

# User Manual

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**Abstract.**

## 1 multiVitamin

multiVitamin is a software package which allows users to perform multiple alignments with graphs. There are two algorithms available for this task. The Bron Kerbosch [1] and the VF2 algorithm [2]. Details on the algorithms can be found in the theory section.

The main function of the package is to align two or more graphs following a binary alignment guide tree containing the "best" alignment procedure. The resulting multiple graph alignment is represented as one single graph itself. Next to this main function, there is also the possibility to show all co-optimals of an alignment between 2 graphs or to view graphical representations of graphs. Sounds exciting, so let's get started!

## 2 Installation

Clone the repo from github with

```
$ git clone https://github.com/mk36fyvy/multivitamin.git
```

Navigate to the directory containing the setup.py file

```
$ cd [yourDir]/multivitamin/multivitamin
```

and type

```
$ pip3 install .
```

Done, now you can run multiVitamin in the command shell. You can test if the installation was successful by typing

```
$ multiVitamin -h
```

## 3 Graph Format

A good first step is to familiarize yourself with the graph format. A graph file is basically a text file, we use the extension '.graph'. It is also used interiorly to identify graph files as such.

The file itself is structured as follows (see below for an example):

Every graph consists of nodes and edges. Every node has an id, which is a unique integer (handled as a String internally) for each node. Optionally, nodes can be labelled. The id and label are separated by a semicolon (;) in the '.graph'-file. The edges are two connected nodes and represented by the ids of the two nodes separated by a semicolon.

Now, let's walk through the overall structure of the '.graph'-file:

- There is a possibility to insert comment lines by preceding comments with `//` or writing `;comment` after the comment. These lines will be ignored by the parser. They can be written anywhere in the first paragraph.
- The first line indicates the author of the graph. This line has to start with the word "author" to be recognized by the parser.
- The second and third lines show the number of nodes and edges respectively. The parser only sees two consecutive integers at the end of the lines, so it is important that the number of nodes is provided before the number of edges.
- The next three lines are booleans that indicate whether the nodes / edges are labelled and if the graph is directed. It is important to stick to the order indicated in the example below.
- The blank line indicates the start of the node section. It is important to choose an integer as node id. The label can contain any symbol, except for `;`.
- The node section is separated from the edge section again by a blank line. If the `Directed graph` option is set to `False`, the edges are considered undirected. Internally, every edge's reverse is then added to the graph.

Let's look at a small example graph for illustration. The graph has 6 nodes which are all labelled with a 'c' and 6 edges which are not labelled. Your graphs need to have exactly this format to use this package. Especially the blank lines are important. You can find a decent number of further examples at [multivitamin/graph\\_examples](#). Now that you are familiar with the graph format, we can take a look at how to use the package.

```

AUTHOR: Michel K.
#nodes;6
#edges;6
Nodes labelled;True
Edges labelled;False
Directed graph;False

1;c
2;c
3;c
4;c
5;c
6;c

1;2
1;6
2;3

```

```
3;4
4;5
5;6
```

## 4 Graph alignment basic case

Consider the basic use without any optional flags for the start. You do not have to be in the directory of the package to use multiVitamin . To get an overview of all the flags you can type

```
$ multiVitamin -h
```

If you are experienced with command-line tools the information provided by multivitamin -h might be sufficient to get you started.

CHANGE THIS TO UPDATED VERSION

*%multiVitamin - A multiple alignment tool for graphs*

optional arguments:

```
-h, --help show this help message and exit
-a ALGORITHM, --algorithm ALGORITHM
                        choose an alignment-algorithm (BK or VF2) (default: BK)
                        Warning: VF2 is only suited if there is true graph-
                        subgraph isomorphism!
-s, --save-all save all the graphs produced during the alignment
                        The graphs are saved as "[newick].graph"
-t, --save-shorter save an additional version of the alignment graph with
                        much shorter node ids
-g, --save-guide save the guide tree in Newick-format as "tree.txt"
```

required arguments:

these arguments are mutually exclusive

```
-m FILES [FILES ...] provide .graph files for multiple alignment. '.' is a
                        valid input
-c COOPT [COOPT ...] provide *2* graphs which will be aligned. Co-optimals
                        will be saved in ./results.
-v VIEW get a visual representation of *1* given graph
```

There are three mutually exclusive flags. One of them is always required to make multiVitamin work. For the start we look at the case of a multiple alignment of graphs, which will be the standard use case. The flag for that is -m. The other flags are explained later. Just type "multitVitamin -m ;pathToGraphFile1.graph; ;pathToGraphFile2; ;optionalMoreGraphFiles;" in the command line. This aligns the given graphs with the Bron Kerbosch algorithmn. The

```

---GRAPH ABBREVIATIONS-----
ac1 >> acetylsalicylic_acid
be1 >> benzene
be2 >> benzoic_acid
sa1 >> salicylic_acid

---ALIGNMENT-----
*NODES (ID, LABEL, NEIGHBOURS)
be1:6.be2:5.sa1:5.ac1:5 'c' ( be1:1.be2:4.sa1:4.ac1:4, be1:5.be2:6.sa1:6.ac1:6 )
be1:3.be2:2.sa1:2.ac1:2 'c' ( be1:2.be2:3.sa1:3.ac1:3, be1:4.be2:1.sa1:1.ac1:1 )
be1:5.be2:6.sa1:6.ac1:6 'c' ( be1:4.be2:1.sa1:1.ac1:1, be1:6.be2:5.sa1:5.ac1:5 )
be1:1.be2:4.sa1:4.ac1:4 'c' ( be1:2.be2:3.sa1:3.ac1:3, be1:6.be2:5.sa1:5.ac1:5 )
be1:4.be2:1.sa1:1.ac1:1 'c' ( be1:3.be2:2.sa1:2.ac1:2, be1:5.be2:6.sa1:6.ac1:6 )
be1:2.be2:3.sa1:3.ac1:3 'c' ( be1:1.be2:4.sa1:4.ac1:4, be1:3.be2:2.sa1:2.ac1:2 )

*EDGES (ID, LABEL)
( be1:3.be2:2.sa1:2.ac1:2, be1:2.be2:3.sa1:3.ac1:3 ) ''
( be1:1.be2:4.sa1:4.ac1:4, be1:2.be2:3.sa1:3.ac1:3 ) ''
( be1:3.be2:2.sa1:2.ac1:2, be1:4.be2:1.sa1:1.ac1:1 ) ''
( be1:6.be2:5.sa1:5.ac1:5, be1:1.be2:4.sa1:4.ac1:4 ) ''
( be1:6.be2:5.sa1:5.ac1:5, be1:5.be2:6.sa1:6.ac1:6 ) ''
( be1:5.be2:6.sa1:6.ac1:6, be1:4.be2:1.sa1:1.ac1:1 ) ''

---NEWICK TREE-----
(benzene,(benzoic_acid,(acetylsalicylic_acid,salicylic_acid)))
*****
Successfully created the directory /home/nax/results
All files will be saved here.
Saved graph as be1-be2-ac1-sa1.graph
Saved graph id abbreviations as graph abbreviations.txt

```

Fig. 1: A sample output of a multiple graph alignment

resulting command line seems confusing at first put follows a clear structure and provides all the important information about the alignment.” in the command line. This aligns the given graphs with the Bron Kerbosch algorithmn. The resulting command line seems confusing at first put follows a clear structure and provides all the important information about the alignment. In order to see from which graph each node of the alignent origins, we use abbreviations of the names of the graph input files. A dictionary of the abbreviations of the original names of the graph files is shown first in the command line output. The you see the nodes of the alignment. Each line corresponds to one node of the alignment. A node in an alignment is represented by all nodes that got mathed to eachother. The aligned node is represented as follows: The id of th enode consits of the graph id (more specific of the abbreviation of the graph id) of the graph where the node origins from. This is followed by a colon, followed by the id of the original node. after a dot comes the next aligned node. After that follows a tab and the label of the aligned node (if the graphs were labelled). After another tab the neighbors of the node are listed. The next line shows the same for the next node of the alignment.

The next section shows the edges of the aligned graph. The format follows the format of the edges. It basically shows to nodes seperated by a comma and a tab.

Additionally to the command line output multiVitamin creates a result folder. This folder contains the aligned graph as .graph file. The first line of the .graph file shows the Newik tree of the mulitple alignment. This file could theoretically be used for further computation with multitVitamin. The results folder contains also a .txt file of the abbreviations of the graph files. These abbreviations are used to indicate in the alignment which node origins from which graph.

## 4.1 Graph Alignment with the Vf2

Let us still consider the basic use case with the required flag -m. This flag can be combined with all optional flags (see section advanced use). One of those optional flags is -a which specifies the algorithm that will be used for the alignment.

In order to use the VF2 algorithm for the alignment you have to use the flag -a with "vf2" as parameter in addition to the specification of the input files. So the command would be "multivitamin -f jpathToGraphFile1.graph i jpathToGraphFile2 i joptionalMoreGraphFiles i" -a vf2" This option is only suited for subgraph graph isomorphisms. So the smaller graph must always fit entirely in the bigger graph. Despite this restriction the Vf2 is able to align graphs based on the label of the node ids. So a structural isomorphism is not enough in this case, but the nodes that could aligned structurally must also fulfill some kind of semantic condition. This is useful to align chemical molecules that are represented as graphs. This condition can be determined by the user in the custom.py

### 4.1.1 Custom.py

The custom.py file can be found in the multivitamin directory of the root folder. This file contains a function the check\_semantics function, which by default returns always true. The code of the custom.py file looks as follows:

```
[numbers=left,xleftmargin=5mm]
def check_semantics( n, m ):
    '''This function is used in VF2 algorithm to decide,
    whether aligning to nodes is allowed from the
    label point of view.
    In the sample example below, two nodes will only
    be accepted as a legal matching, if the two
    labels are exactly the same.

    Insert your scoring logic below:'''

    # if n.label == m.label:
    #     return True
    # else:
    #     return False

    return True

def get_results_dir():
    '''define, where the result files will be saved.
    Default is a dir named 'results' in the current
    working directory'''

    return "/results"
```

If you untoggle the single comments and out-comment the return True, you get a simple

semantic check that only allows nodes to be aligned that have the same label. However you can go as crazy as you want with it.

Besides the `check_semantics` method you can also specify the path where your results should be saved. Just specify your preferred path as a string as return value from `get_results_dir()`. Before we jump into the supplementary classes, which you might want to modify for a highly sophisticated `check_semantics` function, we want to take a look at the other flag options. Note: A change in the supplementary classes is not recommended.

## 5 Advanced Usage

To make multiVitamin work you need one (exactly one) of the following flags: `-m`, `-c`, `-v`. The `-m` option has been discussed in depth in the previous sections. It can be used with every optional flag.

The `-v` flag allows for graphical representation of our `.graph` files. You need to provide a graph file. In most cases this will be the graph of the multiple alignment. Use: `"multiVitamin -v jfile.graph"` It is not compatible with any of the optional flags.

The `-c` flag works only for exactly two input files. It gives all the co-optimal alignments of the two given graphs. There exist cases where two graphs have more than one can be aligned in different, yet optimal ways. The `-c` flag calculates all these cases. Since `-c` only works with 2 graphs the optional flags are not helpful. Only the `-a` flag could be used to use the `vf2` algorithm. Note that this leads to high computation costs. Let us consider the optional flags now. If you want to save the guide tree in an extra file use `-g`. If you not only want the final multiple alignment graph, but also all alignments in between use `-s`. If you have three graphs for example `graph1`, `graph2` and `graph3`. Let us assume multiVitamin gives us a graph which multiple alignment is represented by this Newick tree: `((graph1,graph2),graph3)`. Now you with the option `-s` you will also get the alignment of `graph1` and `graph2` as a graph file.

## 6 Basic Classes

In order to implement the Bron Kerbosch and `Vf2` algorithm, we used three classes. A graph class, a node class and an edge class. In general the classes should not be altered by the user. However for the sophisticated user there may be cases in which he or she wants to make adjustments to these classes. Therefore we will give a brief overview of them. Figure 2 shows simple UML diagrams of our base classes. The most important methods and attributes will be explained.

### 6.1 Node

Our node objects are seen in the context of a graph. Thus every node contains information about its neighbours. Therefore a node has a `neighbours` attribute which is a set of Node objects. In case of directed graphs one can distinguish between in and out neighbours. Thus a node object has a separate set for these two kinds of neighbours. This is useful in the `Vf2` algorithm. The `mult_id` attribute gives an option to name the node in a multiple alignment.

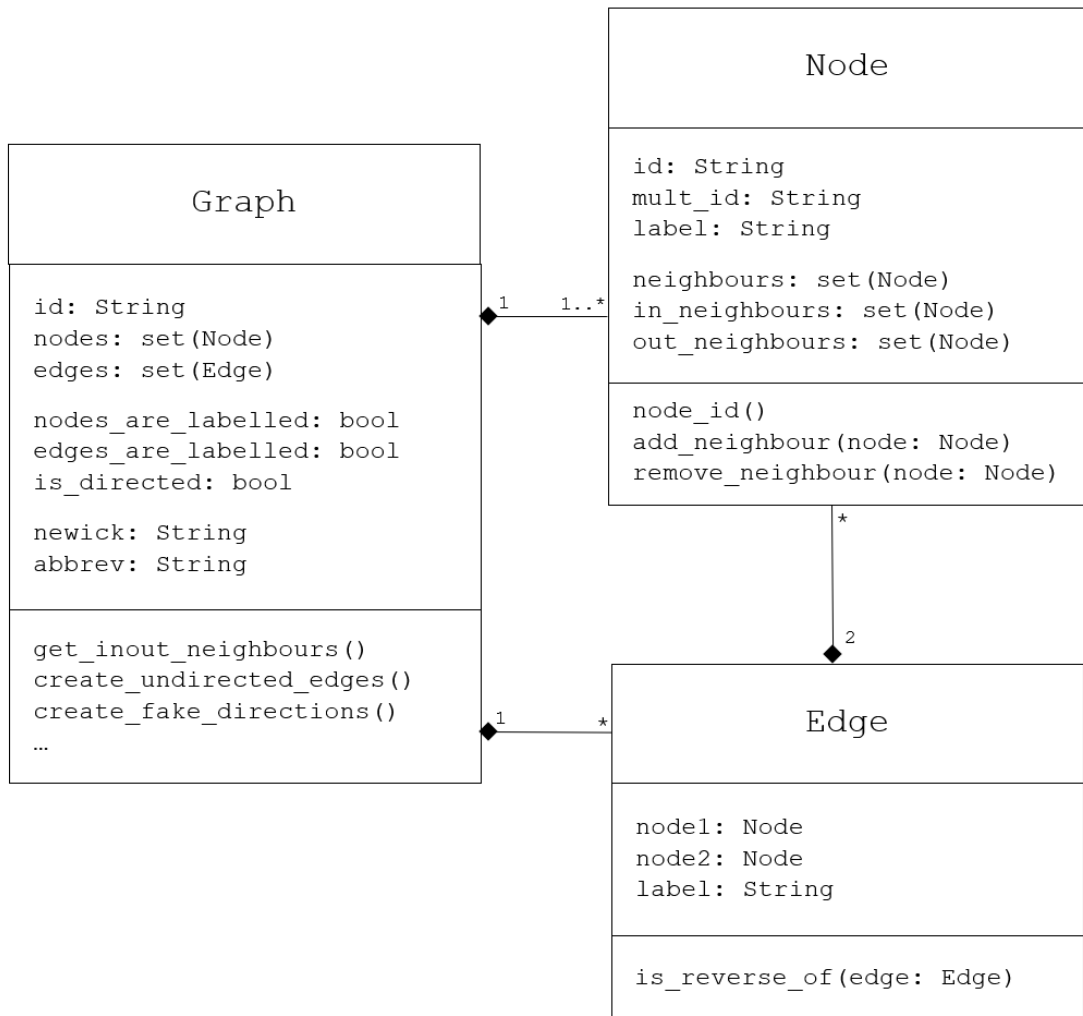


Fig. 2: A UML class diagram of the base classes

### 6.1.1 Edge

An Edge consists of two node objects and an optional label. For each Edge can be checked if it is the reverse of another edge. This only makes sense for directed graphs.

## 6.2 Graph Class

Besides the obvious attributes of nodes, edges and an id, a graph object also has an abbreviation which is TODO. Since the Vf2 only works with directed graphs, we gave our graphs the possibility to give the graph directions. Consider the Edge A-B. The `create_fake_directions` method now points node A to B and node B to A.

The method `create_undirected_edges` fills the edge set of the Graph with help of the neighbour set of the node class. The method `get_inout_neighbours` fills the in and outneighbour sets of the node objects in the node set.

All classes have built-in print methods for more readable output as well as built-in methods to compare the objects. So you can determine which graph is larger by the amount of nodes in the nodes sets and also determine if two graphs are equal based on their id.

Nodes can be also compared by its id and edges by the ids of their nodes.

## 7 Theoretical Foundations of the Algorithms

### 7.1 Bron Kerbosch

The Bron Kerbosch algorithm solves the problem of finding a maximal common induced subgraphs of two graphs G and H indirectly. This is done by finding the maximal clique in the modular product of G and H. The set of nodes of the modular product of G and H is the cartesian  $N(G) \times N(H)$ , with N being the set of all vertices of the corresponding graph. Any nodes  $(u,v)$  and  $(x,y)$ , with  $u,x \in G$  and  $v,y \in H$  in the modular product are adjacent if:

- u and x were neighbours in G and y and x were neighbours in H
- u and x were not neighbours in G and y and x were not neighbours in H

Before giving two graphs to the Bron Kerbosch to align them we need to build the modular product of these graphs. The rest of our implementation is straight forward:

```
def bk_pivot ( self, r, p, x ):
    if not any ( [p, x] ):
        self.results.append(r)
        return r
    pivot = self.find_max_pivot( p, x )
    for v in p[:]:
        if v in pivot.neighbours:
            continue
        r_ = r | {v}
        x_ = x & v.neighbours
```



```

p_ = [n for n in v.neighbours if n in p ]
self.bk_pivot ( r_, p_, x_ )
p.remove(v)
x.add(v)

```

The goal of the Bron Kerbosch ,is as mentioned, to find the maximal clique in the modular product. A clique is a subgraph where every node is connected to every other node (=complete graph). A maximal clique is a clique so that no node can added to the subgraph such that the subgraph remains complete.

At the beginning r and x are empty sets and theset p that contains all nodes of the modular product of the two input graphs. All nodes in p are candiates for extending the clique.

## 7.2 VF2

## 8 Profiling

To get an idea which algorithm is better sutied for which kind of graph (size, structure), we did a profiling study. Therefore we used three different kind of graphs: random graphs, wheel graphs and tree graphs. The graphs where generate with the help of the python package networkx []. We did the profiling for the follwoing number of nodes: (3,8,13,18,23,XX TODO). In case of the random graphs we created 10 different random graphs for each node count and took the average time per node count. The following figures show the results of our profiling study.