Project Report: Física Teórica I

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Identification and classification of information-processing building blocks on genetic regulatory network

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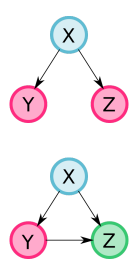
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Abstract

Fibration building blocks of a information flow network represent the sets of nodes that are symmetric with respect to the processing of information, that is, the sets of nodes that process equivalent information. Here, we have reproduced important results concerning the identification and classification of the fibration building blocks of directed networks, constructed from real network data. More specifically, using the transcriptional regulatory network data of the Escherichia Coli bacteria, we quantify the clusters of nodes that synchronously process equivalent information and then we classify these clusters, called network fibers, based on its specific topological features. This way, in order to consistently present the obtained results, in this report we first give a brief description of the theory concerning the graph fibration morphism and its main definitions related to information flow symmetries. Next, we detail the methods adopted to correctly identify and classify the network fibers. More specifically, to establish an optimal framework, I show the implementation details of the Minimal Balanced Coloring algorithm used to find the corresponding fibers in the network, presenting a slightly improvement for the algorithm complexity and its implementation. At last, showing the proper methods for data preparation, I describe the results obtained concerning the fiber statistics for the specific case of the Escherichia Coli regulatory network, to properly compare with the recent results presented at Morone et. al. (2019).

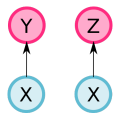
1 Brief Introduction

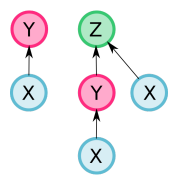
This statement requires citation [1]; this one does too [2] [3] [4] [5]. Lorem ipsum dolor sit amet, consectetur adipiscing elit. Aenean dictum lacus sem, ut varius ante dignissim ac. Sed a mi quis lectus feugiat aliquam. Nunc sed vulputate velit. Sed commodo metus vel felis semper, quis rutrum odio vulputate. Donec a elit porttitor, facilisis nisl sit amet, dignissim arcu. Vivamus accumsan pellentesque nulla at euismod. Duis porta rutrum sem, eu facilisis mi varius sed. Suspendisse potenti. Mauris rhoncus neque nisi, ut laoreet augue pretium luctus. Vestibulum sit amet luctus sem, luctus ultrices leo. Aenean vitae sem leo.



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2 Coarsest Refinement Partitioning Algorithm

In order to identify the correct distribution of network fibers we have to find a efficient procedure to split the network in different disjoint partitions, in which all nodes from a same partition should receive information from the same other partitions. To do that, we treat our problem as the same of find the coarsest relational refinement partitioning for the given network. In this section, we detail the optimal algorithm used to find this coarsest partitioning in a context of graph fibrations and show the relative simple implementation of this method.

2.1 Algorithm description

The algorithm used in this project is a slightly modified version of the algorithm presented by Paige and Tarjan [4], having a time complexity of $O(M \log N)$, being very efficient for sparse networks. This algorithm has the same runtime order than the algorithm from Cardon and Crushemore [2], but it has a simpler implementation and smaller prefactors, exhibiting a better approach to our problem. Even though the algorithm applies for general situations, here we give the details of the algorithm to its application for a network, so we introduce the necessary definitions within this context.

A network G is completely defined as G(V,E) by the sets of nodes V and of the connections E. The network has then N=|V| nodes that can have connections with one another defined by the set E containing M=|E| ordered pair of nodes, denoting the directed connections between the network nodes. Over the network, we can define a graph partition P of V as a set of pairwise disjoints subsets of V whose union is all V

$$P = \bigcup_{i} P_{i} \tag{1}$$

where P_i are the elements of the partition P, in which we call it blocks. Within this context, if we consider an additional graph partition Q that has the property that all the block of Q is contained in a block of P, we say that Q is a refinement

partition of P.

Considering a block B in P, we say that the block B is stable with respect with a set S if or all elements of B connects an element of S or none element of B points to any element of S. By that we define that a graph partition P of V is stable if it is stable with respect with its own blocks. Having defined that, the graph coarsest partition problem is that of finding, for a given set of connections E and initial partition P over V, the coarsest stable refinement of P.

Considering all that, for the proper identification of the fibers on the network we have to construct a stable graph partition that is equivalent to the coarsest (minimal number of blocks) refinement of the network with respect to the information flow passing through each node. For that, we need to extend the concept of stability over the partition for one that accounts for the information received by each block of the partition. Thus, in order to identify the clusters with isomorphic input-tree for all their nodes, we require that the partition should be not only stable but input-tree stable. That means that for a subset $S \subseteq V$, a graph partition P over the network G(V, E) is input-tree stable with respect to S if for all the block $B \in P$, the equality

$$|E^{-1}(\{x\}) \cap S| = |E^{-1}(\{y\}) \cap S| \tag{2}$$

is satisfied for all the elements $x, y \in B$.

Find a set S in which the current partition P is input-tree unstable. Replace P by the output of I-split(S,P). Guarantees that the set S or unions of used sets never be used again.

Since the finest partitioning possible is the one in which every node is itself

a block, the refinement step can be proceeded at most N-1 times, guaranting that the algorithm terminates with the correct answer, since stability is inherited by the refinement process. However, to guarantees that the algorithm has a optimal runtime we have to find a efficient way to select the appropriate sets to the refinement step, without choosing repeated sets. Fortanately, this can be easily done for the construction of a input-tree stable partition.

Given a set S of nodes from the network G and a given input-tree unstable graph partition Q, the blocks $B \in Q$ that are input-tree unstable with respect to S can be splitted in several blocks B_j to have a stable input-tree for S. Then, each splitted block will have the property defined by

$$B_j = \{ x \in B : |E^{-1}(\{x\}) \cap S| = j \}$$
(3)

where the number of splitted block must be larger than one to a proper split process take place. Then, for each unstable block $B \in Q$ all the splitted block, except the largest one, can be put a queue to be used ahead in the algorithm as a refinement set. This ensures that none repeated sets or union of repeated sets can be used during the algorithm execution.

Having said all that, the complete algorithm to find the correct network fibers of a network G(V,E) consists in initializing a graph partition Q over all nodes from V except the nodes that do not receive information from any other node, in which each one of these will be defined as isolated fibers already in the beginning of the algorithm. The partition Q is defined as one block containing all the other nodes in the network. The algorithm maintains a queue L of possible refinement sets, initially containing the single block of Q and all the isolated blocks defined at the beginning. Then, we proceed as

Remove from L its first set S. Replace Q by the I-split(S, Q). Whenever a block $B \in Q$ splits into two or more nonempty blocks, add all but the largest to the back of L.

And this process is repeated until the queue L is empty. At this point, the resulted partition Q represent the coarsest input-tree stable partitioning of the network G(V,E), where each block represents a network fiber with all its nodes having isomorphic input-trees.

2.2 Data preparation and algorithm implementation

We apply the above algorithm in the genetic regulatory network of the *Escherichia Coli* bacteria. We obtain the genetic E. Coli network through its transcriptional

regulatory interactions data, where each gene is regulated by a transcription factor protein. Since a transcription factor production in the cell is regulated by a gene, we can define a directed connection between two genes if a gene regulates the production of a transcriptional factor, which it regulates another gene. Since a transcription factor can be either an activator(positive) or repressor(negative), or even behaves as both(dual), the links between genes can carry different types of messages. Because of that, it is important that the partitioning algorithm accounts the type of message to construct appropriate input-trees for the network fibers partition. Therefore, for a proper application of the algorithm on the *Escherichia Coli* genetic network, we label each node uniquely, either as number or string names, and also each link with the type of connection between genes.

Considering this context, we explain now how the algorithm should proper be implemented to have an optimal perfomance concerning its runtime complexity. After discussing the data structures necessary to deal correctly with data, we design an algorithm recipe to show all the main steps of the process explained above. Also, anyone can acess our own personal implementation for genetic regulatory networks in the link https://github.com/higorsmonteiro/fiberblocks on github platform.

Given a directed network G(V,E) to be partitioned in fibers, it is very important for the correctness of the algorithm that the nodes that do not receive any information, that is, the nodes that do not have any inward connections, be preprocessing as isolated blocks. This means that the initial graph partition P is divided in two different partition P' e P'', the first one contained all the single-node blocks containing the solitaire nodes v in which $|E^{-1}(\{v\})| = 0$, and the second one containing initialing a single block containing all the other nodes w in which $|E^{-1}(\{w\})| \ge 1$. The importance of this preprocessing is to guarantee that solitaire nodes do not be put on the same fibers during the refinement steps. Even though the blocks of P' are used as refinement sets, P' is not used by the algorithm. Thus, the final partition is the union of P' and the result of the refinement partitioning of P''.

After the above preprocessing we define the data structures for the partition Q for its blocks B. A partition is a doubly linked list of blocks, which allows that deletion of blocks be made in constante time O(1) as long we have the block memory address during the procedure. A block is just a structure containing indexing data along with a doubly linked list containg all the nodes that belongs to it. Together with a queue of blocks L, these constructions are the main data structures necessary for an efficient implementation of the refinement partitioning algorithm. At the beginning of the algorithm, we enqueue all the blocks of P'

e P'' to L. Then, we start the algorithm initializing a partition Q = P'' and by removing the top element set S of L, then we apply the I-split(S, Q) to identify the input-tree unstable blocks of Q and split them into input-tree stable blocks with respect to S. This way, all the splitted blocks are push to the end of L, with exception the largest resulted blocks for each splitted block. As we have mentioned, the algorithm terminates when there is no more sets in L. Even though this is the whole algorithm, the implementation of the I-split function might not be so straightforward, since a given block B can be splitted into an arbitrary number of blocks. In respect to that below we propose the following implementation construction to the splitting function:

```
Algorithm 1: I-split (S, \overline{Q}, L)
   Input : A set S and a partition \bar{Q}
   Output: Input tree stable partition \bar{Q} with respect to S
1 Initialize U as an empty partition;
2 for \forall B \in \bar{Q} do
        if \exists \{w_i, w_j\} \subseteq B : |E^{-1}(\{w_i\}) \cap S| \neq |E^{-1}(\{w_j\}) \cap S| then
            push B to \bar{U};
            Initialize \bar{T} as an empty partition;
            for \forall w_i \in B do
                 if \exists X \in \bar{T} : |E^{-1}(\{w_i\}) \cap S| = X(E) then
                      insert w_i in X;
 8
                 else
                      create a new block X;
10
                      insert w_i in X;
                      X(E) \leftarrow |E^{-1}(\{w_i\}) \cap S|;
                      push block X to \bar{T}
13
                 end
14
            end
15
        end
16
        enqueue all blocks X \in \overline{T} in L, except the largest one;
17
        push all blocks X \in \bar{T} to \bar{Q};
19 end
20 delete all blocks B \in \bar{U} in \bar{Q}
```

In the algorithm recipe above, our *I-split* function receives a set S and the partition \bar{Q} as input and returns a input-tree stable \bar{Q} with respect to S as output. Besides its list of nodes, a block X has another attribute accessed as X(E), which

receives, during the splitting process, the number of inward links coming from the current refinement set S. This way, a node w belonging to the block to be splitted can be put in the block $X \in \bar{T}$ that has the same attribute value, like stated in the conditional expression at line 7 of the algorithm above.

Finally, considering all the discussion above, we can explicit the complete algorithm in just a few steps.

Algorithm 2: Coarsest Refinement Graph Partitioning

```
Input: A network G(V, E)

1 Initialize S as an empty set;

2 Initialize L as an empty queue;

3 Initialize \bar{P}, \bar{P}', \bar{P}'', \bar{Q} as empty partitions;

4 B' = \{\{w\} \in V : |E^{-1}(\{w\})| = 0\};

5 B'' = \{v \in V : |E^{-1}(\{v\})| \geq 1\};

6 push all B' to \bar{P}';

7 push B'' to \bar{P}'';

8 enqueue B'' \in \bar{P}'' to L;

9 enqueue all blocks B' \in \bar{P}' to L;

10 \bar{Q} \leftarrow \bar{P}'';

11 while |L| \neq 0 do

12 |S \leftarrow dequeue(L);

13 |\bar{Q} \leftarrow I\text{-input}(S, \bar{Q}, L)|

14 end
```

A concrete implementation of the algorithm in a programming language is presented at the link https://github.com/higorsmonteiro/fiberblocks together with *E. Coli* prepared data and another examples. Further information about the application of the codes in other genetic regulatory network data can be found at the same link given.

Results

Using the algorithm above, we applied it to a set of networks examples before its proper application on the *Escherechia Coli* genetic regulatory network data. The reason for that is merely to show that the not just the algorithmic approach chosen is consistent but the written code per si is correctly implementated. For each network, we have a adjacency list containing all the directed connections between the nodes and for each connection we have a string defining the type of regulation

of a specific link, where here, considering the genetic regulation context, the types can be positive, negative or dual. The first three examples are all small networks, containing at most 48 nodes, and so all can be used to test the correctness of the algorithm applied. For that three examples we show the network drawing and their fibers distribution identified as the result of the algorithm.

The first example is showed at the figure 1. This network is disconnected, containing two main weakly connected components, and contains only positive regulation between the interactions between the genes, showing the simpler case where the connections between nodes in a network are all of the same type. In this case, we find five non-trivial fibers, that is, fibers with size larger than one.

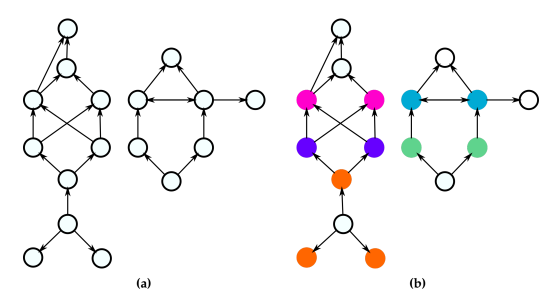
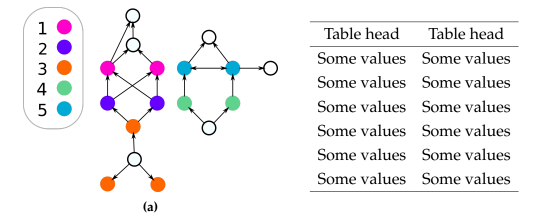


Figure 1

At the end of the algorithm, all the fibers identified are input-tree stable with respect to all fibers in the network, including the fiber itself. Then, labelling each fiber by an index or color we can then construct the fibration building blocks through the calculation of the fundamental class number n and through the subclass number l. Again, a fibration building block is defined as the nodes inside the fiber as well as the l external nodes that directly regulates at least one node of the current fiber. By that, we calculate the fundamental class number n_k of the fiber k by constructing the adjacency matrix A_{ij}^k of its fibration building block and then calculating the largest eigenvalue of it. The value of n quantifies the information loops of the fibration building blocks and can be either a integer or fractal golden ration.

Labelling the resulted fiber distribution, we obtain the following fibration classification for each fibration buuilding block of the network containing more

than one node:



The second example exhibits a connected component containing N=21 nodes, where all the three type of regulation are present. For this network we find the total of eleven fibers, where just six are fibers containing more than one node in it. As we can notice, for two arbitrary nodes be in the same fiber they must not just receive the same number of inward connections but the same number of each type of inward connections, this way receive the equivalent information from the rest of the network. We can see this for the example of the fibers 4 e 5, where even though both receives two links from the same central node, the fibers receives different types of information from that node, meaning that they are not the equivalent fibers.

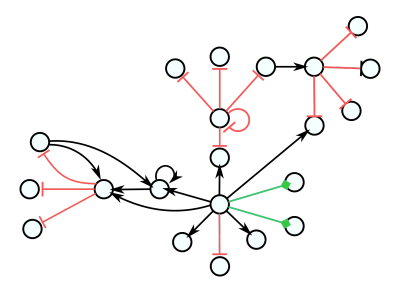
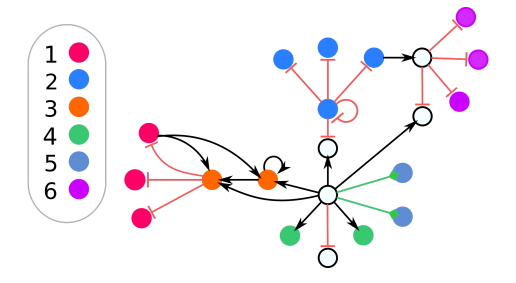


Table head	Table head
Some values	Some values



If we consider the fiber 3, besides the inner autoregulation loop we have a loop information going outside the fiber and going back through an external regulator inside the fiber 1. In cases like this the branching ratio of the fiber has a fractal value represented by a golden ratio ϕ_d resulted a loop information represented by a fibonacci sequence. This way, fiber 3 can be classified by the values $|\phi_d, l=2\rangle$ where the fiber has two external regulator nodes. The fibration classification for the non-trivial fibers is showed in the table ??.

Before applying the algorithm for the whole network, we chose, for the finality of visualization and testing, to apply first the algorithm for a weakly connected component of the *E. Coli* network. The same network is showed at the supplementary information from [1]. The fiber distribution found is consistent with the one presented at [1], presenting the total of fifteen non-trivial fiber groups as shown in figure 3.

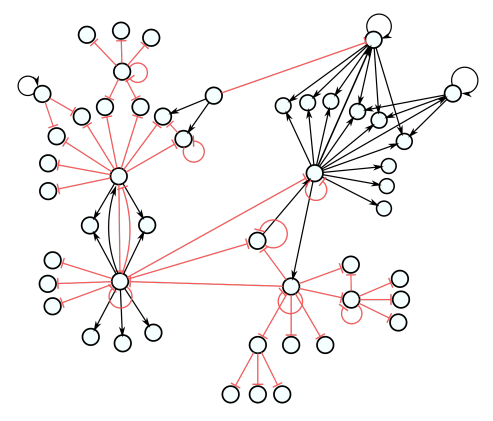
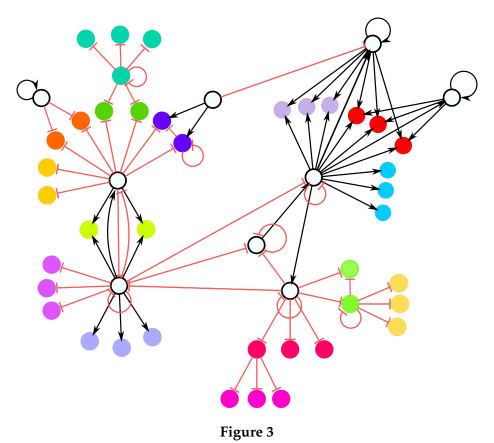


Figure 2



The connected component showed at the figure 2 is a representation of a component in the whole regulatory network of the *Escherichia Coli* bacteria and the resulted fibers shows just a portion of the whole fibers pattern. In the table X we give the classification for each fibration building block obtained for the whole bacteria network, including of course the fibers presented in the figure 3. In total, we have found 84 non-trivial fiber groups containing a total of 462 nodes, obtaining a characteristic fiber statistics for the *Escherichia Coli* network data. To guarantee the correctness of the fiber distribution found we calculate after the application of the refinement partitioning algorithm the information received for each node inside a fiber and compares if each one process equivalent information.

The statistics concerning the number of each fibration block on the network is given in the table X, showing describe the pattern of the statistics of the table that concerns the bacteria fiber statistics.

3 Conclusion

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