

Reaction Motifs in Metabolic Networks

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Abstract. The classic view of metabolism as a collection of metabolic pathways is being questioned with the currently available possibility of studying whole networks. Novel ways of decomposing the network into modules and motifs that could be considered as the building blocks of a network are being suggested. In this work, we introduce a new definition of motif in the context of metabolic networks. Unlike in previous works on (other) biochemical networks, this definition is not based only on topological features. We propose instead to use an alternative definition based on the functional nature of the components that form the motif. After introducing a formal framework motivated by biological considerations, we present complexity results on the problem of searching for all occurrences of a reaction motif in a network, and introduce an algorithm that is fast in practice in most situations. We then show an initial application to the study of pathway evolution.

1 Introduction

Network biology is a general term for an emerging field that concerns the study of interactions between biological elements [2]. The term *molecular interaction networks* may designate several types of networks depending on the kind of molecules involved. Classically, one distinguishes between gene regulatory networks, signal transduction networks and metabolic networks. Protein-protein interaction networks represent yet another type of network, but this term is rather linked to the techniques (such as Yeast-2-hybrid) used to produce the data and covers possibly several biological processes (including, for example, the formation of complexes and phosphorylation cascades) [16].

One of the declared objectives of network biology (or systems biology in general) is whole cell simulation [9]. However, dynamic simulation requires knowledge on reaction mechanisms such as the kinetic parameters describing a Michaelis-Menten equation. Besides the fact that such knowledge is often unavailable or unreliable, the study of the static set of reactions that constitute metabolism is equally important, both as a first step towards introducing dynamics, and in itself. Indeed, such static set represents not what is happening at a given time in a given cell but instead the capabilities of the cell, including capabilities the cell does not use. A careful analysis of this set of reactions for a given organism, alone or in comparison with the set of other organisms, may also help to arrive at a better understanding on how metabolism evolves. It is this

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set we propose to study in this paper. More precisely, in the following sections, the term “metabolism” should be understood as the static set of reactions involved in the synthesis and degradation of small molecules. Regulation information is not taken into consideration for now. It may be added in a later step, as the “software” running on the “hardware” of a metabolic network [15].

A major issue concerning the study of biochemical networks is the problem of their organisation. Several attempts have been made to decompose complex networks into parts. These “parts” have been called modules or motifs, but no definition of such terms seems to be completely satisfying.

Modules have first been mentioned by Hartwell *et al.* [6] who outline the general features a module should have but provide no clear definition for it. In the context of metabolic networks, a natural definition of modules could be based on the partition of a metabolic network into the metabolic pathways one can find in databases: modules would thus be the pathways as those have been established. The advantage of this partition, and thus of modules representing pathways, is that it reflects the way metabolism has been discovered experimentally (starting from key metabolites and studying the ability of an organism to synthesize or degrade them). The drawback is that it is not based on objective criteria and therefore is not universal (indeed, the number of metabolic pathways and the frontiers between them vary from one database to the other).

Several attempts to give systematic and practical definitions have been made using graph formalisms [14,10,5] and constraint-based approaches [11]. Graph-based methods range from a simple study of the local connectivity of metabolites in the network [14] to the maximisation of a criterion expressing modularity (number of links within modules) [5]. The only information used in these methods is the topology of the network. In the case of constraint-based approaches, the idea is quite different. First, a decomposition of the network into functional sets of reactions is performed (by analysis of the stoichiometric matrix [12]) and then modules are defined from the analysis of these functional states. The result is not a partition in the sense that all reactions might not be covered and a single reaction might belong to several modules.

Unlike the definition of module, the notion of motif has not been studied in the context of metabolic networks. In general, depending on what definition is adopted for modules and motifs, there is no clear limit between the two notions besides the difference in size. In the context of regulatory networks, motifs have been defined as small, repeated and perhaps evolutionary conserved subnetworks. In contrast with modules, motifs do not function in isolation. Furthermore, they may be nested and overlapping [22]. This definition refers to general features that regulatory motifs are believed to share but it provides no practical way to find them. A more practical definition has been proposed, still in the context of gene regulatory networks (and other types of non-biological networks such as the web or social networks). These are “network motifs” and represent patterns of interconnections that recur in many different parts of a network at frequencies much higher than those found in randomized networks [17]. This definition is purely topological and disregards the nature of the components in a motif. It assumes that the local topology of the network is sufficient to model function (which is understood here as the dynamic behaviour of the motif). This assumption seems ac-

ceptable when studying the topology of the internet and may also hold when analysing gene regulatory networks, but it appears not adapted to metabolic networks. In a static context, a topological definition of motif seems indeed inappropriate as similar topologies can give rise to very different functions.

In the definition of motif we introduce, the components of the network play the central part and the topology can be added as a further constraint only. This is the main biological contribution of this paper.

Its main algorithmical contribution comes from the fact that the definition of motif we adopt leads to new questions. Indeed, if searching for “purely” topological motifs may be formally modelled as a subgraph isomorphism problem, this no longer applies when searching for motifs where the features describing the components are the important elements and topology is initially indifferent (connectivity only is taken into account). Observe that the problem we address is different from pathway alignment because we wish to go beyond the notion of pathway in order to study the network as a whole. Moreover, in [19] and [13], the pathways are modelled as, respectively, chains and trees to simplify the problem. This simplification may seem reasonable in the case of a pathway alignment, it is no longer so in the case of general networks.

The paper addresses complexity issues related to this new definition of a graph motif, providing hardness results on the problem, and then presents an exact algorithm that is fast in practice for searching for such motifs in networks representing the whole metabolism of an organism. The paper ends with an initial application of the algorithm to the formulation of hypotheses on the evolution of pathways.

2 Preliminaries

2.1 Data

The metabolic network analysed in this work was obtained from the PATHWAY database from KEGG [8]. Data describing reactions, compounds and enzymes were downloaded and stored locally using a relational database management system (postgreSQL). The KEGG database contains metabolic data concerning 209 sequenced organisms. The network we built from such data is therefore a consensus of our current knowledge on the metabolisms of all those organisms. As a consequence, sequences of reactions present in the network may have been observed in no organism. To avoid this configuration, one can “filter” the consensus network by an organism of interest, keeping only in the dataset reactions catalysed by enzymes the organism is considered to be able to synthesize. We adopt a different strategy by choosing to perform our motif search on the consensus network and to possibly filter the results in a second step, allowing for easier comparative analysis between organisms.

Moreover, we use an additional information present in KEGG: the notion of primary/secondary metabolites. Indeed, in the KEGG reference pathway diagrams (maps), only primary metabolites are represented and connect reactions together, whereas secondary metabolites are not drawn (even though they participate in the reaction). A typical example of a secondary metabolite is the ATP molecule in an ATP-consuming reaction. (Observe that, unlike the notion of ubiquitous compound [14], the notion of primary/secondary metabolite is relative to a reaction.) Keeping all metabolites in the

network leads to the creation of artefactual links between reactions and the bias introduced can lead to inaccurate results such as considering metabolic networks as small-world networks as shown in [3]. Withdrawing secondary metabolites may not be the best strategy to adopt, but it represents a simple way of avoiding this bias.

2.2 Graph Models

Several formal models have been in use to study metabolic networks. The choice of a formal model seems to depend mainly on the nature of the hypotheses one wishes to test (qualitative or quantitative, static or dynamic) and on the size of the network under study. Differential equations seem well adapted to study the dynamic aspects of very small networks whereas graphs enable the static study of very large networks.

Between these two ends of the spectrum, semi-quantitative models have been proposed. For example, Petri nets allow for the simulation and dynamical analysis of small networks [21], while constraint-based models provide a mathematical framework enabling to decompose the network into functional states starting only from information on stoichiometry and making the assumption that the network is at steady-state [12].

As our goal is to deal with large networks and work with the least possible *a priori*, graph models seem appropriate. In previous genome-scale studies [7], graphs have been used mainly for topological analyses regardless of the nature of their components (reactions, compounds and enzymes). We propose to enrich the graph models and take into consideration some of the features of such components.

Formally, a graph G is defined as a pair (V, E) , with V a set of *vertices* and $E \subseteq V \times V$ a set of *edges*. The edges represent the relations between the vertices and may be directed or undirected. The vertices and edges of the graph can be labelled.

The most intuitive graph representation of a metabolic network is provided by a bipartite graph. A bipartite graph has two types of vertices which in the context of metabolic networks represent, respectively, reactions and chemical compounds. The compound graph is a compact version of the bipartite graph where only compound vertices are kept and information on the reactions is stored as edge labels. The reaction graph is the symmetric representation of a compound graph (*i.e.*, reaction vertices are kept and information on the compounds is stored as edge labels). Directed versions of these graphs can be drawn expressing the irreversibility of some reactions. The information concerning the reversibility of reactions is generally not well-known. Indeed, contradictions may be found within a same database. We therefore consider this information as uncertain and, in an initial step, assume that all reactions are reversible. This apparently strong hypothesis seems preferable than considering a reaction as irreversible when it actually is reversible (leading to a loss of information).

In the following sections, we denote by C a finite set of labels, which we refer as *colours*, that correspond to reaction labels. Also, we assume the graph $G = (V, E)$ is undirected and that we are given, for each vertex, a set of colours from C . Reversibility and edge labels will not be used. If needed, one can use them in a later step.

2.3 Motif Definition

We define a motif using the nature of the components it contains.

Definition 1. A motif is a multiset of elements from the set C of colours.

As mentioned earlier, we choose in this definition not to introduce any constraint on the order of the reactions nor on topology. This choice is motivated by the wish to explore the network with the least possible *a priori* information on what we are searching for. Topology and order of the reactions can be used later as further constraints. The advantage of this strategy is that the impact of each additional constraint can then be measured.

2.4 Occurrence Definition

Intuitively, an occurrence is a connected set of vertices labelled by the colours of the motif. For a precise definition, let R be a set of vertices of G and let M be a motif of the same size as R . Let $H(R, M)$ denote the bipartite graph whose set of vertices is $R \cup M$ and where there is an edge between a vertex v of R and a vertex c of M if and only if v has c as one of its colours.

Definition 2. Definition of an exact occurrence of a motif

An exact occurrence of a motif M is a set R of vertices of G such that $H(R, M)$ has a perfect matching and R induces a connected subgraph of G .

If one is strict on the relation of similarity between colours (colours are considered the same only if they are identical), the risk is to find a single occurrence, or none, of any given motif in the network [3]. Moreover, since studying the evolution of what the graph G represents is one of our main objectives, it seems relevant to allow for flexibility in the search for occurrences of a motif.

With this in mind, we introduce a function S (detailed later) that assigns, to each pair c_i, c_j in $C \times C$, a score which measures the similarity between c_i and c_j . Two colours are considered similar if this score is superior to a threshold s . We then adapt our definition of exact occurrence by modifying $H(R, M)$ in the following way. There will be an edge between a vertex v in R and a colour c in M if and only if there exists a colour c' of v such that the value of $S(c', c) \geq s$. Further, we generalise this to the case where the threshold s is different for every element c in M . The latter is motivated by the idea that some elements in the motif we are searching for may be more crucial than others. Observe that these considerations are independent of the definition of S that is discussed in the next section.

Another type of flexibility can then be added, that allows for gaps in the occurrences. By this we mean, roughly, allowing the occurrence to have more vertices just to achieve the connectivity requirement. These extra vertices are not matched to the elements of the motif. Two types of control on the number of gaps are considered: local and global. Intuitively, a local gap control policy bounds the maximum number of consecutive gaps allowed between a pair of matched vertices of R . A global control policy bounds the total number of gaps in an occurrence.

This leads to the following definition of an approximate occurrence of a motif, where we denote by G_R the subgraph of G induced by a set R of vertices of G .

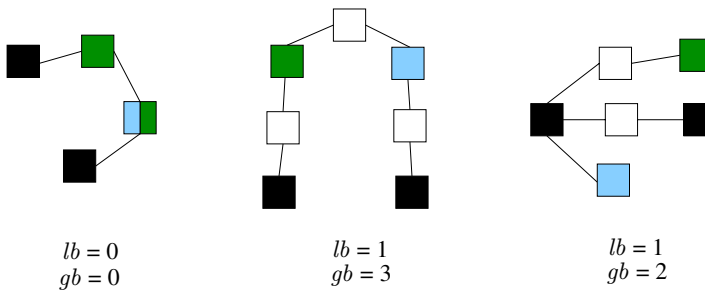


Fig. 1. Subgraphs induced by occurrences for the motif {black, black, dark grey, light grey}.

Definition 3. Definition of an approximate occurrence of a motif

Let lb and gb be the local and global gap control bounds and let M be a motif. For each c in M , let s_c be a number. An approximate occurrence of M (with respect to lb , gb and the thresholds s_c) is any minimal set R of vertices of G that has a subset R' that satisfies the following conditions:

1. the bipartite graph $H(M \cup R', E_H)$ with $E_H = \{\{c, v\} \in M \times R' \mid \text{there exists a colour } c' \text{ of } v \text{ such that } S(c', c) \geq s_c\}$ contains a perfect matching;
2. for each subset B of R' such that $B \neq \emptyset$ and $R' \setminus B \neq \emptyset$, the length of a shortest path in G_R between B and $R' \setminus B$ is at most lb ;
3. $|R| - |R'| \leq gb$.

The minimality requirement on the set R avoids uninteresting approximate occurrences that are simple copies of other occurrences with extra vertices connected to them.

Observe that when no gaps are allowed then $R = R'$ and condition 2 means simply that G_R is connected. An example is given in Figure 1.

2.5 Reaction Similarity

We now discuss function S for the problem of metabolic networks and reaction motifs in such networks. Various functions of different nature may be used. We present here two possible ways to define S .

The first one is based on alignment. Indeed, in order to compare reactions, which is what function S is used for, one can compare the enzymes that catalyse these reactions by performing an alignment of their sequences (or structures). An element of C would then be a protein sequence (or structure). The function S assigns a sequence (or structure) alignment score and s is a user-defined threshold that has to be met to consider the sequences (structures) similar. In the case of whole networks, sequences are preferable since many structures are not known.

The second example is the one we adopt in this paper. It is based on a hierarchical classification of enzymes developed by the International Union of Biochemistry and Molecular Biology (IUBMB) [1]. It consists in assigning to each enzyme a code with 4 numbers expressing the chemistry of the reaction it catalyses. This code is known as the

enzyme's EC number (for Enzyme Commission Number). The first number of the EC number can take values in $[1 \dots 6]$, each number symbolizing the 6 broad classes of enzymatic activity. (1. Oxidoreductase, 2. Transferase, 3. Hydrolase, 4. Lyase, 5. Isomerase, 6. Ligase.) Then each of the three remaining numbers of the EC number provides additional levels of detail. For example, the EC number 1.1.1.1 refers to an oxidoreductase (1) with CH-OH as donor group and NAD⁺ as acceptor group.

An element of C is in this case an EC number. The function S then assigns a similarity score between two EC numbers that corresponds to the index of the deepest level down to which they remain identical. For example, $S(1.1.1.2, 1.1.1.3) = 3$. Two EC numbers are considered similar if their similarity score is above a user-defined cut-off value s in $[0 \dots 4]$. The advantage of this definition of similarity between colours, *i.e.*, reactions, is that it is more directly linked to the notion of function. Reactions compared with this measure are likely to be functionally related (and possibly evolutionarily related also).

3 Algorithmics

3.1 Hardness Results

The formal problem we address is the following:

Search Problem. Given a motif M and a labelled undirected graph G , find all occurrences of M in G .

As mentioned earlier, this problem is different from subgraph isomorphism because the topology is not specified for the motif.

For this problem, we may assume the graph is connected and all vertices have colours that appear in the motif. Otherwise, we preprocess the graph throwing away all the vertices having no colour appearing in the motif and solve the problem in each component of the resulting graph.

A natural variant of the Search Problem consists in, given a motif and a labelled graph, deciding whether the motif occurs in the graph or not. As before, we may assume the graph is connected, all vertices are labelled with colours and all colours appear in the motif. It is easy to see this decision version of the Search Problem is in NP. We show next that it is NP-complete even if G is a tree, which implies that the Search Problem is NP-complete for trees. For the following proof, we consider the version where no gaps are allowed.

NP-Complete for Trees. We have the following proposition.

Proposition 1. *The Search Problem is NP-complete even if G is a tree.*

Proof. We present a reduction from EXACT COVER BY 3-SETS (X3C):

INSTANCE: Set X with $|X| = 3q$ and a collection \mathcal{C} of 3-element subsets of X .

QUESTION: Does \mathcal{C} contain an exact cover for X , *i.e.*, a subcollection $\mathcal{C}' \subseteq \mathcal{C}$ such that every element of X occurs in exactly one member of \mathcal{C}' ?

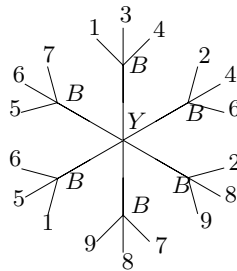


Fig. 2. Tree T and its labels for $X = \{1, \dots, 9\}$ and $\mathcal{C} = \{\{1, 3, 4\}, \{2, 4, 6\}, \{2, 8, 9\}, \{7, 8, 9\}, \{1, 5, 6\}, \{5, 6, 7\}\}$. The motif M in this case is $\{Y, B, B, B, 1, \dots, 9\}$.

Let $X = \{1, \dots, 3q\}$ and $\mathcal{C} = \{C_1, \dots, C_n\}$ be an instance of X3C. The instance for the decision version of the Search Problem consists of a motif $M = \{Y, B, \dots, B, 1, \dots, 3q\}$, where B appears q times in M , and a tree T as follows. (See Figure 2 for an example.) There are four vertices in T for each i , $1 \leq i \leq n$, three of them are leaves in T , each one labelled by one of the elements of C_i . The fourth vertex, named r_i , is adjacent to the three leaves and has colour B . Besides these $4n$ vertices, there is only one more vertex in T , which is labelled Y and is adjacent to each r_i . This completes the description of the instance. Clearly it has size polynomial in the size of X and \mathcal{C} .

To complete the reduction, we need to argue that the motif M occurs in T if and only if there is a subcollection \mathcal{C}' of \mathcal{C} such that each element of X occurs exactly in one member of \mathcal{C}' .

Suppose there is such a \mathcal{C}' . Clearly $|\mathcal{C}'| = q$. Let R be the set of vertices of T consisting of the vertex labelled Y and the four vertices of each C in \mathcal{C}' . The subgraph of T induced by R is connected. Also, in R , there is a vertex labelled Y , q vertices labelled B (one for each C in \mathcal{C}') and one labelled by each element in X (because of the property of \mathcal{C}'). That is, R is an occurrence of M in T .

Now, suppose there is an occurrence of M in T , that is, there is a set R of $1 + 4q$ vertices of T that induces a connected subgraph of T and has a vertex labelled by each of the colours in M . Let \mathcal{C}' consist of the sets C_i in \mathcal{C} whose vertex r_i in T is in R . Let us prove that each element of X appears in exactly one of the sets in \mathcal{C}' . First, note that the vertex labelled Y is necessarily in R , because it is the only one labelled Y and there is a Y in M . Then, as R induces a connected graph, a leaf from a set C_i is in R if and only if r_i is also in R . But R must contain exactly q vertices labelled B . Consequently, $|\mathcal{C}'| = q$ and, as R must contain $1 + 4q$ vertices, all three leaves of each C in \mathcal{C}' must be in R , and these are all vertices in R . As R must contain a vertex labelled after each element in X , there must be exactly one set in \mathcal{C}' containing each element in X . \square

Fixed Parameter Tractability. This problem is fixed-parameter tractable with parameter k . Indeed, a naive fixed-parameter algorithm consists in generating all possible topologies for the input motif M , and then searching for each topology by using a subtree isomorphism algorithm. Since it is enough to generate all possible tree topologies for M , the number of topologies to consider depends (exponentially) on k only, and subtree isomorphism is polynomial in the size of both the motif M and the tree T where

M is sought. This reasoning is not valid anymore when the motif must be searched in a general graph G as subgraph isomorphism is NP-complete even when the motif is a tree [4].

General Complexity Results. Table 1 summarizes the complexity of the Search Problem for various types of motifs and graphs. As mentioned, it is enough to consider that our motifs are trees (or paths). This is because topology is indifferent (only connectivity matters).

By *fixed* in the Table, we mean that the colours of the vertices in a path (respectively tree) are fixed, otherwise (*i.e.* path/tree *not fixed*) we mean that we are searching for a path (respectively tree) with the given vertex colours but do not care in what order they appear, provided they all appear.

Motifs that are paths are already hard problems for general graphs G . This can be shown by a reduction from the Hamiltonian path problem.

Table 1. Complexity results for the motif Search Problem

| MOTIF \ TYPE OF GRAPH | | path | tree | graph |
|-----------------------|-----------|------------|-------------------------|-------------|
| path | fixed | polynomial | polynomial | NP-complete |
| | not fixed | polynomial | polynomial | NP-complete |
| tree | fixed | — | polynomial | NP-complete |
| | not fixed | — | NP-complete, FPT in k | NP-complete |

Since the instances we have to consider in the case of metabolic networks are relatively small (3184 vertices and 35284 edges for the network built from the KEGG Pathway database), it is possible to solve the problem exactly, provided some efficient pruning is applied. This is described in the next section.

3.2 Exact Algorithm

Version with no Gaps. We now present an exact algorithm which solves the Search Problem. We first explain it for the simple case where the gap parameters lb and gb are set to 0 and then we show how it can be extended to the general case.

Let M be the motif we want to search for. A very naive algorithm would consist in systematically testing all sets R of k vertices as candidates for being an occurrence, where $k = |M|$. For R to be considered an occurrence of M , the subgraph induced by R must be connected and there must be a perfect matching in the bipartite graph $H(R, M)$ that has an edge between $r \in R$ and $c \in M$ if and only if c is similar to one of the colours at vertex r . The search space of all combinations of k vertices among the n vertices in G is huge. We therefore show two major pruning ideas arising from the two conditions that R has to fulfill to be validated as an occurrence of M .

The connectivity condition can be checked by using a standard method for graph traversal, such as breadth first search (BFS). In our case, a BFS mixed with a backtracking strategy is performed starting from each vertex in the graph. At each step of the search, a subset of the vertices in the BFS queue is marked as part of the candidate

set R . The queue, at each step, contains only marked vertices and neighbours in G of marked vertices. Also, there is a pointer p to the last marked vertex in the queue. At each step, there are two cases to be analysed: either there are k vertices marked or not. If there are k vertices marked, we have a candidate set R at hand. We submit R to the test of the colouring condition, described below, and we backtrack to find the next candidate set. If there are less than k vertices marked, then there are two possible cases to be analysed: either p is pointing to the last vertex in the queue or not. If p is not pointing to the last vertex in the queue, we move p one position ahead in the queue, mark the next vertex and queue its neighbours that are not in the queue already (checking the latter can be done in constant time by adding a flag to each vertex in the original graph). Then we repeat, that is, start a new step. If, on the other hand, p is pointing to the last vertex in the queue, then we backtrack. The backtracking consists of unmarking the vertex pointed to by p , unqueueing its neighbours that were added when it was marked, moving p to the previous marked vertex in the queue and starting a new step. (If no such vertex exists, the search is finished.) Next we describe the test of the colouring condition.

Given a candidate set R , one can verify the colouring condition by building the graph H and checking whether it has a perfect matching or not. In fact, we can apply a variation of this checking to a partial set R , that is, we can, while constructing a candidate set R , be checking whether the corresponding graph H has or not a complete matching. The latter is a matching that completely covers the partial candidate set R . If there is no such matching, we can move the search ahead to the next candidate set. This verification can be done in constant time using additional data structures that are a constant time the size of the motif.

Extra optimisations can also be added. For instance, instead of using every vertex as a seed for the BFS, we can use only a subset of the vertices: those coloured by one of the colours from the motif, preferably the less frequent in the graph.

Allowing for Gaps. Allowing for local but not global gaps (*i.e.*, setting $lb > 0$ and $gb = \infty$) can easily be done by performing the lb -transitive closure of the initial graph G and applying the same algorithm as before to the graph with augmented edge set. The p -transitive closure of a graph G for p a positive integer is the graph obtained from G by adding an edge between any two vertices u and v such that the length l of a shortest path from u to v in the original graph satisfies $1 < l \leq p$. The p -transitive closure can be done at the beginning of the algorithm or on the fly. In the latter case, when a next vertex is added to the queue, instead of queueing its neighbours only, all vertices at distance at most p from it are queued (if they are not already in the queue) where by distance between any two vertices we mean the number of vertices other than these two in a shortest path between them.

Allowing for global gaps as well as local ones is more tricky. The reason is that an unmarked vertex can be put in the queue because of many different marked vertices. When backtracking in the queue at any step in the algorithm, unmarked vertices that have been queued only because of the marked vertex v that is being dequeued can be safely eliminated from the queue. Unmarked vertices $\{v_i\}$ that were queued because of the vertex being dequeued *and* of at least one other marked vertex will remain (somewhere) in the queue. Therefore, in order to correctly account for the global number of

gaps introduced so far in the current occurrence, one must consider all the remaining marked vertices that implied the queuing of $\{v_i\}$. Extra information must be kept to locate in constant time the unmarked vertices $\{v_i\}$ and to update the global count of gaps. This information can be kept in a balanced tree of size proportional to $k = |M|$ associated with each queued unmarked vertex u' . Each node in the tree corresponds to a marked vertex u that could have led to the queuing of u' and is labelled by the distance from u' to u (this distance is at most lb). Keeping, updating and using the extra information adds a multiplicative term in $O(k \log k)$ to the time complexity of the algorithm, which seems reasonable.

On average, searching for all occurrences of a motif of size 4 with no gaps and threshold $s = 3$ takes 8 microseconds of CPU time on a Pentium 4 (CPU 1.70 GHz) with 512 Mb of memory.

4 Application

The approach we propose, and have described in the previous sections, should enable both to generate some hypotheses on the evolution of metabolic pathways, and to analyse global features of the whole network.

We start by presenting a case study motivated by trying to understand how metabolic pathways evolve. We do not directly answer this question, which is complex and would be out of the scope of this paper. Instead, we give a first example of the type of evolutionary question people have been asking already and have addressed in different, often semi-manual ways in the past [20], and that the algorithm we propose in this paper might help treat in a more systematic fashion.

As in [20], one is often interested in a specific pathway, and, for instance, in finding whether this pathway can be considered similar to other pathways in the whole metabolic network thus suggesting a common evolutionary history. The metabolic pathway we chose as example is valine biosynthesis. Focusing on the last five steps of the pathway, we derived a motif $M = \{1.1.1.86, 1.1.1.86, 4.2.1.9, 2.6.1.42, 6.1.1.9\}$ and performed the search for this motif using initially a cut-off value s of 4 for the similarity score between two EC numbers (that is, between two reaction labels). With this cut-off value, the motif was found to occur only once. (see Figure 3).

From this strictly defined motif, we then relaxed constraints by first lowering the cut-off value s from 4 to 3 and then setting the gap parameters to 1 (motif denoted by M'). Additional occurrences were found. Three of them particularly drew our attention (see Figure 3).

The first one corresponds to the five last steps of the isoleucine biosynthesis. The second one corresponds to the five last steps of the leucine biosynthesis. Together, they suggest a common evolutionary history for the biosynthesis pathways of valine, leucine and isoleucine.

An interesting point concerning the second occurrence is the fact that the order of the reactions is not the same as in the other pathways. This occurrence would not have been found if we had used a definition of motif where the order was specified.

Finally, the third occurrence that drew our attention was formed by reactions from both the biosynthesis of valine and a distinct metabolic pathway, namely the biosynthesis of Panthotenate and CoA. This latter case illustrates a limit of our current general

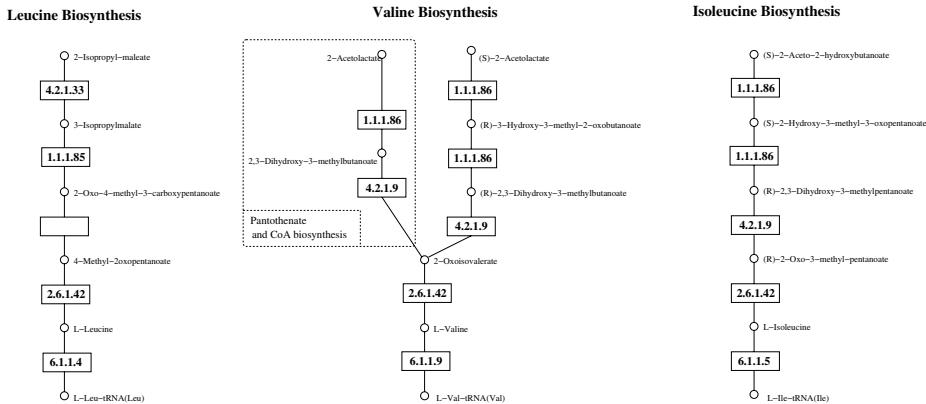


Fig. 3. Bipartite representation of the results obtained when searching for the following motif : $M' = \{1.1.1, 4.2.1, 2.6.1.42, 6.1.1\}$ with local and global gap bounds set to 1. The empty box in the leucine biosynthesis represents a spontaneous reaction.

way of thinking about metabolism: frontiers between metabolic pathways as defined in databases are not tight. If we had taken such frontiers into account, we would not have found this occurrence that overlaps two different pathways. Yet such occurrence can be given a biological meaning: it can be seen as a putative alternative path for the biosynthesis of valine.

To complement this analysis, one should add that the results presented in this section hold for 125 organisms in KEGG among which *S. cerevisiae* and *E. coli*.

Intrigued by the potential importance of inter-pathway occurrences, we computed their proportion in the general case of a randomly chosen motif. By systematically testing all motifs of size 3 and 4 (with cut-off values set to 3), we found that, on average, a motif of size 3 (respectively 4) has 74% (respectively 92%) of its occurrences that are inter-pathway occurrences. All inter-pathway occurrences may not represent biologically meaningful chemical paths but the proportions above suggest that a lot of information may be lost when studying pathways and not networks.

5 Conclusion

In this paper, we presented a novel definition of motif, called a “reaction motif”, in the context of metabolic networks. Unlike previous works, the definition of motif is focused on reaction labels while the topology is not specified. Such novel definition raises original algorithmic issues of which we discuss the complexity in the case of the problem of searching for such motifs in a network. To demonstrate the utility of our definition, we show an example of application to the comparative analysis of different amino-acid biosynthesis pathways. This work represents a first step in the process of exploring the building blocks of metabolic networks. It seems promising in the sense that, with a simple definition of motif, biologically meaningful results are found.

We are currently working on an enriched definition of motif that will take into account information on input and output compounds. The current definition already en-

ables to discover regularities in the network. Enriched definitions should enable to test more precise hypotheses.

In this paper, we used a particular formalism for analysing a metabolic network through the identification of motifs. Other formalisms have been employed or could be considered. As J. Stelling indicated in his review of 2004 [18], each formalism gives a different perspective and confronting them seems to be a promising way of getting at a deeper understanding of such complex networks.

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