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Methods for community detection in multi-layer networks

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Introduction

Complex network theory is an evolution of graph theory, whose origin traces back to Euler's famous publication of 1736 on the "Seven Bridges of Königsberg" [1]. Since then a lot has been learned about graphs and their mathematical properties. Networks have emerged as effective tools for modelling and analysing complex systems of interacting entities. Graphs arise naturally in many disciplines, such as social networks (which arise via offline and/or online interactions) [2], information networks (i.e. hyperlinks between webpages on the World Wide Web) [3], infrastructure networks (i.e. transportation routes between cities) [4], and biological networks (i.e. metabolic interactions between cells or proteins, food webs) [5].

One of the most relevant issue of graphs representing real systems is the identification of communities, or clustering, i.e. the organization of vertices in clusters or modules, with more edges connecting vertices of the same group and fewer edges joining vertices of different groups. Community detection in large networks is potentially very useful. Nodes belonging to the same community might have other properties in common. Society offers a variety of possible group organizations: families, working and friendship circles, villages, towns, nations. In recent years, the spread of Internet has led to the development of virtual communities, which live on the Web. Social communities have been studied for a long time [2], but they also occur in many networked systems from biology, computer science, engineering, economics, etc. For example, biological systems are organized hierarchically with networks of communities interacting at various levels, from ecosystems to networks of synaptic connections between neurons, up to gene and metabolic networks. In protein-protein interaction networks, communities correspond to group proteins having the same specific function within the cell [5], in the graph of the World Wide Web they may represent groups of pages dealing with the same topics [6], in metabolic networks they may be related to functional modules [7]. Communities can have concrete applications. For example, clustering Web clients who have similar interests and are geographically close could be served by a dedicated server, improving the performance of services provided on the World Wide Web [3]; or clusters of large graphs can be used to create data structures and handle navigational queries, like path searches [8].

Community detection is important also to classify vertices, according to their structural position in the modules. So, vertices with a central position in their clusters, i.e. sharing many edges with the other nodes of the group, may have an important function of control and stability within the group; vertices lying at the boundaries between clusters are important for leading the relationships between communities.

Another important aspect related to community structure is the hierarchical organization. Real networks are usually composed by communities including smaller communities, which in turn include smaller communities, etc. For example, the human body is composed by organs, organs by tissues, tissues by cells, and so on.

Community detection is a very hard problem and not yet satisfactorily solved, despite the extensive studies in the literature. Many algorithms have been developed, however they are designed for single-layer networks. This assumption is almost always a simplification, which can lead to misleading results. Recent advances in the study of networked systems have shown that the interconnected world is composed of networks that are coupled to one another through different layers, where each layer represents one of many possible types of interactions. For example, individuals have multiple relationships in social networks, such as economic, political, and financial. In biology, protein interaction networks consist of seven distinct layers that account for different genetic and physical interactions. Analysing multi-layer networks is of great importance because many interesting patterns cannot be obtained by analysing single-layer networks. For example, in multi-layer cancer networks where each layer corresponds to a specific clinical stage, a community represents a biological pathway that is critical for cancer diagnosis and therapy. However, it is hard to extract communities in multi-layer networks because of two reasons. First, multi-layer communities cannot be easily quantified because analyses on these multi-layer networks remain lacking. Second, complexity of multiple networks poses a challenge on finding algorithms for identifying communities in the multi-layer case. Despite these difficulties, great efforts have been devoted to the extraction of multi-layer communities. Most of the current algorithms either reduce multi-layer networks into a single-layer network or extend the algorithms for single-layer networks by using consensus clustering. However, these algorithms are criticized for their low accuracy because they either cannot preserve the community structure in compressed networks or ignore the connection among various layers. To overcome these problems, we must simultaneously take into account multiple layers.

In this thesis project, we propose multiple algorithms for community detection in multi-layer networks. The algorithms are based on the popular Louvain heuristic method for single-layer networks [9], which is a locally greedy modularity-increasing sampling process over the set of partitions.

The work is organized as follows.

Chapter 1 presents a broad description of the community detection problem. Firstly, we introduce some primary notions of graph theory and then we focus on the main problems that arise from community detection.

Chapter 2 contains a summary of the most relevant methods related to the study of community detection. In the first Section we focus on methods for single-layer graphs. In the second Section we study algorithms proposed for the multi-layer case.

Chapter 3 is the main contribution of this thesis and it presents our algorithms in detail. The first algorithm is called Louvain extension and the second one is named Louvain Multiobjective. Both of them try to extend the Louvain heuristic method to the multi-layer case. We present different variants of these two algorithms.

Chapter 4 shows some results obtained testing the methods. We did some tests on both artificial and real world networks to better compare the performances of the algorithms.

Finally, Chapter 5 reports some conclusions taking into consideration both the formulation and the experimentation of the methods.

Contents

Chapter 1

Problem Statement

Graph theory is extremely useful as representation of a wide variety of systems in different areas, such as biology [5], sociology [2], technology [3], and many others. All these networks can be studied as graphs, thus graph analysis has become crucial to understand the features of these systems.

A graph G is composed by a pair of sets (N, E) where N is the set of nodes and E is the set of edges and it is a subset of all the possible pairs of nodes in V.

The *adjacent matrix* of a graph is a square $|N| \times |N|$ matrix A such that its element A_{ij} is one when there is an edge from vertex i to vertex j , and zero otherwise.

In this work we only consider undirected graphs and just a single edge between nodes is allowed.

Many complex systems are composed of coupled networks through different layers, rather than just one, where each layer represents one of many possible types of interactions. They are called multi-layer networks.

We represent a multi-layer graph with k layers through a sequence $(N_s, E_s)_{s=1,\dots,k}$ where N_s and E_s are respectively the set of nodes and the set of edges of layer s.

Connected to a multi-layer graph, we can consider a set of adjacent matrices $(A_s)_{s=1,\dots,k}$ where k is the number of layers and A_s is the adjacent matrix of layer s.

We focus our attention on multi-layer graphs where just edges vary between layers, thus each node is present in all layers, and there are not edges between nodes of different layers.

Community detection has attracted significant attention during the recent years. The goal of community detection is to partition vertices of a graph into densely-connected components (i.e. the so called communities).

The main problems of graph clustering are the concepts of community and partition, which are not rigorously defined, although they are intuitive concepts.

Intuitively, we get the notion that a community must have more edges between the nodes of the community than edges linking nodes of the community with the rest of the graph. However, there are many alternative definitions of community.

Local definitions consider communities as separate entities and evaluate them independently of the rest of the graph. Global definitions, despite the local ones, consider communities as an integral part of the graph, which cannot be studied regardless of the rest. There are many global criteria to find communities, some of them use some global properties just indirectly, incorporating them in an algorithm that shows communities at the end. Some definitions are based on vertex similarity. The main idea is that a community is a group of nodes similar to each other, so they compute the similarity between each pair of nodes and put most similar nodes in the same community.

Another problem of community detection is the concept of partition. A partition is a division of a graph in clusters, such that each node belongs to just one of them. In real systems we can have overlapping communities, where a node can belong to more communities. However, we don't consider this case in our work. The main problem is to distinguish a good partition from a bad one. For this reason, we need a quantitative criterion to measure the goodness of a graph clustering. A quality function is a function that assigns a number to each partition of a graph, thus one can sort them by their value and identify the best one. Nevertheless, the answer depends on the quality function and on the community concept that are used. This problem of quantifying the value of partitions becomes even more complicated in the multi-layer case, because a given partition can be very good for one layer but very bad for another.

In this work we rely on the idea that a graph has a community structure if it is different from a random graph. In fact, a random graph should not have a community structure, since any two nodes have the same probability to be adjacent or not. A null model is used as term of comparison, to verify if whether a graph shows a community structure or not. For this reason, it maintains some structural features of the original graph. This concept is the basis of the definition of modularity, a quality function where a subgraph is a community if the number of edges inside the subgraph exceeds the expected number of internal edges that the same subgraph would have in the null model. Modularity can be written as follows

$$
Q = \frac{1}{2m} \sum_{i,j} (A_{ij} - P_{ij}) \delta(C_i, C_j)
$$
\n(1.1)

where the sum runs over all pairs of vertices, A is the adjacency matrix, m the total number of edges of the graph, P_{ij} represents the expected number of edges between vertices i and j in the null model, and C_i is the community of node *i*. The function δ yields one if vertices i and j are in the same community $(C_i=C_j)$, zero otherwise.

In the literature have been suggested several versions of modularity and null model. The most popular null model has been proposed by Newman and Girvan, where edges are linked at random, under the constraint that the expected degree of each vertex of the null model stays the same as the degree of the correspondent node in the original graph [10]. So, the final expression of modularity reads

$$
Q = \frac{1}{2m} \sum_{i,j} \left(A_{ij} - \frac{k_i k_j}{2m} \right) \delta(C_i, C_j)
$$
 (1.2)

where k_i is the degree of node i (the sum of all the edges incident to i).

Large positive values of modularity indicate good partitions. The modularity of the whole graph, taken as a single community, is zero, and it is always smaller than one, and can be negative as well. For instance, the partition in which each vertex is a community is always negative. This implies that, if there are no partitions with positive modularity, the graph has no community structure. Another feature of modularity is that the gain due to the movement of a node from one community to another can be easily calculated.

Unfortunately, modularity has got some limits. The main problem is the so-called resolution limit [11], that may prevent to find community that are small with respect to the graph as a whole, even if they are cliques. Thus, when the partition with maximum modularity includes whole, even if they are chques. Thus, when the partition with maximum modularity includes clusters with total degree of the order of \sqrt{m} or smaller, where m is the number of total edges of the graph, one cannot distinguish if the clusters are single communities or if they are combined together. The reason of the resolution limit is the definition of the null model, where we assume that each node can interact with every other node, but more reasonably each node interacts just with a part of the graph. This limit has a large impact in real applications.

To overcome this problem, one could recursively apply modularity optimization on each community of the obtained clustering [11] [12]. However, we don't have a stopping criteria and each cluster uses a different null model, since communities can be of different sizes.

Good et al. [13] discovered that modularity values are very close to the global maximum, nevertheless partitions with high modularity are not necessarily similar to each other and the global maximum is impossible to reach.

Another disadvantage of modularity is that can be applied just to single-layer graphs. In this thesis project we hence propose some models that take advantage of this modularity function and try to extend it to the multi-layer case. All the models are based on the Louvain heuristic method for single-layer networks [9], which is a locally greedy modularity-increasing sampling process over the set of partitions. The most intuitive idea, already studied in the literature, is to use the modularity average on the layers. Thus, we propose two methods that take into account variance of modularity on the layers, in addition to the average. We present also a more sophisticated filter type algorithm. We focus on the multiobjective aspect of the problem and maintain just the modularity vectors that are not dominated according to a suitably developed Pareto search. We implemented these methods in Matlab and we performed some experiments both on artificial and real world networks.

Chapter 2

Related Work

In this Chapter, we present a brief review of the literature related to the community detection problem. In the first Section 2.1, we start by describing the classical approaches for single-layer graphs, following Fortunato [14]. Traditional algorithms include graph partitioning, hierarchical, partitional and spectral clustering; modern methods include divisive algorithms, modularity based methods, spectral algorithms, dynamic algorithms, methods based on statistical inference, non-negative matrix factorization, nonlinear spectral algorithms and total variation approaches. In Section 2.2 we present some algorithms for multi-layer graphs: first we introduce community detection algorithms in two-layer graphs, later detection algorithms that can support multi-layer graphs containing more than or equal to two layers.

2.1 Community Detection in Single-Layer Graphs

Many community detection approaches have been proposed for single-layer graphs. Fortunato conducted a survey on this topic [14]. He presents:

- *traditional clustering methods* divided into: graph partitioning, hierarchical, partitional and spectral clustering;
- *modern methods*, divided into categories based on the type of approach: divisive algorithms, modularity based methods, spectral algorithms, dynamic algorithms, methods based on statistical inference. In this section we add some more recent methods: non-negative matrix factorization, nonlinear spectral algorithms and total variation approaches.

We do not present algorithms to find overlapping communities, multiresolution and hierarchical techniques because they are not connected to our work.

2.1.1 Traditional clustering methods

Traditional clustering methods include graph partitioning, hierarchical, partitional and spectral clustering.

Graph partitioning

Graph partitioning aims to divide the vertices in an established number of groups of predefined size, such that the cut size, i.e. the number of edges running between clusters, is minimal. Specifying the number of clusters of the partition is necessary. The graph partitioning problem is NP-hard, however several algorithms have good results, even if their solutions are not necessarily optimal [15]. Many algorithms perform a bisection of the graph, and partitions into more than two clusters are usually attained by iterative bisectioning [16]. The well-known max-flow mincut theorem by Ford and Fulkerson [17], that states that the minimum cut between any two vertices s and t of a graph carries the maximum flow that can be transported from s to t across the graph, has been used to determine minimal cuts from maximal flows in clustering algorithms [18] [19]. We usually do not have information about the community structure of a graph, in such cases this procedure is not useful, and one must make some unjustified assumptions about the number and size of the clusters.

Hierarchical clustering

The vertices of a graph can be grouped at different levels, with small clusters included within large clusters, which are in turn included in larger clusters, and so on. In order to reveal the multilevel structure of the graph, one may use hierarchical clustering algorithms [20]. Hierarchical clustering is very common in social network analysis, biology, engineering, marketing. As the base of hierarchical clustering methods there is the definition of a similarity measure between vertices. After a measure is chosen, one computes the similarity for each pair of vertices, obtaining the similarity matrix. Hierarchical clustering techniques can be classified in two categories: agglomerative algorithms, in which clusters are iteratively merged if their similarity is sufficiently large, and divisive algorithms, in which clusters are iteratively split by removing edges connecting vertices with low similarity. Hierarchical clustering has the advantage that a preliminary knowledge on the number and size of the clusters is not required. However, it does not give us a way to discriminate between the many partitions obtained by the procedure.

Partitional clustering

Partitional clustering is another common class of methods to find communities in a graph. The number of clusters is preassigned, say k. Each vertex is a point of a metric space and a distance measure is defined between them. The distance is a measure of dissimilarity between vertices. The purpose is to partition the points in k clusters so to maximize/minimize a given cost function based on distances between points. The most popular partitional technique in the literature is k-means clustering $[21]$, where the cost function is the total intra-cluster distance, or squared error function. The limitations of partitional clustering is both that the number of clusters must be specified at the beginning, as for the graph partitioning algorithms, and that the embedding in a metric space for some graphs can be not natural.

Spectral clustering

Supposed to have a pairwise similarity function S defined between a set of n objects, which is symmetric and non-negative. Spectral clustering includes all methods and techniques that partition the set into clusters by using the eigenvectors of matrices, like S itself or other matrices derived from it. The objects could be points in some metric space, or the vertices of a graph. Spectral clustering consists of a transformation of the initial set of objects into a set of points in space, whose coordinates are elements of eigenvectors: the set of points is then clustered via standard partitional techniques, like k-means clustering. The first contribution on spectral clustering was a paper by *Donath and Hoffmann* [22], who used the eigenvectors of the adjacency matrix for graph partitions. In the same year, Fiedler [23] used the eigenvectors of the Laplacian matrix, by far the most used matrix in spectral clustering.

2.1.2 Modern clustering methods

Modern clustering methods include divisive algorithms, modularity based methods, spectral algorithms, dynamic algorithms, methods based on statistical inference, non-negative matrix factorization, nonlinear spectral algorithms and total variation approaches.

Divisive algorithms

The problem of divisive algorithms consists in detecting the edges that connect vertices of different communities and remove them, so that the clusters get disconnected from each other. The critical point is to find a property of intercommunity edges that could identify them. Divisive methods just perform hierarchical clustering on the graph at study, so they do not introduce new techniques.

The most popular algorithm is that proposed by Girvan and Newman [24] [10]. Edges are selected according to the values of measures of edge centrality, estimating the importance of edges according to some property or process running on the graph. Girvan and Newman study in deep the concept of betweenness, which is a variable expressing the frequency of the participation of edges to a process. Many modifications of this method have been proposed, like the algorithm by Holme et al. where vertices, rather than edges, are removed [25]; the algorithm proposed by Pinney and Westhead, that is able to find overlapping communities [26]; the method designed by Estrada based on the concept of communicability between nodes [27].

Modularity based methods

Newman-Girvan modularity [28], originally introduced to define a stopping criterion for the algorithm of Girvan and Newman, has rapidly become the most used quality function. Modularity optimization is an NP-complete problem, however there are currently several algorithms able to find good approximations of the modularity maximum in a reasonable time. We concentrate on clustering methods that require modularity, directly and/or indirectly: *greedy techniques*, simulated annealing, extremal optimization, spectral optimization.

The first *greedy technique* to maximize modularity was designed by *Newman* [29]. It is an agglomerative hierarchical clustering method, where groups of vertices are successively joined to form larger communities such that modularity increases after the merging. At the beginning, all vertices of the graph are put in different communities. Edges are not initially present, they are added one by one during the procedure. An edge is chosen such that this partition gives the maximum increase of modularity with respect to the previous configuration. A lot of modifications of this method have been proposed [30], [31], [32].

A different greedy approach is the Louvain method, introduced by Blondel et al. [9]. Initially, each vertices is a community. The first step consists of a sequential sweep over all vertices. Given a vertex i, one computes the gain in weighted modularity coming from putting i in the community of its neighbour j and picks the community of the neighbour that yields the largest increase of modularity, as long as it is positive. At the end of the sweep, one obtains the first level partition. In the second step communities are replaced by supervertices. Two supervertices are connected if there is at least an edge between vertices of the corresponding communities and the weight of the edge between the supervertices is the sum of the weights of the edges between the represented communities at the lower level. The whole procedure is repeated iteratively. At some point, the algorithm stops because modularity cannot increase anymore.

Simulated annealing [33] is a probabilistic procedure for global optimization used in different fields and problems. It performs an exploration of the space of possible states, looking for the global optimum of a function F , say its maximum. Transitions from one state to another occur with probability 1 if F increases after the change, otherwise they occur with a probability $exp(\beta \Delta F)$, where ΔF is the decrease of the function and β is an index of stochastic noise, which increases after each iteration. Simulated annealing was first used for modularity optimization by Guimerá et al. [34]. Its standard implementation [7] is composed by two types of moves: local moves, where a single vertex is shifted from one cluster to another randomly; global moves, consisting of mergers and splits of communities. The method can potentially come very close to the true modularity maximum, but it is slow.

Extremal optimization (EO) is a heuristic search procedure proposed by Boettcher and Percus [35], in order to achieve an accuracy comparable with simulated annealing, but with a gain in computer time. It is based on the optimization of local variables, expressing the contribution of each unit of the system to the global function at study. Duch and Arenas [36] used this technique for modularity optimization. Modularity can be written as a sum over the vertices: the local modularity of a vertex is the value of the corresponding term in this sum. Dividing the local modularity of the vertex by its degree, we obtain a fitness measure for each vertex. In this way the measure is normalized and does not depend on the degree of the vertex. At the beginning, the vertices are divided randomly into two groups of the same size. At each iteration, the vertex with the lowest fitness is shifted to the other cluster. The move changes the partition, so the local fitnesses of many vertices need to be recalculated. The process stops when the global modularity cannot be improved any more. After the bipartition, each cluster is considered as a graph on its own and the procedure is repeated, as long as the global modularity increases for the partitions found.

Spectral optimization aims to optimize modularity using the eigenvalues and eigenvectors of the modularity matrix. One can maximize modularity via spectral clustering, by replacing the Laplacian matrix with the modularity matrix [10] [29].

In the most recent literature on graph clustering, several modifications and extensions of modu-

larity can be found, motivated by specific classes of clustering problems or graphs that one may want to analyse. For instance, due to the so-called resolution limit of the modularity function, Traag at al. compared the network to a constant factor, instead of a random null model as usual [37].

Spectral algorithms

Spectral properties are used to find partitions, as in the case of spectral clustering, which considers the eigenvectors of Laplacian matrix, or in the optimization of modularity, which uses the eigenvectors of the modularity matrix.

The algorithm proposed by *Donetti and Muñoz* uses the eigenvectors of the Laplacian matrix [38]. It turns vertices into points of a metric space, using the eigenvectors components as coordinates, because they are close for vertices in the same community.

Also the algorithm designed by Alves uses spectral properties of the Laplacian matrix [39]. Here effective conductances for pairs of vertices is computed, looking at the graph as an electric network with edges of unit resistance.

Capocci et al. used the eigenvectors of a right stochastic matrix, that should have similar properties as the Laplacian [40].

Yang and Liu proposed a recursive bisectioning procedure, which uses the spectral properties of the adjacency matrix [41].

In recent times, Fasino and Tudisco studied in deep spectral properties of modularity matrices, that are related to the community detection problem [42] [43].

Dynamic algorithms

Dynamic methods use processes running on the graph, focusing on *spin-spin interactions*, random walks and *synchronization*.

One of most popular spin model in statistical mechanics, is the Potts model [44]. It describes a system of spins that can be in q different states. If Potts spin variables are assigned to the vertices of a graph with community structure, and the interactions are between neighbouring spins, the modules could be likely recovered from like-valued spin clusters of the system, as there are many more interactions inside communities than outside.

Random walks [45] can also be useful to find communities. If a graph has a strong community structure, a random walker spends a long time inside a community due to the high density of internal edges and consequent number of paths that could be followed.

Synchronization [46] is an emergent phenomenon occurring in systems of interacting units and is present in nature, society and technology. In a synchronized state, the units of the system are always in the same or similar state. Synchronization has also been applied to find communities in graphs. If oscillators are placed at the vertices, with initial random phases, and have nearest-neighbour interactions, oscillators in the same community synchronize first, whereas a full synchronization requires a longer time. So, if one follows the time evolution of the process, states with synchronized clusters of vertices can be stable and durable, so they can be easily individuated.

Methods based on statistical inference

Statistical inference [47] has the intent to deduce properties of data sets, starting from a set of observations and model hypotheses. If the data set is a graph, the model, based on hypotheses about connections between nodes, has to fit the actual graph topology. These methods try to find the best fit of a model to the graph, where the model assumes that vertices have some sort of classification, based on their connectivity patterns.

Generative models adopt Bayesian inference [48], in which the best fit is obtained through the maximization of a likelihood. Bayesian inference uses observations to estimate the probability that a given hypothesis is true. It is based on two elements: the evidence, expressed by the information one has about the system (e.g. through measurements), and a statistical model with parameters.

Block modelling [49] is a popular approach in statistics and social network analysis to decompose a graph in classes of vertices with common properties. Vertices are grouped in classes of equivalence. There are two main definitions of topological equivalence for vertices: structural equivalence [50], in which vertices are equivalent if they have the same neighbours; regular equivalence [51] [52], in which vertices of a class have similar connection patterns to vertices of the other classes.

Model selection [53] try to find models which are both simple and good at describing a system/process. To select a model, there is not one defined way, but some heuristics.

Information theory has also been used to detect communities in graphs. Ziv et al. [54] have designed a method in which the information contained in the graph topology is compressed such to preserve some predefined information. This is the underlying philosophy of the information bottleneck method [55].

Non-negative Matrix Factorization

Non-negative Matrix Factorization (NMF) [56] [57] has been applied to many applications such as clustering and classification. It provides a linear representation of non-negative data in high dimensional space with the product of two non-negative matrices. Some papers explicitly include the notion of sparseness, improving the found decompositions [58] [59]; some others incorporate discriminant constraints in the decomposition [60]. Some recent research works suggest that data of many applications in a high dimensional Euclidean space are usually embedded in a low dimensional manifold [61]. To explore the local structure on the low dimensional manifold, papers have proposed Locality Preserving NMF and Neighbourhood Preserving NMF, which add constraints between a point and its neighbours [62] [63].

Nonlinear Spectral algorithms and Total Variation approaches

Modularity optimization is an NP-complete problem. We have seen that many algorithms, such as spectral clustering methods or non-negative matrix factorization (NMF) methods, relax the discrete optimization space into a continuous one to obtain an easier optimization procedure. However, in general the solution of the relaxed continuous problem and that of the discrete NPhard problem can be very different. A new set of algorithms obtains tighter relaxations, taking

idea from the image processing literature. They are based on the concept of total variation (TV), which favours the formation of sharp indicator functions in the continuous relaxation. These functions equal one on a subset of the graph, zero elsewhere and exhibit a non-smooth jump between these two regions. At the beginning, total variation techniques had a recursive bi-partitioning procedure to handle more than two classes. Later, Bühler and Hein and Bresson et al. proposed two methods that do not rely on a recursive procedure [64] [65].

In 2013, Hu et al. showed that modularity optimization is equivalent to minimizing a particular non-convex total variation based functional over a discrete domain [66] and, in 2018, Boyd et al. showed that this equivalence states for a convex total variation based functional [67]. Both algorithms assume that the number of communities is known. This process is called non-linear exact relaxation of the modularity function.

In recent times, Tudisco et al. and Tudisco and Higham proposed instead nonlinear spectral methods [68] [69].

Finally, in 2020, Cristofari et al. introduced the modularity total variation (TVQ) and showed that its box-constrained global maximum coincides with the maximum of the original discrete modularity function [70].

2.2 Community Detection in Multi-Layer Graphs

In contrast to the community detection problem in single graphs, new challenges arise for community detection in multi-layer graphs. It is natural to detect multi-layer communities by extending the algorithms for single-layer community detection. The most popular approaches that have been employed for this extension can be grouped into two strategies: the first one reduces the multi-layer networks into a single-layer network and then applies single-layer network algorithms to obtain the communities in the collapsed network [71]; whereas the second strategy obtains the communities for each layer applying single-layer network algorithms, and then combines the obtained communities by using consensus clustering [72]. However, these algorithms are criticized for their low accuracy because they either cannot preserve the community structure in compressed networks or ignore the connection among various layers. Another problem is connected to noise, in fact these methods usually suppose that each layer is informative but in real networks some of them are just noise. To overcome these problems, algorithms must simultaneously take into account multiple layers. Thus, there is necessity to develop effective algorithms for community detection in multi-layer networks, rather than by simply extending the available single-layer network algorithms.

In their work, Kim and Lee [73] divide community detection algorithms for two-layer graphs, from detection algorithms that can support multi-layer graphs containing more than or equal to two layers. We follow this subdivision in our review.

2.2.1 Community detection in two-layer graphs

In this subsection, we introduce community detection algorithms in two-layer graphs. All algorithms described in this section can only support two-layer graphs and mostly consider structural and attribute information.

Cluster Expansion

Li et al. [74] proposed a hierarchical community detection algorithm based on both relations and textual attributes using the cluster expansion idea. Initially the algorithm quinckly finds the centers as seed of communities, then it expands the centers into the communities. The algorithm is composed by four steps: core probing, core merging, affiliation, and classification.

Matrix Factorization

 Qi et al. [75] proposed a community detection algorithm based both on link structure and edge content using the Edge-Induced Matrix Factorization (EIMF). The key point of this algorithm is the use of edge content for the community detection process, which can be useful when nodes interact with multiple communities, since it can help distinguishing between the different interactions of nodes. This algorithm firstly takes into consideration just the link structure, then it incorporates also the edge content.

Unified Distance

Zhou et al. [76] proposed a community detection algorithm, called SA-Cluster, based on both structural and attribute similarities using a unified distance measure. The main contributions of SA-Cluster are a unified distance measure to take simultaneously into account both structural and attribute similarities and a weight self-adjustment method to control the degree of importance of structural and attribute similarities.

Model-Based Method

Xu et al. [77] proposed a model-based community detection method based on both structural and attribute aspects of a graph. The main idea of this approach is the use of a probabilistic model that fuses both structural and attribute information instead of an artificial distance measure. The algorithm firstly constructs the probabilistic model and then a variational approach to solve it.

Pattern Mining

Silva et al. [78] proposed a community detection algorithm based on structural correlation pattern mining, called SCPM. The key point of SCPM is to reveal the connection between vertex attributes and dense subgraphs using both frequent itemset mining and quasi-clique mining. Here, a dense subgraph is defined by a quasi-clique.

Graph Merging

Ruan et al. [79] proposed a community detection approach, called CODICIL, to combine structural and attribute information using the graph merging process. The main contribution of this algorithm is to delate noise in the link structure using content information.

2.2.2 Community detection in multi-layer graphs

In this section, we introduce community detection algorithms that can support multi-layer graphs containing more than or equal to two layers.

Matrix Factorization

Tang et al. [80] and Dong et al. [81] proposed graph clustering algorithms for multi-layer graphs based on matrix factorization. The key point of these two algorithms is to fuse different information by extracting common factors from multiple layers, which may then be used by general clustering methods. Tang et al. [80] approximates adjacency matrices while Dong et al. [81] approximates graph Laplacian matrices.

Pattern Mining

Zeng et al. [82] proposed a subgraph mining algorithm for finding quasi-cliques that appear on multiple layers with a frequency above a given threshold. The main contribution of this algorithm is to find cross-graph quasi-cliques that are frequent, coherent, and closed. Generally, the cross-graph quasi-clique has been defined as a set of vertices belonging to a quasi-clique that appears on all layers and must be the maximal set [83].

Spectral Methods

Some methods try to extend spectral clustering to multi-layer graphs. In general, these algorithms aim to define a graph operator that contains all the information of the multi-layer graph such that the eigenvectors corresponding to the smallest eigenvalues are informative about the clustering structure. These methods usually rely on some sort of arithmetic mean, for example the Laplacian of the average adjacency matrix or the average Laplacian matrix [84]. Further examples are the work of Zhou and Burges, which defines a multiple cut graph, which is good on average while it may not be the best on single graphs [85], and the algorithm designed by Chen and Hero, that performs convex aggregation of layers based on graph noise models [86].

Other Approaches

Some approaches adopt Bayesian inference [48], in which certain hypotheses about connections between nodes are made to find the best fit of a model to the graph through the optimization of a suitable likelihood [87].

Other methods try to propose an extension of Newman's modularity [28] and connected null models. For instance, the method proposed by *Wilson et al.* finds overlapping communities and proves consistency in a suitable multi-layer stochastic block model [88].

A co-training approach is proposed by Kumar and Daumé [89], where the algorithm aims to find a consistent clustering under the main assumption that all the layers are informative, so each single layer has a piece of meaningful information from its own perspective. Kumar et al. [90] concentrated on this approach under the notion of co-regularization.

Chapter 3

Presentation of the Methods

In this Chapter, we present some methods for community detection in multi-layer networks. The algorithms are based on the Louvain heuristic method for single-layer graphs [9], that we mentioned in Chapter 2 describing modularity based methods.

The natural extension of this method to the multi-layer case, already studied in the literature, is to locally maximize the modularity average on the layers during phase 1, instead of the modularity of a single layer. In Section 3.1, we suggest two variants that, in the selection criteria of the algorithm during phase 1, take into account the variance of modularity on the layers, in addition to the average. This is due to the fact that in real networks we can have two cases: the informative case, where there are all informative layers, so each single layer has a piece of meaningful information from its own perspective, but we can also have the noisy case, where there are some noisy layers, which give us wrong information about communities.

In Section 3.2, we present a more sophisticated filter type algorithm. We are studying a multiobjective optimization problem, so we decide to memorize all the possible moves in a list and then we maintain just the modularity vectors that are not dominated according to a suitably developed Pareto search. We use different functions as criterion for handling the list, in particular, we use the average on modularity of the layers, and the convex combination of the average and the variance, thus getting two different approaches. Notice that these methods with unit length of the list correspond respectively to the algorithms described before.

These algorithms coincide with the Louvain method on single layer graphs.

Matlab codes of the methods are available in the Appendix.

3.1 Louvain Expansion

In this Section, we present the *Louvain Expansion* method for community detection in multilayer networks. It is based on the Louvain method for single-layer graphs [9] and tries to expand it to the multi-layer case.

The input of the algorithm is a multi-layer graph G with k layers. We suppose to have multilayer graphs where just edges vary between layers, thus each node is present in all layers, and

there are not edges between nodes of different layers. Each layer is an undirected graph and just a single edge between nodes is allowed. The output of the algorithm is a final assignment of nodes to communities.

The algorithm is composed of two phases that are repeated iteratively.

At the beginning of the first phase, each node forms a community. The algorithm calculates some values related to this first partition, s.a. Q_s the modularity of the clustering in layer s for $s=1,..,k$ and a defined function F connected to them. We consider as neighbours of a node i the union of the neighbours of node i in the various layers. Then the algorithm starts a loop, where each node is considered in order (for this reason the indexing of the nodes changes the output of the algorithm). Call i the node taken into consideration and C_i its community. The algorithm removes node *i* from its community C_i and calculates the modularity gain $\Delta Q1_i$ on each layer. For each neighbour j of node i (if the community of node j has not already been considered), the method includes i in the community of node j, called C_j , and calculates the corresponding modularity gain $\Delta Q_{i\rightarrow j}$ on each layer. Now the algorithm calculates the gain $\Delta F_{i\rightarrow j}$ of the defined function F , for changing the community of node i. Among all the positive gains of function F , which therefore give an increase of the function, the algorithm selects the highest one $\Delta F_{i\rightarrow j*}$ and put node *i* into the corresponding community C_{j*} . The algorithm recalculates the modularity and the function values corresponding to this new partition. This first phase stops when it finds a local maxima of the function, i.e. when no individual move can improve the function value.

The second phase remains unchanged respect to the original method for single-layer graphs. The algorithm constructs a reduced network, where each community becomes a node, such that all-singleton partition has the same value of modularity as the partition that we identified at the end of the first phase. To do so, the weights of the links between the new nodes are given by the sum of the weight of the links between nodes in the corresponding two communities, and links between nodes of the same community lead to self-loops for the community.

The algorithm then iterates the whole procedure, until the heuristic converges, i.e. until phase 2 induces no further changes.

Look at Algorithm 1 for a pseudocode of the Louvain Expansion method.

Note that the output of the algorithm depends on the order of the nodes. Therefore, the way in which we index the new communities at the end of phase 2 is important. It seems that the ordering of the nodes does not influence significantly the final modularity, however it appears to affect the computational time. In the literaturate, this aspect is still unclear. We decide to order the nodes due to the community size.

The Louvain heuristic is very popular for its simplicity and efficiency. Part of the algorithm's efficiency results from the fact that the modularity can be calculated iteratively during the procedure. At the beginning of phase 1, the method calculates the modularity from scratch. In Chapter 1 we proposed a formula to calculate modularity of an unweighted graph (equation (1.2)), however during phase 2 the algorithm creates a reduced graph with weighted edges,

Algorithm 1 Louvain Expansion

Input: G multi-layer graph Output: C final assignment of nodes to communities

```
1: function LOUVAIN EXPANSION(G)2:
 3: repeat
 4: PHASE 1
 5: Initialize:
 6: C \leftarrow initial partition, where each vertex of graph G is a community
 7: Q \leftarrow \text{modularity vector of the initial partition}8: F \leftarrow function value of the initial partition
 9: NB \leftarrow neighbour nodes vector
10: repeat
11:
12: for each node i do
13: remove node i from its community
14: \Delta Q1_i \leftarrow modularity gain for removing node i from its community
15:
16: for each node j that is neighbour of node i do
17: insert node i into community of node j18: \Delta Q2_{i\rightarrow j} \leftarrow modularity gain for inserting i into community of j
19: \Delta F_{i \to j} \leftarrow function gain for changing the community of node i
20: end for
21: \Delta F_{i\rightarrow j^*} \leftarrow best function gain
22:
23: if \Delta F_{i\rightarrow j^*} > 0 then
24: move node i into community of node j^*25: Q \leftarrow \text{modularity vector of the new partition}26: F \leftarrow function value of the new partition
27: end if
28: end for
29:
30: until no improved clustering found.
31:
32: PHASE 2
33: G \leftarrow reduced graph where each community of partition C is a node
34:
35: until no improved clustering found.
36: return C
37: end function
```
therefore we show also the corresponding formula for weighted graphs

$$
Q_{\rm w} = \frac{1}{2w} \sum_{i,j} \left(W_{ij} - \frac{s_i s_j}{2w} \right) \delta(C_i, C_j) \tag{3.1}
$$

where the sum runs over all pairs of vertices, W is the incidence matrix, w the sum of the weights of all the edges of the graph, s_i is the strength of node i (the sum of the weights of all the edges incident to i), and C_i is the community of node i. The function δ yields one if vertices i and j are in the same community $(C_i=C_i)$, zero otherwise.

From now we suppose to work with a unweighted graph, however each formula can be easily extended to the weighted case.

To calculate modularity, the algorithm actually uses this formula, that is equivalent to the equation (1.2),

$$
Q = \sum_{c \in C} \frac{|E(c)|}{m} - \left(\frac{\sum_{i \in N} k_i}{2m}\right)^2
$$
 (3.2)

where $C=(C_1,...,C_n)$ is the clustering, $m=|E|$ is the number of edges, $|E(c)|$ is the sum of all the links between nodes in C and k_i is the degree of node i .

During the loop, the algorithm calculates the gain in modularity in an easy way. The gain ΔQ_1 obtained by moving a node i from its community c can easily be computed by

$$
\Delta Q I_i = \frac{\sum_{tot} k_i}{2m^2} - \frac{k_i^2}{2m^2} - \frac{k_{i,in}}{m}
$$
\n(3.3)

where \sum_{tot} is the sum of weights of the links incident to node in c, k_i is the degree of node i, $k_{i,in}$ is the sum of weights of the links from i to nodes in c, m is the sum of weights of all the links inside the network.

The following expression is used in order to evaluate the change of modularity $\Delta Q2_i$ when an isolated node i is moved into the community c of node j

$$
\Delta Q2_{i\to j} = \frac{\sum_{i,in}}{m} - \frac{\sum_{tot} \cdot k_i}{2m^2}
$$
\n(3.4)

where $\sum_{i,in}$ is the sum of weights of the links inside community c, \sum_{tot} is the sum of weights of the links incident to node in c, k_i is the degree of node i, m is the sum of weights of all the links inside the network.

Thus, if a node i has changed community, the algorithm calculates the modularity of the new partition just adding to the initial modularity value the gains obtained above, rather than calculate it from scratch. Thanks to this fact, the Louvain heuristic is extremely fast.

In our method, it's reasonable to use a function F that can include the information of the multiple layers and that can be calculated iteratively.

The most intuitive idea to extend the Louvain heuristic from the single-layer to the multi-layer case, already studied in the literature, is to take as function F the average of modularity on the layers

$$
M_Q = \frac{\sum_{s=1}^{k} Q_s}{k} \tag{3.5}
$$

where k is the number of layers and Q_s is the modularity of layer s. The gain of this function can be calculated easily as follow

$$
\Delta M_Q = \frac{\sum_{s=1}^k \Delta Q_s}{k} \tag{3.6}
$$

where k is the number of layers and ΔQ_s is the gain on modularity of layer s. We refer to this method with *community-average* (ComA). Matlab code of this method is available in Appendix A.

In real networks, different situations arise. We study two cases: the informative case, where each single layer has a piece of meaningful information from its own perspective, and the noisy case, where there are some noisy layers, which give us wrong information about communities. In order to analyse better these two situations, we propose two functions that take into account sample variance of modularity on the layers

$$
V_Q = \frac{\sum_{s=1}^{k} (Q_s - M_Q)^2}{k - 1}
$$
\n(3.7)

where k is the number of layers, Q_s is the modularity on layer s and M_Q is the average of modularity on the layers.

We suggest to take a convex combination of the average and the variance of modularity on the layers. In particular, we study these two function

$$
F_{-} = (1 - \gamma)M_Q - \gamma V_Q \tag{3.8}
$$

$$
F_{+} = (1 - \gamma)M_Q + \gamma V_Q \tag{3.9}
$$

where M_Q is the average of modularity on the layers, V_Q is the variance of modularity on the layers and $\gamma \in [0, 1]$.

The idea behind is that: for the informative case we would like to maximize the average and minimize the variance of modularity on the layers, instead in the noisy case we want to maximize both the values.

The gain of both the functions can be easily calculated during the algorithm respectively in these ways

$$
\Delta F_{-} = (1 - \gamma)\Delta M_Q - \gamma \left(V_{\Delta Q} + \frac{2}{k - 1} (Q - M_Q)^t (\Delta Q - \Delta M_Q) \right) \tag{3.10}
$$

$$
\Delta F_{+} = (1 - \gamma)\Delta M_Q + \gamma \Big(V_{\Delta Q} + \frac{2}{k - 1}(Q - M_Q)^t(\Delta Q - \Delta M_Q)\Big) \tag{3.11}
$$

where k is the number of layers, Q is a vector in which the entrance s is the initial modularity of layer s, M_Q is the initial average of modularity on the layers, ΔQ is a vector in which the entrance s is the gain in modularity of layer s, ΔM_Q is the gain of the average of modularity on the layers (that coincides with $M_{\Delta Q}$ the average of ΔQ , as shown in equation (3.6)), $V_{\Delta Q}$ is the variance of ΔQ , calculated as follows

$$
V_{\Delta Q} = \frac{\sum_{s=1}^{k} (\Delta Q_s - \Delta M_Q)^2}{k - 1}
$$
\n(3.12)

We refer to the method that uses the function F_{-} as *community-variance-minus* (ComV-) and to the method that uses the function F_+ as *community-variance-plus* (ComV+). Matlab codes of these methods are available in Appendix B and Appendix C.

All the methods that we have introduced coincide with the original Louvain algorithm if applied to single-layer graphs.

3.2 Louvain Multiobjective

In this Section, we present the *Louvain Multiobjective* method for community detection in multilayer networks, that is more sophisticated respect to the one described in Section 3.1. The algorithm is a filter type method that takes into account the multiobjective nature of the problem. It is based on the Louvain method for single-layer graphs [9] and tries to expand it to the multi-layer case. The basic idea is to not decide just one community to put in node i during phase 1, but to follow more case studies.

The problem of maximizing modularity over multiple layers is a problem of multiobjective optimization. We consider a vector $Q=(Q_s)_{s=1,..,k}$ where k is the number of layers, and we want to maximize simultaneously all its entries. If there are no conflicts between the entries, the straightforward optimal solution of the problem is obtained solving separately k optimization problems. However, this is not what usually happens in real networks. In multiobjective optimization there is not a unique way to define the concept of optimality, since there is not a total order for \mathbb{R}^k . Each partial order defines a different definition of optimality. We adopt a definition that was proposed for the first time by Edgeworth in 1881 and later revised by Vilfredo Pareto in 1896 [91]. All the definitions are referred to a maximisation problem.

Definition 1. Given two vectors z^1 and $z^2 \in \mathbb{R}^k$, z^1 dominates z^2 according to Pareto, and we write $z^1 \geq_P z^2$, if

$$
z_i^1 \geq z_i^2
$$
 for each index $i=1,..,k$ and
\n $z_j^1 > z_j^2$ for at least one index $j=1,..,k$.

This binary relation induces a partial order over \mathbb{R}^k . Thus we can give the definition of optimality according to Pareto.

Definition 2. A vector $z^* \in \mathbb{R}^k$ is Pareto optimal if there is not other vectors $z \in \mathbb{R}^k$ such that $z^* \leq_P z$.

The Pareto front is the set of all Pareto optimals.

We define a filter as a list of vectors such that no vector dominates the others.

The input of the Louvain Multiobjective algorithm is a multi-layer graph G with k layers. We suppose to have multi-layer graphs where just edges vary between layers, thus each node is present in all layers, and there are not edges between nodes of different layers. Each layer is an

undirected graph and just a single edge between nodes is allowed. The output of the algorithm is a final assignment of nodes to communities.

The algorithm is composed of two phases that are repeated iteratively.

At the beginning of the first phase, each node forms a community. The algorithm calculates some values related to this first partition, s.a. Q_s the modularity of the clustering in layer s for $s=1,..,k$ and a defined function F connected to them. The method inserts the initial partition and the corresponding modularity vector in a filter L . We consider as neighbours of a node i the union of the neighbours of node i in the various layers. Then the algorithm starts a loop, where each node is considered in order (for this reason the indexing of the nodes changes the output of the algorithm). Call i the node taken into consideration. For each partition in filter L, the method does the following procedure. Call C_i the community of node i in the considered partition. The algorithm removes node i from its community C_i and calculates the modularity gain ΔQ_1 on each layer. For each neighbour j of node i (if the community of node j has not already been considered), the method includes i in the community of node j, called C_j , and calculates the corresponding modularity gain $\Delta Q2_{i\rightarrow j}$ on each layer. Now the algorithm calculates the gain $\Delta F_{i\to j}$ of the defined function F, for changing the community of node *i*. If the function gain is positive, which therefore gives an increase of the function, the algorithm memorises the partition and the corresponding modularity vector in the filter. More precisely, the new modularity vector is added to L only if it is not dominated and, if this condition is verified, the method delates from the list all the modularity vectors that are now dominated by the new one. When the method has finished doing this procedure on each element of the initial filter, it checks the length of the new list L. If the filter is too long compared to a previously decided length h , the filter is cut removing the partitions with the least function values, until it is of the required length. This first phase stops when no moves change the filter. At this point, the method selects the partition of the filter with the maximum value of the function.

The second phase remains unchanged respect to the original method for single-layer graphs. The algorithm constructs a reduced network, where each community becomes a node, such that all-singleton partition has the same value of modularity as the partition that we identified at the end of the first phase. To do so, the weights of the links between the new nodes are given by the sum of the weight of the links between nodes in the corresponding two communities, and links between nodes of the same community lead to self-loops for the community.

The algorithm then iterates the whole procedure, until the heuristic converges, i.e. until phase 2 induces no further changes.

Look at Algorithm 2 for a pseudocode of the Louvain Multiobjective method.

Note that, also in this method, the output depends on the order of the nodes. We decide to order the nodes due to the community size to study the informative case, instead to index them following the initial order to study the noisy case, because this case has a higher computational time.

The method uses a filter because we want to get closer to the Pareto front. However, considering all the case studies would be too expensive in terms of computational time. For this reason, the method cuts the list to a length h using as a criterion a function F .

Algorithm 2 Louvain Multiobjective

Input: G multi-layer graph Output: C final assignment of nodes to communities

function LOUVAIN MULTIOBJECTIVE (G)

```
repeat
    PHASE 1
    Initialize:
         C \leftarrow initial partition, where each vertex of graph G is a community
         Q \leftarrow modularity vector of the initial partition
         F \leftarrow function value of the initial partition
         NB \leftarrow neighbour nodes vector
         L \leftarrow (C,Q) filter
    repeat
        for each node i do
            L_{old} \leftarrow Lfor each element (C,Q) in L_{old} do
                remove node i from its community
                \Delta Q1_i \leftarrow modularity gain for removing node i from its community
                for each node j that is neighbour of node i do
                    insert node i into community of node j\Delta Q2_{i\rightarrow j} ← modularity gain for inserting i into community of j
                    \Delta F_{i\rightarrow j} ← function gain for changing the community of node i
                    if \Delta F_{i\rightarrow j} > 0 then
                        move node i into community of node j: C_{i\rightarrow j} new partition
                        Q_{i\rightarrow j} ← modularity vector of the new partition
                        if Q_{i\rightarrow j} not dominated by any element of the filter L then
                            L \leftarrow L \cup (C_{i\rightarrow j}, Q_{i\rightarrow j})L \leftarrow L \setminus (elements dominated by (C_{i\rightarrow j}, Q_{i\rightarrow j}))
                        end if
                    end if
                end for
            end for
            if length(L) > h then
                remove from L the partitions with the least function values, until it is of length hend if
        end for
    until no changes of filter L
    (C^*,Q^*) \leftarrow element of the filter with the maximum value of the function
    PHASE 2
    G \leftarrow reduced graph where each community of partition C^* is a node
```
until no improved clustering found. return C end function

We study three variants of this method, taking as function F : the average of modularity on layers M_Q described in (3.5), and the functions $F₊$ and $F₊$ defined in (3.8) and (3.9). We refer to these algorithms respectively as multi-average (MultiA), multi-variance-minus (MultiV-), multi-variance-plus (MultiV+).

Matlab codes of these methods are available respectively in Appendix D, Appendix E and Appendix F.

As we have seen in Section 3.1, the algorithm calculates the modularity and the function values in a iterative and easy way.

Note that this method coincides with the method described in Section 3.1 when the filter has a unit length. Therefore, it is equal to the original Louvain algorithm if applied to single-layer graphs.

Chapter 4

Experimentation

We implemented the methods described in Chapter 3 using Matlab, starting from a available Matlab code for the Louvain heuristic for single layer graphs [92]. The Matlab codes are reported in the Appendix. We tested the algorithms using a machine equipped with i7 processor with 1.8 GHz and 16 GB of ram memory. The tests were performed both on artificial networks and on real world networks.

In this Chapter, we show the obtained results. Section 4.2 presents the results achieved on artificial graphs, and Section 4.3 reports the algorithm outputs obtained on real datasets. Preliminarily, in Section 4.1 we focus on how evaluate the partitions obtained as results of the methods, to later compare them. In particular, we used the accuracy and the Normalized Mutual Information.

4.1 Evaluation

In order to compare the results of the algorithms described in Chapter 3, we need to evaluate the partitions obtained as output. In the literature, various ways have been proposed on how to measure the performance of an algorithm for community detection. However, this question has not been fully resolved yet. We worked under the main assumption of knowing the community structure of the graph, that we call standard partition. On the other hand, we refer to the partitions obtained by the algorithms as predicted partitions. We evaluated the results of our algorithms through two values: the accuracy (AC) and the Normalized Mutual Information (NMI). To calculate both these values, we use indirectly the confusion matrix.

The *confusion matrix* T is a table that allows visualization of the performance of an algorithm. It has a row for each community of the standard partition and a column for each community of the predicted partition. The element T_{ij} represents the number of nodes in the j community of the predicted partition that are in the i community of the standard partition. The sum of the i row (resp. column) gives us the total number of nodes of the i community of the standard (resp. predicted) partition. The values on the diagonal are the nodes that are in the same community in both the partitions, instead the values off-diagonal are the nodes placed in the wrong community.

One disadvantage of the confusion matrix is that it depends on the labels of the communities.

The first way adopted to measure the results of the algorithms is the *accuracy* (Ac) , i.e. the percentage of nodes placed in the right community. First of all, we calculated the number of nodes that are placed in the right community, using the confusion matrix and labelling the communities appropriately. We then considered the percentage of the nodes placed in the right community respect to the total number of nodes, in order to possibly compare results reffered to graphs of different size.

We evaluated the output partitions obtained by the algorithms also through the *Normalized* Mutual Information (NMI), because we supposed that the community structures were known. Mutual Information is widely used in physics, statistics, and machine learning as a tool for comparing different labellings of a set of objects. In network science it is perhaps the standard measure for quantifying the performance of community detection algorithms. However, it can give inaccurate answers under certain conditions, s.a. when communities have different size. For this reason, Danon et al. proposed a normalization of this measure, called Normalized Mutual Information [93]. Given the standard partitioning C^* and the obtained partitioning C, the Normalized Mutual Information can be calculated by the following formula

$$
NMI(C^*, C) = \frac{2 \cdot I(C^*, C)}{H(C^*) + H(C)}
$$
\n(4.1)

where $H(C)$ is the entropy of partition C and $I(C^*, C)$ is the Mutual Information between C^* and C. This is defined by

$$
I(C^*, C) = H(C^*) - H(C^*|C)
$$
\n(4.2)

where $H(C^*|C)$ is the conditional entropy of C^* respect to C. We calculated the NMI using the confusion matrix T, through the formula

$$
NMI(C^*, C) = \frac{-2 \cdot \sum_{i=1}^{|C|} \sum_{i=1}^{|C^*|} T_{ij} \cdot \log\left(\frac{T_{ij}t}{T_i \cdot T_j}\right)}{\sum_{i=1}^{|C|} T_i \cdot \log\left(\frac{T_i}{t}\right) + \sum_{j=1}^{|C^*|} T_{.j} \cdot \log\left(\frac{T_{.j}}{t}\right)}
$$
(4.3)

where $t = \text{sum}(T)$ is the number of nodes, T_i is the sum of row i of T and $T_{i,j}$ is the sum of column j of T .

NMI(C^*, C) is a number between 0 and 1. It is equal to 1 when the two partitions are identical and equal to 0 when the two partitions are totally independent, s.a. when C is the single community that includes the whole graph.

A nice feature of NMI is that it is invariant under permutations of the labels of the communities.

4.2 Artificial Networks

We tested the algorithms described in Chapter 3 on artificial networks. In this Section, we show and compare the obtained results.

The Stochastic Block Model (SBM) is a generative model for graphs showing certain clusters structures through the parameters p_{in} and p_{out} . These parameters represent the edge probabilities: given nodes v_i and v_j the probability of observing an edge between them is p_{in} (resp. p_{out}), if v_i and v_j belong to the same (resp. different) cluster.

We analysed two different settings: in the first one all layers have the same class structure, in the second setting one layer is informative and the remaining layers are just noise. We set $p_{in} \gg p_{out}$ on the informative layers and $p_{in} = p_{out}$ on the noisy layers.

In particular, we created networks with 4 communities of 125 nodes each and with $k = 2, 3$ layers, by fixing $p_{in} = 0.1$ and varying p_{out} . In the noisy layers, we fixed $p_{in} = p_{out} = 0.1$.

For each case, we report the results in a table. Each row corresponds to a method, which is indicated in the first column. We studied the Louvain Multiobjective models for length of the filter $h = 2, 3$ and, in the definition of function $F₋$ in equation (3.8) and $F₊$ in equation (3.9), we set $\gamma = 0.1, 0.3, 0.5$. The ratio between p_{in} and p_{out} changes between columns and it is specified in the second row. For each method and ratio between p_{in} and p_{out} , the tables report the accuracy (Ac) in percentage, the Normalized Mutual Information (NMI) and the execution time in seconds (Cpu) of the corresponding output. The best performances are marked with bold fonts and gray background and second best performances with only gray background. We show the results also using bar plots. In the tables and in the bar plots, we report the average of the values on 10 runs. We summarize the results of the multiple runs through some boxplots.

4.2.1 Informative case

In the informative case all layers have the same community structure, so each single layer has a piece of meaningful information.

For this case, we compared the models *community-average* (ComA) and *community-variance*minus (ComV-) (Section 3.1), multi-average (MultiA) and multi-variance-minus (MultiV-) (Section 3.2). The idea behind is that for the informative case we would like to maximize the average and minimize the variance of modularity on the layers.

Table 4.1 shows the average results for the informative case on graphs with $k = 2$ layers. Figure 4.1 and Figure 4.2 represent the Accuracy and the NMI of the results in some bar plots. We summarize the results of the 10 runs in the boxplots in Figure 4.3 and Figure 4.4.

 p_{in} is fixed equal to 0.1 and p_{out} varies according to the ratio between p_{in} and p_{out} equal to 3, 2.5, 2, 1.5.

The results get worse as the ratio between p_{in} and p_{out} decreases, in fact intuitively the communities are less defined when p_{out} is close to p_{in} . For this reason, the execution times behave the same way, increasing as the ratio decreases. In particular, the Louvain Multiobjective methods are slower than the others.

All the methods perform very good in the first two cases with $p_{in}/p_{out} = 3, 2.5, a$ little less well in the third case with $p_{in}/p_{out} = 2$ and get worst in the last situation with $p_{in}/p_{out} = 1.5$. The algorithms show similar results unless for $p_{in}/p_{out} = 2$, where the better performances are obtained by MultiV- with $h = 2, \gamma = 0.3$ and 0.5.

We studied in depth the more critical interval [2.5, 2], sampling it. Table 4.2 shows the performances to vary of $p_{in}/p_{out} = 2.6, 2.4, 2.2, 2$. The methods still perform approximately the same way. Figure 4.5 and Figure 4.6 represent the Accuracy and the NMI of the results in some bar plots. We summarize the results of the 10 runs in the boxplots in Figure 4.7 and Figure 4.8.

Table 4.3 shows the average results for the informative case on graphs with $k = 3$ layers. Figure 4.9 and Figure 4.10 represent the Accuracy and the NMI of the results in some bar plots. We summarize the results of the 10 runs in the boxplots in Figure 4.11 and Figure 4.12.

 p_{in} is fixed equal to 0.1 and p_{out} varies according to the ratio between p_{in} and p_{out} equal to 3, 2.5, 2, 1.5.

Also in this case, the results get worse as the ratio between p_{in} and p_{out} decreases, and the execution times behave the same way. In particular, the Louvain Multiobjective methods are slower than the others.

All the methods perform very good and similar for values of $p_{in}/p_{out} = 3, 2.5, 2$, but unfortunately all of them get worst in the last case with $p_{in}/p_{out} = 1.5$.

We analysed better the more interesting interval [2, 1.5] in Table 4.4, taking $p_{in}/p_{out} = 2, 1.9$, 1.8, 1.7. The algorithms obtain almost the same results for the first two values; in the third case MultiV- with $h = 2$, $\gamma = 0.1$ and ComA achieve the best results; finally in the last case MultiA with $h = 3$ definitely outperforms all the other methods. Figure 4.13 and Figure 4.14 represent the Accuracy and the NMI of the results in some bar plots. We summarize the results of the 10 runs in the boxplots in Figure 4.15 and Figure 4.16.

Compared to the case with 2 layers, all the methods give best results. In fact, in the informative case, have multiple layers correspond to have more information. Nevertheless, execution times are higher for graphs with 3 layers.

In general, in the informative case on graphs with both 2 and 3 layers: when the community structure is well defined, all the algorithms perform well; when it is more confused, they give different outputs, but there is not a method that always dominated all the others.

4.2.2 Noisy case

In the noisy case just one layer is informative, so it has a community structure, and all the other layers are just noise, so they give wrong information about the clustering.

For this case, we compared the models community-average (ComA) and community-varianceplus (ComV+) (Section 3.1), multi-average (MultiA) and multi-variance-plus (MultiV+) (Section 3.2). The idea behind is that for the noisy case we would like to maximize both the average and the variance of modularity on the layers.

Table 4.5 shows the average results for the noisy case on graphs with $k = 2$ layers. Figure 4.17 and Figure 4.18 represent the Accuracy and the NMI of the results in some bar plots. We
INFORMATIVE CASE: $k=2$												
		$p_{\rm in}/p_{\rm out}$ =3		$p_{\rm in}/p_{\rm out} = 2.5$			$p_{\rm in}/p_{\rm out}$ =2			$p_{\rm in}/p_{\rm out}$ =1.5		
	Ac	NMI	Cpu	Ac	NMI	Cpu	Ac	NMI	Cpu	Ac	NMI	Cpu
ComA	99.74	0.989	2.52	99.04	0.962	3.98	75.04	0.575	6.96	21.84	0.045	5.36
MultiA	99.76	0.990	140.72	98.84	0.955	214.42	74.22	0.577	254.66	21.08	0.049	287.80
$h=2$												
MultiA	99.80	0.992	147.78	98.92	0.957	328.00	78.20	0.620	425.17	21.58	0.047	451.14
$h=3$												
$ComV-$	99.76	0.990	2.86	98.84	0.955	5.05	75.44	0.588	8.03	21.02	0.043	7.14
$\gamma = 0.1$												
$ComV-$	99.76	0.990	3.46	98.76	0.952	5.35	73.86	0.577	7.77	21.00	0.041	6.90
$\gamma = 0.3$												
$ComV-$	99.74	0.990	6.05	98.74	0.952	5.66	74.74	0.591	7.41	21.46	0.042	7.51
$\gamma = 0.5$												
MultiV-	99.76	0.990	163.26	98.80	0.954	248.27	77.08	0.609	262.40	21.44	0.050	292.21
$\gamma = 0.1, h = 2$												
MultiV-	99.80	$\bf 0.992$	170.17	98.96	0.959	446.78	75.72	0.588	421.59	22.08	$\bf 0.051$	437.92
$\gamma = 0.1$, h=3												
MultiV-	99.80	0.992	169.90	98.98	0.960	250.46	81.64	$\bf 0.656$	262.39	21.10	0.045	295.12
$\gamma = 0.3, h = 2$												
MultiV-	99.72	0.988	197.79	98.96	0.959	311.20	76.70	0.593	435.59	20.90	0.043	452.44
$\gamma = 0.3, h = 3$												
MultiV-	99.78	0.991	106.79	98.74	0.951	228.23	80.72	0.642	266.18	21.40	0.051	287.89
$\gamma = 0.5, h = 2$												
MultiV-	99.76	0.990	185.08	98.92	0.958	331.87	76.84	0.579	419.15	21.70	0.046	460.25
$\gamma = 0.5, h = 3$												

Table 4.1: Experiments in the informative case on artificial graphs with $k=2$ layers. Notation: best performances are marked with bold fonts and gray background and second best performances with only gray background.

Figure 4.1: Accuracy of the experiments in the informative case on artificial graphs with $k=2$ layers

Figure 4.2: NMI of the experiments in the informative case on artificial graphs with $k=2$ layers

Figure 4.3: Accuracy of the experiments in the informative case on artificial graphs with $k=2$ layers

Figure 4.4: NMI of the experiments in the informative case on artificial graphs with k=2 layers

INFORMATIVE CASE: $k=2$												
		$p_{\rm in}/p_{\rm out}$ =2.6		$p_{\rm in}/p_{\rm out} = 2.4$			$p_{\rm in}/p_{\rm out}$ =2.2				$p_{in}/p_{out}=2$	
	Ac	NMI	Cpu	Ac	NMI	Cpu	Ac	NMI	Cpu	Ac	NMI	Cpu
ComA	98.90	0.956	3.13	98.32	0.936	3.99	93.28	0.833	4.15	75.04	0.575	6.96
MultiA	98.94	0.975	139.63	98.30	0.934	197.19	96.78	0.888	219.62	74.22	0.577	254.66
$h=2$												
MultiA	98.96	0.958	227.07	98.42	0.939	276.52	97.16	0.895	295.49	78.20	0.620	425.17
$h=3$												
$ComV-$	98.90	0.956	4.20	98.32	0.935	4.90	97.26	0.898	4.98	75.44	0.588	8.03
$\gamma = 0.1$												
$ComV-$	98.92	0.956	4.29	98.26	0.933	4.92	97.00	0.891	5.06	$73.86\,$	0.577	7.77
$\gamma = 0.3$												
$ComV-$	98.94	0.957	4.59	98.44	0.940	4.65	97.20	0.896	4.83	74.74	0.591	7.41
$\gamma = 0.5$												
MultiV-	98.92	0.956	144.55	98.46	0.941	200.38	96.70	0.880	203.30	77.08	0.609	262.40
$\gamma = 0.1, h = 2$												
MultiV-	98.96	0.958	216.71	98.36	0.937	286.79	97.06	0.890	296.29	75.72	0.588	421.59
$\gamma = 0.1$, h=3												
MultiV-	99.00	0.959	180.89	98.30	0.935	192.75	96.52	0.885	200.45	81.64	0.656	262.39
$\gamma = 0.3, h = 2$												
MultiV-	98.94	0.957	240.51	98.32	0.936	287.64	95.86	0.874	295.01	76.70	0.593	435.59
$\gamma = 0.3, h = 3$												
MultiV-	99.00	0.959	144.67	98.42	0.939	188.69	96.98	0.888	199.50	80.72	0.642	266.18
$\gamma = 0.5, h = 2$												
MultiV-	99.00	0.959	231.16	98.20	0.931	280.32	96.82	0.886	297.55	76.84	0.579	419.15
$\gamma = 0.5, h = 3$												

Table 4.2: Experiments in the informative case on artificial graphs with $k=2$ layers. Notation: best performances are marked with bold fonts and gray background and second best performances with only gray background.

Figure 4.5: Accuracy of the experiments in the informative case on artificial graphs with $k=2$ layers

Figure 4.6: NMI of the experiments in the informative case on artificial graphs with $k=2$ layers

Figure 4.7: Accuracy of the experiments in the informative case on artificial graphs with $k=2$ layers

Figure 4.8: NMI of the experiments in the informative case on artificial graphs with k=2 layers

INFORMATIVE CASE: $k=3$												
		$p_{\rm in}/p_{\rm out}$ =3		$p_{\rm in}/p_{\rm out} = 2.5$			$p_{\rm in}/p_{\rm out} = 2$				$p_{\rm in}/p_{\rm out}{=}1.5$	
	Ac	NMI	Cpu	Ac	NMI	Cpu	Ac	NMI	Cpu	Ac	NMI	Cpu
ComA	99.90	0.996	5.11	99.84	0.993	5.42	98.80	0.952	5.42	25.56	0.076	7.20
MultiA	99.90	0.996	231.03	99.84	0.993	264.99	98.80	0.952	275.88	24.38	0.079	325.63
$h=2$												
MultiA	99.90	0.996	358.93	99.84	0.993	408.75	98.80	0.952	417.78	26.20	0.076	477.31
$h=3$												
$ComV-$	99.90	0.996	5.94		99.84 0.993	6.19	98.72	0.949	6.31	25.36	0.072	10.13
$\gamma = 0.1$												
$ComV-$	99.90	0.996	6.09	99.84	0.993	6.11	98.74	0.950	6.36	27.48	0.093	9.75
$\gamma = 0.3$												
$ComV-$	99.90	0.996	5.67	99.84	$\bf{0.993}$	5.75	98.72	0.949	6.29	29.00	0.096	8.68
$\gamma = 0.5$												
MultiV-	99.90	0.996	231.36		99.84 0.993 271.00		98.70	0.948	277.56	24.24	0.071	312.35
$\gamma = 0.1, h = 2$												
MultiV-	99.88	0.995	352.77		99.84 0.993	402.17	98.84	0.953	426.00	27.88	0.092	507.82
$\gamma = 0.1, h = 3$												
MultiV-	99.90	0.996	273.54	99.84	0.993 266.89		98.86		0.954 231.57	25.00	0.076	334.74
$\gamma = 0.3$, h=2												
MultiV-	99.90	0.996	354.30	99.82	0.992	406.74	98.78	0.952	425.64	24.44	0.072	509.90
$\gamma = 0.3$, h=3												
MultiV-	99.92	0.997	234.06	99.84	$\bf 0.993$	268.40	98.80	0.952	277.64	24.76	0.072	337.01
$\gamma = 0.5, h = 2$												
MultiV-	99.90	0.996	346.97	99.84	0.993	396.08	98.82	0.953	422.18	26.98	0.104	521.42
$\gamma = 0.5, h = 3$												

Table 4.3: Experiments in the informative case on artificial graphs with k=3 layers. Notation: best performances are marked with bold fonts and gray background and second best performances with only gray background.

Figure 4.9: Accuracy of the experiments in the informative case on artificial graphs with $k=3$ layers

Figure 4.10: NMI of the experiments in the informative case on artificial graphs with k=3 layers

Figure 4.11: Accuracy of the experiments in the informative case on artificial graphs with k=3 layers

Figure 4.12: NMI of the experiments in the informative case on artificial graphs with k=3 layers

INFORMATIVE CASE: $k=3$												
		$p_{\rm in}/p_{\rm out}$ =2		$p_{\rm in}/p_{\rm out}$ =1.9			$p_{\rm in}/p_{\rm out}{=}1.8$				$p_{\rm in}/p_{\rm out}$ =1.7	
	Ac	NMI	Cpu	Ac	NMI	Cpu	Ac	NMI	Cpu	Ac	NMI	Cpu
ComA	98.80	0.952	5.42	98.02	0.924	6.04	96.20	0.865	6.59	61.88	0.440	7.56
MultiA	98.80	0.952	233.60	98.16	0.929	249.74	90.72	0.804	275.88	65.76	0.467	293.36
$h=2$												
MultiA	98.80	0.952	357.67	98.10	0.927	396.72	84.92	0.742	417.78	85.64	$\bf 0.684$	440.85
$h=3$												
$ComV-$	98.72	0.949	6.23	97.76	0.915	6.31	96.02	0.859	7.80	69.76	0.528	9.81
$\gamma = 0.1$												
$ComV-$	98.74	0.950	6.36	97.90	0.920	7.07	89.36	0.791	8.47	52.44	0.342	9.20
$\gamma = 0.3$												
$ComV-$	98.72	0.949	6.29	98.04	0.925	7.27	94.84	0.838	8.76	68.78	0.499	10.45
$\gamma = 0.5$												
MultiV-	98.70	0.948	238.61	98.02	0.923	266.97	95.88	0.858	277.56	58.94	0.390	303.89
$\gamma = 0.1, h = 2$												
MultiV-	98.84	0.953	369.92	98.18	0.929	404.17	96.30	$\bf0.868$	426.00	70.76	0.512	475.77
$\gamma = 0.1$, h=3												
MultiV-	98.86	0.954	231.57	98.18		0.929 265.74	93.06	0.819	298.45	69.80	0.530	312.52
$\gamma = 0.3, h = 2$												
MultiV-	98.78	0.952	365.66	97.90	0.919	425.64	89.24	0.784	437.12	63.30	0.460	449.65
$\gamma = 0.3, h = 3$												
MultiV-	98.80	0.952	241.91	98.10	0.926	267.20	96.10	0.863	275.88	53.40	0.349	277.64
$\gamma = 0.5, h = 2$												
MultiV-	98.82	0.953	356.77	97.96	0.922	392.99	84.18	0.718	422.18	64.54	0.439	437.57
$\gamma = 0.5, h = 3$												

Table 4.4: Experiments in the informative case on artificial graphs with k=3 layers. Notation: best performances are marked with bold fonts and gray background and second best performances with only gray background.

Figure 4.13: Accuracy of the experiments in the informative case on artificial graphs with $k=3$ layers

Figure 4.14: NMI of the experiments in the informative case on artificial graphs with k=3 layers

Figure 4.15: Accuracy of the experiments in the informative case on artificial graphs with k=3 layers

Figure 4.16: NMI of the experiments in the informative case on artificial graphs with k=3 layers

summarize the results of the 10 runs in the boxplots in Figure 4.19 and Figure 4.20.

 p_{in} is fixed equal to 0.1 and p_{out} varies according to the ratio between p_{in} and p_{out} equal to 3, 2.5, 2.

The results get worse as the ratio between p_{in} and p_{out} decreases, in fact intuitively the communities are less defined when p_{out} is close to p_{in} . For this reason, the execution times behave the same way, increasing as the ratio decreases. In particular, the Louvain Multiobjective methods are slower than the others.

ComV+ outperforms all the other methods in the first two cases with $p_{in}/p_{out} = 3, 2.5$. All the algorithm gives bad results in the last case with $p_{in}/p_{out} = 2$.

Table 4.6 shows the average results for the noisy case on graphs with $k = 3$ layers. Figure 4.21 and Figure 4.22 represent the Accuracy and the NMI of the results in some bar plots. We summarize the results of the 10 runs in the boxplots in Figure 4.23 and Figure 4.24.

 p_{in} is fixed equal to 0.1 and p_{out} varies according to the ratio between p_{in} and p_{out} equal to 3, 2.5, 2, 1.5.

Also in this case, the results get worse as the ratio between p_{in} and p_{out} decreases, and the execution times behave the same way. In particular, the Louvain Multiobjective methods are slower than the others.

MultiV+ with $\gamma = 0.5$, $h = 3$ achieves the best result in the first situation with $p_{in}/p_{out} = 3$. All the algorithms obtained bad results in all the other cases.

Compared to the case with 2 layers, all the methods give worst results. In fact, in the noisy case, just the first layer is informative and all the others are noise, so the noise increases as the number of layer increases. Moreover, execution times are higher for graphs with $k = 3$ layers.

In general, in the noisy case on graphs with both 2 and 3 layers, almost all the methods overcome ComA, and the best performances are obtained by the methods that take into consideration the variance of modularity on the layers as well as the average, giving them the same weight in equation (3.9).

All the algorithms achieve better results in the informative case then the noisy one. Moreover, execution times are lower in the informative case than in the noisy case. In fact, in the informative case each layer has a piece of meaningful information, instead in the noisy case some layers can give wrong information.

4.3 Real World Networks

In this Section, we compare the performances of the proposed approaches on real networks. We considered three real datasets: 3sources [94], BBCSports [95] and Wikipedia [96]. We used the corresponding layer matrices provided by Mercado et al. [97], which have used a similarity measure and have considered the unweighted version of the symmetric k-nearest neighbour graph (i.e. they have taken the k neighbours with highest correlation). Also in this case, we knew the community structure of the graphs.

3sources dataset corresponds to new articles of BBC, Reuters and Guardian. It produces a

NOISY CASE: $k=2$												
		$p_{\rm in}/p_{\rm out} = 3$			$p_{\rm in}/p_{\rm out}$ =2.5		$p_{\rm in}/p_{\rm out}$ =2					
	Aс	NMI	Cpu	Aс	NMI	Cpu	Ac	NMI	Cpu			
ComA	49.42	0.318	7.34	27.00	0.109	7.35	22.46	0.054	8.01			
MultiA	58.68	0.369	255.28	29.42	0.116	289.26	21.06	0.053	312.98			
$h=2$												
MultiA	47.06	0.289	226.51	30.22	0.128	383.26	21.62	0.056	483.20			
$h=3$												
$ComV+$	56.84	0.382	7.03	27.36	0.102	8.32	21.30	0.053	9.00			
$\gamma = 0.1$												
$ComV+$	54.56	0.343	5.56	29.36	0.128	9.52	21.50	0.054	9.99			
$\gamma = 0.3$												
$ComV+$	75.86	0.581	9.19	35.34	0.173	9.62	21.88	0.058	10.41			
$\gamma = 0.5$												
$MultiV+$	56.30	0.368	260.59	32.74	0.142	281.59	20.78	0.054	300.20			
$\gamma = 0.1, h = 2$												
$MultiV+$	58.14	0.370	379.66	29.30	0.126	386.27	22.08	0.060	439.44			
$\gamma = 0.1, h = 3$												
$MultiV+$	59.22	0.396	190.88	28.36	0.126	254.34	21.30	0.052	269.15			
$\gamma = 0.3, h = 2$												
$MultiV+$	57.54	0.394	211.16	30.78	0.144	379.35	21.36	0.057	525.10			
$\gamma = 0.3, h = 3$												
$MultiV+$	62.36	0.432	164.39	32.62	0.157	252.06	24.02	0.067	283.00			
$\gamma = 0.5, h = 2$												
$MultiV+$	65.02	0.466	318.62	31.98	0.154	407.74	22.12	0.067	474.81			
$\gamma = 0.5, h = 3$												

Table 4.5: Experiments in the noisy case on artificial graphs with $k=2$ layers. Notation: **best** performances are marked with bold fonts and gray background and second best performances with only gray background.

Figure 4.17: Accuracy of the experiments in the noisy case on artificial graphs with k=2 layers

Figure 4.18: NMI of the experiments in the noisy case on artificial graphs with k=2 layers

Figure 4.19: Accuracy of the experiments in the noisy case on artificial graphs with k=2 layers

Figure 4.20: NMI of the experiments in the noisy case on artificial graphs with k=2 layers

NOISY CASE: $k=3$												
		$p_{\rm in}/p_{\rm out} = 3$			$p_{\rm in}/p_{\rm out}$ =2.5		$p_{\rm in}/p_{\rm out}$ =2					
	Aс	NMI	Cpu	Aс	NMI	Cpu	Ac	NMI	Cpu			
ComA	31.08	0.140	6.61	23.32	0.074	7.15	21.48	0.046	8.23			
MultiA	36.54	0.183	198.97	23.18	0.069	242.43	20.72	0.045	280.91			
$h=2$												
MultiA	35.40	$\,0.169\,$	366.69	21.76	0.066	431.68	21.86	0.054	441.05			
$h=3$												
$ComV+$	37.36	0.196	8.67	23.92	0.085	9.97	20.34	0.044	13.69			
$\gamma = 0.1$												
$ComV+$	38.68	0.219	8.58	25.94	0.095	9.57	20.76	0.045	11.57			
$\gamma = 0.3$												
$ComV+$	52.64	0.328	9.88	25.52	0.092	10.34	21.32	0.055	12.94			
$\gamma = 0.5$												
$MultiV+$	36.62	0.188	245.82	22.84	0.070	293.56	21.06	0.037	331.74			
$\gamma = 0.1, h = 2$												
$MultiV+$	32.32	0.151	348.44	23.56	0.077	363.38	21.52	0.051	440.29			
$\gamma = 0.1, h = 3$												
$MultiV+$	42.10	0.227	225.27	23.70	0.082	260.47	21.30	0.047	277.16			
$\gamma = 0.3, h = 2$												
$MultiV+$	36.00	0.189	373.14	23.16	0.069	418.36	21.20	0.042	439.75			
$\gamma = 0.3$, h=3												
$MultiV+$	48.02	0.306	253.78	27.08	0.102	292.30	21.48	0.057	309.10			
$\gamma = 0.5, h = 2$												
$MultiV+$	59.48	0.402	362.97	24.44	0.099	408.00	20.80	0.049	442.13			
$\gamma = 0.5, h = 3$												

Table 4.6: Experiments in the noisy case on artificial graphs with $k=3$ layers. Notation: **best** performances are marked with bold fonts and gray background and second best performances with only gray background.

Figure 4.21: Accuracy of the experiments in the noisy case on artificial graphs with k=3 layers

Figure 4.22: NMI of the experiments in the noisy case on artificial graphs with k=3 layers

Figure 4.23: Accuracy of the experiments in the noisy case on artificial graphs with k=3 layers

Figure 4.24: NMI of the experiments in the noisy case on artificial graphs with k=3 layers

muti-layers graph with 169 nodes, 3 layers, and 6 communities of size 56, 21, 11, 18, 51, 12. BBCSports is a dataset of sports articles of BBC and corresponds to a multi-layer graph with 544 nodes, 2 layers and 5 communities of size 62, 104, 193, 124, 61.

The last dataset, called Wikipedia, reports Wikipedia articles and gives a multi-layer graph with 693 nodes, 2 layers and 10 communities of size 34, 88, 96, 85, 65, 58, 51, 41, 71, 104.

We analysed the informative and the noisy settings. In the noisy case, we kept the first layer and we added to all the other layers a matrix, generated by the Stochastic Block Model with $p_{in} = p_{out} = 0.05$. For each dataset, we tested the algorithms on 10 runs.

For each of the two cases, we report the results in a table. Each row corresponds to a method, which is indicated in the first column. We studied the Louvain Multiobjective models for length of the filter $h = 2, 3$ and, in the definition of function F in equation (3.8) and $F₊$ in equation (3.9) , we set $\gamma = 0.1, 0.3, 0.5$. The names of the datasets are reported in the second row. For each real dataset we show the accuracy (Ac) in percentage, the Normalized Mutual Information (NMI) and the execution time in seconds (Cpu) of the corresponding output. The best performances are marked with bold fonts and gray background and second best performances with only gray background. We show the results also using bar plots. In the noisy case, in the tables and in the bar plots, we report the average of the values on 10 runs. We summarize the results of the multiple runs through some boxplots.

4.3.1 Informative case

In the informative case, all layers have the same community structure, so each single layer has a piece of meaningful information.

For this case, we compared the models *community-average* (ComA) and *community-variance*minus (ComV-) (Section 3.1), multi-average (MultiA) and multi-variance-minus (MultiV-) (Section 3.2). The idea behind is that for the informative case we would like to maximize the average and minimize the variance of modularity on the layers.

Table 4.7 shows the results for the informative case. Figure 4.25 and Figure 4.26 represent the Accuracy and the NMI of the results in some bar plots.

All the methods perform very good on the first two datasets 3sources and BBCSports. ComA performs the worst results. The best performances are achieved by MultiV- with $h = 2$,

 $\gamma = 0.5$ in the first dataset and by MultiV- with $h = 3$, $\gamma = 0.1$ in the second one. In the Wikipedia dataset the results are slightly worse and the highest outputs are obtained by MultiA with $h = 3$, ComV- with $\gamma = 0.3$ and ComA.

Execution times increase with increasing number of nodes and are higher for the Louvain Multiobjective methods.

4.3.2 Noisy case

In the noisy case, we kept the first layer and we added to all the other layers a matrix, generated by the Stochastic Block Model with $p_{in} = p_{out} = 0.05$. For each dataset, we tested the algorithms on 10 runs.

For this case, we compared the models community-average (ComA) and community-varianceplus (ComV+) (Section 3.1), multi-average (MultiA) and multi-variance-plus (MultiV+) (Section 3.2). The idea behind is that for the noisy case we would like to maximize both the average and the variance of modularity on the layers.

Table 4.8 shows the average results for the noisy case. Figure 4.27 and Figure 4.28 represent the Accuracy and the NMI of the results in some bar plots. We summarize the results of the 10 runs in the boxplots in Figure 4.29 and Figure 4.30.

All the methods perform well on the first two datasets 3sources and BBCSports. The worst results are obtained by algorithms that use function F_{+} with $\gamma = 0.3$ and 0.5. Unfortunately, all the methods give bad results for the Wikipedia dataset.

All the algorithms achieve better results in the informative case than the noisy one. Moreover, execution times are lower in the noisy case than in the informative case. In fact, in the informative case each layer has a piece of meaningful information, in fact in the noisy case some layers can give wrong information.

As seen in the informative case, execution times increase with increasing number of nodes and are higher for the Louvain Multiobjective methods.

INFORMATIVE CASE												
		3sources			BBCSport			Wikipedia				
	Ac	NMI	Cpu	Ac	NMI	Cpu	Ac	NMI	Cpu			
ComA	85.80	0.749	0.21	74.82	0.753	2.09	55.56	0.544	4.02			
MultiA	86.39	0.765	3.59	89.89	0.825	51.44	52.53	0.521	136.54			
$h=2$												
MultiA	86.98	0.773	5.30	79.04	0.791	85.77	55.84	0.546	207.34			
$h=3$												
$ComV-$	85.80	0.749	0.25	82.35	0.789	2.54	54.83	0.520	5.87			
$\gamma = 0.1$												
$ComV-$	85.80	0.749	0.26	75.18	0.751	2.40	41.70	0.285	5.43			
$\gamma = 0.3$												
$ComV-$	86.98	0.781	0.24	83.27	0.798	2.69	37.52	0.285	8.94			
$\gamma = 0.5$												
MultiV-	86.39	0.765	3.69	84.38	0.784	56.48	54.55	0.503	144.25			
$\gamma = 0.1, h = 2$												
MultiV-	86.98	0.773	5.60	90.44	0.837	89.38	54.40	0.520	265.18			
$\gamma = 0.1, h = 3$												
MultiV-	86.39	0.765	3.64	84.56	0.787	56.22	44.30	0.337	167.55			
$\gamma = 0.3, h = 2$												
MultiV-	88.76	0.805	5.97	80.51	0.787	70.36	44.30	0.369	264.88			
$\gamma = 0.3, h = 3$												
MultiV-	88.76	0.812	4.13	80.88	0.784	50.78	37.09	0.266	178.91			
$\gamma = 0.5, h = 2$												
MultiV-	86.98	0.775	8.12	86.40	0.816	82.10	36.65	0.279	207.46			
$\gamma = 0.5, h = 3$												

Table 4.7: Experiments in the informative case on real datasets. Notation: best performances are marked with bold fonts and gray background and second best performances with only gray background.

Figure 4.25: Accuracy of the experiments in the informative case on real datasets

Figure 4.26: NMI of the experiments in the informative case on real datasets
NOISY CASE									
	3sources			BBCSport			Wikipedia		
	Aс	NMI	Cpu	Ac	NMI	Cpu	Ac	NMI	Cpu
ComA	75.50	0.706	1.01	81.18	0.749	4.49	17.59	0.065	7.84
MultiA	76.80	0.715	14.48	81.84	0.752	128.82	17.30	0.067	397.11
$h=2$									
MultiA	77.04	0.717	22.32	83.25	0.757	192.67	17.34	0.068	452.51
$h=3$									
$ComV+$	77.63	0.717	1.20	80.74	0.739	5.29	17.29	0.063	8.79
$\gamma = 0.1$									
$ComV+$	74.73	0.698	1.28	71.10	0.686	4.97	17.17	0.061	9.22
$\gamma=0.3$									
$ComV+$	69.53	0.661	1.09	65.09	0.648	5.03	16.26	0.058	9.07
$\gamma = 0.5$									
$MultiV+$	77.87	0.717	15.00	81.27	0.739	129.70	17.84	0.066	302.08
$\gamma = 0.1, h = 2$									
$\overline{\text{MultiV}} +$	74.14	0.698	23.33	77.67	0.728	190.31	17.71	0.066	453.98
$\gamma = 0.1, h = 3$									
$MultiV+$	74.20	0.699	14.69	74.03	0.698	122.65	17.55	0.063	317.09
$\gamma = 0.3, h = 2$									
$MultiV+$	75.33	0.703	21.85	74.23	0.699	192.38	17.30	0.062	441.93
$\gamma = 0.3, h = 3$									
$MultiV+$	67.81	0.658	14.25	65.42	0.648	125.40	16.70	0.058	320.44
$\gamma = 0.5, h = 2$									
$MultiV+$	69.59	0.661	21.05	63.14	0.644	187.00	17.11	0.058	450.51
$\gamma = 0.5, h = 3$									

Table 4.8: Experiments in the noisy case on real datasets. Notation: best performances are marked with bold fonts and gray background and second best performances with only gray background.

Figure 4.27: Accuracy of the experiments in the noisy case on real datasets

Figure 4.28: NMI of the experiments in the noisy case on real datasets

Figure 4.29: Accuracy of the experiments in the noisy case on real datasets

Figure 4.30: NMI of the experiments in the noisy case on real datasets

Chapter 5

Conclusions

In this thesis project we presented multiple methods for community detection in multi-layer graphs.

Most of the current algorithms either reduce multi-layer networks into a single-layer network or extend the algorithms for single-layer networks by using consensus clustering. However, these algorithms are criticized for their low accuracy because they either cannot preserve the community structure in compressed networks or ignore the connection among various layers. To overcome these problems, we tried to simultaneously take into account multiple layers.

The algorithms that we proposed are based on the Louvain heuristic method, that is a popular algorithm for community detection in single-layer graphs. It is an iterative procedure composed by two phases. During phase 1, it uses as criterion the modularity function, that is the most popular quality function for measuring the goodness of partitions.

The most intuitive idea to extend this work to the multi-layer case, already studied in the literature, is to use the average of modularity on the layers.

The first method that we proposed, called Louvain Expansion, instead of considering just the modularity average, it takes into account the modularity variance.

The second algorithm, called Louvain Multiobjective, is more sophisticated and it is a filter type method, in fact it maintains just the modularity vectors that are not dominated according to a suitably developed Pareto search.

We suggested different versions of these methods to better analyse two situations: the informative case, where each layer has the same community structure, and the noisy case, where instead just some layers present a community structure and all the others are noise.

We implemented all the algorithms using Matlab.

The methods have been tested on both artificial and real world networks. In both cases, we studied the informative and the noisy situation. We compared the results of the different algorithms calculating the accuracy and the Normalized Mutual Information.

We generated artificial networks with 2 and 3 layers using the Stochastic Block Model. In the informative case all the algorithms behave almost the same way unless in few occasions; instead in the noisy case the method already proposed in the literature, that considers just the average of modularity on the layers, is outdated by almost all the other methods. In particular, the best performances are achieved by the algorithms that consider the modularity variance as well as the modularity average giving them the same weight.

We tested our methods also on three real datasets from the literature. Also in this case we verified that our proposed approaches are competitive with the already proposed method.

For time reasons, we did not test our models against state-of-the-art algorithms and this would be an additional step for providing robust results.

For further research in future studies, we could improve the models using a filter also in the phase 2 and trying to nominate the communities in different ways, s.a. randomizing the labels at the beginning of phase 1.

We should perform an in depth computational analysis with different values of h length of the list and γ weight of variance in the objective function.

A further goal is to test the methods on different type networks. For instance, we could use artificial graphs with different number of nodes, communities of various size, or with more than 3 layers. We should also test the algorithms on real networks from different fields.

A further idea is to evaluate and compare the outputs of the methods through the use of different tools.

Appendix A

Matlab code of the community-average method (Section 3.1)

```
1 % community - average
2\frac{9}{6}3 % Inputs :
4 % M : adjacent matrix
5 % s : 1 = Recursive computation
6 % : 0 = Just one level computation
7 \frac{9}{6}8 % Output :
9 % COMTY , structure with the following information
10 % for each level i :
11 % COMTY .COM{i} : vector of community partition
12 % COMTY . SIZE {i} : vector of community sizes
13 % COMTY .MOD{i} : vector of modularities of clustering on the layers
14 % COMTY . Average (i) : average of modularity on the layers
15 % COMTY . Niter (i) : Number of iteration before convergence
16 % ending :
17 function [COMTY ending] = community_average (M,z)
18 if nargin < 1
19 error ('not enough argument');
20 end
21
22 if nargin < 2
23 Z = 1;
24 end
25
26 S = size (M);
27 \text{ N} = S(1);
28 if length(S) == 3
```

```
29 \text{ k} = S(3);
30 else k = 1;
31 end
32
33 ending = 0;
34
35 for s=1: k
36 M2 (:,:,s) =M (:,:,s) -diag (diag (M (:,:,s)));
37 end
38
39 % total weight of the graph
40 for s=1:k41 if z == 142 M(:,: ,s) = M(:,: ,s) + diag(diag(M(:,: ,s)));43 end
44 m\{s\} = sum(sum(M(:,: , s)))/2;45 end
46
47 Niter = 0; % number of iterations
48
49 if (sum(cellfun((@x)x>0,m)) == 0) | N == 150 ending = 1;
51 COMTY = 0;
52 return
53 end
54
55 % Delate layers with m=0
56 M(:,:,(cellfun(\mathcal{Q}(x)x == 0, m)))=[];
57 \text{ k} = \text{k} - \text{sum} (\text{cellfun} (\mathbb{Q}(x) x == 0, m));
58
59 COM = 1: S(1); % Community of node i
60
61 for s=1:kK(:,: ,s) = sum(M(:,: ,s)); % K(i) = sum of wieght incident to node i
63 SumTot(:,:,s) = sum(M(:,:,s)); %SumTot(i) = sum of weight incident to
           nodes in community i
64 SumIn(:,:,s) = diag(M(:,:,s))'; %SumIn(i)= sum of weight inside
          community i (loops)
65 %At the beginning each node is a
                                            community
66 Q{s} = compute_model = {f} (COM, M(:, : , s));
67 end
68 Average = sum([Q{\:};}])/k;
69
70 % Neighbor {j}{s} neighbor of node j in layer s
71 for j=1:N72 for s=1: k
73 temp=M2(:,:,s);
74 Neighbor<sub>-j</sub>{s} = find (temp(j,:));
```

```
75 end
76 Neighbor {j}= Neighbor_j ;
77 end
78
79 gain = 1;
80 while (gain == 1)81 gain = 0;
82 for i=1: N
83 C_i = COM(i);84 NB=unique (cat (2, Neighbor\{i\}\{:\}));
85 G = zeros(1, N); % Gain vector
86 best increase = -inf:
87 Cnew = Ci;
88 COM(i) = -1;
89 % remove i from its community
90 for s=1: k
91 CNi = (COM == Ci); %list of nodes in Ci community, without i
92 Ki_in_i {s} = sum (M(i,CNi ,s) ) ; %sum of weights between i and Ci
93 % Gain of modularity
94 GQ_i {\text{is}} = K(:,i,s) * SumTot(:,Ci,s) / (2 * (m{s}^(2))) - Ki_in_i {s} / m{s}s \} - ((K(:,i,s)) \hat{(2)}) / (2 * (m[s]^{(2)}));95 % Recalculate values
96 SumTot(:, Ci, s) = SumTot(:, Ci, s) - K(:, i, s); % weights incident toCi community
97 SumIn(:,Ci,s) = SumIn(:,Ci,s) - 2*sum(M(i,CNi,s)) - M(i,i,s); %weights community i
98 end
99 for j=1: length(NB)100 Cj = COM (NB(j));
101 if (G(C_i)) == 0)102 CNj = (COM==Cj); % nodes in community Cj, without j
103 for s=1:k
104 Ki_in_j{s} = sum(M(i,CNj,s)); %sum of weights between i and
                 C<sub>j</sub>105 GQ_j{s} = Ki_in_j{s}/m{s} - (K(:,i,s)*SumTot(:,Cj,s))/(2*(m{
                 s}^(2))); % gain deltaQ if I put isolated node i in Cj
106 end
107 %gain average
108 for s=1:k
109 DQ_j{s} = GQ_i{s} + GQ_j{s};110 end
111 G(C_i) = (sum ([DQ_i[{:}]]) / k);112 if G(Cj) > best_increase
113 best_increase = G(Cj);
114 DQ_j_t = DQ_j;\mathtt{Cnew\_t} \ = \ \mathtt{Cj} \ ;116 end
117 end
118 end
```

```
119 if best_increase > -10^(-15)
120 Cnew = Cnew_t;
121 Average = Average + best_increase;
122 for s=1:k
123 Q{s} = Q{s} + DQ_j_t(s;124 end
125 end
126 % Recalculate
127 Ck = (COM = = Cnew);
128 for s=1:k129 SumIn(:,Chew, s) = SumIn(:,Chew, s) + 2*sum(M(i, Ck, s)) + M(i, i, s);130 SumTot (:, Cnew, s) = SumTot (:, Cnew, s) + K(:, i, s);
131 end
132 COM(i) = Cnew;
133 if (Cnew \tilde{c} = Ci)
134 gain = 1;
135 end
136 end
137 Niter = Niter + 1;
138 end
139 Niter = Niter - 1;
140 [COM] = reindex_com (COM) ;
141 COMTY.COM\{1\} = COM;
142 COMTY .MOD {1} = [Q {:}];
143 COMTY . Average (1) = Average ;
144 COMTY. Niter (1) = Niter;
145
146
147 % Perform part 2
148 if (z == 1)149 Mnew = M;
150 Mold = Mnew:
151 COMcur = COM;
152 COMfull = COM;
153 j = 2;
154 while 1
155 Mold = Mnew ;
156 S2 = size (Mold);
157 Nnode = S2(1);
158
159 COMu = unique (COMcur);
_{160} Ncom = length (COMu);
161 ind_com = sparse (Ncom, Nnode);
162 ind_com_full = sparse (Ncom, N);
163 for p=1: Ncom
164 ind = find (COMcur==p);
165 ind_com(p,1:length(ind)) = ind;
166 end
167 for p=1: Ncom
```

```
168 ind = find (COMfull == p);
169 ind_com_full (p, 1: length (ind) ) = ind;170 end
171 Mnew = [];
172 for s=1:k
173 Mnew (:,:,s) = zeros (Ncom, Ncom); %new matrix (each node is a
             community )
174 for m=1: Ncom
175 for n=m: Ncom
176 ind1 = ind_com(m, :);
177 \quad \text{ind2 = ind\_com(n, :)};178 % weights of edges between communities
179 Mnew (m, n, s) = sum(sum (Mold (ind1 (ind1 > 0), ind2 (ind2 > 0), s))) ) ;
180 Mnew (n,m,s) = \text{sum}(\text{sum}(1,nd1(\text{ind}1>0),ind2(\text{ind}2>0),s)) ) ;
181 end
182 end
183 end
184 [COMt e] = community_average (Mnew, 0);
185 if (e^{\pi} = 1)186 COMfull = sparse (1, N);
187 COMcur = COMt. COM\{1\};
188 for p=1: Ncom
189 ind1 = ind_{\texttt{com\_full}}(p, :);190 COMfull (ind1 (ind1 >0)) = COMcur (p);
191 end
_{192} [COMfull2] = reindex_com (COMfull);
193 COMTY.COM{j} = COMfull2;
194 COMTY. MOD{j} = COMt. MOD{1};
195 COMTY . Average (j) = COMt . Average (1) ;
196 COMTY. Niter (j) = COMt. Niter;
197 Ind = (COMfull2 == COMTY.COM{j-1});
198 if (sum (Ind) == length (Ind))
199 return;
200 end
201 else
202 return ;
203 end
204 j = j + 1;
205 end
206 end
207 end
208
209 % Re - index community partition by size
210 function [C Ss] = reindex_com ( COMold )
_{211} C = sparse (1, length (COMold));
212 COMu = unique (COMold);
213 S = sparse (1, length(COMu));
```

```
214 for l=1:length (COMu)
215 S(1) = length (CDMold (CDMold == CDMu(1)));
216 end
217 [SS \quad INDS] = sort(S, 'descend');
218 for l=1: length ( COMu )
219 C(COMold = COMu(INDs(1))) = 1;220 end
221 end
222
223 % Compute modulartiy
224 function MOD = compute_modularity (C, Mat)
225
226 S = size (Mat);
227 \quad N = S(1);
228 m = sum(sum(Mat))/2; %total weight
229 MOD = 0;
230 COMu = unique (C);
231
232 for j=1: length ( COMu )
233 Cj = (C== COMu (j) ) ; % list of nodes in Cj
234 Ec = sum (sum (Mat (Cj, Cj))); %sum of weights between nodes in Cj
235 Et = sum (sum (Mat (Cj, :))); %sum of weights of nodes incident in nodes
            of Cj
236 if Et >0
237 MOD = MOD + Ec/(2*m)-(Et/(2*m))^2;
238 end
239 end
240 end
```
Appendix B

Matlab code of the community-variance-minus method (Section 3.1)

```
1 % community - variance - minus
2\frac{9}{6}3 % Inputs :
4 % M : weight matrix
5 % lambda : weight of variance in function for cut filter
6 \text{ % } z : 1 = \text{Recursive computation}7 % : 0 = Just one level computation
8 %
9 % Output :
10 % COMTY , structure with the following information
11 % for each level i :
12 % COMTY .COM{i} : vector of community partition
13 % COMTY . SIZE {i} : vector of community sizes
14 % COMTY .MOD{i} : vector of modularities of clustering on the layers
15 % COMTY . Average (i) : average of modularity on the layers
16 % COMTY . Niter (i) : Number of iteration before convergence
17 \frac{9}{6}18 function [COMTY \tending] = community\_variance\_minus (M, lambda)19
20 if nargin \leq 121 error ('not enough argument');
22 end
23
24 if nargin < 2
25 error ('not lambda defined');
26 end
27
28 if nargin < 3
```

```
29 z = 1;
30 end
31
32 S = size(M);33 N = S(1) ;
34 if length (S) == 335 k = S(3);
36 else k = 1;
37 end
38
39 ending = 0;
40
41 for s=1:k42 M2 (:,:,s) = M(:,:,s) - diag (diag (M(:,:,s)));
43 end
44
45 % total weight of the graph
46 for s=1: k
47 if \; z == 148 M(:, :, s) = M(:, :, s) + diag(diag(M(:, :, s)));
49 end
50 m\{s\} = sum(sum(M(:,: , s)))/2;51 end
52
53 Niter = 0; % number of iterations
54
55 if (sum(cellfun((@(x)x>0,m)) == 0) | N == 156 ending = 1;
57 COMTY = 0;
58 return ;
59 end
60
61 % Delate layers with m=0
62 M(:,:, (cellfun(@(x)x == 0, m))) = [];
63 k = k - sum ( cellfun (Q(x)x == 0, m) );
64
65 COM = 1: S(1); % Community of node i
66
67 for s=1: k
68 K(:,: ,s) = sum(M(:,: ,s)); % K(i) = sum of wieght incident to node i
69 SumTot(:,:,s) = sum(M(:,:,s)); %SumTot(i) = sum of weight incident to
           nodes in community i
70 SumIn(:,:,s) = diag(M(:,:,s))'; %SumIn(i)= sum of weight inside
          community i (loops)
71 %At the beginning each node is a
                                           community
72 \qquad Q\{s\} = \text{compute\_modularity}(COM, M(:, :, s));
73 end
74 Average = sum([Q{\:};}])/k;
```

```
75 if k == 176 Variance = 0;
77 else
78 Variance = (sum(([Q\{:\}) - Average) . (2))) / (k-1); % variance
79 end
80 Function = (1-lambda) * Average - lambda * Variance; %function
81
82 % Neighbor {j}{s} neighbor of node j in layer s
83 for j=1: N
84 for s=1: k
85 temp=M2(:,:,s);
86 Neighbor_j {s} = find ( temp (j ,:) ) ;
87 end
88 \text{Neighbour} {j}=Neighbor_j;
89 end
90
91 gain = 1;
92 while (gain == 1)
93 gain = 0;
94 for i=1: N
95 Ci = COM (i) ;
96 NB = unique (cat (2, Neighbour {i} {:}) );
97 G = zeros(1, N); % Gain vector
98 best_increase = -inf;
99 Cnew = Ci;
100 COM(i) = -1;
101 % remove i from its community
102 for s=1: k
103 CNi = (COM == Ci); %list of nodes in Ci community, without i
104 Ki_in_i {s} = sum (M(i,CNi ,s) ) ; %sum of weights between i and Ci
105 % Gain of modularity
106 GQ_i {s} = K(:,i,s) * SumTot(:,Ci,s) /(2*(m{s}^(2))) - Ki_in_i{s}/m{
              s \} - ((K(:,i,s))^{(2)}) / (2 * (m[s]^{(2)}));107 %Recalculate values
108 SumTot(:,Ci,s) = SumTot(:,Ci,s) - K(:,i,s); % weights incident to
               Ci community
109 SumIn(:,Ci,s) = SumIn(:,Ci,s) - 2* sum(M(i,CNi,s)) - M(i,i,s); %
              weights community i
110 end
111 for j=1: length (NB)
112 C j = COM (NB (j));
113 if (G(C_i)) == 0)114 CNj = (COM==Cj); % nodes in community Cj, without j
115 for s=1:k
116 Ki_in_j {s} = sum(M(i, CNj, s)); %sum of weights between i and
                  Ci
117 GQ_{j} \{s\} = Ki_{in_{j}} \{s\} / m\{s\} - (K(:,i,s) * SumTot(:,Cj,s)) / (2 * (m\{s\} + N))s}<sup>^</sup>(2))); % gain deltaQ if I put isolated node i in Cj
118 end
```

```
119 % variance gain
120 if \ k == 1121 GV_j = 0;
122 else
123 for s=1:k
124 DQ{s} = GQ_i{s} + GQ_j{s}; % gains
125 end
126 M_DQ = sum ([DQ{:}]) /k; % gain average
127 GV_j = ((\text{sum}([DQ{\:}:)] - M_DQ) .^(2)) ) / (k-1) + (2/(k-1)) *sum (sum (([Q{\:}] - Average) + ([DQ{\:}] - M_DQ))) ; % gainvariance
128 end
129 % % gain of the function
130 G(Cj) = (1-\lambda) * M_DQ - \lambdambda * GV_{-j};131 if G(Cj) > best_increase
132 best_increase = G(Cj); % gain function
133 Q_t = DQ; % gain of modularity;
134 M_t = M_DQ; % gain average
135 V_t = G V_j; \% gain of variance136 Cnew_t = Cj;
137 end
138 end
139 end
_{140} if best_increase > -10^(-15)
141 Cnew = Cnew_t;
142 for s=1:k
143 Q{s} = Q{s} + Q_t{s};144 end
145 Average = Average + M_t;
146 Variance = Variance + V_t;
147 Function = Function + best_increase ;
148 end
149 % Recalculate
150 Ck = (COM = = Cnew);
151 for s=1:k
152 SumIn (:, Cnew, s) = SumIn (:, Cnew, s) + 2* sum (M(i, Ck, s)) + M(i, i, s);
153 SumTot (:, Cnew, s) = SumTot (:, Cnew, s) + K(:, i, s);
154 end
155 COM(i) = Cnew;
156 if ( Cnew ~= Ci )
157 gain = 1;
158 end
159 end
160 Niter = Niter + 1;
161 end
162 Niter = Niter - 1;
_{163} [COM] = reindex_com(COM);
164 COMTY. COM{1} = COM;
165 COMTY. MOD\{1\} = [Q\{\cdot\}];
```

```
166 COMTY . Average (1) = Average ;
167 COMTY. Variance (1) = Variance;
168 COMTY. Function (1) = Function;
169 COMTY. Niter (1) = Niter;
170
171 % Perform part 2
172 if (z == 1)173 Mnew = M;
174 Mold = Mnew;
175 COMcur = COM;
176 COMfull = COM:
177 i = 2;
178
179 while 1
180 Mold = Mnew;
181 S2 = size (Mold);
182 Nnode = S2(1);
183
184 COMu = unique (COMcur);
185 Ncom = length (COMu);
186 ind_com = sparse (Ncom, Nnode);
187 ind_com_full = sparse (Ncom, N);
188 for p=1: Ncom
189 ind = find (COMcur==p);
190 ind\_com(p, 1: length (ind)) = ind;191 end
192 for p=1: Ncom
193 ind = find (COMfull == p);
194 ind_com_full (p, 1: length (ind) ) = ind;195 end
196 Mnew = [];
197 for s=1:k
198 Mnew (i, j, s) = zeros (Ncom, Ncom); %new matrix (each node is a
             community )
199 for m=1: Ncom
200 for n=m: Ncom
201 ind1 = ind_{\text{com}}(m, :);202 ind2 = ind_com(n, :);
203 % weights of edges between communities
204 Mnew (m,n,s) = sum ( sum ( Mold (ind1 (ind1 > 0), ind2 (ind2 > 0), s)
                      ) ) ;
205 Mnew (n,m,s) = sum (sum (Mold (ind1 (ind1 >0),ind2 (ind2 >0),s))) ) ;
206 end
207 end
208 end
209 [ COMt e] = community_variance_minus (Mnew ,lambda ,0) ;
210 if (e^{\sim} = 1)211 COMfull = sparse (1, N);
```

```
212 COMcur = COMt. COM\{1\};
213 for p=1: Ncom
214 ind1 = ind_com_full(p, :);
215 COMfull (ind1 (ind1 >0)) = COMcur (p);
216 end
217 [COMfull2] = reindex_com(COMfull);
218 COMTY.COM{j} = COMfull2;
219 COMTY. MOD{j} = COMt. MOD{1};
220 COMTY . Average (j) = COMt . Average (1) ;
221 COMTY . Variance (j) = COMt . Variance (1);
222 COMTY . Function (j) = COMt . Function (1);
223 COMTY. Niter (i) = COMt. Niter :
224 Ind = (COMfull2 == COMTY.COM{j-1});
225 if (sum(Ind) == length(Ind))226 return ;
227 end
228 else
229 return ;
230 end
231 j = j + 1;
232 end
233 end
234 end
235
236 % Re - index community partition by size
237 function [C Ss] = reindex_com(COMold)
238 C = sparse (1, length (CDMold));
239 COMu = unique ( COMold ) ;
240 S = sparse (1, length(COMu));
241 for l=1:length (COMu)
S(1) = \text{length}( \text{CDMold}( \text{CDMold} == \text{CDMu}(1)));
243 end
244 [Ss INDs] = sort (S, 'descend');
245 for l=1:length (COMu)
246 C(COMold == COMu (INDs(1)) = 1;
247 end
248 end
249 % Compute modulartiy
250 function MOD = compute_modularity (C, Mat)
251
252 S = size (Mat);
253 \text{ N} = S(1);
254 m = sum (sum (Mat) ) / 2; % total weight
255 MOD = 0;
256 COMu = unique (C);
257
258 for j=1: length ( COMu )
259 Cj = (C == COMu(j)); %list of nodes in Cj
260 \, Ec = sum (sum (Mat (Cj, Cj))); %sum of weights between nodes in Cj
```

```
261 Et = sum (sum (Mat (Cj ,:) ) ) ; %sum of weights of nodes incident in nodes
           of Cj
262 if Et >0
263 MOD = MOD + Ec/(2*m) -(Et/(2*m)) ^2;
264 end
265 end
266 end
```
Appendix C

Matlab code of the community-variance-plus method (Section 3.1)

```
1 % community - variance - plus
2\frac{9}{6}3 % Inputs :
4 % M : weight matrix
5 % lambda : weight of variance in function for cut filter
6 \text{ % } z : 1 = \text{Recursive computation}7 % : 0 = Just one level computation
8 %
9 % Output :
10 % COMTY , structure with the following information
11 % for each level i :
12 % COMTY .COM{i} : vector of community partition
13 % COMTY . SIZE {i} : vector of community sizes
14 % COMTY .MOD{i} : vector of modularities of clustering on the layers
15 % COMTY . Average (i) : average of modularity on the layers
16 % COMTY . Niter (i) : Number of iteration before convergence
17 \frac{9}{6}18 function [COMTY \end{equation} = community_variance_plus (M, \text{lambda}, z)19
20 if nargin \leq 121 error ('not enough argument');
22 end
23
24 if nargin < 2
25 error ('not lambda defined');
26 end
27
28 if nargin < 3
```

```
29 z = 1;
30 end
31
32 S = size(M);33 N = S(1) ;
34 if length (S) == 335 k = S(3);
36 else k = 1;
37 end
38
39 ending = 0;
40
41 for s=1:k42 M2 (:,:,s) = M(:,:,s) - diag (diag (M(:,:,s)));
43 end
44
45 % total weight of the graph
46 for s=1: k
47 if \; z == 148 M(:, :, s) = M(:, :, s) + diag(diag(M(:, :, s)));
49 end
50 m\{s\} = sum(sum(M(:,: , s)))/2;51 end
52
53 Niter = 0; % number of iterations
54
55 if (sum(cellfun((@(x)x>0,m)) == 0) | N == 156 ending = 1;
57 COMTY = 0;
58 return ;
59 end
60
61 % Delate layers with m=0
62 M(:,:, (cellfun(@(x)x == 0, m))) = [];
63 k = k - sum ( cellfun (Q(x)x == 0, m) );
64
65 COM = 1: S(1); % Community of node i
66
67 for s=1: k
68 K(:,: ,s) = sum(M(:,: ,s)); % K(i) = sum of wieght incident to node i
69 SumTot(:,:,s) = sum(M(:,:,s)); %SumTot(i) = sum of weight incident to
           nodes in community i
70 SumIn(:,:,s) = diag(M(:,:,s))'; %SumIn(i)= sum of weight inside
          community i (loops)
71 %At the beginning each node is a
                                           community
72 \qquad Q\{s\} = \text{compute\_modularity}(COM, M(:, :, s));
73 end
74 Average = sum([Q{\:};}])/k;
```

```
75 if k == 176 Variance = 0;
77 else
78 Variance = (sum(([Q\{:\}) - Average) . (2))) / (k-1); % variance
79 end
80 Function = (1-lambda) * Average + lambda * Variance; %function
81
82 % Neighbor {j}{s} neighbor of node j in layer s
83 for j=1: N
84 for s=1: k
85 temp=M2(:,:,s);
86 Neighbor_j {s} = find ( temp (j ,:) ) ;
87 end
88 \text{Neighbour} {j}=Neighbor_j;
89 end
90
91 gain = 1;
92 while (gain == 1)
93 gain = 0;
94 for i=1: N
95 Ci = COM (i) ;
96 NB = unique (cat (2, Neighbour {i} {:}) );
97 G = zeros(1, N); % Gain vector
98 best_increase = -inf;
99 Cnew = Ci;
100 COM(i) = -1;
101 % remove i from its community
102 for s=1: k
103 CNi = (COM == Ci); %list of nodes in Ci community, without i
104 Ki_in_i {s} = sum (M(i,CNi ,s) ) ; %sum of weights between i and Ci
105 % Gain of modularity
106 GQ_i {s} = K(:,i,s) * SumTot(:,Ci,s) /(2*(m{s}^(2))) - Ki_in_i{s}/m{
              s \} - ((K(:,i,s))^{(2)}) / (2 * (m[s]^{(2)}));107 %Recalculate values
108 SumTot(:,Ci,s) = SumTot(:,Ci,s) - K(:,i,s); % weights incident to
               Ci community
109 SumIn(:,Ci,s) = SumIn(:,Ci,s) - 2* sum(M(i,CNi,s)) - M(i,i,s); %
              weights community i
110 end
111 for j=1: length (NB)
112 C j = COM (NB (j));
113 if (G(C_i)) == 0)114 CNj = (COM==Cj); % nodes in community Cj, without j
115 for s=1:k
116 Ki_in_j {s} = sum(M(i, CNj, s)); %sum of weights between i and
                  Ci
117 GQ_{j} \{s\} = Ki_{in_{j}} \{s\} / m\{s\} - (K(:,i,s) * SumTot(:,Cj,s)) / (2 * (m\{s\} + N))s}<sup>^</sup>(2))); % gain deltaQ if I put isolated node i in Cj
118 end
```

```
119 % variance gain
120 if \ k == 1121 GV_j = 0;
122 else
123 for s=1:k
124 DQ{s} = GQ_i{s} + GQ_j{s}; % gains
125 end
126 M_DQ = sum ([DQ{:}]) /k; % gain average
127 GV_j = ((\text{sum}([DQ{\:}:)] - M_DQ) .^(2)) ) / (k-1) + (2/(k-1)) *sum (sum (([Q{\:}] - Average) + ([DQ{\:}] - M_DQ))) ; % gainvariance
128 end
129 % % gain of the function
130 G(Cj) = (1-\lambda) * M_DQ + \lambda131 if G(Cj) > best_increase
132 best_increase = G(Cj); % gain function
133 Q_t = DQ; % gain of modularity;
134 M_t = M_DQ; % gain average
135 V_t = G V_j; \% gain of variance136 Cnew_t = Cj;
137 end
138 end
139 end
_{140} if best_increase > -10^(-15)
141 Cnew = Cnew_t;
142 for s=1:k
143 Q{s} = Q{s} + Q_t{s};144 end
145 Average = Average + M_t;
146 Variance = Variance + V_t;
147 Function = Function + best_increase ;
148 end
149 % Recalculate
150 Ck = (COM = = Cnew);
151 for s=1:k
152 SumIn (:, Cnew, s) = SumIn (:, Cnew, s) + 2* sum (M(i, Ck, s)) + M(i, i, s);
153 SumTot (:, Cnew, s) = SumTot (:, Cnew, s) + K(:, i, s);
154 end
155 COM(i) = Cnew;
156 if ( Cnew ~= Ci )
157 gain = 1;
158 end
159 end
160 Niter = Niter + 1;
161 end
162 Niter = Niter - 1;
_{163} [COM] = reindex_com(COM);
164 COMTY. COM{1} = COM;
165 COMTY. MOD\{1\} = [Q\{\cdot\}];
```

```
166 COMTY . Average (1) = Average ;
167 COMTY. Variance (1) = Variance;
168 COMTY. Function (1) = Function;
169 COMTY. Niter (1) = Niter;
170
171 % Perform part 2
172 if (z == 1)173 Mnew = M;
174 Mold = Mnew;
175 COMcur = COM;
176 COMfull = COM:
177 i = 2;
178
179 while 1
180 Mold = Mnew;
181 S2 = size (Mold);
182 Nnode = S2(1);
183
184 COMu = unique (COMcur);
185 Ncom = length (COMu);
186 ind_com = sparse (Ncom, Nnode);
187 ind_com_full = sparse (Ncom, N);
188 for p=1: Ncom
189 ind = find (COMcur==p);
190 ind\_com(p, 1: length (ind)) = ind;191 end
192 for p=1: Ncom
193 ind = find (COMfull == p);
194 ind_com_full (p, 1: length (ind) ) = ind;195 end
196 Mnew = [];
197 for s=1:k
198 Mnew (i, j, s) = zeros (Ncom, Ncom); %new matrix (each node is a
             community )
199 for m=1: Ncom
200 for n=m: Ncom
201 ind1 = ind_{\text{com}}(m, :);202 ind2 = ind_com(n, :);
203 % weights of edges between communities
204 Mnew (m,n,s) = sum ( sum ( Mold (ind1 (ind1 > 0), ind2 (ind2 > 0), s)
                      ) ) ;
205 Mnew (n,m,s) = sum (sum (Mold (ind1 (ind1 >0),ind2 (ind2 >0),s))) ) ;
206 end
207 end
208 end
209 [ COMt e] = community_variance_plus (Mnew ,lambda ,0) ;
210 if (e^{\sim} = 1)211 COMfull = sparse (1, N);
```

```
212 COMcur = COMt. COM\{1\};
213 for p=1: Ncom
214 ind1 = ind_com_full(p, :);
215 COMfull (ind1 (ind1 >0)) = COMcur (p);
216 end
217 [COMfull2] = reindex_com(COMfull);
218 COMTY.COM{j} = COMfull2;
219 COMTY. MOD{j} = COMt. MOD{1};
220 COMTY . Average (j) = COMt . Average (1) ;
221 COMTY . Variance (j) = COMt . Variance (1);
222 COMTY . Function (j) = COMt . Function (1);
223 COMTY. Niter (i) = COMt. Niter ;
224 Ind = (COMfull2 == COMTY.COM{j-1});
225 if (sum(Ind) == length(Ind))226 return ;
227 end
228 else
229 return ;
230 end
231 j = j + 1;
232 end
233 end
234 end
235
236 % Re - index community partition by size
237 function [C Ss] = reindex_com(COMold)
238 C = sparse (1, length (CDMold));
239 COMu = unique ( COMold ) ;
240 S = sparse (1, length(COMu));
241 for l=1:length (COMu)
S(1) = \text{length}( \text{CDMold}( \text{CDMold} == \text{CDMu}(1)));
243 end
244 [Ss INDs] = sort (S, 'descend');
245 for l=1:length (COMu)
246 C(COMold == COMu (INDs(1)) = 1;
247 end
248 end
249
250 % Compute modulartiy
251 function MOD = compute_modularity (C,Mat)
252
253 S = size (Mat);
254 \text{ N} = S(1);
255 m = sum (sum (Mat) ) /2; % total weight
256 MOD = 0;
257 COMu = unique (C);
258
259 for j=1: length ( COMu )
260 Cj = (C == CDMu(j)); %list of nodes in Cj
```

```
261 Ec = sum (sum (Mat (Cj ,Cj) ) ) ; %sum of weights between nodes in Cj
262 Et = sum (sum (Mat (c_j, ...)); %sum of weights of nodes incident in nodes
           of Cj
263 if Et >0
264 MOD = MOD + Ec/(2*m)-(Et/(2*m))^2;
265 end
266 end
267 end
```
Appendix D

Matlab code of the multi-average method (Section 3.2)

```
1 %multi - average
2\frac{9}{6}3 % Inputs :
4 % M : vector of weight matrix of each layer
5 % h : length of the filter
6 \text{ % } z : 1 = \text{Recursive computation}7 % : 0 = Just one level computation
8 %
9 % Output :
10 % L the filter with the following information
11 % for each element l:
12 % 1{1}=Q cell with modularity of each layer
13 % 1{2}={COMcur COMfull COMindex} communities in the current graph, in
      the
14 % original graph , communities in the current graph reindexed
15 % l {3}= SumTot (sum of weights of the links incident to nodes in a
      community )
16 % l {4}= SumIn (sum of weights of the links inside a community )
17 \% 1{5}= Average (average)
18 %
19
20
21 function [L \text{ ending}] = \text{multi\_average}(M, h, z)22 if nargin < 1
23 error ('not enough argument');
24 end
25 if nargin < 2
26 error ('not h defined');
27 end
28 if nargin < 3
29 z = 1;
```

```
30 end
31
32 S = size(M);33 \text{ N} = S(1);
34 if length (S) = 335 k = S(3);
36 else k = 1;
37 end
38
39 ending = 0;
40
41 for s=1:k42 M2 (:,:,s) = M(:,:,s) - diag (diag (M(:,:,s)));
43 end
44
45 % total weight of the graph
46 for s=1: k
47 if \; z == 148 M(:, :, s) = M(:, :, s) + diag(diag(M(:, :, s)));
49 end
50 m{s} = sum(sum(M(:,: , s)))/2;51 end
52
53 Niter = 0; % number of iterations
54
55 % Calculation of the beginning values
56 COM\{1\} = 1:S(1); % current community %At the beginning each node is a
      community
57 COM\{2\} = 1:S(1); % original graph commuity
58 COM{3} = 1:S(1); % community reindexed
59 %COM(i)= Community of node i
60
61 for s=1:kK(:,: ,s) = sum(M(:,: ,s)); % K(i) = sum of wieght incident to node i
63 SumTot(:,:,s) = sum(M(:,:,s)); %SumTot(i) = sum of weight incident to
           nodes in community i
64 SumIn(:,:,s) = diag(M(:,:,s))'; %SumIn(i)= sum of weight inside
          community i (loops)
65 %At the beginning each node is a
                                           community
66 Q{s} = compute_model{andularity(COM{1}, M(:,:, s))};67 end
68 % average
69 Average = sum([Q{\:};}] ) / k;
70
71 % filter
72 L = \{ [ {Q}, {COM}, {SumTot}, {SumIn}, Average] \};73 %If no edges in any layer or just one node
74 if (sum(cellfun((\mathfrak{C}(x)x>0, m)) == 0) | N == 1 \% logical or
```

```
75 ending = 1;
76 else
77
78 % Delate layers with m=0
79 \text{ M}(:,:,(cellfun(\mathcal{Q}(x)x == 0, m)))=[];
80 k = k - sum ( cellfun (Q(x)x == 0, m) );
81
82 % Neighbor {j}{s} neighbor of node j in layer s
83 for j=1: N
84 for s=1: k
85 temp=M2(:,:,s);
86 Neighbor_j {s} = find ( temp (j ,:) ) ;
87 end
88 Neighbor {j}= Neighbor_j ;
89 end
90
91
92 GAIN = 1;
93 while ( GAIN == 1) % Stop when putting nodes in other communities do not
      increase the modularity
94    L_old=L;
95 L_new=L;
96 for i=1:N %for each node, in this order
97 L_o = L;
98 for 1=1: length (L_o) %for each situation in the filter
99 % delate the point from the filter
100 L_new_0 = L_new;101 index = cellfun (\mathcal{C}(x) isequal (x, L_o{1}), L_new, 'UniformOutput'
                , 1) ;
102 L_new (index) = [];
103 % step1
104 [L_new, U] = step_1 \left( \{L_0\{1\}\{1\}\}, \{L_0\{1\}\{2\}\}, L_0\{1\}\{3\}, L_0\{1\}}{4} , L_o {l}{5} ,k, Neighbor ,K,M,N,m,L_new ,h,i) ;
105 %if no new point , insert again the initial point in the filter
106 if U == 0\verb|L_new=L_new_0|;108 end
109 end
110 L_new = cut_filter (L_new, h, k);
\mathtt{l}=\mathtt{L\_new} \; ;112 end
113 % Check if nothing happened
114 if length (L_old) "=length (L_new)
115 GAIN = 1;
116 else
117 C_01d = [];
118 C_{\text{new}} = [];
119 for l=1:length (L_old)
R_old = [];
```

```
121 R new = [];
122 for s=1:kR_old = [R_old L_old{1}{1}{s}];
124 R_new = [R_new L_new{1}{1}{s};
125 end
126 C_{old} = [C_{old} ; R_{old}];
127 C_new = [C_new; R_new];128 end
129 C = \text{ismembertol}(C_old, C_new, 10^(-6)) ;130 if size(C, 1) * size(C, 2) == sum(sum(C))131 GAIN=0;
132 end
133 end
134 Niter = Niter + 1;
135 end
136 end
137
138 % Perform part 2
139 if (z == 1)140 L = cut_fitter(L,1,k); % one final element141 Mnew=M; % new weight matrix of this iteration
142 Mold = Mnew ; % old weight matrix of iteration befor
143 COMcur = L {1}{2}{3}; % communities in the graph of this iteration
144 COMfull = L {1}{2}{3}; % communities in the original graph
145 LL=L;
146 j=2; % number of pass (step1+step2)
147 S=1;148 while S
149 L_old=LL;
150 Mold = Mnew;
151 S2 = size (Mold); % number of nodes
152 Nnode = S2(1);
153
154 COMu = unique ( COMcur ) ; % list of communities without repetitions
             and in sorted order
155 Ncom = length ( COMu ) ; % number of communities
156 ind_com = sparse (Ncom , Nnode ) ; % zero matrix with a row for each
             community and a column for each node in this configuration
157 ind_com_full = sparse (Ncom ,N) ; % zero matrix with a row for each
             community and a column for each node of the original graph
158
159 for p=1: Ncom % for each community
160 ind = find (COMcur==p);
161 ind_com(p, 1: length(ind)) = ind;
162 end
163 for p=1: Ncom
164 ind = find (COMfull == p);
165 ind_com_full (p, 1: length (ind) ) = ind;166 end
```

```
167 Mnew=[];
168 for s=1:k
169 Mnew (:,:,s) = sparse (Ncom, Ncom); %new matrix (each node is
              a community )
170 for m=1: Ncom
171 for n=m: Ncom
172 ind1 = ind_{\text{1}}(m, :);173 ind2 = ind2 com(n, :);
174 % weights of edges between communities
Mnew(m,n,s) = sum(sum(Mold(ind1(cind1>0),ind2(cind2>0))), s)) ) :
176 Mnew (m,m,s) = sum (sum (Mold (ind1 (ind1 > 0) ,ind2 (ind2 > 0))), s) ) ) :
177 end
178 end
179 end
180 % apply first step to this new matrix but z=0 without recursive
181 [LLL e] = multi_average (Mnew, h, 0);
182 if isempty (LLL)
183 L=LLL;
184 return
185 else
186 LL = cut_finter(LLL, 1, k); %one final element
187 \%if (e \tilde{=} 1)
188 COMfull = sparse (1, N); % communities of the original graph
189 COMcur = LL{1}{2}{3}; % communities
190 for p=1: Ncom
191 ind1 = ind_com_full (p, :); % nodes in community p in the
                 original graph
192 COMfull (ind1 (ind1 > 0) = COMcur (p); % community now of
                 node i
193 end
194 [COMfull] = reindex_com (COMfull);
195 LL\{1\}\{2\}\{2\}= COMfull;
196 % communities do not change
197 if isequal (L_old {1}{2}{2}, LL {1}{2}{2})
198 % return not just this value , but all the value of the
                 last list
199 % calculate COMfull for each element of the last list
200 for l=1: length (LLL)
201 COMfull = sparse (1 ,N) ; % communities of the original
                    graph
202 COMcur = LLL{1}{2}{3}; % communities
203 for p=1:Ncom204 ind1 = ind_com_full (p ,:) ; % nodes in community p
                        in the original graph
205 COMfull (ind1 (ind1>0)) = COMcur (p); % community
                        now of node i
206 end
```

```
207 [COMfull] = reindex_com (COMfull);
208 LLL {1} {2} {2} = COMfull;
209 end
_{210} L = LLL;
211 S=0;
212 end
213 j = j + 1; % start another pass
214 tEnd = toc;
215 if tEnd > 1200
216 L = \{\}:
217 return
218 end
219 end
220 end
221 end
222 end
223224 % Compute step_1
225 function [L,U] = step_1 (Q,COM ,SumTot ,SumIn , Average ,k, Neighbor ,K,M,N,m,L,
      h,i)
226
227 U=0; %if U=1 add at least an element to the list, otherwise put again
      the initial point in the filter
228 COMcur=COM{1}{1}; % current community
229 Ci = COMcur (i) ; % community of node i
230
231 NB= unique (cat (2, Neighbour {i}{:}));
232
233 SumTot2=SumTot:
234 SumIn2=SumIn;
235
236 % remove i from its community
237 for s=1:k238 COMcur2 = COMcur;
239 \text{CMcur2(i)} = -1;240 CNi = ( COMcur2 == Ci) ; % list of nodes in Ci community , without i
241 Ki_in_i {s} = sum (M(i,CNi ,s) ) ; %sum of weights between i and Ci
GQ_i is = (K(:,i,s) * SumTot2(:,Ci,s)) / (2 * (m{s}^c)(2)) - Ki_in_i {s} / m{s}\} - ((K(:,i,s)) \hat{(2)}) / (2 * (m\{s\} \hat{(2)}));243 % Recalculate values
244 SumTot2(:,Ci,s) = SumTot2(:,Ci,s) - K(:,i,s); % weights incident to
         Ci community
245 SumIn2(:,Ci,s) = SumIn2(:,Ci,s) - 2* sum(M(i,CNi,s)) - M(i,i,s); \frac{9}{6}weights community i
246 end
247
248 G = sparse(1, N);249 for j=1: length(NB)250 SumTot3=SumTot2;
```
```
251 SumIn3=SumIn2;
252 Cj = COMcur (NB(j) ) ; % community of node j
253 if (G(Cj) == 0) %If not tried with another node of community Cj yet
254 G (C<sub>j</sub>) =1;
255 for s=1:k
256 COMcur3 = COMcur;
257 COMcur3(i) = -1;
258 %I put i in the community of j for each layer
259 CNj = ( COMcur3 == Cj) ; % nodes in community Cj , without j
260 Ki_in_j{s} = sum(M(i,CNj,s)); %sum of weights between i and
                  Ci
261 GQ_j{s} = Ki_in_j{s}/m{s} - (K(:,i,s)*SumTot3(:,Cj,s))/(2*(m
                  \{s\}^(2)) ; % gain deltaQ if I put isolated node i in Cj
262 % Recalculate
263 SumTot3(i, C_j, s) = SumTot3(i, C_j, s) + K(i, i, s);
264 SumIn3(:,\text{Cj},\text{s}) = SumIn3(:,\text{Cj},\text{s}) + 2* \text{sum}(M(i,\text{CNj},\text{s})) + M(i,i,s) ;
265 end
266 COMcur4=COMcur;
267 COMcur4 (i) = Cj;
268 COM\{1\}\{1\}= COMcur4 ;
269 [COMcur4] = reindex_com(COMcur4);
270 COM {1} {3} = COMcur4 ;
271 for s=1:k272 DQ{s} = GQ_i{s} + GQ_j{s}; % gain modularity
273 end
274 GA_j = sum ([DQ{:}])/k; % gain average
275 if GA_{-j} > -10^(-14)276 A_j = Average + GA_j;
277 % modularity
278 for s=1:k
279 Q_{j} \{s\} = Q \{1\} \{s\} + DQ \{s\};280 end
281 [L] = add_to_finter (\lbrace Q_i \rbrace, COM, SumTot3, SumIn3, A_i, L, k);282 U=1;
283 end
284 end
285 end
286 end
287
288 %add a point to the filter
289 function [L] = add_to_filter (QQ ,COM ,SumTot ,SumIn , Average ,L,k)
290
291 if length (L) z = 0292
293 COMcur = COM {1}{1}; % current community
294
295 %1. Check if the new point is dominated by a point in the filter
296 DD=0;
```

```
297 \quad 1=1;298 while (DD == 0) & 1 \leq -\operatorname{length}(L)299 D=0;
300 s=1;
301 while (D == 0) & s <= k302 if QQ {1}{ s} >L{l }{1}{ s}
303 D = 1;
304 end
305 S=S+1;306 end
307 if D %the new point is not dominated by l
308 DD =1;
309 end
310 1=1+1;311 end
312 if DD % the new point is not dominated
313 DDD=0;
314 1=1;315 while (DDD == 0) & l <= length (L)
316 D=1;
317 S=1;
318 while D & s \le k319 if abs (L{1}{t}{s} - Q{t}{s}) > 10^(-14)320 D = 0;
321 end
322 s = s + 1;
323 end
324 if D
325 DDD = 1;
326 end
327 1 = 1 + 1;
328 end
329
330 if DDD ==0
331
332 %2. Check if the other points in the filter are dominated by the new
      point %
333
334 for l=1: length (L)
335 D=1;
336 S=1;337 while D & s \le k338 if L{1}{s} > QQ{1}{s} + 10^(-14)339 D = 0;
340 end
341 S=S+1;
342 end
343 if D %l is dominated by the new point so I remove it from the
           filter
```

```
344 L{1}=[];
345 end
346 end
347
348 empties =( cellfun ( @isempty ,L) ) ;
349 L(empties) = [];
350 L{end +1}=[ QQ ,COM ,SumTot ,SumIn , Average ]; %Add the new point to the
         filter
351
352 end
353 end
354 else
355 L{end +1}=[ QQ ,COM ,SumTot ,SumIn , Average ]; %Add the new point to the
           filter
356 end
357 end
358
359 %cut the filter
360 function [L] = cut_fitter(L,h,k)361
362 % calculate the function for every element of the list
363 if length (L) >h
364 for l=1: length (L)
365 % function
366 Average {l} = L{l }{5};
367 end
368
369 % remuve the situations in the filter with the lower values
370 for t=1: (\text{length} (L) - h)371 [mn,idx]=min([Average{:}]);
372 v=find ([Average {: }] == mn); %if more element with same average values,
         delate the last one ( because with h=1 I have same result as
         cluster_jl_average_variance )
373 idx_t = v(\text{end});
374 Average {idx_t}=Inf;
375 L{idx_t}=[];
376 end
377
378 empties = find (cellfun (@isempty, L));
379 L( empties ) = [];
380 end
381 end
382
383 % Compute modulartiy
384 function MOD = compute_modularity (C,Mat)
385
386 S = size (Mat);
387 \quad N = S(1);
388
```

```
389 m = sum(sum(Mat))/2; % total weight390
391 MOD = 0;
392 COMu = unique (C); %list of communities without repetiotions and in
      sorted order
393 %for each community calculate modularity and then sum all together
394 for j=1: length (CDMu)395 Cj = (C== COMu (j) ) ; % list of nodes in Cj
% faster then C j = find (C== COMu (j))
397 Ec = sum (sum (Mat (Cj, Cj))); %sum of weights between nodes in Cj
\%Mat(C_j, C_j) submatrix
399 Et = sum (sum (Mat (C<sub>j</sub>,:))); %sum of weights of nodes incident in nodes
           of Cj
400 if Et >0
401 MOD = MOD + Ec/(2*m)-(Et/(2*m))<sup>-2</sup>;
402 end
403 end
404
405 end
406
407 % Re-index community partition by size
408 function [C Ss] = reindex_com ( COMold )
409 C = sparse (1, length (CDMold));
410 COMu = unique (COMold);
411 S = sparse(1, length(COMu));
412 for l=1: length ( COMu )
413 S(1) = \text{length}(COMold(COMold == COMu(1)));414 end
415 [Ss INDs] = sort (S, 'descend');
416 for l=1: length (COMu)
417 C(COMold == COMu(INDs(1))) = 1;418 end
419 end
420 %Re - index community partition no by size but by initial order
421 % function [C] = reindex_com (COMold)
422 %C = sparse (1 , length ( COMold )); % vector with a index for each node
423 % COMu = unique (COMold); % unique (v) = same data of v but not repetitions
      and in sorted order
424 % COMolds = sort ( COMold );
425 % COMu = COMolds ([ true ; diff ( COMolds (:)) >0]);
426 %for l=1:length (COMu)
427 % C(COMold = COMu(1)) = 1; %Riname the communities
428 %end
429 %end
```
Appendix E

Matlab code of the multi-variance-minus method (Section 3.2)

```
1 %multi - variance - minus
2 \frac{9}{6}3 % Inputs :
4 % M : vector of weight matrix of each layer
5 \text{ % } z : 1 = \text{Recursive computation}6 % : 0 = Just one level computation
7 % Output :
8 % L the filter with the following information
9 % for each element l:
10 % l {1}= Q cell with modularity of each layer
11 % 1{2}={COMcur COMfull COMindex} communities in the current graph, in
       the
12 % original graph, communities in the current graph reindexed
13 % l {3}= SumTot (sum of weights of the links incident to nodes in a
      community )
14 % l {4}= SumIn (sum of weights of the links inside a community )
15 % l {5}= Average ( average )
16 \t% 16 = Function (funtion)
17 \frac{9}{6}18
19
20 function [L \text{ ending}] = \text{multi\_variance\_minus}(M, \text{lambda}, h, z)21
22 if nargin < 1
23 error ('not enough argument');
24 end
25 if nargin < 2
26 error ('not lambda defined');
```

```
27 end
28 if nargin < 3
29 error ('not h defined');
30 end
31 if nargin < 4
32 z = 1;
33 end
34
35 S = size (M);
36 \text{ N} = S(1);
37 if length (S) ==3
38 \text{ k} = S(3) ;
39 else k = 1;
40 end
41
42 ending = 0;
43
44 for s=1: k
45 M2 (:,:,s) = M (:,:,s) - diag (diag (M (:,:,s)));
46 end
47
48 % total weight of the graph
49 for s=1: k
50 if z==151 M(:, :, s) = M(:, :, s) + diag(diag(M(:, :, s)));
52 end
53 m{s} = sum(sum(M(:,: , s)))/2;54 end
55
56 Niter = 0; % number of iterations
57
58 % Calculation of the beginning values
59 COM\{1\} = 1:S(1); % current community %At the beginning each node is a
      community
60 COM\{2\} = 1:S(1); % original graph commuity
61 COM\{3\} = 1:S(1); % community reindexed
62 %COM(i)= Community of node i
63
64 for s=1: k
65 K(:,:,s) = sum(M(:,:,s)); % K(i) = sum of weight incident to node i66 SumTot(:,:,s) = sum(M(:,:,s)); %SumTot(i)= sum of weight incident to
           nodes in community i
67 SumIn(:,:,s) = diag(M(:,:,s))'; %SumIn(i)= sum of weight inside
          community i (loops)
68 %At the beginning each node is a
                                           community
69 Q{s} = compute_modelarity (COM{1}, M(:,:, s));
70 end
71 % average
```

```
72 Average = sum([Q{\:}] ) / k;
73 % variance
74 if k == 175 Variance = 0;
76 else
77 Variance = (sum(([Q{\:}:)] - Average) . (2)) ) / (k-1) ;78 end
79 % function
80 Function = (1-lambda) * Average - lambda * Variance;
81
82 % filter
83 L = \{ [0], [COM], SumTot, SumIn, Average, Function] \};
84 %If no edges in any layer or just one node
85 if (sum(cellfun((@(x)x)>0,m)) ==0) | N == 1 %| logical or86 ending = 1;
87 else
88
89 % Delate layers with m=0
90 M(:,:,(cellfun(\mathcal{Q}(x)x == 0, m)))=[];
91 k = k - sum(cellfun((@x)x == 0, m));92
93 % Neighbor {j}{s} neighbor of node j in layer s
94 for j=1: N
95 for s=1: k
96 temp=M2(:,:,s);
97 \text{Neighbour}_j\{s\} = \text{find}(\text{temp}(j,:));
98 end
99 Neighbor {j}= Neighbor_j ;
100 end
101
102
103 GAIN = 1:
104 while ( GAIN == 1) % Stop when putting nodes in other communities do not
      increase the modularity
105 L_old=L;
106 L_new=L;
107 for i=1:N %for each node, in this order
108 L_0 = L;109 for l=1: length (L_o) %for each situation in the filter
110 % Melate the point from the filter
111 L_new_o=L_new;
112 index = cellfun(Q(x) isequal(x, L_0{1}), L_new, 'UniformOutput'
                 , 1) ;
113 L_new(intdev) = [];
114 % step1
115 [L_new, U] = step_1 \left( \{L_0\{1\}\{1\}\}, \{L_0\{1\}\{2\}\}, L_0\{1\}\{3\}, L_0\{1\}}{4} , L_o {l}{5} , L_o {l}{6} ,k, Neighbor ,K,M,N,m,L_new ,h,lambda ,
                 i) ;
116 %if no new point , put again the initial point in the filter
```

```
117 if U=-0118 L_new=L_new_o;
119 end
120 end
121 L_new = cut_filter (L_new, h, k, lambda);
122 L=L_new;
123 end
124 % Check if nothing happened
125 if length (L_old) = length (L_new)126 GAIN = 1;
127 else
128 C old = [];
129 C new = [];
130 for l=1:length (L_old)131 R_{\text{ol}}d = [];
132 R_new=[];
133 for s=1:k
134 R_old = [R_old L_old{1}{1}{s}];
135 R_new = [R_new L_new{1}{1}{s}136 end
137 C_01d = [C_01d ; R_01d];138 C_new = [C_new; R_new];139 end
140 C = \text{ismembertol}(C\_old, C\_new, 10^(-6));
141 if size(C, 1) * size(C, 2) == sum(sum(C))142 GAIN = 0;
143 end
144 end
145 Niter = Niter + 1;
146 end
147 end
148
149 % Perform part 2
150 if (z == 1)151 L = cut_fitter(L,1,k); % one final element152 Mnew =M; % new weight matrix of this iteration
153 Mold = Mnew ; % old weight matrix of iteration befor
154 COMcur = L {1}{2}{3}; % communities in the graph of this iteration
155 COMfull = L {1}{2}{3}; % communities in the original graph
156 LL=L;
157 j=2; \frac{0}{2} number of pass (step1+step2)
158 S=1;
159 while S
160 L_old=LL;
161 Mold = Mnew;
162 S2 = size (Mold); % number of nodes
163 Nnode = S2(1);
164
165 COMu = unique ( COMcur ) ; % list of communities without repetitions
```

```
and in sorted order
166 Ncom = length (COMu); % number of communities
167 ind_com = sparse (Ncom, Nnode); % zero matrix with a row for each
            community and a column for each node in this configuration
168 ind_com_full = sparse (Ncom ,N) ; % zero matrix with a row for each
            community and a column for each node of the original graph
169
170 for p=1: Ncom % for each community
171 ind = find (COMcur == p);
172 ind_com (p ,1: length (ind ) ) = ind ; %in row p put the indeces of
                the nodes in community p in this configuration
173 end
174 for p=1: Ncom
175 ind = find (COMfull ==p);
176 ind_com_full (p ,1: length (ind ) ) = ind ; %in row p put the
               indeces of the nodes in community p in the original graph
177 end
178 Mnew=[];
179 for s=1:k
180 Mnew (:,:,s) = sparse (Ncom, Ncom); %new matrix (each node is
               a community )
181 for m=1: Ncom
182 for n=m: Ncom
183 ind1 = ind_{\text{1}}(m, :);184 ind2 = ind2 com(n, :);
185 % weights of edges between communities
186 Mnew (m,n,s) = sum ( Mold (ind1 (ind1 >0), ind2 (ind2 >0)
                      ,s) ) ) ;
187 Mnew (n,m,s) = sum ( Mold (ind1 (ind1 >0), ind2 (ind2 >0)
                      ,s) ) ) ;
188 end
189 end
190 end
191 % apply first step to this new matrix but z=0 without recursive
192 [LLL e] = multi_variance_minus (Mnew, lambda, h, 0);
193 if isempty (LLL)
194 L=LLL;
195 return
196 else
197 LL = cut_fitter(LLL,1,k); %one final element
198 COMfull = sparse (1, N); % communities of the original graph
199 COMcur = LL{1}{2}{3}; % commuities200 for p=1:Ncom201 ind1 = ind_com_full (p, :); % nodes in community p in the
                   original graph
202 COMfull ( ind1 (ind1 >0) ) = COMcur (p) ; % community now of
                   node i
203 end
204 [COMfull] = reindex_com (COMfull);
```

```
205 LL \{1\} \{2\} \{2\} = COMfull ;
206 % communities do not change
207 if isequal (L_old {1}{2}{2}, LL {1}{2}{2})
208 % return not just this value , but all the value of the
                   last list
209 % calculate COMfull for each element of the last list
210 for l=1: length (LLL)
211 COMfull = sparse (1, N); % communities of the original
                      graph
212 COMcur = LLL{1}{2}{3}; % communities
213 for p=1: Ncom
214 ind1 = ind_com_full (p ,:) ; % nodes in community p
                          in the original graph
215 \text{CDMfull}(\text{ind1}(\text{ind1} > 0)) = \text{CDMcur(p)}; % community
                         now of node i
216 end
217 [COMfull] = reindex_com (COMfull);
218 LLL\{1\}{2}{2}= COMfull;
219 end
220 L = LLL;
221 S=0;
222 end
223 j = j + 1; % start another pass
224 tEnd = toc;
225 if tEnd > 1200
226 L = \{\};
227 return
228 end
229 end
230 end
231 end
232 end
233
234 % Compute step_1
235 function [L,U] = step_1 (Q,COM ,SumTot ,SumIn , Average , Function ,k, Neighbor ,K
     ,M,N,m,L,h,lambda ,i)
236
237 U=0; %if U=1 add at least an element to the list, otherwise insert again
      the initial point in the filter
238 COMcur=COM{1}{1}; % current community
239 Ci = CDMcur(i); % community of node i240
241 NB=unique (cat (2, Neighbor\{i\}\{:\}));
242
243 SumTot2 = SumTot ;
244 SumIn2=SumIn;
245
246 % remove i from its community
247 for s=1:k
```

```
248 COMcur2 = COMcur;
249 COMcur2(i) = -1;
250 CNi = ( COMcur2 == Ci) ; % list of nodes in Ci community , without i
251 Ki_in_i{s} = sum(M(i,CNi,s)); %sum of weights between i and Ci
252 GQ_i{s} = (K(:, i, s) * SumTot2(:, Ci, si, s)) / (2 * (m{s}^(2))) - Ki_in_i{s}/m{s}\} - ((K(:,i,s)) \hat{(2)}) / (2 * (m\{s\} \hat{(2)}));253 % Recalculate values
254 SumTot2(:,Ci,s) = SumTot2(:,Ci,s) - K(:,i,s); % weights incident to
          Ci community
255 SumIn2(:,\text{Ci},s) = \text{SumIn2}(:,\text{Ci},s) - 2*\text{sum}(M(i,\text{CNi},s)) - M(i,i,s);weights community i
256 end
257
258 G = sparse(1, N);259 for j=1: length (NB)
260 SumTot3=SumTot2;
261 SumIn3=SumIn2;
262 Cj = COMcur (NB(j) ) ; % community of node j
263 if (G(Cj) == 0) %If not tried with another node of community Cj yet
264 G (C<sub>j</sub>) =1;
265 for s=1:k
266 COMcur3 = COMcur;
267 COMcur3(i) = -1;
268 % Insert i in the community of j for each layer
269 CNj = ( COMcur3 == Cj) ; % nodes in community Cj , without j
270 Ki_in_j{s} = sum(M(i,CNj,s)); %sum of weights between i and
                  C<sub>j</sub>271 GQ_{j} \{s\} = Ki_{in_{j}} \{s\} / m\{s\} - (K(:,i,s) * SumTot3(:,Cj,s)) / (2 * (m{s }^(2) ) ) ; % gain deltaQ if I put isolated node i in Cj
272 %Recalculate
273 SumTot3(:, Cj, s) = SumTot3(:, Cj, s) + K(:, i, s);274 SumIn3(:, Cj, s) = SumIn3(:, Cj, s) + 2*sum(M(i, CNj, s)) + M(i, i, j)s) ;
275 end
276 COMcur4=COMcur;
277 COMcur4 (i) = Cj;
278 COM\{1\}\{1\}= COMcur4 ;
279 [COMcur4] = reindex_com(COMcur4);
280 COM\{1\}\{3\} = COMcur4 ;
281
282 % variance
283 if k == 1284 GV_j = 0;285 else
286 for s=1:k
287 DQ{s} = GQ_i{s} + GQ_j{s}; % gains
288 end
289 M_DQ = sum([DQ{\:}])/k; % gain average
290 GV_j = (( \text{sum} ((DQ {\:}:\}) - M_DQ) .^(2) )) / (k-1) ) + (2/(k-1) ) * \text{sum} (
```

```
sum (([Q{1}{:}] - Average) .* ([DQ{:}] - M_DQ))); %gain variance
291 end
292 GF_j = (1-lambda * M_DQ - lambda * GV_j;
293 if GF_j > -10^(-14)P_1 = Function + GF_1;
295 % modularity
296 for s=1:k
297 Q_j\{s\} = Q_{1}\{1\}\{s\} + DQ_{s};298 end
299 % average
300 A_i = Average + M_DQ;
301 [L] = add_to_filter ({Q_i} , COM, SumTot3, SumIn3, A<sub>-</sub>j, F<sub>-</sub>j, L, k);
302 U=1;
303 end
304 end
305 end
306 end
307
308 %add a point to the filter
309 function [L] = add_to_filter (QQ ,COM ,SumTot ,SumIn , Average , Function ,L,k)
310
311 if length (L) \tilde{ } = 0
312
313 COMcur=COM{1}{1}; % current community
314
315 %1. Check if the new point is dominated by a point in the filter
316 DD=0;
317 \quad 1=1;318 while (DD==0) & l <= length (L)
319 D=0;
320 S=1;321 while (D == 0) & s <= k322 if QQ{1}{s} > L{1}{1}{s}323 D = 1;
324 end
325 S=S+1;
326 end
327 if D %the new point is not dominated by l
328 DD = 1;
329 end
330 1=1+1;331 end
332 if DD % the new point is not dominated
333 DDD = 0;
334 1=1;335 while (DDD == 0) & l <= length (L)
336 D=1;
337 S=1;
338 while D & s <= k
```

```
339 if abs (L{1}{t}{s} - QQ{1}{s} > 10^(-14)
340 D = 0;
341 end
342 s = s + 1;
343 end
344 if D
345 DDD = 1;
346 end
347 <br> 1 = 1 + 1;348 end
349
350 if DDD ==0
351
352 %2. Check if the other points in the filter are dominated by the new
       point %
353
354 for l=1: length (L)
355 D=1;
356 s=1;
357 while D & s \le k358 if L{1}{1}{s} > QQ{1}{s} + 10^(-14)359 D = 0;
360 end
361 S=S+1;362 end
363 if D %l is dominated by the new point so I remove it from the
             filter
364 L{1}=[;
365 end
366 end
367
368 empties = (cellfun (@isempty, L);
369 L(empties) = [];
370 L{end +1}=[ QQ ,COM ,SumTot ,SumIn , Average , Function ]; %I add the new point
        to the filter
371
372 end
373 end
374 else
375 L{end +1}=[ QQ ,COM ,SumTot ,SumIn , Average , Function ]; %I add the new
         point to the filter
376 end
377 end
378
379 %cut the filter
380 function [L] = cut_filter (L,h,k,lambda)381
382 % calculate the function for every element of the list
383 if length (L) >h
```

```
384 for l=1:length (L)385 % function
386 Function \{1\} = L\{1\}\{6\};387 end
388
389 % remuve the situations in the filter with the lower values
390 for t=1: (\text{length} (L) - h)391 [mn,idx]=min ([Function {:}]);
392 v=find ([Function {: }] == mn); %if more element with same average values,
         delate the last one ( because with h=1 I have same result as
         cluster_jl_average_variance )
393 idx t = v(end) ;
394 Function { idx t } = Inf;
395 L{idx_t}=[];
396 end
397
398 empties = find ( cellfun ( @isempty ,L) ) ;
399 L(empties) = [];
400 end
401 end
402
403 % Compute modulartiy
404 function MOD = compute_modularity (C, Mat)
405
406 S = size (Mat);
407 \quad N = S(1);
408
409 m = sum (sum(Mat))/2; %total weight
410
411 MOD = 0;
412 COMu = unique (C) ; % list of communities without repetiotions and in
       sorted order
413 %for each community calculate modularity and then sum all together
414 for j=1: length(COMu)415 Cj = (C == COMu(j)); %list of nodes in Cj
416 Ec = sum (sum (Mat (Cj, Cj))); %sum of weights between nodes in Cj
417 Et = sum (sum (Mat (Cj,:))); % sum of weights of nodes incident in nodes
            of Cj
418 if Et >0
419 MOD = MOD + Ec/(2*m)-(Et/(2*m))<sup>-2</sup>;
420 end
421 end
422
423 end
424
425 % Re-index community partition by size
426 function [C Ss] = reindex_com ( COMold )
427 C = sparse (1, length (CDMold) );
428 COMu = unique (COMold);
```

```
429 S = sparse(1, length(COMu));
430 for l=1: length ( COMu )
S(1) = \text{length}(COMold(COMold == COMu(1)));
432 end
433 [Ss INDs] = sort (S, 'descend');
434 for l=1: length ( COMu )
435 C(COMold == COMu(INDs(1))) = 1;436 end
437 end
438 %Re-index community partition not by size but by initial order
439 % function [C] = reindex_com (COMold)
440 %C = sparse (1 , length ( COMold )); % vector with a index for each node
441 % COMu = unique ( COMold ); % unique (v)= same data of v but not repetitions
       and in sorted order
442 % COMolds = sort ( COMold );
443 % COMu = COMolds ([ true ; diff ( COMolds (:)) >0]);
444 %for l=1: length ( COMu )
445 % C(COMold = COMu(1)) = 1; %Riname the communities
446 %end
447 %end
```
Appendix F

Matlab code of the multi-variance-plus method (Section 3.2)

```
1 %multi - variance - plus
2\frac{9}{6}3 % Inputs :
4 % M : vector of weight matrix of each layer
5 % h : length of the filter
6 % lambda : weight of variance in function for cut filter
7 % z : 1 = Recursive computation
8 % : 0 = Just one level computation
9 % Output :
10 % L the filter with the following information
11 % for each element l:
12 % 1{1}=Q cell with modularity of each layer
13 % 1{2}={COMcur COMfull COMindex} communities in the current graph, in
      the
14 % original graph, communities in the current graph reindexed
15 % l {3}= SumTot (sum of weights of the links incident to nodes in a
      community )
16 % l {4}= SumIn (sum of weights of the links inside a community )
17 \frac{\%}{\%} 1{5}=Average (average)
18 % 1{6}= Function (funtion)
19 %
20
21
22 function [L ending] = multi_variance_plus (M, lambda, h, z)
23
24 if nargin < 1
25 error ('not enough argument');
26 end
```

```
27 if nargin < 2
28 error ('not lambda defined');
29 end
30 if nargin < 3
31 error ('not h defined');
32 end
33 if nargin < 4
34 z = 1;
35 end
36
37 S = size(M):
38 \text{ N} = S(1):
39 if length (S) = 3k = S(3);
41 else k = 1;
42 end
43
44 ending = 0;
45
46 for s=1:k47 M2 (:,:,s) = M (:,:,s) - diag (diag (M (:,:,s)));
48 end
49
50 % total weight of the graph
51 for s=1: k
52 if z==153 M(:, :, s) = M(:, :, s) + diag(diag(M(:, :, s)));
54 end
55 m{s} = sum(sum(M(:,: , s)))/2;56 end
57
58 Niter = 0; % number of iterations
59
60 % Calculation of the beginning values
61 COM \{1\} = 1: S(1); % current community %At the beginning each node is a
      community
62 COM\{2\} = 1:S(1); % original graph commuity
63 COM{3} = 1: S(1); % community reindexed
64 %COM(i)= Community of node i
65
66 for s=1: k
67 K(:,:,s) = sum(M(:,:,s)); % K(i)= sum of wieght incident to node i
68 SumTot(:,:,s) = sum(M(:,:,s)); %SumTot(i)= sum of weight incident to
           nodes in community i
69 SumIn(:,:,s) = diag(M(:,:,s))'; %SumIn(i)= sum of weight inside
          community i (loops)
70 %At the beginning each node is a
                                          community
71 Q{s} = compute_modularity (COM{1}, M(:,:,s));
```

```
72 end
73 % average
74 Average = sum([Q{\f : }]) / k;
75 % variance
76 if k == 177 Variance = 0;
78 else
79 Variance = (sum(([Q{\:}:)] - Average) . (2))) / (k-1);
80 end
81 % function
82 Function = (1-lambda) * Average + lambda * Variance;
83
84 l<sup>6</sup>\frac{1}{2}filter
85 L = \{ [\{ Q \}, \{ COM \}, \text{SumTot}, \text{SumIn}, \text{Average}, \text{Function} ] \};86 %If no edges in any layer or just one node
87 if (sum(cellfun((@x)x)>0,m)) == 0) | N == 1 % | logical or88 ending = 1;
89 else
90
91 % Delate layers with m=0
92 M(:,:,(cellfun(\mathcal{O}(x)x==0,m)))=[];
93 k = k - sum(cellfun((@x)x == 0, m));94
95 % Neighbor {j}{s} neighbor of node j in layer s
96 for j=1: N
97 for s=1: k
98 temp=M2(:,:,s);
99 \text{Neighbour}_j\{s\} = \text{find}(\text{temp}(j,:));
100 end
101 Neighbor {j}=Neighbor_j;
102 end
103
104
105 GAIN = 1;
106 while ( GAIN == 1) % Stop when putting nodes in other communities do not
       increase the modularity
107 % while Niter <=2
108 L_old=L;
109 L_new=L;
110 for i=1:N %for each node, in this order
111 L_0 = L;112 for l=1: length (L_o) %for each situation in the filter
113 % Melate the point from the filter
114 L_new_0 = L_new;115 index = cellfun (@(x) isequal (x,L_o{l}) , L_new , ' UniformOutput '
                  , 1) ;
116 L_new(intdev) = [];
117 % step1
118 [L_new, U] = step_1 \left( \{L_0\{1\}\{1\}\}, \{L_0\{1\}\{2\}\}, L_0\{1\}\{3\}, L_0\{1\}
```

```
{}_{1}^{1}{4}, L_o{1}{5}, L_o{1}{6}, k, Neighbor, K, M, N, m, L_new, h, lambda,
              i) ;
119 119 %if no new point, insert again the initial point in the filter
120 if \, U == 0\verb|L_new=L_new_0|;122 end
123 end
124 L_new = cut_filter (L_new, h, k, lambda);
125 L=L_new;
126 end
127 % Check if nothing happened
128 if length (L_old) \tilde{} = length (L_new)
129 GAIN = 1;
130 else
131 C_01d = [];
132 C_new = [];
133 for l=1:length (L_old)
134 R_old = [];
135 R_new = [];
136 for s=1:k
137 R_old = [R_old L_old(1){1}S];138 R_new = [R_new L_new{1}{1}{s};
139 end
C_old = [C_old ; R_old];
141 C_new = [C_new; R_new];142 end
143 C = \text{ismembertol}(C_old, C_new, 10^(-6));
144 if size(C, 1) * size(C, 2) == sum(sum(C))145 GAIN=0;
146 end
147 end
148 Niter = Niter + 1;
149 tEnd = toc;
150 if tEnd > 500
151 L = { };
152 return
153 end
154 end
155 end
156
157 % Perform part 2
158 if (z == 1)159 L = cut_fitter(L,1,k); % one final element160 Mnew=M; % new weight matrix of this iteration
161 Mold=Mnew; %old weight matrix of iteration befor
162 COMcur = L {1}{2}{3}; % communities in the graph of this iteration
163 COMfull = L{1}{2}{3}; % communities in the original graph
164 LL=L;
165 j=2;% number of pass (step1+step2)
```

```
166 S=1;
167 while S
168 L_old=LL;
169 Mold = Mnew;
170 S2 = size (Mold); % number of nodes
171 Nnode = S2(1);
172
173 COMu = unique (COMcur); %list of communities without repetitions
            and in sorted order
174 Ncom = length (COMu); % number of communities
175 ind_com = sparse (Ncom, Nnode); % zero matrix with a row for each
            community and a column for each node in this configuration
176 ind_com_full = sparse (Ncom ,N) ; % zero matrix with a row for each
            community and a column for each node of the original graph
177
178 for p=1: Ncom % for each community
179 ind = find (COMcur == p);
180 ind_com (p ,1: length (ind ) ) = ind ; %in row p insert the indeces
                of the nodes in community p in this configuration
181 end
182 for p=1:Ncom183 ind = find (COMfull == p);
184 ind_com_full (p ,1: length (ind ) ) = ind ; %in row p insert the
               indeces of the nodes in community p in the original graph
185 end
186 Mnew = [];
187 for s=1:k
188 Mnew (:,:,s) = sparse (Ncom, Ncom); %new matrix (each node is
               a community )
189 for m=1: Ncom
190 for n=m : Ncom191 ind1 = ind com (m , :) ;
192 ind2 = ind_com(n, :);
193 % weights of edges between communities
194 Mnew (m,n,s) = sum ( Mold (ind1 (ind1 >0), ind2 (ind2 >0)
                      ,s) ) ) ;
195 Mnew (n,m,s) = sum (sum (Mold (ind1 (ind1>0), ind2 (ind2>0)
                      ,s) ) ) ;
196 end
197 end
198 end
199 % apply first step to this new matrix but z=0 without recursive
200 [LLL e] = multi_variance_plus (Mnew, lambda, h, 0);
201 if isempty (LLL )
202 L=LLL;
203 return
204 else
205 LL = cut_fitter(LLL,1,k); %one final element
206 COMfull = sparse (1, N); \% communities of the original graph
```

```
207 COMcur = LL{1}{2}{3}; % communities
208 for p=1: Ncom
209 ind1 = ind_com_full (p ,:) ; % nodes in community p in the
                  original graph
210 COMfull ( ind1 (ind1 >0) ) = COMcur (p) ; % community now of
                  node i
211 end
212 [COMfull] = reindex_com (COMfull);
213 LL \{1\}{2}{2}= COMfull;
214 % communities do not change
215 if i \equiv 2i \pmod{L_0} if i \equiv 1 \{2\} \{2\}, \text{LL} \{1\} \{2\} \{2\})
216 % return not just this value , but all the value of the
                  last list
217 % calculate COMfull for each element of the last list
218 for l=1: length (LLL)
219 COMfull = sparse (1 ,N) ; % communities of the original
                      graph
220 COMcur = LLL{1}{2}{3}; % communities
221 for p=1:Ncom222 ind1 = ind_com_full (p, :); % nodes in community p
                         in the original graph
223 COMfull (ind1 (ind1 > 0) = COMcur (p) ; % community
                         now of node i
224 end
225 [COMfull] = reindex_com (COMfull);
226 LLL\{1\}\{2\}\{2\}= COMfull;
227 end
228 L = LLL;
229 S=0;
230 end
231 j = j + 1; % start another pass
232 tEnd = toc;
233 if tEnd > 600
234 L = {};
235 return
236 end
237 end
238 end
239 end
240 end
241
242 % Compute step_1
243 function [L,U] = step_1 (Q,COM ,SumTot ,SumIn , Average , Function ,k, Neighbor ,K
     ,M,N,m,L,h,lambda ,i)
244
245 U=0; %if U=1 add at least an element to the list, otherwise insert again
      the initial point in the filter
246 COMcur = COM {1}{1}; % current community
247 Ci = COMcur(i); % community of node i
```

```
248
249 NB=unique (cat (2, Neighbor\{i\}\{:\}));
250
251 SumTot2=SumTot;
252 SumIn2=SumIn;
253
254 % remove i from its community
255 for s=1: k
256 COMcur2 = COMcur;
257 COMcur2(i) = -1;
258 CNi = ( COMcur2 == Ci) ; % list of nodes in Ci community , without i
259 Ki in i {s} = sum (M(i, CNi, s)) ; %sum of weights between i and Ci
260 GQ_i {S} = (K(:,i,s) * SumTot2(:,Ci,s)) / (2 * (m{s}^c(2))) - Ki_in_i {s} / m{s}\} - ((K(:,i,s)) \hat{(2)}) / (2 * (m[s] \hat{(2)}));261 % Recalculate values
262 SumTot2 (: ,Ci ,s) = SumTot2 (: ,Ci ,s) - K(: ,i,s) ; % weights incident to
          Ci community
263 SumIn2(:,Ci,s) = SumIn2(:,Ci,s) - 2* sum(M(i,CNi,s)) - M(i,i,s); \frac{\%}{\%}weights community i
264 end
265
266 G = sparse(1, N);267 for j=1: length(NB)268 SumTot3=SumTot2;
269 SumIn3=SumIn2;
270 Cj = COMcur(NB(j)); % community of node j
271 if (G(C_j) == 0) %If have not tried with another node of community Cj
           yet
272 G (C<sub>j</sub>) =1;
273 for s=1:k
274 COMcur3 = COMcur;
275 COMcur3(i) = -1;
276 %I put i in the community of j for each layer
277 CNj = ( COMcur3 == Cj) ; % nodes in community Cj , without j
278 % faster then CNj = find ( COMcur == Cj)
279 Ki_in_j{s} = sum(M(i,CNj,s)); %sum of weights between i and
                  C_1280 GQ_j{s} = Ki_in_j{s}/m{s} - (K(:,i,s)*SumTot3(:,Cj,s))/(2*(m
                  {s }^(2) ) ) ; % gain deltaQ if I put isolated node i in Cj
281 % Recalculate
282 SumTot3(:,Cj, s) = SumTot3(:,Cj, s) + K(:,i, s);283 SumIn3(:, Cj, s) = SumIn3(:, Cj, s) + 2*sum(M(i, CNj, s)) + M(i, i,s) ;
284 end
285 COMcur4=COMcur;
286 COMcur4 (i) = Cj;
287 COM \{1\} {1} = COMcur4 ;
288 [COMcur4] = reindex_com (COMcur4);
289 COM {1} {3} = COMcur4 ;
```

```
291 % variance
292 if k == 1293 GV_j = 0;294 else
295 for s=1:k
296 DQ{s} = GQ_ii{s} + GQ_j{s}; % gains
297 end
298 M_DQ = sum (DQ {\cdot}; ) \wedge k; % % gain average
299 GV_j = (( \text{sum} ((DQ {\:}:\) - M_DQ) .^{})(2) ) ) / (k-1) + (2/(k-1)) * \text{sum} (k-1)sum ((Q{1}{f}:)-Average) * ([DQ{f}:}] - M_DQ)); %gain variance
300 end
301 GF_j = (1 - \lambda m b da) * M_D Q + \lambda m b da * G V_i;302 if GF_i > -10^(-14)303 \quad F_j = \text{Function} + \text{GF}_j;304 % modularity
305 for s=1:k
306 Q_j{s} = Q{1}{s} + DQ{s};307 end
308 % average
309 A_{-}j = Average + M_DQ;
310 [L] = add_to_filter ({Q_i} , COM, SumTot3, SumIn3, A<sub>-</sub>j, F<sub>-</sub>j, L, k);
311 U=1;
312 end
313 end
314 end
315 end
316
317 %add a point to the filter
318 function [L] = add_to_filter (QQ ,COM ,SumTot ,SumIn , Average , Function ,L,k)
319
320 if length (L) z = 0321
322 COMcur = COM {1}{1}; % current community
323
324 %1. Check if the new point is dominated by a point in the filter
325 DD=0;
326 \quad 1=1;327 while (DD==0) & l <= length (L)
328 D=0;
329 S=1;330 while (D == 0) & s <=k_{331} if QQ{1}{s} > L{1}{1}{s}332 D=1;
333 end
334 S=S+1;
335 end
336 if D %the new point is not dominated by l
337 DD = 1;
```

```
338 end
339 1=1+1;340 end
341 if DD % the new point is not dominated
342 DDD=0;
343 1=1;344 while (DDD == 0) & l <= length (L)
345 D=1;
346 s=1;
347 while D & s \le k348 if abs (L{1}{1}{s} - QQ{1}{s} > 10^(-14)
349 D=0;
350 end
351 s = s + 1;
352 end
353 if D
354 DDD = 1;
355 end
356 1 = 1 + 1;357 end
358
359 if DDD ==0
360
361 %2. Check if the other points in the filter are dominated by the new
       point %
362
363 for l=1: length (L)
364 D=1;
365 S=1;
366 while D & s <=k
367 if L{1}{1}{s} > QQ{1}{s} + 10^(-14)368 D = 0;
369 end
370 S=S+1;371 end
372 if D %l is dominated by the new point so I remove it from the
           filter
373 L{1}=[];
374 end
375 end
376
377 empties = (cellfun (@isempty,L));
378 L(empties) = [];
379 L{end+1}=[QQ, COM, SumTot, SumIn, Average, Function]; %I add the new point
       to the filter
380
381 end
382 end
383 else
```

```
384 L{end +1}=[ QQ ,COM ,SumTot ,SumIn , Average , Function ]; %I add the new
           point to the filter
385 end
386 end
387
388 %cut the filter
389 function [L] = cut_filter (L,h,k, lambda )
390
391 % calculate the function for every element of the list
392 if length (L) >h
393 for l=1: length (L)
394 % function
395 Function \{1\} = L\{1\}\{6\};396 end
397
398 % remuve the situations in the filter with the lower values
399 for t=1: (\text{length} (L) - h)_{400} [mn, idx] = min ([Function {: }]);
401 v=find ([Function \{\cdot\}] ==mn); %if more element with same average values,
         delate the last one ( because with h=1 I have same result as
         cluster_jl_average_variance )
_{402} idx_t=v(end);
403 Function {idx_t}=Inf;
404 L\{idx_t\} = [];
405 end
406
407 empties = find (cellfun (@isempty, L));
408 L( empties ) = [];
409 end
410 end
411
412 % Compute modulartiy
413 function MOD = compute_modularity (C, Mat)414
415 S = size (Mat);
416 N = S(1);
417
418 m = sum (sum(Mat))/2; %total weight
419
420 MOD = 0;
421 COMu = unique (C); %list of communities without repetiotions and in
       sorted order
422 %for each community I calculate modularity and then sum all together
423 for j=1: length(COMu)424 Cj = (C == COMu(j)); %list of nodes in Cj
425 Ec = sum (sum (Mat (Cj, Cj))); %sum of weights between nodes in Cj
426 Et = sum (sum (Mat (Cj ,:) ) ) ; %sum of weights of nodes incident in nodes
            of Cj
427 if Et >0
```

```
428 MOD = MOD + Ec/(2*m)-(Et/(2*m))<sup>2</sup>;
429 end
430 end
431
432 end
433
434 % Re-index community partition by size
435 function [C Ss] = reindex_com ( COMold )
436 C = sparse(1, length(COMold));437 COMu = unique (COMold);
438 S = sparse (1, length(COMu));
439 for l=1: length (COMu)
440 S(1) = length (COMold (COMold == COMu(1)));
441 end
_{442} [Ss INDs] = sort (S, 'descend');
443 for l=1: length ( COMu )
444 C(COMold == COMu (INDs (1))) = 1;
445 end
446 end
447 %Re-index community partition not by size but by initial order
448 % function [C] = reindex_com ( COMold )
449 %C = sparse (1 , length ( COMold )); % vector with a index for each node
450 % COMu = unique ( COMold ); % unique (v)= same data of v but not repetitions
       and in sorted order
451 % COMolds = sort ( COMold );
452 % COMu = COMolds ([ true ; diff ( COMolds (:)) >0]);
453 \quad \% for l=1:l ength (COMu)
454 % C(COMold = COMu(1)) = 1; %Riname the communities
455 %end
456 %end
```
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