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About

This document contains examples on using some functions of CBioInfCpp.h library.

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Written by Sergey Chernouhov (chernouhov@rambler.ru).

Code for all Exercises is also represented at StudyingCBioInfCpp.cpp.

Exercise 1. Srings

Reading strings from a file, finding the shortest superstring (using greedy algorithm). Generating reverse complement of a string, calculating its GC-content, computing Hamming distance between two strings. Align 2 strings (edit distance alignment). Finding in a string all its substrings that have edit distance <=1 to some given string.

First of all we should to include CBioInfCpp to our code. Place the file CBioInfCpp.h to the folder where others libs to be included are. Or write the whole path to CBioInfCpp.h after #include.

So our code will start its building as follows:

```
#include <CBioInfCpp.h>
// Including CBioInfCpp
// Подключаем библиотеку CBioInfCpp
using namespace std;
int main()
{
   return 0;
}
```

We are ready. Let's read a collection of strings from a file. To do so one should download Inl(strings).txt to the project's folder or, another way, to any place (in this case one should write the whole path to the file for ifstream).

We shall read strings to std::vector DataS. The results of doing this exercise we shall out both to screen and to file Outl(strings).txt. So le's start. And to check the strings' reading we may using VectorCout (to out the result to screen) and VectorFout (to out to file):

```
#include <CBioInfCpp.h>
// Including CBioInfCpp
// Подключаем библиотеку CBioInfCpp
using namespace std;
int main()
```

```
{
     vector <string> DataS;
     // Container for strings to be read from file In1(strings).txt
     DataS.clear();
     ifstream fin1 ("In1(strings).txt");
     // If you place file not to "standart place" please write the entire
     ofstream fout1 ("Out1(strings).txt"); // File to write results for the
     Exercise 1.
     StringsRead(fin1, DataS); //Reading strings to vector DataS
     VectorCout(DataS); // Out DataS to screen
     VectorFout(DataS, fout1); // Out dataS to file
     fin1.close();
     fout1.close();
     return 0;
}
I hope, all it's ok. Now let's add two empty strings to our DataS. And after
we are going to try the function ShortestSuperstring (it finds the shortest
superstring using greedy algorithm). Let's do it.
int main()
     vector <string> DataS;
     // Container for strings to be read from file In1(strings).txt
     DataS.clear();
     ifstream fin1 ("In1(strings).txt");
     // If you place file not to "standard place" please write the entire
     path
     ofstream fout1 ("Out1(strings).txt"); // File to write results for the
     Exercise 1.
     StringsRead(fin1, DataS); //Reading strings to vector DataS
     VectorCout(DataS); // Out DataS to screen
     VectorFout(DataS, fout1); // Out DataS to file
     DataS.push back(""); //adding empty string
     DataS.push back(""); //another one
     string ShortestSuperstring = ShortSuperstringGr(DataS);
     // Generating shortest supersring using greedy algorithm
     cout<<"ShortestSuperstring: "<<ShortestSuperstring<<endl; // out it to</pre>
     fout1<<"ShortestSuperstring: "<<ShortestSuperstring<<endl; // out it</pre>
     too file
     fin1.close();
     fout1.close();
     return 0;
}
See it? And now some bioinformatics. We'll find reverse complement (the
function rp), compute GC-content, and transcribe DNA to RNA:
int main()
{
     vector <string> DataS;
     // Container for strings to be read from file In1(strings).txt
     DataS.clear();
     ifstream fin1 ("In1(strings).txt");
```

```
path
      ofstream fout1 ("Out1(strings).txt"); // File to write results for the
      Exercise 1.
      StringsRead(fin1, DataS); //Reading strings to vector DataS
      VectorCout(DataS); // Out DataS to screen
      VectorFout(DataS, fout1); // Out DataS to file
      DataS.push back(""); //adding empty string
      DataS.push back(""); //another one
      string ShortestSuperstring = ShortSuperstringGr(DataS);
      // Generating shortest superstring using greedy algorithm
      cout<<"ShortestSuperstring: "<<ShortestSuperstring<<endl; // out it to</pre>
      fout1<<"ShortestSuperstring< "<<ShortestSuperstring<<endl; // out it</pre>
      too file
      cout<<"Reverse complement of ShortestSuperstring:"</pre>
      <<rp>(ShortestSuperstring) <<endl;</p>
      // generating and printing to screen reverse complement of
      ShortestSuperstring
      fout1<<"Reverse complement of ShortestSuperstring: "</pre>
      <<rp>(ShortestSuperstring) <<endl;</p>
      // generating and printing to file reverse complement of
      ShortestSuperstring
      cout<<"GC-content of Reverse complement of ShortestSuperstring: "</pre>
      <<qcDRNA (rp(ShortestSuperstring))<<endl;
      // calculating GC-content of this reverse complement (to screen)
      fout1<<"GC-content of Reverse complement of ShortestSuperstring: "</pre>
      <<qcDRNA(rp(ShortestSuperstring))<<endl; // calculating GC-content of
      this reverse complement (to file)
      string RNAofShortestSuperstring;
      RNAfromDNA (ShortestSuperstring, RNAofShortestSuperstring); // now let's
      generate RNA string upon ShortestSuperstring and write it to
      RNAofShortestSuperstring
      cout<<"RNAofShortestSuperstring: "<<RNAofShortestSuperstring<<endl;</pre>
      fout1<<"RNAofShortestSuperstring: "<<RNAofShortestSuperstring<<endl;</pre>
      fin1.close();
      fout1.close();
      return 0;
}
Now it's time: (1) to evaluate Hamming distance between ShortestSuperstring
    RNAofShortestSuperstring, (2) to cut the last 2 symbols from
RNAofShortestSuperstring then
                                    align RNAofShortestSuperstring
RNAofShortestSuperstring (Edit Distance Alignment). Furthermore, we'll find
in the string ShortestSuperstring all its substrings, that have edit distance
to "ACA" <=1 (function FindMutatedRepeatsED).
int main()
      vector <string> DataS;
      // Container for strings to be read from file In1(strings).txt
      DataS.clear();
      ifstream fin1 ("In1(strings).txt");
      // If you place file not to "standard place" please write the entire
      ofstream fout1 ("Out1(strings).txt"); // File to write results for the
      Exercise 1.
```

// If you place file not to "standard place" please write the entire

```
StringsRead(fin1, DataS); //Reading strings to vector DataS
VectorCout(DataS); // Out DataS to screen
VectorFout(DataS, fout1); // Out DataS to file
DataS.push back(""); //adding empty string
DataS.push back(""); //another one
string ShortestSuperstring = ShortSuperstringGr(DataS);
// Generating shortest superstring using greedy algorithm
cout<<"ShortestSuperstring: "<<ShortestSuperstring<<endl; // out it to</pre>
fout1<<"ShortestSuperstring: "<<ShortestSuperstring<<end1; // out it</pre>
too file
cout<<"Reverse complement of ShortestSuperstring:"</pre>
<<rp><<rp (ShortestSuperstring) <<endl;</p>
\ensuremath{//} generating and printing to screen reverse complement of
ShortestSuperstring
fout1<<"Reverse complement of ShortestSuperstring: "</pre>
<<rp><<rp (ShortestSuperstring) <<endl;</p>
// generating and printing to file reverse complement of
ShortestSuperstring
cout<<"GC-content of Reverse complement of ShortestSuperstring: "</pre>
<<gcDRNA (rp(ShortestSuperstring))<<endl;
// calculating GC-content of this reverse complement (to screen)
fout1<<"GC-content of Reverse complement of ShortestSuperstring: "</pre>
<<qcDRNA(rp(ShortestSuperstring))<<endl; // calculating GC-content of
this reverse complement (to file)
string RNAofShortestSuperstring;
RNAfromDNA (ShortestSuperstring, RNAofShortestSuperstring); // now let's
generate RNA string upon ShortestSuperstring and write it to
RNAofShortestSuperstring
cout<<"RNAofShortestSuperstring: "<<RNAofShortestSuperstring<<endl;</pre>
fout1<<"RNAofShortestSuperstring: "<<RNAofShortestSuperstring<<endl;</pre>
let's calculate Hamming distance between ShortestSuperstring and
RNAofShortestSuperstring
cout<<"Hamming Distance between ShortestSuperstring and</pre>
RNAofShortestSuperstring: "<<HmDist(ShortestSuperstring,
RNAofShortestSuperstring) << endl;</pre>
fout1<<"Hamming Distance between ShortestSuperstring and</pre>
RNAofShortestSuperstring: "<<HmDist(ShortestSuperstring,
RNAofShortestSuperstring) << endl;
RNAofShortestSuperstring.pop back();
// Deleting the last symbol from RNAofShortestSuperstring
RNAofShortestSuperstring.pop back();
// Another time
string sr1, sr2;
// These strings will contain results of Alignment (minimizing Edit
Distance)
cout<<"Edit Distance between ShortestSuperstring and</pre>
RNAofShortestSuperstring: "<<EditDistA(ShortestSuperstring,
RNAofShortestSuperstring, sr1, sr2) << endl;
fout1<<"Edit Distance between ShortestSuperstring and</pre>
RNAofShortestSuperstring: "<<EditDistA(ShortestSuperstring,
RNAofShortestSuperstring, sr1, sr2) << endl;
cout<<"Edit Distance Alignment: "<<endl<<sr1<<endl<<sr2<<endl;</pre>
fout1<<"Edit Distance Alignment: "<<endl<<sr1<<endl<<sr2<<endl;</pre>
```

```
// Now let's find in the string ShortestSuperstring all its substrings
      that have edit distance to "ACA" <=1.
     std::set <std::pair <int, int>> Result;
      // The function FindMutatedRepeatsED returns results as set of pairs
     there the first one is a starting position of substring in the string,
     and the second is its lenght
     FindMutatedRepeatsED ("ACA", ShortestSuperstring, 1, Result); //done
     cout<<"Substrings of ShortestSuperstring that have EditDistance to</pre>
     string ACA <=1 :"<<endl;</pre>
      fout1<<"Substrings of ShortestSuperstring that have EditDistance to
     string ACA <=1 :"<<endl;</pre>
      for (auto it=Result.begin(); it!=Result.end(); it++)
              cout<<ShortestSuperstring.substr((*it).first,</pre>
             *it).second) <<endl; //outing substrings to screen
             fout1<<ShortestSuperstring.substr((*it).first,</pre>
              (*it).second) << endl; //outing substrings to file</pre>
          }
      fin1.close();
      fout1.close();
      return 0;
}
```

Exercise 2. Reading graphs from file, outing graphs to screen and to file, finding Eulerian Cycle

If we start from the very beginning we should include CBioInfCpp as it has been said above. Then we'll have this:

```
Koд:
#include <CBioInfCpp.h>
// Including CBioInfCpp
// Подключаем библиотеку CBioInfCpp
using namespace std;
int main()
{
    return 0;
}
```

See the result in Outl(strings).txt.

Lel's read from files: unweighted graph A1, weighted graph (weights are int) A2, weighted graph (weights are double) A3. Note that A1 and A2 may be read to vector <int> and A3 - to pair < vector<int>, vector<double>> A3. Also note that numbers assigned to vertices may be negative too.

To start we should download files that contain these graphs (function UWGraphRead for unweighted graph, function WGraphRead for weighted graph). We should place these files to the project's folder or, another way, to any other place (in this case one should write the whole path to the file for ifstream). These are required files:

InUnWeightedGraph.txt; // An unweighted graph is here (edge list format)

```
InWeightedGraphIntegers.txt // A weighted graph is here (edge list format,
weights are integers)
InWeightedGraphDoubles.txt // A weighted graph is here (edge list format,
weights are double)
After we may check ourselves using functions GraphCout (to out graph to
screen) and GraphFout (to out graph to file).
#include <CBioInfCpp.h>
// Including CBioInfCpp
// Подключаем библиотеку CBioInfCpp
using namespace std;
int main()
      ifstream fin2 1 ("InUnWeightedGraph.txt"); // An unweighted graph is
      here (edge list format)
      ifstream fin2 2 ("InWeightedGraphIntegers.txt"); // A weighted graph
      is here (edge list format, weights are integers)
      ifstream fin2 3 ("InWeightedGraphDoubles.txt"); // A weighted graph is
      here (edge list format, weights are double)
      ofstream fout2 ("Out2(Graphs1).txt"); // File to write results for the
      Exercise 2.
      vector <int> A1; //Container for unweighted graph A1
      UWGraphRead(fin2 1, A1);
      cout<<"Here is unweighted graph A1: "<<endl;</pre>
      GraphCout(A1, false); //Out graph to screen, as it is unweighted the
      second parameter should be "false"
      fout2<<"Here is unweighted graph A1: "<<endl;</pre>
      GraphFout(A1, false, fout2); //Out graph to file, as it is unweighted
      the second parameter should be "false"
      vector <int> A2; //Container for weighted graph A2, weights are
      integers
      WGraphRead(fin2 2, A2);
      cout<<"Here is weighted graph A2: "<<endl;</pre>
      GraphCout(A2, true); //Out graph to screen, as it is weighted the
      second parameter should be "true"
      fout2<<"Here is weighted graph A2: "<<endl;</pre>
      GraphFout(A2, true, fout2); //Out graph to file, as it is weighted the
      second parameter should be "true"
      pair < vector<int>, vector<double>> A3; //Container for unweighted
      graph A3, weights are double
      WGraphRead(fin2 3, A3);
      cout<<"Here is weighted graph A3: "<<endl;</pre>
      GraphCout(A3); //Out graph to screen, as it is set by pair A3 it may be
      weighted only
      fout2<<"Here is weighted graph A3: "<<endl;</pre>
      GraphFout(A3, fout2); //Out graph to file, as it is set by pair A3 it
      may be weighted only
      fout2.close();
      fin2_1.close();
      fin2_2.close();
      fin2 3.close();
   return 0;
}
```

Now we know how to read graphs and to out them to screen/ file. As the A1, A2, A3 differs one from another by edge's weight only, their Eulerian cycles/ paths should be the same, aren't they? Let's check it using EPathDGraph (note that it considers input graphs as <u>directed</u>). Also note that EPathDGraph will fill for us 2 vectors, i.e. vector R (Eulerian cycle (in this case the function returns 1)/ path (and in this - it returns 2) itself) and vector I (it will contain all isolated vertices). If there is no Eulerian cycle/ path or input data are incorrect the function returns -1 and empty R and I.

We should also remember that vertices may have negative numbers assigned.

```
#include <CBioInfCpp.h>
// Including CBioInfCpp
// Подключаем библиотеку CBioInfCpp
using namespace std;
int main()
      ifstream fin2 1 ("InUnWeightedGraph.txt"); // An unweighted graph is
      here (edge list format)
      ifstream fin2 2 ("InWeightedGraphIntegers.txt"); // A weighted graph
      is here (edge list format, weights are integers)
      ifstream fin2 3 ("InWeightedGraphDoubles.txt"); // A weighted graph is
      here (edge list format, weights are double)
      ofstream fout2 ("Out2(Graphs1).txt"); // File to write results for the
      Exercise 2.
      vector <int> A1; //Container for unweighted graph A1
      UWGraphRead(fin2 1, A1);
      cout<<"Here is unweighted graph A1: "<<endl;</pre>
      GraphCout(A1, false); //Out graph to screen, as it is unweighted the
      second parameter should be "false"
      fout2<<"Here is unweighted graph A1: "<<endl;</pre>
      GraphFout(A1, false, fout2); //Out graph to file, as it is unweighted
      the second parameter should be "false"
      vector <int> A2; //Container for weighted graph A2, weights are
      integers
      WGraphRead(fin2 2, A2);
      cout<<"Here is weighted graph A2: "<<endl;</pre>
      GraphCout(A2, true); //Out graph to screen, as it is weighted the
      second parameter should be "true"
      fout2<<"Here is weighted graph A2: "<<endl;</pre>
      GraphFout(A2, true, fout2); //Out graph to file, as it is weighted the
      second parameter should be "true"
      pair < vector<int>, vector<double>> A3; //Container for unweighted
      graph A3, weights are double
      WGraphRead(fin2 3, A3);
      cout<<"Here is weighted graph A3: "<<endl;</pre>
      GraphCout(A3); //Out graph to screen, as it is set by pair A3 it may be
      weighted only
      fout2<<"Here is weighted graph A3: "<<endl;</pre>
      GraphFout(A3, fout2); //Out graph to file, as it is set by pair A3 it
      may be weighted only
      vector <int> R; //Container for Eulerian cycle/path to be found
      vector <int> I; //Container for isolated vertices to be found
      cout<<EPathDGraph (A1, R, false, I)<<endl; //as A1 is unweighted we</pre>
      should set 3d parameter as "false"
      fout2<<EPathDGraph (A1, R, false, I)<<endl;</pre>
      // EPathDGraph returns value "1" if Eulerian cycle has been found or
      value "2" if Eulerian path has been found
```

```
// or "-1" together with empty R and Isolates if no cycle/ path found.
      //Note that EPathDGraph considers graps as directed ones.
      cout<< "Eulerian cycle of graph A1: "<<endl;</pre>
      VectorCout(R); // vector R - to screen
      cout<< "Isolated vertices of graph A1: "<<endl;</pre>
      VectorCout(I); // vector I - to screen
      fout2<< "Eulerian cycle of graph A1: "<<endl;</pre>
      VectorFout(R, fout2); // vector R - to file
      fout2<< "Isolated vertices of graph A1: "<<endl;</pre>
      VectorFout(I, fout2); // vector I - to file
      cout<<EPathDGraph (A2, R, true, I)<<endl; //as A1 is unweighted we</pre>
      should set 3d parameter as "true"
      fout2<<EPathDGraph (A2, R, true, I)<<endl;</pre>
      // EPathDGraph returns value "1" if Eulerian cycle has been found or
      value "2" if Eulerian path has been found
      // or "-1" together with empty R and Isolates if no cycle/ path found.
      cout<< "Eulerian cycle of graph A2: "<<endl;</pre>
      VectorCout(R); // vector R - to screen
      cout<< "Isolated vertices of graph A2: "<<endl;</pre>
      VectorCout(I); // vector I - to screen
      fout2<< "Eulerian cycle of graph A2: "<<endl;</pre>
      VectorFout(R, fout2); // vector R - to file
      fout2<< "Isolated vertices of graph A2: "<<endl;</pre>
      VectorFout(I, fout2); // vector I - to file
      cout<<EPathDGraph (A3, R, I)<<endl; //as A3 is set by pair it may be</pre>
      only weighted
      fout2<<EPathDGraph (A3, R, I)<<endl;</pre>
      // EPathDGraph returns value "1" if Eulerian cycle has been found or
      value "2" if Eulerian path has been found
      // or "-1" together with empty R and Isolates if no cycle/ path found.
      cout<< "Eulerian cycle of graph A3: "<<endl;</pre>
      VectorCout(R); // vector R - to screen
      cout<< "Isolated vertices of graph A3: "<<endl;</pre>
      VectorCout(I); // vector I - to screen
      fout2<< "Eulerian cycle of graph A3: "<<endl;</pre>
      VectorFout(R, fout2); // vector R - to file
      fout2<< "Isolated vertices of graph A3: "<<endl;</pre>
      VectorFout(I, fout2); // vector I - to file
      fout2.close();
      fin2 1.close();
      fin2 2.close();
      fin2 3.close();
   return 0;
The result of Ex.2 we may see in Out2(Graphs1).txt
```

Exercise 3. Checking if 2 graphs are isomorphic; finding all subgraphs that are isomorphic to given template graph

Now let's turn to (1) the graph isomorphism problem and (2) the problem of finding of all subgraphs in a graph A that are isomorphic to given template graph B. CBioInfCpp have the function **SubGraphsInscribed** for it. Some info on evolution of this function and its testing is here:

https://github.com/chernouhov/CBioInfCpp-0-/tree/master/TestsGraphIsomorphismInfo

https://github.com/chernouhov/CBioInfCpp-0/tree/master/TestsIsomorphicSubGraphsFinding

https://doi.org/10.24108/preprints-3111977

```
So let's start as above:

Koд:
#include <CBioInfCpp.h>
// Including CBioInfCpp
// Подключаем библиотеку CBioInfCpp
using namespace std;
int main()
{
    return 0;
}
```

We should read "more larger" unweigted graph A from a file InA.txt and template graph B1 from InB1.txt. As we have already known we should use the function UWGraphRead for it. But before we should download these files to the project's directory or to some other place (in this case one should write the whole path to the file for ifstream).

Also we'll create graph B2 as follows. First let the graph B2 = B1 and then let's add integer 2 to all numbers that are assigned to each vertex of B2 (we may do so using the function RenumVGraph). So B2 will be isomorphic to B1 and we will check it too.

```
#include <CBioInfCpp.h>
// Including CBioInfCpp
// Подключаем библиотеку CBioInfCpp
using namespace std;
int main()
      vector <int> A; // Container for unweighted graph A ("more large one")
      is here (edge list format)
      vector <int> B1; // Container for unweighted graph B1 is here (edge
      list format)
      vector <int> B2; // Container for unweighted graph B2 is here (edge
      list format)
      ifstream fin3 A ("InA.txt"); // Unweighted graph A is here (edge list
      ifstream fin3 B ("InB1.txt"); // Unweighted graph B1 is here (edge list
      format)
      ofstream fout3 ("Out3(Graphs2).txt"); // File to write results for the
      Exercise 3.
      UWGraphRead(fin3 A, A); // Reading unweighted graph A from a file
      UWGraphRead(fin3 B, B1); // Reading unweighted graph B1 from a file
      B2=B1; // Let's graph B2=B1
      RenumVGraph(B2,2, false); // Rename all vertices of B2: let's add 2 to
      their numbers; the 3d parameter is set as false as graph is unweighted.
      cout<<"Here is unweighted graph B1: "<<endl;</pre>
      GraphCout(B1, 0); //Out graph to screen, as it is unweighted the second
      parameter should be "false" i.e. 0
      cout<<"Here is unweighted graph B2: "<<endl;</pre>
```

```
GraphCout(B2, 0); //Out graph to screen, as it is unweighted the second
      parameter should be "false" i.e. 0
      fout3<<"Here is unweighted graph B1: "<<endl;</pre>
      GraphFout(B1, 0, fout3); //Out graph to file, as it is unweighted the
      second parameter should be "false" i.e. 0
      fout3<<"Here is unweighted graph B2: "<<endl;</pre>
      GraphFout(B2, 0, fout3); //Out graph to file, as it is unweighted the
      second parameter should be "false" i.e. 0
    return 0;
}
Now let's check if B1 is isomorphic to B2. We'll check it for two cases:
graphs are directed and graphs are undirected. We may set this condition via
bool "directed" as a parameter of the function SubGraphsInscribed.
The function SubGraphsInscribed returns the number of subgraphs nave been
found; the subgraphs themselves will be contained here in set<vector <int>>
SubGraphs. If input graphs are isomorphic it will find the only subgraph,
i.e. the "larger" graph itself.
So, let's try:
#include <CBioInfCpp.h>
// Including CBioInfCpp
// Подключаем библиотеку CBioInfCpp
using namespace std;
int main()
      vector <int> A; // Container for unweighted graph A ("more large one")
      is here (edge list format)
      vector <int> B1; // Container for unweighted graph B1 is here (edge
      list format)
     vector <int> B2; // Container for unweighted graph B2 is here (edge
      list format)
      ifstream fin3 A ("InA.txt"); // Unweighted graph A is here (edge list
      ifstream fin3 B ("InB1.txt"); // Unweighted graph B1 is here (edge list
      format)
      ofstream fout3 ("Out3(Graphs2).txt"); // File to write results for the
      Exercise 3.
      UWGraphRead(fin3 A, A); // Reading unweighted graph A from a file
      UWGraphRead(fin3 B, B1); // Reading unweighted graph B1 from a file
      B2=B1; // Let's graph B2=B1
      RenumVGraph(B2,2, false); // Rename all vertices of B2: let's add 2 to
      their numbers; the 3d parameter is set as false as graph is unweighted.
      cout<<"Here is unweighted graph B1: "<<endl;</pre>
      GraphCout(B1, 0); //Out graph to screen, as it is unweighted the second
     parameter should be "false" i.e. 0
      cout<<"Here is unweighted graph B2: "<<endl;</pre>
      GraphCout(B2, 0); //Out graph to screen, as it is unweighted the second
     parameter should be "false" i.e. 0
      fout3<<"Here is unweighted graph B1: "<<endl;</pre>
      GraphFout(B1, 0, fout3); //Out graph to file, as it is unweighted the
      second parameter should be "false" i.e. 0
      fout3<<"Here is unweighted graph B2: "<<endl;</pre>
      GraphFout(B2, 0, fout3); //Out graph to file, as it is unweighted the
      second parameter should be "false" i.e. 0
      set<vector <int>> SubGraphs; // Here will be subgraphs found
```

```
bool directed = true; //lets consider both graphs as directed ones
      cout<<SubGraphsInscribed (B1, B2, SubGraphs, directed, 0, 1, 0)<<endl;</pre>
      //trying to find in B1 all subgraphs that are isomorphic to B2
      fout3<<SubGraphsInscribed (B1, B2, SubGraphs, directed, 0, 1, 0)<<endl;</pre>
      // it should be 1 as they are isomorphic themselves
      for (auto it = SubGraphs.begin(); it!= SubGraphs.end(); it++) // now
      let's out all found (sub) graphs:
        cout<<"isomorphic (sub)graph found (directed graph case):"<<"</pre>
       "<<(*(it)).size()/2<<" edges"<<endl; // to screen
       GraphCout(*(it), false);
        fout3<<"isomorphic (sub)graph found: (directed graph case)"<<"</pre>
       "<<(*(it)).size()/2<<" edges"<<endl; // to file
        GraphFout(*(it), false, fout3);
      }
      directed = 0; //lets consider both graphs as undirected ones
      cout<<SubGraphsInscribed (B1, B2, SubGraphs, directed, 0, 1, 0)<<endl;</pre>
      fout3<<SubGraphsInscribed (B1, B2, SubGraphs, directed, 0, 1, 0)<<endl;</pre>
      for (auto it = SubGraphs.begin(); it!= SubGraphs.end(); it++)
        cout<<"isomorphic (sub)graph found (undirected graph case):"<<"</pre>
       "<<(*(it)).size()/2<<" edges"<<endl;
        GraphCout(*(it),false);
        fout3<<"isomorphic (sub)graph found (directed graph case):"<<"</pre>
       "<<(*(it)).size()/2<<" edges"<<endl;
        GraphFout(*(it), false, fout3);
    return 0;
}
Now we may find in the graph A all its subgraphs that are isomorphic to
template graph B1. We should consider the case both graphs are directed (8
subgraphs will be found) and the case both graphs are undirected (24
subgraphs will be found).
Here is the code for the whole Ex.3:
#include <CBioInfCpp.h>
// Including CBioInfCpp
// Подключаем библиотеку CBioInfCpp
using namespace std;
int main()
      vector <int> A; // Container for unweighted graph A ("more large one")
      is here (edge list format)
      vector <int> B1; // Container for unweighted graph B1 is here (edge
      list format)
      vector <int> B2; // Container for unweighted graph B2 is here (edge
      list format)
     ifstream fin3 A ("InA.txt"); // Unweighted graph A is here (edge list
      ifstream fin3 B ("InB1.txt"); // Unweighted graph B1 is here (edge list
      ofstream fout3 ("Out3(Graphs2).txt"); // File to write results for the
      Exercise 3.
```

```
UWGraphRead(fin3 A, A); // Reading unweighted graph A from a file
UWGraphRead(fin3 B, B1); // Reading unweighted graph B1 from a file
B2=B1; // Let's graph B2=B1
RenumVGraph(B2,2, false); // Rename all vertices of B2: let's add 2 to
their numbers; the 3d paremeter is set as false as graph is unweighted.
cout<<"Here is unweighted graph B1: "<<endl;</pre>
GraphCout(B1, 0); //Out graph to screen, as it is unweighted the second
parameter should be "false" i.e. 0
cout<<"Here is unweighted graph B2: "<<endl;</pre>
GraphCout(B2, 0); //Out graph to screen, as it is unweighted the second
parameter should be "false" i.e. 0
fout3<<"Here is unweighted graph B1: "<<endl;</pre>
GraphFout(B1, 0, fout3); //Out graph to file, as it is unweighted the
second parameter should be "false" i.e. 0
fout3<<"Here is unweighted graph B2: "<<endl;</pre>
GraphFout(B2, 0, fout3); //Out graph to file, as it is unweighted the
second parameter should be "false" i.e. 0
set<vector <int>> SubGraphs; // Here will be subgraphs found
bool directed = true; //lets consider both graphs as directed ones
cout<<SubGraphsInscribed (B1, B2, SubGraphs, directed, 0, 1, 0)<<endl;</pre>
//trying to find in B1 all subgraphs that are isomorphic to B2
fout3<<SubGraphsInscribed (B1, B2, SubGraphs, directed, 0, 1, 0)<<endl;</pre>
// it should be 1 as they are isomorphic themselves
for (auto it = SubGraphs.begin(); it!= SubGraphs.end(); it++) // now
let's out all found (sub)graphs:
 cout<<"isomorphic (sub)graph found (directed graph case):"<<"</pre>
 "<<(*(it)).size()/2<<" edges"<<endl; // to screen
 GraphCout(*(it),false);
  fout3<<"isomorphic (sub)graph found: (directed graph case)"<<"</pre>
 "<<(*(it)).size()/2<<" edges"<<endl; // to file
  GraphFout(*(it), false, fout3);
}
directed = 0; //lets consider both graphs as undirected ones
cout<<SubGraphsInscribed (B1, B2, SubGraphs, directed, 0, 1, 0)<<endl;</pre>
fout3<<SubGraphsInscribed (B1, B2, SubGraphs, directed, 0, 1, 0)<<endl;</pre>
for (auto it = SubGraphs.begin(); it!= SubGraphs.end(); it++)
  cout<<"isomorphic (sub)graph found (undirected graph case):"<<"</pre>
 "<<(*(it)).size()/2<<" edges"<<endl;
  GraphCout(*(it), false);
  fout3<<"isomorphic (sub)graph found (directed graph case):"<<"</pre>
 "<<(*(it)).size()/2<<" edges"<<endl;
  GraphFout(*(it), false, fout3);
//now let's find in A all subgraphs that are isomorphic to graph B1
//first of all let's out them
cout<<"Here is unweighted graph A: "<<endl;</pre>
GraphCout(A, 0); //Out graph to screen, as it is unweighted the second
parameter should be "false" i.e. 0
cout<<"Here is unweighted graph B1: "<<endl;</pre>
```

```
GraphCout(B1, 0); //Out graph to screen, as it is unweighted the second
parameter should be "false" i.e. 0
fout3<<"Here is unweighted graph A: "<<endl;</pre>
GraphFout(A, 0, fout3); //Out graph to file, as it is unweighted the
second parameter should be "false" i.e. 0
fout3<<"Here is unweighted graph B1: "<<endl;</pre>
GraphFout(B1, 0, fout3); //Out graph to file, as it is unweighted the
second parameter should be "false" i.e. 0
// directed graph case
directed = true; //lets consider both graphs (A and B1) as directed
ones
cout<<SubGraphsInscribed (A, B1, SubGraphs, directed, 0, 1, 0)<<"</pre>
subraphs of A that are isomorphic to B1 is found: "<<endl; //trying to
find in A all subgraphs that are isomorphic to {\tt B1}
fout3<<SubGraphsInscribed (A, B1, SubGraphs, directed, 0, 1, 0)<<"</pre>
subraphs of A that are isomorphic to B1 is found:"<<endl;
for (auto it = SubGraphs.begin(); it!= SubGraphs.end(); it++) // now
let's out all found (sub) graphs:
 cout<<"isomorphic (sub)graph found (directed graph case):"<<"</pre>
 "<<(*(it)).size()/2<<" edges"<<endl; // to screen
 GraphCout(*(it), false); //Out graph to screen, as it is unweighted
 the second parameter should be "false"
 fout3<<"isomorphic (sub)graph found: (directed graph case)"<<"</pre>
 "<<(*(it)).size()/2<<" edges"<<endl; // to file
 GraphFout(*(it), false, fout3); //Out graph to file, as it is
 unweighted the second parameter should be "false"
cout<<SubGraphs.size()<<" subraphs of A that are isomorphic to B1 is</pre>
found"<<endl;
fout3<<SubGraphs.size()<<"subraphs of A that are isomorphic to B1 is</pre>
found"<<endl;</pre>
// undirected graph case
directed = 0; //lets consider both graphs (A and B1) as undirected ones
cout<<SubGraphsInscribed (A, B1, SubGraphs, directed, 0, 1, 0)<<"</pre>
subraphs of A that are isomorphic to B1 is found:"<<endl; //trying to
find in A all subgraphs that are isomorphic to B1
fout3<<SubGraphsInscribed (A, B1, SubGraphs, directed, 0, 1, 0)<<"</pre>
subraphs of A that are isomorphic to B1 is found:"<<endl;
for (auto it = SubGraphs.begin(); it!= SubGraphs.end(); it++)
 cout<<"isomorphic (sub)graph found (undirected graph case):"<<"</pre>
 "<<(*(it)).size()/2<<" edges"<<endl;
 GraphCout(*(it), false); //Out graph to screen, as it is unweighted
 the second parameter should be "false"
 fout3<<"isomorphic (sub)graph found (directed graph case):"<<"</pre>
 "<<(*(it)).size()/2<<" edges"<<endl;
 GraphFout(*(it), false, fout3); //Out graph to file, as it is
 unweighted the second parameter should be "false"
cout<<SubGraphs.size()<<" subraphs of A that are isomorphic to B1 is</pre>
found"<<endl;
fout3<<SubGraphs.size()<<"subraphs of A that are isomorphic to B1 is</pre>
found"<<endl;</pre>
```

```
return 0;
}
The result of Ex.3 we may see in Out3(Graphs2).txt
```