Encoded Expansion: An Efficient Algorithm to Discover Identical String Motifs

Aqil M. Azmi*, Abdulrakeeb Al-Ssulami

Department of Computer Science, College of Computer & Information Sciences, King Saud University, Riyadh, Saudi Arabia

Abstract

A major task in computational biology is the discovery of short recurring string patterns known as motifs. Most of the schemes to discover motifs are either stochastic or combinatorial in nature. Stochastic approaches do not guarantee finding the correct motifs, while the combinatorial schemes tend to have an exponential time complexity with respect to motif length. To alleviate the cost, the combinatorial approach exploits dynamic data structures such as trees or graphs. Recently (Karci (2009) Efficient automatic exact motif discovery algorithms for biological sequences, Expert Systems with Applications 36:7952–7963) devised a deterministic algorithm that finds all the identical copies of string motifs of all sizes ≥2 in theoretical time complexity of O(nL+L^2), and a space complexity of O(nL^2), where n is the length of the input sequence and L is the length of the longest possible string motif. In this paper, we present a significant improvement on Karci’s original algorithm. The algorithm that we propose reports all identical string motifs of sizes ≥2 that occur at least τ times. Our algorithm starts with string motifs of size 2, and at each iteration it expands the candidate string motifs by one symbol throwing out those that occur less than τ times in the entire input sequence. We use a simple array and data encoding to achieve theoretical worst-case time complexity of O(nL), and a space complexity of O(n). Encoding of the substrings can speed up the process of comparison between string motifs. Experimental results on random and real biological sequences confirm that our algorithm has indeed a linear time complexity and it is more scalable in terms of sequence length than the existing algorithms.

Citation: Azmi AM, Al-Ssulami A (2014) Encoded Expansion: An Efficient Algorithm to Discover Identical String Motifs. PLoS ONE 9(5): e95148. doi:10.1371/journal.pone.0095148

Editor: Junwen Wang, The University of Hong Kong, Hong Kong

Received October 14, 2013; Accepted March 24, 2014; Published May 28, 2014

Copyright: © 2014 Azmi, Al-Ssulami. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This work was supported by a special fund of the Research Center of the College of Computer and Information Sciences (CCIS) at King Saud University. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: aqil@ksu.edu.sa

Introduction

There has been a considerable interest in motif discovery from both computer scientists and computational biologists. Motifs or binding sites are significant for understanding the mechanism behind regulating gene expressions. From a biological sequence analysis perspective, the significant pattern is the substring that is either over-represented or under-represented in a biological sequence. Therefore, the main problem is to identify the most or the rarest recurring patterns. Some of the methods depend on comparing the biological sequence with the background sequences to discover exceptional motifs. Those methods require generating specific length background sequences randomly. The problem of deciding whether the generated sequences respect the motif constraints, the number of motif occurrences, the length of motif, is NP-Complete [1]; essentially meaning that it is computationally hard for all practical purposes. To solve this problem we require an exponential time complexity, though space complexity may not be exponential.

Over the years, many algorithms were developed to discover and report motifs; and most of these algorithms were either stochastic or combinatorial in nature [2]. The stochastic algorithms such as Expectation Maximization (EM) [3], take a set of input sequences, the motif length, and an initial guess for the motif. This guess is generated either randomly or supplied by the user. The algorithm returns a probabilistic model of the consensus pattern, or the motif. EM assumes that there is a single motif occurring in each input sequence. It is possible that EM fails to return the correct motif, that is if we were unfortunate in picking a good starting point. Subsequently newer algorithms came into existence that extended EM, e.g. Gibbs Sampling [4,5,6,7,8,9], and Multiple EM for Motif Elicitation (MEME) algorithm [10,11,12,13,14,15]. On the other hand, the combinatorial approaches are very expensive because they exhaustively generate and search for each possible permutation of a given length making them impractical for motif sizes over 10 [16]. Some of the algorithms that fall into this category includes: Weeder [17], MotifEnumerator [18], Seeder [19], the algorithm by Marschall and Rahmann [20], VINE [21], PairMotif [22], and PairMotif+. Few of these algorithms resort to smart pruning to reduce the search space [17,19,23].

Most of the above algorithms rely on dynamic data structure to process the data. When it comes to the insertion and deletion operations, the dynamic data structures are very efficient. However, traversing dynamic data structures is less efficient than doing so on the static data structures as the data might be scattered all around [24,25]. Static data structures, e.g. arrays, are preferred if the algorithm was properly crafted. Our goal is to devise an efficient algorithm that discovers all the identical string motifs and does not rely on dynamic data structure thus saving on the memory requirements. The algorithm must be highly efficient and
Algorithm to Discover Identical String Motifs

Proposed Algorithm

Let $\Sigma$ denote the set of finite symbols (alphabet), we define $\Sigma^* = \{w \mid w \in \Sigma^* \text{ and } w \in \Sigma \}$ where $\Sigma^* = \epsilon$ (empty string) to be a set composed of elements each of length $k$. The $\Sigma^*$ denotes the set of all strings formed using the symbols in $\Sigma$. In our case the alphabet $\Sigma = \{A,C,G,T\}$ contains the four bases for the DNA biological sequence, so that any DNA sequence belongs to the language $\Sigma^*$. In this work we assume that all the indices start from 0. Let $S$ be a string of length $n$, $S = s_0s_1s_2 \ldots s_{n-1} \in \Sigma^*$. Substrings of $S$ which start at position $p$ and are of length $k$ are denoted $S_{p:k} = s_ps_{p+1} \ldots s_{p+k-1}$ where $0 \leq p \leq n-k$.

Our objective is: given the sequence $S$, find all the identical string motifs of lengths up to $L(2 \leq L < n/\tau)$, where all motifs of length $L + 1$ appears less than $\tau$ times in $S$. A motif of length $L$ is a substring that is repeated at least $\tau$ times in different positions in $S$.

Our algorithm proceeds as follows. We start by generating the pair bases ($2$-lets CanMotifs) and their starting positions. For example, if $S =$ TATAC and $\tau = 2$, then the list is $\{(TA, 0), (AT, 1), (TA, 2), (AC, 3)\}$ where the tuple stands for (CanMotif, starting position). The algorithm encodes the $2$-lets CanMotifs using the codes in Table 1, resulting in $\{(12, 0), (3, 1), (12, 2), (1, 3)\}$. Next the encoded sequence is sorted. Sorting is achieved by going over the list sixteen times. For our example, the list after the sorting will be $\{(1, 3), (3, 1), (12, 0), (12, 2)\}$. Then we delete all the entries where the encoded CanMotif occurred less than $\tau$ times resulting in $\{(12, 0), (12, 2)\}$. What remains is a list of all $2$-lets motifs. The algorithm will proceed to the next stage only if it was successful in discovering motif(s) at the current stage.

We start the next stage by generating an encoded list of all $3$-lets CanMotifs. The list is generated by augmenting every occurrence of $2$-lets motifs (out of the previous stage) with one nucleotide to its right in the input sequence $S$. Going back to our previous example we have only one $2$-lets motif, TA. The generated $3$-lets CanMotifs are $TAT$ and $TAC$ which will be encoded (more on that later). Once the encoded list is sorted we delete all CanMotifs that occur less than the threshold $\tau$. The remainder is a list of all $3$-lets motifs. The process continues discovering larger motifs and stops when there are no more motifs, or if they occur less than $\tau$ times. There are three aspects of the algorithm that will be covered in depth including the generation of the CanMotifs, the encoding scheme; and finally sorting and discovery of the motifs. For our ensuing discussion we would like to denote $\Phi(\cdot)$ for the set of $k$-lets CanMotifs.

Generating CanMotifs

In our algorithm we use $k$-lets motifs to generate the list of $k+1$-lets CanMotifs. Since there are only sixteen $2$-lets CanMotifs, we generate them all and this scheme is applied for $k \geq 2$. This is done by augmenting each occurrence of the $k$-lets motif with one right nucleotide. Figure 1 illustrates our scheme.

There are two issues that need to be addressed: (1) why did we not use $k$-lets CanMotifs to generate the list of $k+1$-lets CanMotifs; and (2) what difference (if any) does it make if we augment the $k$-lets motifs with one left nucleotide instead of one right nucleotide in our algorithm.

**Theorem 1.** It is redundant to use $k$-lets CanMotifs over $k$-lets motifs to generate the $k+1$-lets CanMotifs.

**Proof 1.** It suffices to show that a $k+1$-lets motif cannot be derived from a non-motif substring of length $k$. Let $S_{j}^{(k)}$ be a substring of length $k$ of the string $S$. Assume that $S_{j}^{(k)}$ is not a motif. Therefore there are no substrings $S_{i}^{(k)}$, $j \neq i$ in $S$ which equals $S_{j}^{(k)}$. Now, in the best case, both substrings share at most $k-1$ symbols, otherwise they will be equal and $S_{j}^{(k)}$ is a motif. Augmenting $S_{j}^{(k)}$ and $S_{i}^{(k)}$ with a single symbol each gives us at best a $k$-lets motif which is not what we are looking for.

A consequence of the above is a faster algorithm since the set of $k$-lets motifs is much smaller than the set of $k$-lets CanMotifs. For
example, consider the sequence, AACTGCTACTT. There are 10 2-lets CanMotifs: AA, AC, CT, TG, GC, CT, TA, AC, CT and TT, but only two 2-lets motifs: AC and CT. These 2-lets motifs will be augmented by one right nucleotide to form the set of 3-lets CanMotifs.

**Theorem 2.** Augmenting the \( k \)-lets motifs in either direction (left or right) to generate the list of \( k+1 \)-lets CanMotifs will yield the same set of \( k+1 \)-lets motifs.

**Proof 2.** Note that a CanMotif may or may not yield a motif, and that two different CanMotifs cannot yield the same motif. It does not matter if left and right augmentations result in a different set of CanMotifs, the important thing is that they both result in identical motifs. Suppose that we have two identical substrings of length \( k \), \( S_{i}^{(k)} \sim S_{j}^{(k)} \). Let \( \varphi \) represent this \( k \)-lets motif. Note that \( s_{i-1} \) is the left nucleotide of \( S_{i}^{(k)} \) and \( s_{i+k} \) is its right nucleotide, for \( S_{j}^{(k)} \) is \( s_{j-1} \) and \( s_{j+k} \) respectively. Augmenting the \( k \)-lets motif \( \varphi \) with the left nucleotide results in the following \( k+1 \)-lets CanMotifs: \( \{ s_{i-1} S_{i}^{(k)}, s_{i-1} S_{i}^{(k)} \} = \{ s_{i-1}^{(k+1)}, s_{i-1}^{(k+1)} \} \), while augmenting \( \varphi \) with the right nucleotide results in the \( k+1 \)-lets CanMotifs: \( \{ S_{i}^{(k)} S_{j}^{(k)}, S_{j}^{(k)} S_{j}^{(k)} \} \). Depending on whether the left and the right nucleotides are the same or not, we have four different cases:

- Case \( s_{i-1} \neq s_{j-1} \) and \( s_{i+k} \neq s_{j+k} \). Here neither augmentation of the \( k \)-lets motif \( \varphi \) will yield a \( k+1 \)-lets motif.
- Case \( s_{i-1} = s_{j-1} \) and \( s_{i+k} \neq s_{j+k} \) (Figure 2). Since \( S_{i-1}^{(k)} = S_{j-1}^{(k)} \) we have another \( k \)-lets motif, \( \varphi' \). To shorten the argument we will assume that \( s_{i-2} \neq s_{j-2} \) and \( s_{i+k-1} \neq s_{j+k-1} \). Now the left augmentation of the \( k \)-lets motif \( \varphi \) will be the same as the right augmentation of the \( k \)-lets motif, \( \varphi' \). In other words both will yield the same \( k+1 \)-lets motif. Certainly, the right augmentation of \( \varphi \) and the left augmentation of \( \varphi' \) will yield a \( k+1 \)-lets CanMotif, one which will not result in a motif.
- Case \( s_{i-1} \neq s_{j-1} \) and \( s_{i+k} = s_{j+k} \). Argument similar to above.

### Table 1. Encoding of pair bases.

| AA | AG | AC | CA | CG | CT | GA | GG | GC | TT |
|----|----|----|----|----|----|----|----|----|----|
| 0  | 1  | 2  | 3  | 4  | 5  | 6  | 7  | 8  | 9  |
| 10 | 11 | 12 | 13 | 14 | 15 |

This is the full list of all 2-lets candidate motifs (CanMotifs).

**Figure 1.** Each occurrence of \( k \)-lets motif is augmented with right nucleotide to form the \( k+1 \)-lets CanMotif.

doi:10.1371/journal.pone.0095148.g001

**Figure 2.** Left augmentation of \( \varphi \) and the right augmentation of \( \varphi' \) yield the same \( k+1 \)-lets motif.

doi:10.1371/journal.pone.0095148.g002
CanMotifs is a method for identifying motifs in a sequence, and in general the maximum number of unique motifs that can be discovered is $16^2$ for two-letter motifs, $64^3$ for three-letter motifs, and so on. This algorithm is designed to handle motifs of any size, as a result the above encoding is always used.

The basic idea is to map a $k$-letter CanMotif $\varphi$ to a single integer. However, if we insist on a unique encoding for each different motif, we will have an integer overflow. We want our algorithm to avoid a situation where the encoding of the generated CanMotif is the same as that of the encoding of one of the input motifs. The variable $\Delta$ is used to take care of this problem. The test $p \geq n - k$ allows us to reject the rightmost $k$-letter motif because there is no right nucleotide it can be augmented with. The function $\text{NucVal}$ maps a nucleotide to a numerical value $\{A, C, G, T\rightarrow 0, 1, 2, 3\}$. It can be easily shown that no two different CanMotifs have the same encoding.

As we mentioned earlier, one of our concerns was avoiding the integer overflow while computing the encoding of CanMotif. Our solution is to re-number the motifs sequentially starting from 0. The re-encoding scheme is straightforward (Algorithm 2). It goes over the entire encoded motif assigning a new code to each unique motif.

Algorithm 2. Algorithm to re-encode the $k$-letter motifs so it always starts from 0.

**Input.** List of encoded $k$-letter motifs $(z_1, p_1), (z_2, p_2), \ldots, (z_n, p_n)$ in increasing order, i.e. $z_i \leq z_j$ for all $i \leq j$, where the tuple stands for (encoded $k$-letter motif, its starting position).

**Output.** List of re-encoded $k$-letter motifs $(x'_1, p'_1), (x'_2, p'_2), \ldots, (x'_n, p'_n)$.

**Begin.**

1. $\Delta \leftarrow 1 + \max z$ //value of largest encoded motif
2. Loop over all encoded $k$-letter motifs
   - if $p \geq n - k$ then continue //reached boundary
   - $x' \leftarrow \Delta + 2x + \text{NucVal}(z_p + k)$
   - $p' \leftarrow p$
3. End.

The data is encoded using the tuple (encoded CanMotif, its starting position). The starting position helps in recovering the CanMotif and in fetching the right nucleotide which the motif has to be augmented with.

We use Table 1 for encoding 2-letter CanMotifs. For larger CanMotifs, we use Algorithm 1 to generate and encode $k + 1$-letter CanMotifs out of encoded $k$-letter motifs.

We want to avoid a situation where the encoding of the generated CanMotif is the same as that of the encoding of one of the input motifs. The variable $\Delta$ is used to take care of this problem. The test $p \geq n - k$ allows us to reject the rightmost $k$-letter motif because there is no right nucleotide it can be augmented with. The function $\text{NucVal}$ maps a nucleotide to a numerical value $\{A, C, G, T\rightarrow 0, 1, 2, 3\}$. It can be easily shown that no two different CanMotifs have the same encoding.

As we mentioned earlier, one of our concerns was avoiding the integer overflow while computing the encoding of CanMotif. Our solution is to re-number the motifs sequentially starting from 0. The re-encoding scheme is straightforward (Algorithm 2). It goes over the entire encoded motif assigning a new code to each unique motif.

**Algorithm 2.** Algorithm to re-encode the $k$-letter motifs so it always starts from 0.

**Input.** List of encoded $k$-letter motifs $(z_1, p_1), (z_2, p_2), \ldots, (z_n, p_n)$ in increasing order, i.e. $z_i \leq z_j$ for all $i \leq j$, where the tuple stands for (encoded $k$-letter motif, its starting position).

**Output.** List of re-encoded $k$-letter motifs $(x'_1, p'_1), (x'_2, p'_2), \ldots, (x'_n, p'_n)$.

**Begin.**

1. $\text{old_encoding} \leftarrow 0$
2. $\text{new_encoding} \leftarrow 0$
3. for $i \leftarrow 1 \ldots n$
   - if $z_i \neq \text{old_encoding}$
     - $\text{old_encoding} \leftarrow z_i$
     - $\text{new_encoding} \leftarrow \text{new_encoding} + 1$
4. $z_i \leftarrow \text{new_encoding}$
5. End.

Back to our initial example $S = \text{TATAC}$, the encoded 2-letter motifs were $\{(12, 0), (12, 2)\}$ where 12 is the encoding of 2-letter motif, TA (Table 1). The other number in the tuple, 0 and 2 in our case, represents the starting position of the motif. The algorithm returns the generated 3-letter encoded CanMotifs $\{(64, 0), (64, 2)\}$. For this computation, the algorithm sets the value of $\Delta = 13$. So we have two different 3-letter CanMotifs, one whose encoded value is 62 (CanMotif TAC), and the other is 64 (CanMotif TAT). Although we can determine what actual CanMotifs these are (using the starting position and the value of $\Delta$), this is unnecessary.
for our task. The important thing is whether we have other CanMotifs of similar values?

**Sorting and discovering motifs**

Given a list of encoded \( k + 1 \)-lets CanMotifs the simplest way to discover the motifs is by sorting this list. It is a better method than doing an exhaustive search for each CanMotif. But the fastest sorting algorithm has a time complexity \( O(n \log n) \). We, however, will use counting sort, a linear time sorting algorithm.

Using counting sort, we can sort the list of encoded 2-lets CanMotifs using 2 passes only. In the first pass, we count the number of occurrences of each distinct key value. That is, make a histogram of all encoded 2-lets AA, AC, AG, … etc., and in the second pass, we use arithmetic on this count to determine the position of each key value. As counting sort uses key values as indexes into the list, so it is linear in the size of the input list. Thus the cost of this algorithm is \( O(n) \).

For 3-lets and beyond the situation is as follows. Given a sorted list of encoded \( k \)-lets motifs \((k \geq 2)\) we generate a sorted list of encoded \( k + 1 \)-lets CanMotifs. Initially we feed the encoded 2-lets motifs (sorted using counting sort) to get a sorted list of encoded 3-lets CanMotifs. For subsequent values of \( k \) the input is a subset of the output of the previous stage. So if the output is sorted that means the input to the next stage is sorted too. Algorithm 1 already generates the desired encoded CanMotifs; we only need to modify it to output in a sorted form.

Suppose that \( ((x_1,y_1),(x_1,y_2),(x_1,y_3),(x_2,y_1),(x_2,y_2),(x_2,y_3),\ldots) \) is the sorted list of encoded \( k \)-lets motifs, where \( x_1 < x_2 \ldots \) are the encoded motifs and \( p_1,p_2,\ldots \) their starting positions. In generating the encoded \( k + 1 \)-lets CanMotifs, we note that the largest value \( x_1 \) goes to is \( \Delta + 4x_1 + 3 \) while the smallest value \( x_2 \) goes to is \( \Delta + 4x_2 \). We can easily see that the relation \( \Delta + 4x_1 + 3 < \Delta + 4x_2 \) is true for any integer values of \( x_1 < x_2 \). This suggests a naive sorting algorithm where we do 4 passes (a pass for each nucleotide) over each group in array \( B \) and renumber the different motifs starting from 0. Algorithm 2 already generates the desired encoded CanMotifs; we only need to modify it to output in a sorted form.

The algorithm

The complete algorithm is shown as Algorithm 3. All the calculations are done in the array \( B \), where at iteration \( k \) the \( B[i].value \) holds the encoding of the \( k \)-lets which starts at position \( B[i].pos \).

Algorithm 3. The full listing of the identical string motif discovery algorithm. The algorithm automatically keeps discovering larger and larger motifs and stops when there are no more motifs to be found.

**Input.** \( Seq[i].Nucleotide \) the input sequence of size \( SeqLength \), and threshold \( \tau \)

**Output.** Display the full list of identical string motifs

**Begin.**

1. Initialize \( Seq[i].NucVal \) with the corresponding nucleotide value (A, C, G, T→0, 1, 2, 3)

2. Fill \( Seq[i].value \) with the encoding of the pair bases (see Table 1), where \( Seq[i].value \) is the encoding for the pair at \( Seq[i..i+1].Nucleotide \)

3. Count sort \( Seq[i] \) on \( .value \) field in ascending order saving \( B[j].value \) (count \( Seq[i].value \) and \( B[j].pos \)→i)

4. \( k \)→2

5. \( n \)→\( SeqLength \) – 1

6. while \( (n>0) \) do

7. \( \{ \)

8. //remove all motifs that occur less than the threshold \( \tau \)

9. \( i \)→0

10. while \( (i<n) \) do

11. \( \{ \)

12. Let \( \zeta = \) number of entries that has the same \( .value \) as \( B[j].value \)

13. if \( (\zeta < \tau) \) then

14. \( \{ \)

15. Mark \( B[i], B[i+1], \ldots, B[i+\zeta] \) for deletion

16. \( i \)→\( i+\zeta \)

17. \( \} \)

18. Discard all marked entries in array \( B \) by shifting the contents

19. \( ce \)→new size of the array \( B \) //number of occurrences of all \( k \)-lets motifs

20. for \( i=0 \ldots c-1 \) do

21. renumber the different motifs starting from 0

22. \( Last-\text{last}+1 \)-largest motif number

23. \( \text{new} \)→c

24. Output \( k \)-lets motifs and their starting positions

25. \( q \)→0

26. \( k=k+1 \)

27. \( \text{loop} \) over each group in array \( B \) //each group has the same \( .value \)

28. \( \{ \)

29. \( x \)→index of the first element in the group

30. \( \beta \)→index of the last element in the group

31. for \( i=0 \ldots 3 \) do

32. \( \{ \)

33. for \( j=x \ldots \beta \) do

34. \( \{ \)

35. \( xx \)→\( B[j].pos+k-1 \)

36. if \( (xx<\text{SeqLength} \&\& \text{Seq}[x].NucVal = = \tilde{e}) \) then

37. \( \{ \)

38. \( \} \)

39. \( \} \)

40. \( \} \)

41. \( \} \)

42. \( \} \)

43. \( \} \)

44. \( \} \)

45. \( \} \)

46. \( \} \)

47. \( \} \)

48. \( \} \)

49. \( \} \)

50. \( B = \text{tmp} \)

51. \( \} \)

52. End.

The code should be simple to follow. The while-loop at lines 13–21 expect a list of all CanMotifs sorted on their encoding value.
Since the list is sorted, we can determine the occurrence of each motif easily and remove those which occur fewer than the threshold \( t \). We re-encode (re-number) the encodings of motifs so that it always start from 0. The idea is to prevent an integer overflow, see the discussion in (Section CanMotifs encoding scheme.) This is achieved by the loop at line 26 (see Algorithm 2 for the details). Lines 31–49 code the algorithm in Figure 3, which generates a sorted list of encoded \( k \)-lets CanMotifs from a sorted list of \( k \)–lets motifs.

When applying the algorithm on the sample sequence below,

**ATAGACGTAGTATACGTCAGCATGCAG**

using the threshold \( t = 2 \), the algorithm discovers 9 different 2–lets motifs, 5 different 3–lets motifs and two different 4–lets motifs. Figure 4 marks all the 3–lets and 4–lets motifs in the example sequence.

The algorithm discovers and reports all motifs including those that overlap, e.g. TATA (Figure 4). Surely, overlapped motifs may not make sense biologically, but the algorithm does report them. These motifs can be easily discarded through a simple conditional statement that ensures the starting positions of any two \( k \)-lets motifs is at least \( k \) symbols apart.

**Complexity analysis of the algorithm**

For the complexity analysis of the algorithm, we assume the size of the input sequence is \( n \). Starting with computational complexity. Line 1 requires \( n \) operations. We can use a hash table to initialize the \( \text{Seq[ ]}.value \) with the encoding of pair bases (line 3) which costs \( n \) operations. In (Section Sorting and discovering motifs) we have shown that sorting 2–lets CanMotifs (line 5) costs \( 2n \) operations. The while-loop (lines 13–21) is used to mark all CanMotifs if they happen to occur less than the threshold \( t \) times. This can be efficiently implemented using \( 2n \) operations. In the first pass, we just mark those CanMotifs destined for removal with an invalid encoding, say \(-1\), then deleting them (line 22) in the second pass by shifting the contents. A single pass is sufficient to re-number the motifs. The loop to generate the sorted list of encoded

---

**Figure 3. A linear algorithm to generate a sorted list of encoded \( k+1 \)-lets CanMotifs from a sorted list of encoded \( k \)-lets motifs.**

Each group is sorted individually.

doi:10.1371/journal.pone.0095148.g003

---

**Figure 4. All the 3 and 4–lets identical string motifs in the sample sequence.**

doi:10.1371/journal.pone.0095148.g004
CanMotifs (lines 33–49) costs $4n$ operations as shown in (Section Sorting and discovering motifs). The copying in line 50 costs $2n$ operations. Note that the main while-loop (lines 9–51) is repeated $L$ times, where $L$ is the length of the longest motif that occurs not less than the threshold $t$.

The theoretical worst-case time complexity is $4n^2 + 9n(L - 1) = O(nL)$. In reality, the time complex-

### Table 3. Experimental results of running our algorithm on selected sets from the data sets [27,28] using $\tau = 2$.

| Sequences | k-lets | Overlapping | Examples with starting positions | No | Yes |
|-----------|--------|-------------|----------------------------------|----|-----|
| dm02r     | 11     | 2           | ATCCCCAATCCC→748, 760; ATCCCCAATCCC→748, 760 |
|           | 10     | 9           | TCTTGGGCC→670, 1164 |
|           | 9      | 25          | CTGGGGGGG→672, 1534 |
| yst09r    | 23     | 2           | GAAAAAAAAAAAAAAAAAAAAAAA→11659, 13120 |
|           | 17     | 3           | |
|           | 16     | 5           | TGAAAAAAAAAAAAAAAAAAAA→861, 11658 |
|           | 15     | 13          | GTTTCAAGCTGAGG→324, 1319 |
| hm20r     | 42     | 1           | ACTCGGGAGGCTGAGGAGAATCACCTGAGACCCGGAGG→24338, 64502 |
|           | 41     | 3           | GAGACAGCTGCCTGGCAACTGTTGGAAACCCCGTCTCTACTA→4490, 58687 |
| dm01g     | 14     | 2           | CACGCGGAGGAGGAGAAGAATCACCTGAGACCCGGAGG→1372, 1393 |
|           | 13     | 5           | TTATTTATATTT→32, 55 |
|           | 12     | 11          | |
|           | 11     | 28          | |
| mus03g    | 12     | 2           | TCTCCAAATCTA→755, 1103; TCTTGGGAGCT→1575, 2400 |
|           | 11     | 4           | |
|           | 10     | 13          | |
| yst01g    | 14     | 1           | GATCTCAAACAAAA→4985, 6920 |
|           | 13     | 3           | GAACCAAGATG→506, 2239 |
|           | 12     | 12          | CTAAAAGATAA→2611, 4585 |
|           | 11     | 37          | ACCAAAGATGG→508, 2241 |
| hm20m     | 18     | 1           | TGCGCCAGGGCTGGGCTG→34498, 69949 |
|           | 17     | 3           | GCACCAGGCTCCCGCCG→25468, 47769 |
|           | 16     | 7           | CGTCGACGCCCCTCCC→5487, 69910 |
|           | 15     | 17          | AAATGCCACCGAG→35282, 59636 |
|           | 14     | 47          | GCCCTCAGCCCGGC→2385, 26115 |

(a) Count the number of different motifs. For non-overlapping motifs we only consider motifs if their starting position is further apart than their length. (b) The starting position is based on index starting at 0. We followed [26] in treating each of the sequences as a single string. For example, yst09r.fasta is composed of 16 substrings each having 1000 nucleotides. These are merged into a single string with 16000 nucleotides. This set includes real (sequences suffixed ‘r’), generic (sequences suffixed ‘g’), and markov (sequences suffixed ‘m’) data sets. Only larger sized identical string motifs are reported.

doi:10.1371/journal.pone.0095148.t003

### Table 4. Execution time (in seconds) to find identical string motifs of all sizes on an Intel core i5 based PC running at 2.67 GHz with 4 GB RAM.

| Sequence | Size (# nucleotides) | Karci algorithm | Our algorithm |
|----------|----------------------|-----------------|---------------|
| mus06r   | 1500                 | 0.57            | 0.33          |
| dm06r    | 3000                 | 1.77            | 0.40          |
| yst04r   | 7000                 | 9.49            | 0.57          |
| hm26r    | 9000                 | 18.56           | 0.83          |
| yst09r   | 16000                | 53.43           | 1.21          |
| hm01r    | 36000                | 596.43          | 1.70          |
| hm20r    | 70000                | 2225.15         | 2.45          |

doi:10.1371/journal.pone.0095148.t004
plexity is much lower. Note that \( n \) is the initial size of the input sequence, and this \( n \) actually shrinks at each iteration since we are removing all CanMotifs that occur below the threshold.

For space complexity, we have three arrays all of size \( n \). The array \( \text{Seq} \) has three components (\text{Nucleotide}, \text{NucVal} and \text{value}), while arrays \( B \) and \( \text{tmp} \) both have two components each (\text{pos} and \text{value}). The total space requirement is \( 7n = O(n) \).

### Experimental Results and Discussion

For testing purposes, we will conduct experiments using three different data sets. A data set which is extracted from real data set from TRANSFAC [27]; randomly generated sequences of different sizes; and real biological sequences. The first data set is the same set that is used by Karci [26]. This data set is made up of two different sets: real, and synthetic. The real data set is from TRANSFAC; while the synthetic data set (generic and markov), is created out of extracted data from TRANSFAC using two different schemes to randomly place the binding site [28]. The full data set can be downloaded from the site, [http://bio.cs.washington.edu/assessment/download.html](http://bio.cs.washington.edu/assessment/download.html). The Result of running the algorithm on the first data set is in Table 3. This table will help readers verify the accuracy of our algorithm and at the same time compare it with the results in [26].

Next, we do a performance comparison to compare the execution time of our algorithm against the time needed by the algorithm in [26]. As stated earlier, this work is a significant improvement of the work in [26], which claimed a time complexity of \( O(n^2L) \). For the sake of fair comparison we implemented Karci’s algorithm in MS Visual C#, which is the same environment that we used for our algorithm. Table 4 summarizes the execution time of both algorithms running on the same platform, an Intel core i5 processor based PC running at 2.67 GHz with 4 GB of RAM.

For further testing, we generated random sequences over the alphabet \{A, C, G, T\} of various lengths. For each length, we generated ten different random sequences and calculated the execution time (in seconds) to discover all the identical string motifs of lengths not exceeding 40 nucleotides in real biological sequences.

### Table 5. Execution time (in seconds) to discover all the identical string motifs of lengths not exceeding 40 nucleotides in real biological sequences.

| Organism                        | NCBI RefSeq | Size  | Time  |
|---------------------------------|-------------|-------|-------|
| Vaccinia virus                  | NC_006998.1 | 0.19 M| 1.83  |
| Mycoplasma penetrans HF-2       | NC_004432.1 | 1.36 M| 10.38 |
| Lactobacillus acidophilus NCFM  | NC_006814.3 | 1.99 M| 12.93 |
| Methanothermococcus paludicola SANA  | NC_013665.1 | 2.96 M| 19.24 |
| Acidiphilium multivorum AIU301 | NC_015186.1 | 3.75 M| 27.51 |
| Mycobacterium tuberculosis H37Rv| NC_00962.2  | 4.41 M| 28.81 |
| Pectobacterium wasabiae WPP163  | NC_013421.1 | 5.06 M| 31.13 |
| Mesorhizobium opportunstum WSM2073 chromosome | NC_015675.1 | 6.88 M| 42.14 |
| Saccaropolyspora erythrea NRRL 2338 chromosome | NC_009142.1 | 8.21 M| 55.85 |
| Caenorhabditis elegans Bristol N2 chromosome III | NC_003281.10 | 13.78 M| 105.54 |
| Caenorhabditis elegans Bristol N2 chromosome II | NC_003280.10 | 15.28 M| 119.63 |

The size of the sequences is expressed in M (for millions). doi:10.1371/journal.pone.0095148.t005
average time to discover all the identical string motifs of all possible sizes. Figure 5 plots the average time (in seconds) that our algorithm required for finding all the identical motifs. Using MS Excel, the time \( t \) (in seconds) is given by \( t = 8.533 \times 10^{-6}n - 0.962 \), where \( n \) is the length of the sequence. The linear-in-time behavior of the algorithm is apparent.

The final test is on real biological sequences. Our algorithm was able to discover and report all the identical string motifs including some of impractical sizes. These huge motifs may not make sense biologically, but from a string point of view, they are valid identical motifs. For example, for *Mesorhizobium loti* our algorithm found two identical string motifs of size 6357 nucleotides. Table 5 shows the time to discover all identical string motifs of lengths up to 40. We believe this is reasonable, since in reality motifs rarely exceed 40 nucleotides. Let \( n \) be the length of the sequence, then using MS Excel we can calculate the time (in seconds) which is \( t = 7.776 \times 10^{-6}n - 3.8 \).

Given that, the time function for finding all identical string motifs for random sequences as well as real biological sequences is close, this allows us to claim that our algorithm has indeed a complexity that is linear in time.

**Conclusion and Future Work**

In this paper we presented an algorithm that discovers automatically all the identical string motifs in a given sequence. The idea of the algorithm is rooted in [26]; which had a complexity of \( O(n^2 L + L^2) \) in time, and \( O(nL^2) \) in space, where \( n \) is the length of the input sequence and \( L \) is the length of the longest possible motif. Our enhancement improved the complexity to \( O(nL) \) in time and linear in space. We were able to achieve this due to three factors: an encoding scheme for the motifs by which we have eliminated string comparison operation; relying on motifs only to generate a list of candidate motifs of larger size, which helps in placing a cap on the number of motifs to check among; and the usage of a linear algorithm to sort the encoded motifs thereby simplifying the task of identifying motifs. A further enhancement is the introduction of a threshold for the minimum number of occurrences of a motif. Experimental results on random, synthetic, and real biological sequences demonstrate that our algorithm has a time complexity that is linear.

For future work, we seek to enhance the algorithm so that it can discover the planted \((\ell, d)\) motifs, and the degenerate motifs.

**Implementation and Availability**

The program is implemented in MS Visual C# running under Windows operating system. The executable is available for academic use only. It is obtainable through email request.

**Acknowledgments**

We thank one of the reviewers for suggesting a threshold for the minimum number of motifs. All three anonymous reviewers deserve our appreciation for their constructive comments which helped improve this paper.

**Author Contributions**

Conceived and designed the experiments: AAS AMA. Performed the experiments: AAS. Analyzed the data: AAS AMA. Contributed reagents/materials/analysis tools: AAS. Wrote the paper: AMA.

**References**

1. Rivière R, Barth D, Cohen J, Denia A (2008) Shuffling biological sequences with motif constraints. Journal of Discrete Algorithms 6:192–204.
2. Vanea A, Marsan L, Sagot MF (1999) Promoter sequences and algorithmical methods for identifying them. Research in Microbiology 150:779–799.
3. Lawrence CE, Reilly AA (1990) An expectation maximization (EM) algorithm for the identification and characterization of common sites in unaligned biopolymer sequences. Proteins: Structure, Function, and Bioinformatics 7:41–51.
4. Lawrence CE, Autschbach SF, Boguski MS, Liu JS, Neuwald AF, et al (1993) Detecting subtle sequence signals. A Gibbs sampling strategy for multiple alignment. Science 262:208–214.
5. Liu JS, Neuwald AF, Lawrence CE (1995) Bayesian models for multiple local sequence alignment and Gibbs sampling strategies. Journal of the American Statistical Association 90:1156–1170.
6. Thijs G, Marchal K, Lescot M, et al (2002) A Gibbs sampling method to detect overrepresented motifs in the upstream regions of coexpressed genes. Journal of Computational Biology 9:447–464.
7. Siddharthan R, Sigga ED, van Nimwegen E (2005) PhyloGibbs: A Gibbs sampling motif finder that incorporates phylogeny. PLOS Computational Biology 7:e67.
8. Shida K (2006) GibbSF: a Gibbs sampling method for motif discovery with enhanced resistance to local optima. BMC Bioinformatics 7:486.
9. Defrance M, van Helden J (2009) info-gibbs: a motif discovery algorithm that directly optimizes information content during sampling. Bioinformatics 25:2715–2722.
10. Bailey TL, Elkan C (1995) Unsupervised learning of multiple motifs in biopolymers using expectation maximization. Machine Learning 21:51–80.
11. Grundy WN, Bailey TL, Elkan CP (1996) ParaMEME: a parallel implementation and a web interface for a DNA and protein motif discovery tool. Computer Applications in the Biosciences 12:303–310.
12. Bailey TL, Williams N, Misleh C, Li WW (2006) MEME: discovering and analyzing DNA and protein sequence motifs. Nucleic Acids Research 34:W369–W373.
13. Bailey TL, Boden M, Buske FA, Frith M, Grant CE, et al (2009) MEME SUITE: tools for motif discovery and searching. Nucleic Acids Research 37:W202–W208.
14. Bailey TL, Boden M, Whittington T, Machanick P (2010) The value of position-specific priors in motif discovery using MEME. BMC Bioinformatics 11(1):179.
15. Brown P, Baxter L, Hickman R, Beynon J, Moore JD, et al (2013) MEME-LaB: motif analysis in clusters. Bioinformatics 29:1696–1697.
16. GohChThakurta D (2006) Computational identification of transcriptional regulatory elements in DNA sequence. Nucleic Acids Research 34:3583–3598.
17. Patva G, Mereghetti P, Mauri G, Pesole G (2004) Weeder Web: discovery of transcription factor binding sites in a set of sequences from co-regulated genes. Nucleic Acids Research 32:W199–W203.
18. Sze SH, Zhao X (2006) Improved pattern-driven algorithms for motif finding in DNA sequences. Joint Annual RECOMB 2005 Satellite Workshops on Systems Biology and on Regulatory Genomics San Diego, CA, USA, December 2–4, 2005. Lecture Notes in Bioinformatics 4625:198–211.
19. Fauteux F, Blanchette M, Strømvik MV (2000) Seeder: discriminative seeding DNA motif discovery. Bioinformatics 24:2303–2307.
20. Marchall T, Rahmann S (2009) Efficient exact motif discovery. Bioinformatics 25:i356–i364.
21. Huang CW, Lee WS, Hsieh SY (2011) An improved heuristic algorithm for finding motif signals in DNA sequences. IEEE/ACM Transactions on Computational Biology and Bioinformatics 8:959–975.
22. Yu Q, Hao H, Zhang Y, Guo H (2012) PairMotif: A New Pattern-Driven Algorithm for Planted \((\ell, d)\) DNA Motif Search. PLOS ONE 7(10):e40412.
23. Yu Q, Hao H, Zhang Y, Guo H, Guo H (2013) PairMotif+: a fast and effective algorithm for de novo motif discovery in DNA sequences. International Journal of Biological Sciences 9:412–424.
24. Crochemore M, Rytter W (1994) Text algorithms. Oxford University Press.
25. Gusfield D (1997) Algorithms on Strings Trees and Sequences Computer Science and Computational Biology. Cambridge University Press.
26. Karci A (2009) Efficient automatic exact motif discovery algorithms for biological sequences. Expert Systems with Applications 36:7952–7963.
27. Wingender E, Dietze P, Karas H, Knuppel R (1996). TRANSFAC: A database for transcription factors and their DNA binding sites. Nucleic Acids Research 24:230–231.
28. Tompa M, Li N, Bally TL, Church GM, De Moor B, et al (2005) Assessing Computational Tools for the Discovery of Transcription Factor Binding Sites. Nature Biotechnology 23(1):137–144.