# RISOTTO: Fast Extraction of Motifs with Mismatches

Nadia Pisanti<sup>1,\*</sup>, Alexandra M. Carvalho<sup>2,\*\*</sup>, Laurent Marsan<sup>3</sup>, and Marie-France Sagot<sup>4</sup>

<sup>1</sup> Dipartimento di Informatica, University of Pisa, Italy
 <sup>2</sup> INESC-ID, Lisbon
 <sup>3</sup> PRiSM, Université de Versailles St Quentin, France
 <sup>4</sup> INRIA Rhône-Alpes, France

**Abstract.** We present in this paper an exact algorithm for motif extraction. Efficiency is achieved by means of an improvement in the algorithm and data structures that applies to the whole class of motif inference algorithms based on suffix trees. An average case complexity analysis shows a gain over the best known exact algorithm for motif extraction. A full implementation was developed and made available online. Experimental results show that the proposed algorithm is more than two times faster than the best known exact algorithm for motif extraction.

## 1 Introduction

Patterns appearing repeated either inside a same string or over a set of strings are important objects to identify. Such repeated patterns are called *motifs* and their identification is called *motif inference* or *motif extraction*. The area has many potential applications, namely to data compression, natural languages, databases, basically, any activity or research requiring text mining [4]. The field of application that concerns us is molecular biology. The motifs in this case may correspond to functional elements in DNA, RNA or protein molecules, or to whole genes whose sequences are strongly similar. In biological applications, it is mandatory to allow for some mismatches between different occurrences of the same motif. In fact point mutations might have taken place, as well as errors in the sequencing procedure, so that molecules that have the same or related function(s), have no longer identical sequences. This is what makes the problem difficult from the computational point of view. In this paper we propose an exact algorithm for the extraction of motifs with mismatches. In particular, we consider *single* and *structured motifs*, which are motifs composed of several disjoint single motifs placed at given distances from each other. The extraction of structured motifs appears particularly interesting because of its application to the detection of binding sites ([3]). Given a text s, the problem is to find repeated

<sup>\*</sup> Partially supported by PRIN project ALGONEXT.

<sup>\*\*</sup> Partially supported by FCT grant SFRH/BD/18660/2004 and FCT Project FEDER POSI/SRI/47778/2002 BioGrid, and by the FCT grant SFRH/BD/18660/2004.

patterns in s according to some parameters that specify the frequency and the structure required for the motifs. In molecular biology, the text is in general a set of DNA sequence. Several exact, heuristic, and probabilistic algorithms for extracting structured motifs exist. Up to date, the best known exact algorithms for the extraction of single [7] and structured [2] motifs perform well when searching for short motifs. In this paper, we propose an improvement to such algorithms in order to deal with long motifs. The problem of extracting long motifs was first adressed by Pevzner and Sze [6]. They considered a precise version of the motif discovery problem: find all single motifs of length 15 with at most 4 mismatches in 20 sequences of size 600. A general set for this problem deserves attention from the algorithmic point of view because its computational complexity is in the worst case exponential with respect to the number e of mismatches allowed among different occurrences of the same motif. The reason is that, to identify motifs of the required length, there can be an explosion of the number of candidates of intermediate length whose extension has to be attempted. This imposes in practice a limit to the length of the motifs themselves, as in many applications the value of e depends on this length. The improvement introduced in this paper acts exactly in these cases, and hence applies to relatively long motifs, being a way to increase the length of motifs that are detectable in practice.

## 2 Single Motif Extraction

A single motif is a word over an alphabet  $\Sigma$ . Given an error rate e, a motif is said to *e-occur* in a sequence if it occurs with at most *e* letters substitution. The single motif extraction problem takes as input N sequences, a quorum  $q \leq N$ , a maximal number e of mismatches allowed, and a minimal and maximal length for the motifs,  $k_{min}$  and  $k_{max}$ , respectively. The problem consists in determining all motifs that e-occur in at least q input sequences. Such motifs are called *valid*. An efficient exact algorithm for the extraction of single motifs with mismatches has been introduced in [7] and is based on a suffix tree: motifs are considered in lexicographical order starting from the empty word, and they are extended to the right as long as the quorum is satisfied until either a valid motif of maximal length is found (if the  $k_{max}$  length is reached), or the quorum is no longer satisfied. In both cases, a new motif is attempted. Formally, the algorithm ([7])we refer to is sketched in Algorithm 1.. At the beginning ExtractSingleMotif is called on the empty word. The algorithm recursively calls itself for longer motifs built by adding letters (step 4), and considers new ones (step 1) when the extension fails (step 2). A valid motif is spelled out whenever a motif whose

#### Algorithm 1. Single motif extraction

```
ExtractSingleMotif(motif m)
```

```
1. for all \alpha \in \Sigma do
```

2. if  $m\alpha$  is valid then

- 3. **if**  $|m\alpha| \ge k_{min}$  **then** spell out the valid motif  $m\alpha$
- 4. if  $|m\alpha| < k_{max}$  then ExtractSingleMotif $(m\alpha)$

length lies within the required minimal and maximal length is detected (step 3). The order in which motifs are generated corresponds to a depth-first visit of a complete trie  $\mathcal{M}$  (the motif tree) of all words of length  $k_{max}$  on the alphabet  $\Sigma$ . The algorithm does not need to allocate the motif tree. The only memory requirement is for the suffix tree  $\mathcal{T}$ . Assuming that the required length of the motif is k (that is  $k_{min} = k_{max} = k$ ), and that at most e mismatches are allowed, the algorithm has worst case time complexity in  $O(Nn_k\nu(e, k))$ , where  $n_k$  is the number of tree nodes at depth k, and  $\nu(e, k)$  is the number of words that differ in at most e letters from a word m of length k. This value does not depend on m, and it holds that  $\nu(e, k) \leq k^e |\Sigma|^e$ . This upper bound is in practice not tight. Nevertheless, no better bound can be given and therefore the time complexity is linear in the input size, but possibly exponential in the number e of mismatches. Since reasonable values for e are proportional to the value of k, this actually places a practical bound on the length required for the motifs. The goal of this paper is to move this bound.

#### 2.1 Using Maximal Extensibility of Factors

The modification we suggest consists in storing information concerning maximal extensibility in order to avoid trying to extend hopeless motifs. For instance (see Fig. 1), assume that in our virtual depth-first visit of the motif tree, we have found out that motif m can be further extended without losing the quorum up to a length of MaxExt(m) only, the latter representing its maximal extensibility. If later on, we are processing a motif m' that has m as a suffix, then the MaxExt(m) information could be useful, as it applies to m' as well because m' can also be extended with at most MaxExt(m) symbols (and possibly less). In particular, we have that if  $|m'| + MaxExt(m) < k_{min}$ , then we can avoid any further attempt to extend m' as there is no hope to reach length  $k_{min}$  for motifs that have m' as prefix. In Algorithm 1., motifs are considered in lexicographical order by a depth-first (virtual) visit of the motif tree  $\mathcal{M}$ . Every time we stop extending a motif, that is, when we (virtually) backtrack in  $\mathcal{M}$ , it is either because we found a valid motif of the maximal length, or because the quorum is no longer satisfied ( $m\alpha$  does not satisfy the condition at step 2, and we start to consider the next one in lexicographical order). In the first case, m is valid, as are all its prefixes, and  $|m| = k_{max}$ . No information on the maximal extension of m nor of



**Fig. 1.** Example where the extension of m' can be avoided, using MaxExt(m), where m is a suffix of m', because  $|m'| + MaxExt(m) < k_{min}$ 

its prefixes can be of any use because all motifs having a prefix of m as suffix can in general still be extended as much as necessary to reach at least the length  $k_{min}$ . For this reason, we set  $MaxExt(m) = +\infty$ . In the second case, m does not satisfy the quorum while all its prefixes do. For reasons that will be clearer later, we chose to only use the maximal extensibility information of motifs of length up to  $k_{min} - 1$ , hence this case can be subdivided into two subcases. When a motif m cannot be extended anymore and it has not reached the length  $k_{min} - 1$ , we set MaxExt(m) = 0. If the motif has reached a length h between  $k_{min} - 1$ and  $k_{max}$ , we set  $MaxExt(\langle m\alpha|_{k_{min}-1}) = h - (k_{min} - 1)$ , where  $\langle m\alpha|_{k_{min}-1}$  is the prefix of length  $k_{min} - 1$  of  $m\alpha$ . Since it can be that  $MaxExt(\langle m\alpha|_{k_{min}-1})$ had already received some value because a previous extension of  $\langle m\alpha|_{k_{min}-1}$  was interrupted, then we change the value of  $MaxExt(\langle m\alpha|_{k_{min}-1})$  only if we are increasing it, as maximal extensibility of a motif refers to its longest extension. All maximal extensibility values are initially set to -1, hence the first attribution to  $MaxExt(\langle m\alpha|_{k_{min}-1})$  will always increase its value.

In all the cases above, the algorithm does not consider any further extension of m, and backtracks. This backtracking consists in either replacing the last letter  $\sigma_{|m|}$  of m (line 1), or considering a shorter motif which in general shares a prefix with m, if  $\sigma_{|m|}$  was the last letter of the alphabet  $\Sigma$ . In this latter case, the whole subtree rooted at the node spelling  $\sigma_1 \dots \sigma_{|m|-1}$  has been (virtually) completely visited. Thus, we have all the information necessary to set the value of  $MaxExt(\sigma_1 \dots \sigma_{|m|-1})$  according to  $MaxExt(x) = 1 + \max_{\alpha \in \Sigma} MaxExt(x\alpha)$ , for all valid motifs x such that  $|x| < k_{min} - 1$ . If the letter  $\sigma_{|m|-1}$  was the last of the alphabet, then the backtracking goes further. In that case, also the MaxExtinformation concerning the word  $\sigma_1 \ldots \sigma_{|m|-2}$  can be filled in in the same way, and so on. As mentioned before, maximal extensibility information can be used for motifs whose extension is being considered and for which this information could actually prevent some useless attempts. Namely, assume we are trying to extend the motif  $m = \sigma_1, \sigma_2, \ldots, \sigma_{|m|}$ . Obviously, we do not know the value of MaxExt(m) yet, and we know  $MaxExt(\sigma_2,\ldots,\sigma_{|m|})$  only if it lexicographically precedes m, that is, it has already been virtually visited in the motif tree. If this is not the case, we check whether  $MaxExt(\sigma_3,\ldots,\sigma_{|m|})$  is already known, and so on, possibly until the singleton  $\sigma_{|m|}$ . If they are all lexicographically greater than m, then no maximal extension information can be used for m, but if for any of them MaxExt is known and it holds that the maximal possible extension is not enough to reach  $k_{min}$ , then the information is useful as it guarantees that attempting to further extend m is useless.

**Lemma 1.** Let  $w \in \Sigma^*$ . We have  $MaxExt(w) \leq MaxExt(v)$  for each v which is a suffix of w.

A consequence of Lemma 1 is that longer suffixes of m can give us more tight bounds on the maximal extensibility information with respect to shorter ones. Therefore, since we start by checking the longest one, as soon as we find a suffix of m that enables us to state that m is not worth further attempts, then we can stop checking the other (shorter) suffixes. That is, if we find a suffix  $|m\rangle_j = \sigma_j, \ldots, \sigma_{|m|}$  of m, with  $1 < j \leq |m|$ , such that  $MaxExt(|m\rangle_j)$  is not enough for m to reach  $k_{min}$  because  $MaxExt(|m\rangle_j)+|m| < k_{min}$ , then we can quit attempting m and all its extensions, and we can consequently update MaxExt(m). On the other hand, if no suffix  $|m\rangle_j$  of m is such that  $MaxExt(|m\rangle_j)+|m| < k_{min}$ , then the maximal extension does not disallow to reach  $k_{min}$ . In this case, we have to go on trying to extend m even if it might be the case that it will never reach the minimal length. The algorithm for single motif extraction using the maximal extensibility information is presented in Algorithm 2.. For simplicity, we denote in the same way a node x and the word spelled by the path from the root to x. Recall that we use  $\langle m\alpha|_{k_{min}-1}$  to denote the prefix of  $m\alpha$  of length  $k_{min} - 1$ , and  $|x\rangle_{|x|-1}$  to denote the suffix of x of length |x| - 1. Finally, for step 3, recall that we assumed that all maximal extensibility values are initially set to -1.

#### Algorithm 2. Single motif extraction with maximal extensibility information

```
\mathbf{ExtractSingleMotif}(\mathsf{motif}\ m)
```

```
1. for all \alpha \in \Sigma do
 2
      x := m\alpha
       repeat x := |x\rangle_{|x|-1} until (x = root \text{ or } MaxExt(x) \neq -1)
 3.
       if x \neq root and MaxExt(x) + |m\alpha| < k_{min} then
 4.
 5.
          MaxExt(m\alpha) := MaxExt(x)
 6.
          stop spelling m\alpha and continue
 7.
       if m\alpha is valid then
          \mathbf{if} \left| m lpha 
ight| \geq k_{min} \ \mathbf{then} spell out the valid motif
 8
           if |m\alpha| < k_{max} then ExtractSingleMotif(m\alpha)
 9.
          else MaxExt(\langle m\alpha |_{k_{min}-1}) := +\infty
10
11.
        else
          if |m\alpha| < k_{min} then MaxExt(m\alpha) := 0
12.
          else if MaxExt(\langle m\alpha|_{k_{min}-1}) < |m\alpha| - (k_{min}-1) then MaxExt(\langle m\alpha|_{k_{min}-1}) := |m\alpha| - (k_{min}-1)
13
     (k_{min} - 1)
14. if |m| < (k_{min} - 1) then MaxExt(m) := 1 + \max_{\alpha \in \Sigma} MaxExt(m\alpha)
```

#### 2.2 Complexity Analysis

The time complexity of Algorithm 2. remains the same as for Algorithm 1. in the worst case. Nevertheless, the proposed improvement has (very positive) effects on the average case. Next we compute the average ratio between the number of attempted extensions by RISO and RISOTTO for single motif extraction and compute the limit from which RISOTTO performs better than RISO.

Assume that the dataset has r planted random motifs of size t, where each motif can be extracted with at most e mismatches, and that the remaining text is uniformly random. This assumption captures the fact that we want to analyze the ratio between the number of attempted extensions by RISO and RISOTTO in the context of a dataset with highly correlated sequences (meeting the application requirements to biological datasets).

Let  $M_i$  be the random variable that gives the number of extracted motifs of size *i* with at most *e* mismatches for the assumed dataset, where  $0 \le i \le t$ . Clearly, we have that  $P(M_0 = 1) = 1$  and  $P(M_t \ge r) = 1$ . The number of attempted extensions by RISO at level i > 0 (when the recursion step is at level *i*) is given by the random variable  $E_i = M_{i-1}|\Sigma|$ , and the total number of attempted extensions for the extraction of a single motif of size *k* is given by  $R_k = \sum_{i=1}^k E_i$ . On the other hand, RISOTTO will only extend words at level *i* if they fulfill the maximum extensibility requirement. Therefore the number of attempted extensions by RISOTTO at level *i* is given by  $E'_i = M_{i-1}|\Sigma|(1-p(i))$ , where p(i) is the probability of a *i*-word having maximal extensibility information to avoid its extension. Furthermore, the total number of attempted extensions by RISOTTO for the extraction of a single motif of size *k* is  $R'_k = \sum_{i=1}^k E'_i$ .

We conclude that to compute the average value of  $\frac{R'_k}{R_k}$  we need to determine the average of the random variables  $M_i$  and the values p(i), for  $i = 1, \ldots, k$ . We proceed by computing the average values of  $M_i$ . Clearly, a planted motif of size t has t - i + 1 segments of size i (considering overlapping). Observe that the average number of mismatches of the e-occurrences of a motif of size t is

$$\overline{e} = \sum_{j=0}^{e} j \frac{\binom{t}{j} (|\varSigma| - 1)^j}{\nu(e, t)}$$

Hence, if we assume the mismatches to distribute uniformly over the segments, the average number of mismatches of the segments of size i of the e-occurrences is  $\overline{e}_i = \frac{i}{t}\overline{e}$ . Thus, the motifs extracted at level i due to the planted motifs are all the neighbors differing at most  $(e - \overline{e}_i)$  letters from the segments of size i of the planted motifs. Since there are r(t - i + 1) segments of size i, the average number of extracted motifs of size i with at most e mismatches, due to the planted motifs, is

$$\overline{T}_i = |\Sigma|^i \left( \sum_{j=0}^{r(t-i+1)-1} \left( 1 - \frac{\nu(e-\overline{e}_i,i)}{|\Sigma|^i} \right)^j \frac{\nu(e-\overline{e}_i,i)}{|\Sigma|^i} \right).$$

Finally, to determine the average value of  $M_i$ , we need to take into account the motifs extracted from the random part of the text, and so, we have  $\overline{M}_i = \overline{T}_i + (|\Sigma|^i - \overline{T}_i)(1 - \pi_i)$ , where  $\pi_i$  is the probability of a random word of size *i* not being extracted with quorum *q* from a set of *N* sequences. Given that the probability of an *e*-neighbor of a word of size *i* not appearing in a random text of size *n* is

$$\delta(i, e, n) = (1 - 1/|\Sigma|^{i})^{(n-i+1)\nu(e,i)} \approx (1 - 1/|\Sigma|^{i})^{ni^{e}|\Sigma|^{e}},$$

the value of  $\pi_i$  can be computed by the following binomial

$$\pi_i = \sum_{j=0}^{q-1} {\binom{N}{j}} \delta(i, e, n)^{N-j} (1 - \delta(i, e, n))^j.$$

We finalize by computing the probability p(i). Since the probability of a suffix of a random word being lexicographically smaller than the random word is  $\frac{1}{2}$ , we have that

$$p(i) = \sum_{j=1}^{i} \frac{1}{2^j} \gamma_{k-i}$$

where  $\gamma_{k-i}$  is the probability of the suffix of size k-i to have information to avoid the extension. Notice that  $\gamma_{k-i}$  is the probability of the suffix of size k-i not being extended to a size greater than k-1, and is given by

$$\gamma_{k-i} = \pi_{k-i} + (1 - \pi_{k-i})\pi_{k-i+1}^{|\Sigma|} + (1 - \pi_{k-i})(1 - \pi_{k-i+1}^{|\Sigma|})\pi_{k-i+2}^{|\Sigma|^2} + \dots$$
$$= \sum_{j=0}^{i-1} \pi_{k-i+j}^{|\Sigma|^j} \prod_{\ell=1}^j (1 - \pi_{k-i+j-\ell}^{|\Sigma|^{j-\ell}}).$$

To understand when RISOTTO starts to provides a gain over RISO, it is important to look to  $E'_i$  and  $E_i$ . Note that  $E'_i$  will be much smaller than  $E_i$  if p(i) is close to 1. Moreover, as soon as random motifs start to disappear,  $M_{i-1}$  will be larger than  $M_i$ , which happens when  $\pi_i$  is close to 1. Both  $\pi_i$  and p(i) depend tightly of  $\delta(i, e, n)$ , that is, if  $\delta(i, e, n)$  is close to 0, so are  $\pi_i$  and p(i), and if  $\delta(i, e, n)$  is close to 1, so are  $\pi_i$  and p(i). Since  $\delta(i, e, n)$  behaves like a Dirac cumulative function for large values of n, that is, it jumps very fast from 0 to 1, we just need to solve the equation  $\delta(i, e, n) = 1/2$  for the variable i to grasp when RISOTTO starts to be faster than RISO, which is just slightly before the solution. The solution of that equation is the fixed point of the following function

$$f(x) = -\log(1 - \frac{1}{2^{nx^e |\Sigma|^e}}) / \log(|\Sigma|).$$

Given that f(x) is contractive, that is, its derivate function takes values in the interval  $(-1 + \epsilon, 1 - \epsilon)$ , the fixed point can be computed by iterating f over an initial value. Finally, notice that the fixed point increases with the values of e, n and  $\Sigma$ . With the previous analysis, we have all the machinery necessary for computing the ratio between the expected number of attempted extensions between RISO and RISOTTO, as well as, from which point RISOTTO performs



Fig. 2. Left: Ratio between the expected number of extensions attempted by RISOTTO and RISO (cf Fig. 3 to compare theoretical with experimental results obtained in the same set). Right: Ratio between performance of RISOTTO and RISO.

better than RISO. As an example, the ratio between the expected number of extensions attempted by RISOTTO and RISO for a dataset consisting of N = 100 sequences of size n = 1000 where we planted r = 1 motif of size t = k = 5..20, with up to e = 2 mismatches, and quorum q = 100, is given in Fig. 2(left). For the dataset considered, the fixed point for f(x) is x = 10.6616.

## 3 Structured Motif Extraction

A structured motif is a pair (m, d) where  $m = (m_i)_{1 \le i \le p}$  is a p-tuple of single motifs and  $d = (d_{min_i}, d_{max_i})_{1 \le i \le p}$  is a (p-1)-tuple of pairs, denoting p-1 intervals of distance between the p single motifs. Each element  $m_i$  of a structured motif is called a box and its minimal and maximal length denoted by  $k_{min_i}$  and  $k_{max_i}$ , respectively. The structured motif extraction problem takes as parameters N input sequences, a quorum  $q \leq N$ , p maximal error rates  $(e_i)_{i < 1 < p}$  (one for each of the p boxes), p minimal and maximal lengths  $(k_{min_i})_{i \leq 1 \leq p}$  and  $(k_{max_i})_{i \leq 1 \leq p}$  (one for each of the p boxes), and p-1 intervals of distance  $(d_{min_i}, d_{max_i})_{i < 1 < p-1}$ (one for each pair of consecutive boxes). Given these parameters, the problem consists in searching for the contents of the boxes, that is the motifs, that have the structure defined by the parameters above and that satisfy the quorum. The algorithm for single motif extraction introduced in [7] is the ancestor of others [2,5] that infer structured motifs. The optimisation introduced in this paper can be applied to any of them. In a few words, the algorithm first builds the factor tree  $\mathcal{T}$  of the input sequences, then it searches for all valid motifs of length at least  $k_{min}$  and up to  $k_{max}$  (as in [7]) and, after updating the data structure (see [2] for details), checks whether there is a second valid motif (again as in [7]) with the required interval between them. The algorithm is described by Algorithm 3. for p = 2, where i indicates whether we are dealing with the first or the second box, and  $\lambda$  is the empty word.

### Algorithm 3. Structured motif extraction

### 3.1 Using Maximal Extensibility of Factors

In the case of structured motifs, the maximal extensibility information for the first box of a motif should be updated as described in Sect. 2.1. However, any failure in attempting to extend a motif during the search of a second box cannot update any value of MaxExt because it refers only to parts of the text that follow a specific first box at a specific distance. In fact, when a first box  $m_1$  of a structured motif is fixed at any given step, the maximal extensibility information

Algorithm 4. Structured motif extraction with maximal extensibility information

**ExtractStructuredMotif**(motif m, box i) 1. for all  $\alpha \in \Sigma$  do if i = 1 or  $e_2 \leq e_1$  then 2. 3  $x := m\alpha$ 4. while  $(x \neq root \text{ or } MaxExt(x) = -1) x := |x\rangle_{|x|-1}$ if  $x \neq root$  and  $MaxExt(x) + |m\alpha| < k_{min_i}$  then 5. if i = 1 then  $MaxExt(m\alpha) := MaxExt(x)$ 6. stop spelling  $m\alpha$  and continue 7. 8. if  $m\alpha$  is valid then 9. if  $|m\alpha| \ge k_{min_i}$  then if i = 2 then spell out the valid motif 10. else follow box-links and update  $\mathcal{T}$  to ExtractStructuredMotif( $\lambda, 2$ ) 11. 12. if  $|m\alpha| < k_{max_i}$  then ExtractStructuredMotif $(m\alpha, i)$ 13. else if i = 1 then  $MaxExt(\langle m\alpha |_{k_{min_1}-1}) := +\infty$ 14. else if i = 1 then 
$$\begin{split} & \text{if } |m\alpha| < k_{min_1} \text{ then } MaxExt(m\alpha) := 0 \\ & \text{else if } MaxExt(\langle m\alpha|_{k_{min_1}-1}) < |m\alpha| - (k_{min_1}-1) \text{ then } MaxExt(\langle m\alpha|_{k_{min_1}-1}) := 0 \end{split}$$
15. 16  $|m\alpha| - (k_{min_1} - 1)$ 17. if i = 1 and  $|m| < (k_{min_1} - 1)$  then  $MaxExt(m) := 1 + \max_{\alpha \in \Sigma} MaxExt(m\alpha)$ 

that concerns the whole sequence is in general an upper bound on the maximal extensibility of fragments of the sequence that are at a given distance from the occurrences of  $m_1$ . Given this observation, a possibility is to use the maximal extensibility information of the first box when searching and trying to extend a second box. Another possibility, while attempting to find a motif for the second box, is to compute and store tighter maximal extensibility information which we can use for the second box being attempted as long as the first box is fixed. In the following, we only address the first alternative, that is, only the first box stores extensibility information. The conditions needed for our optimisation to be applicable in the case of structured motifs may hold even more frequently than in the case of single motifs. In fact, since the search for a valid motif as second box is made after a valid motif for the first box is found, maximal extensibility information may be known also for the whole motif whose extension is attempted and not just for its prefixes. In other words, it may happen that when Algorithm 3. is called with parameters m and 2, the value of MaxExt(m)is already known. Proper suffixes are thus not the only candidates to give useful information when we are trying to find a motif for the second box. The extensibility information can be used as for the case of single motifs except that one has to deal with different error rates among boxes. Indeed,  $e_2$  must be less than or equal to  $e_1$  in order for the extensibility information to be useful for the second box. Otherwise, the maximal extensibility information stored for the first box may be too restrictive, and if it is used, it may cancel the extension of valid motifs. The algorithm for structured motif extraction using the maximal extensibility information is presented in Algorithm 4. Similarly to the case of single motif extraction, the time complexity of Algorithm 4. remains the same as for Algorithm 3. in the worst case, and the improvement proposed accounts only for the average case, as we shall verify in the next section.

### 4 Implementation and Experimental Results

In order to verify the improvement proposed in this paper, a C implementation of the maximal extensibility algorithm, called RISOTTO<sup>1</sup>, was made. The new implementation was tested against a C implementation of the algorithm presented in [2] and called RISO. The results of the experiments we made show a sensible improvement for both single and structured motif extraction when using maximal extensibility information. As we shall see in this section, maximal extensibility may cost some extra space, which is a delicate issue for large datasets, but it can definitely save some hopeless visits, and in general it results very efficient.

We start with some considerations concerning the storage of extensibility information. As we have seen in Sect. 2.1, due to the order in which motifs are considered, we have that only certain subwords of motifs can give useful information concerning maximal extensibility, namely, those that are lexicographically smaller. Since no motif is smaller than itself, we actually only use the MaxExtinformation of motifs that are shorter than the current one, that is, they are proper suffixes. Therefore, since the condition to check is whether or not we can hope to reach the  $k_{min}$  length, then we make use of the MaxExt data only for strings of length at most  $k_{min} - 1$ . Hence, it is not necessary to store this information for motifs that have length  $k_{min}$  or more for the purpose mentioned above. Let us now discuss how much space is required to store the extensibility information until level  $k_{min} - 1$ . We say that a tree is uncompact complete if it is a trie where all possible nodes are present. There is thus no arc whose label contains more than one letter. A previous result [1] makes use of some statistical analysis for stating that a suffix tree of a text of length n is expected to be uncompact complete at the  $log_{|\Sigma|}(n)$  top levels, where  $\Sigma$  is the alphabet of the text. This fact suggests a model to store extensibility information: a static data structure to keep the MaxExt values until level  $log_{|\Sigma|}(n)$ , and a dynamic structure for deeper levels. Since we are interested in the DNA alphabet (composed of the four nucleotides A, C, G, and T), then we have that our suffix tree is uncompact complete at the top  $log_4(n)$  levels where n is the size of the input sequence s. The function  $log_4(n)$  reaches 10 for  $n \approx 10^6$ , it is greater than 11 for  $n = 10^7$ , it is more than 13 for  $n = 10^8$ , and nearly 15 for  $n = 10^9$ . These values correspond to reasonable values for the minimal length  $k_{min}$  of the motif, and they are reached for values n of the text size corresponding to quite big datasets. In the RISOTTO implementation, we took all the observations above into consideration. Since  $k_{min}$  has to be relatively small for our approach to be tractable spacewise, we considered only 1 byte (a char in C) to store MaxExtvalues. In this case, extensibility values must be less than 256, which is quite reasonable. To build a static data structure to store such values until level z, we need z + 1 1-byte arrays, where the *j*-th array has size  $|\Sigma|^j$  with  $0 \le j \le z$ . Therefore, for the case of DNA, the total amount of memory required is  $\frac{4^{z+1}-1}{3}$ bytes. This function gives us values of 1.3MB for z = 10, 5.3MB for z = 11, 5.3MB for z = 10, 5.3MB for z = 11, 5.3MB for z = 10, 5.3MB for z = 11, 5.3MB for z = 10, 5.3MB for z = 11, 5.3MB for z = 10, 5.3MB for z = 10, 5.3MB for z = 11, 5.3MB for z = 10, 5.3MB

 $<sup>^{1}</sup>$  RISOTTO is available at http://algos.inesc-id.pt/~asmc/software/riso.html.

85.3MB for z = 13, and 1.3GB for z = 15. In our experiments, we achieved an optimum trade-off between memory allocation/management and maximal extensibility gain when z = 10. Taking this observation into account, we only allocate values for MaxExt until level  $z = \min\{10, k_{min} - 1\}$ , even for large values of  $k_{min}$ , and disregard deeper levels as well as the dynamic data structure mentioned above. Nevertheless, we allowed this z level to be an implementation parameter. In the end, considering  $z = \min\{10, k_{min} - 1\}$ , RISOTTO requires at most 1.3MB more that RISO for DNA databases, being more than twice faster as we shall see next.

To test maximal extensibility performance we used several randomly generated (with a uniform distribution over the four letters size DNA alphabet) synthetic datasets with planted structured motifs. Each dataset consists of 100 sequences of size 1000 where we planted one motif, possibly structured into several boxes, with 2 mismatches per box. We ran both RISO and RISOTTO requiring a quorum q = 100 and at most 2 mismatches per box so that the output contains at least the planted motif. For each dataset, we made several runs for increasing lengths of the motifs. In particular, given the number of boxes of the structured motifs (in our tests there are p boxes for  $p = 1, \ldots, 4$ ), we have increased the size of the boxes ranging from 5 to 20. As a result, the total motifs size (without counting the gaps) ranges from 5 to 80. For each p (number of boxes), we have plotted in Fig. 3, against the size of the motif (x axis), the



Fig. 3. Ratio between the number of extensions attempted by RISOTTO and RISO (cf Fig. 2(left) to compare theoretical with experimental results obtained in the same set)

ratio between the number of extensions attempted by RISOTTO and those by RISO (y axis). Given than RISOTTO only saves useless attempts, this equals the percentage of saved calls of the recursive procedure. For one box (Fig. 3 top left) we have depicted the results for several runs, while for two, three and four boxes (Fig. 3 top right and bottom) there are one curve for the inference of each box of the structured model. As one would expect, the attempts saved are more when the length of the motif increases and, in particular, the improvement starts when the length of the box is about 10 (this value depends in general from the input sequence and the alphabet size). For one box (see Fig. 3 top left), the number of attempted extension of RISOTTO decreases fast to 40% with respect to RISO (for growing values of the length of the motifs). Even better results, getting as good as attempting only 20% of the extensions of RISO, were achieved when extracting an *i*-th box with  $2 \le i \le p$  (see Fig. 3 (top right and bottom)). Moreover, we present the ratio of speed performance of the computation of RISOTTO with respect to that of RISO. This is shown for all tests together in Fig. 2(**right**) for all possible sizes of the boxes. One can see that the best relative performance is achieved for the first boxes (that is where it is more needed because the search space is very large and noisy), where RISOTTO is up to 2.4 faster than RISO. Finally, in [6] a challenging problem was launched that concerned finding all single motifs of length 15 with at most 4 mismatches in 20 texts of size 600. We ran both RISO and RISOTTO on such instances. We observe a speedup of 1.6 of RISOTTO over RISO. We actually believe that a true challenge should involve texts of larger size. Therefore, we ran tests with the same parameters (length 15 and at most 4 mismatches) on larger input sequences. The results confirm the 1.6 speedup for sequences of length 700 and 800, 1.3 speedup for length 900, and then the speedup decreases, but the time required by RISOTTO is always lower than for RISO.

## References

- 1. J. Allali. Structures d'indexation: les arbres des facteurs. Memoire de maitrise, University of Marne-la-Vallée, 2000.
- A. M. Carvalho, A. T. Freitas, A. L. Oliveira, and M.-F. Sagot. A highly scalable algorithm for the extraction of cis-regulatory regions. In *Proc. APBC'05*, pages 273–282. Imperial College Press, 2005.
- 3. D. Guha-Thakurta and G. D. Stormo. Identifying target sites for cooperatively binding factors. *Bioinformatics*, 17:608–621, 2001.
- 4. M. Lothaire. Applied Combinatorics on words. Cambridge University Press, 2005.
- L. Marsan and M.-F. Sagot. Algorithms for extracting structured motifs using a suffix tree with application to promoter and regulatory consensus identification. J. Comp. Bio., 7:345–360, 2001.
- P. A. Pevzner and S. H. Sze. Combinatorial algorithm for finding subtle signals in dna sequences. In *Proc. ISMB'00*, pages 269–278, 2000.
- M.-F. Sagot. Spelling Approximate repeated or common motifs using a suffix tree. In C.L. Lucchesi and A.V. Moura, editors, *Proc. Latin'98*, volume 1380, pages 111–127, 1998. LNCS.