share any vertices.

7.7 Others

• GraphState.copy(): Return a shallow copy of the GraphState object.

8 More on the graphs and fusion network

For advanced applications, you may need to directly deal with the igraph. Graph class variables in a GraphState object: GraphState.graph, GraphState.unraveled_graph, and GraphState.fusion_network. Here, we present the details on the vertex and edge attributes of these class variables. We assume that the reader is familiar with our strategy in Ref. [1] and the python-igraph library.

8.1 Original graph GraphState.graph

Vertex attributes

- 'name' <str>: Unique label of the vertex.
- (Optional) 'clifford' <str>: Clifford gates applied to the vertex. 'RXd' and 'RZ' indicates the operators $R_X(\pi/2) := \exp\left[-i(\pi/4)\hat{X}\right]$ and $R_Z(-\pi/2) := \exp\left[i(\pi/4)\hat{Z}\right]$, respectively. If it is 'RXd-RZ', it means that $R_X(\pi/2)$ is applied and then $R_Z(-\pi/2)$ is applied.

8.2 Unraveled graph GraphState.unraveled_graph

Vertex attributes

- 'name' <str>: Unique label of the vertex. If the vertex comes from the original graph (namely, if it is not a new vertex created while unraveling), it has the same label as the corresponding vertex in the original graph.
- 'ext_fusion' <None or str>: Name of the other vertex that undergoes an external fusion with the vertex. It is None if the vertex is not involved in any fusions.
- 'clifford' <None or str>: Clifford gates applied to the vertex.

8.3 Fusion network GraphState.fusion_network

Vertex (node) attributes

• 'name' <str>: Unique label of the node. For a vertex 'i' in the fusion network that is connected with two or more edges, its corresponding node group contains the nodes with the names 'i', 'i-1', 'i-2', and so on, where the node 'i' is the seed node of the node group.

- 'seed' <bool>: Whether the node is a seed node or not.
- 'node_group' <str>: Name of the seed node in the same node group.

Edge (link) attributes

- 'name' <str>: Unique label of the link.
- 'kind' <str>: Type of the link. It is one of 'RR', 'LL', and 'RL', which respectively mean root-to-root, leaf-to-leaf, and root-to-leaf.
- 'root_node' <None or str>: Name of the node (connected with the link) whose root qubit is involved in the fusion of the link, if the link has the type of root-to-leaf. It is None if the link has another type.
- 'cliffords' <dict>: Non-trivial Clifford gates applied to the qubits involved in the fusion before the fusion is performed. Its keys are the names of the two nodes connected by the link and its values are the Clifford gates applied to the respective qubits.
- 'order' <int>: Order of the fusion that starts from 1. Fusions with the same order can be performed simultaneously. This attributes is not created before the fusion order is determined by GraphState.calculate_overhead().

References

- [1] S.-H. Lee and H. Jeong, "Graph-theoretical optimization of fusion-based graph state generation," (2023), arXiv:2304.11988 [quant-ph] .
- [2] G. Csardi and T. Nepusz, "The igraph software package for complex network research," InterJournal Complex Systems, 1695 (2006).
- [3] P. Erdős and A. Rényi, "On random graphs I," Publicationes mathematicae 6, 290–297 (1959).
- [4] T. C. Ralph, A. J. F. Hayes, and A. Gilchrist, "Loss-tolerant optical qubits," Phys. Rev. Lett. **95**, 100501 (2005).
- [5] S.-H. Lee, S. Omkar, Y. S. Teo, and H. Jeong, "Parity-encoding-based quantum computing with bayesian error tracking," npj Quantum Inf. 9, 39 (2023).