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INSTITUT NATIONAL DE RECHERCHE EN INFORMATIQUE ET EN AUTOMATIQUE

Finding approximate repetitions under Hamming distance

Roman Kolpakov and Gregory Kucherov

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THÈME 2

A large blue rectangle occupies the lower half of the page. Overlaid on it is a large, light gray stylized 'R'. To the right of the 'R', the words 'Rapport de recherche' are written in a white serif font, with 'Rapport' on the top line and 'de recherche' on the bottom line. A horizontal gray brushstroke underline is positioned below the text.

***Rapport
de recherche***



Finding approximate repetitions under Hamming distance

Roman Kolpakov* and Gregory Kucherov

Thème 2 — Génie logiciel
et calcul symbolique
Projet Adage

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Abstract: The problem of computing tandem repetitions with K possible mismatches is studied. Two main definitions are considered, and for both of them an $O(nK \log K + S)$ algorithm is proposed (S the size of the output). This improves, in particular, the bound obtained in [LS93]. Finally, other possible definitions are briefly analyzed.

Key-words: Algorithm, pattern matching, tandem repeat, approximate repetition

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at Moscow University

Recherche de répétitions approchées sous la distance de Hamming

Résumé : Nous étudions le problème de recherche de répétitions en tandem avec K inégalités. Nous considérons deux définitions principales et pour chacune d'elles nous proposons un algorithme de recherche s'exécutant en temps $O(nK \log K + S)$, où n est la longueur du mot et S est la taille de sortie. Cela améliore en particulier l'algorithme proposé dans [LS93]. Dans la dernière section nous analysons d'autres définitions possibles.

Mots-clés : Algorithme, recherche de motifs, répétition en tandem, répétition approchée

Finding approximate repetitions under Hamming distance

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1 Introduction

Repetitions (periodicities) play a central role in word combinatorics [Lot83, CK97]. On the other hand, repetitions are important from the application perspective. As an example, their properties allow to speed up pattern matching algorithms [GS83, CR95, CH98].

The problem of efficiently identifying repetitions in a given word is one of the classical pattern matching problems [Cro81, Sli83]. A *tandem repeat* or a *square* is a pair of consecutive occurrences of a subword in a word. For example, *baba* is a tandem repeat in word *cbacbabacba*. Since the beginning of 80s [Cro83] it is known that checking whether a word contains *no* tandem repeat (or is *square-free*) can be done in time $O(n)$ (n length of the word). If one wants to find *all* tandem repeats, their number comes into consideration. Word a^n contains $O(n^2)$ tandem repeats. If we restrict ourselves to *primitive*

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squares (i.e. subwords uu where u is not itself a repetition v^k for $k \geq 2$), then a word may contain $O(n \log n)$ of them and this bound is tight. All primitive squares can be found in time $O(n + S)$ where S is their number [Kos94, SG98, KK99a], hence in the worst-case time $O(n \log n)$.

In [KK99b, KK99a], we studied *maximal repetitions* (see also [ML84, Mai89]). Those can be viewed as maximal *runs* of squares [IMS97, SG98], i.e. series of squares of equal length shifted by one letter one with respect to another. For example, *bcbacacacaab* contains a maximal repetition *acacaca* which is a succession of four squares : *acac*, *caca*, *acac*, *caca*. Thus, the set of maximal repetitions can be regarded as an encoding of all tandem repeats in the string. We showed [KK99b] that this encoding is more compact in the worst case, as there are only $O(n)$ maximal repetitions in words of length n . Moreover, all of them can be found in time $O(n)$ [KK99a].

More recently, searching for repetitions in a string received a new motivation, due to the biosequence analysis [Gus97]. Successive occurrences of a fragment often bear important information in DNA sequences and their presence is characteristic for many genomic structures (such as telomer regions for example). From practical viewpoint, satellites and alu-repeats are involved in chromosome analysis and genotyping, and thus are of great interest to genomic researchers. Tools for finding successive repeats are nowadays an obligatory part of integrated systems for analyzing and annotating whole genomes [Ben99]. We refer to [vBSvL⁺97] as an example of biological study of contiguous repetitions in DNA sequences.

The major difficulty in finding biologically relevant repetitions in genomic sequences is a certain variation that must be admitted between the copies of the repeated subword. In other words, biologists are interested in *approximate repetitions* and not necessarily in exact repetitions only. The first natural definition of approximate repetition is an *approximate tandem repeat* which is a subword uv where u and v are within a given distance k and the notion of distance could be one of those usually used in biological applications, such as Hamming distance or edit distance. The problem of finding approximate tandem repeats for both these distances has been studied by G. Landau and J. Schmidt [LS93]. They showed that in case of the Hamming distance (respectively edit distance), all approximate tandem repeats can be found in time $O(nK \log(n/K) + S)$ (respectively $O(nK \log K \log n + S)$), where S is the num-

ber of repeats found. Several other approaches to finding approximate tandem repeats in DNA sequences have been proposed in bioinformatics community – some of them use statistical framework [Ben98, Ben99], some require to specify the size of repeated motif [BW94, RDDD95], some use very general framework and have to make use of some heuristic filtering steps to avoid exponential blow-up [SM98].

This paper deals with finding approximate repetitions using exact combinatorial methods of string matching. We focus on the Hamming distance case when the variability between repeated copies can be only letter replacements. An important motivation is to define structures encoding families of approximate tandem repeats, analogous to maximal repetitions in the exact case. In Section 2, we define two fundamental structures : *globally-defined approximate repetitions* and *runs of approximate tandem repeats*. In Section 3, we show that all globally-defined approximate repetitions can be found in time $O(nK \log K + S)$, where S is their number. In Section 4 we show that the same bound holds for runs of approximate tandem repeats: all of them can be found in time $O(nK \log K + R)$, where R is their number. This result implies, in particular, that all approximate tandem repeats can be found in time $O(nK \log K + T)$ (T their number), improving the $O(nK \log(n/K) + T)$ bound of G. Landau and J. Schmidt for the most interesting case of small K . Finally, in Section 5 we introduce two other possible notions of approximate repetitions and give a brief analysis of their properties.

2 K -mismatch globally-defined repetitions and runs of K -mismatch tandem repeats

Quoting [Ben98], *one difficulty in dealing with (approximate) tandem repeats is accurately defining them*. Even if we concentrate only on mismatches, as it is the case in this paper, different definitions of approximate repetitions can be thought of. Here we introduce two basic notions of approximate repetitions. Other definitions will be discussed in Section 5.

We start by recalling briefly some facts about exact repetitions. The period of a word $w[1 : n]$ is the minimal natural number p such that $w[i] = w[i + p]$ for all $1 \leq i, i + p \leq n$. The ratio n/p is called the *exponent* of w . A *repetition*

is any word with the exponent greater or equal to 2 [KK99b]. A *tandem repeat*, or a *square*, is a word which is a catenation of another word with itself. Equivalently, a tandem repeat is a repetition the exponent of which is an even natural number. In the case when the exponent is equal to 2, the tandem repeat (square) is called *primitive*. The following proposition is well-known (see [Lot83]).

Proposition 2.1 *A word $r[1 : n]$ is a repetition of period $p \leq n/2$ if and only if one of the following conditions holds:*

- (i) $r[1..n - p] = r[p + 1..n]$, and p is the minimal number with this property,
- (ii) any subword of r of length $2p$ is a tandem repeat, and p is the minimal number with this property.

When considering repetitions as subwords of a bigger word, the notion of maximality turns out to be very useful: a repetition is *maximal* iff it cannot be extended (by one letter) to the right or left while keeping the same period. Formally, given a word $w[1 : n]$ and a subword $w[i..j]$ which is a repetition of exponent $e \geq 2$, this repetition is called *maximal* if the period of both $w[i..j + 1]$ (provided that $j < n$) and $w[i - 1..j]$ (provided that $i > 1$) is strictly larger than e . For example, *acaabaababc* contains repetition (tandem repeat) *aabaab* which is not maximal, as the *a* which follows it respects the periodicity. On the other hand, *aabaaba* occurs as a maximal repetition. Maximal repetitions were studied in [Mai89, KK99b, KK99a, SG98].

We now turn to defining approximate repetitions. Similar to the exact case, the basic notion here is the approximate tandem repeat. Assume $h(\cdot, \cdot)$ is the Hamming distance between two words of equal length, that is $h(w_1, w_2)$ is the number of mismatches (letter differences at corresponding positions) between w_1 and w_2 . For example, $h(baaacb, bcabcb) = 2$.

Definition 2.2 *A word $\alpha = \alpha'\alpha''$, such that $|\alpha'| = |\alpha''|$, is called a K -mismatch tandem repeat iff $h(\alpha', \alpha'') \leq K$. Reusing the terminology of the exact case [KK99a], we call number $p = |\alpha'| = |\alpha''|$ the period of α , and words α' , α'' left and right root of α respectively.*

We now want to define a more global structure which would be able to capture “long approximate periodicities”, generalizing repetitions of arbitrary

exponent in the exact case. As opposed to the exact case, Conditions (i)-(ii) of Proposition 2.1 generalize to different notions of approximate repetition. Condition (i) gives rise to the strongest of them:

Definition 2.3 A word $r[1 : n]$ is called a K -mismatch globally-defined repetition of period p , $p \leq n/2$, iff $h(r[1..n-p], r[p+1..n]) \leq K$.

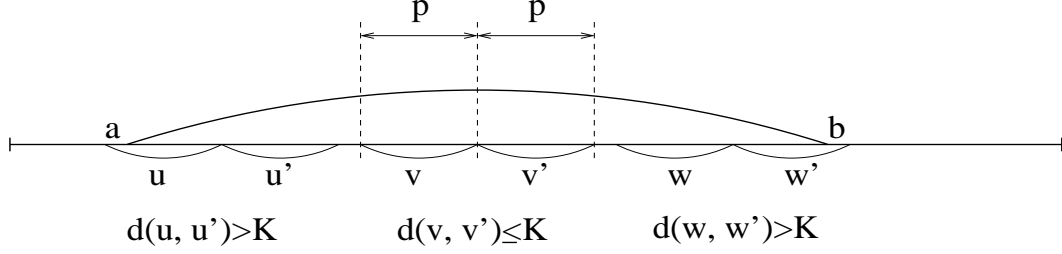
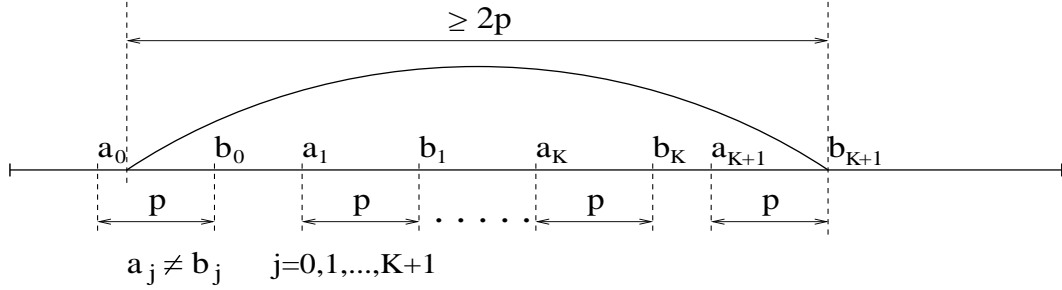
Equivalently, $r[1 : n]$ is a K -mismatch globally-defined repetition of period p , if the number of i such that $r[i] \neq r[i+p]$ is at most K . For example, *abaa abba cbba cb* is a 2-mismatch globally-defined repetition of period 4. *abc abc abc abb abc abc abc abc* is a 1-mismatch globally-defined repetition of period 3 but *abc abc abc abb abc abc abc abb* is not.

Another viewpoint, expressed by Condition (ii) of Proposition 2.1, considers a repetition as an encoding of squares it contains [IMS97, SG98]. Projecting this to the approximate case, we come up with the notion of *run of approximate tandem repeats*:

Definition 2.4 A word $r[1 : n]$ is called a run of K -mismatch tandem repeats of period p , $p \leq n/2$, iff for every $i \in [1..n-2p+1]$, subword $\alpha = r[i..i+2p-1] = r[i..i+p-1]r[i+p..i+2p-1]$ is a K -mismatch tandem repeat of period p .

Similarly to the exact case, when we are looking for approximate repetitions occurring in a word, it is natural to consider *maximal* approximate repetitions. These are repetitions extended to the right and left as far as possible provided that the corresponding definition is still verified. Note that the notion of maximality applies to both definitions of approximate repetition considered above : in both cases we can extend a repetition to the right/left as long as the obtained subword remains a repetition according to the considered definition. Throughout this paper we will be *always* interested in maximal repetitions, without mentioning it explicitly. Note that for both notions of approximate repetitions defined above, the maximality requirement implies that if $w[i : j]$ is a repetition of period p in $w[1 : n]$, then $w[j+1] \neq w[j+1-p]$ (provided $j < n$) and $w[i-1] \neq w[i-1+p]$ (provided $i > 1$)¹. Furthermore, if $w[i : j]$ is a maximal globally-defined repetition, it contains *exactly* K mismatches

¹For one type of repetitions defined in Section 5 this will not be the case, and we will add this condition explicitly.

Figure 1: Maximal run of K -mismatch tandem repeatsFigure 2: Maximal K -mismatch globally-defined repetition

$w[l] \neq w[l+p]$, $i \leq l, l+p \leq j$, unless the whole word w contains less than K mismatches (to simplify the presentation, we always exclude this latter case from consideration).

Figure 1 illustrates the definition of (maximal) run of K -mismatch tandem repeats, and Figure 2 that of (maximal) K -mismatch globally-defined repetitions.

Example 2.5 *The following Fibonacci word contains three runs of 3-mismatch tandem repeats of period 6. They are shown in regular font, in positions aligned with their occurrences. Two of them are identical, and contain each four 3-mismatch globally-defined repetitions, shown in italic for the first run only. The third run is a 3-mismatch globally-defined repetition in itself.*

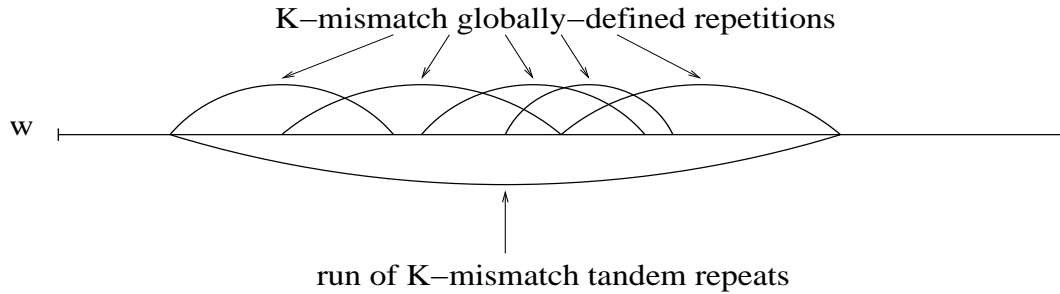


Figure 3: Extension relation

```

010010 100100 101001 010010 010100 1001
      10010 100100 101001
      10010 100100 10
      0010 100100 101
      10 100100 10100
      0 100100 101001
                1001 010010 010100 1
                  10 010100 1001

```

In general, each K -mismatch globally-defined repetition is a subword of a run of K -mismatch tandem repeats. On the other hand, a run of tandem repeats in a word is the union of all globally-defined repetitions it contains. We say then that the notion of run of K -mismatch tandem repeats *extends* that of K -mismatch globally-defined repetition (see Figure 3).

However, a run of tandem repeats may contain as many as a linear number of globally-defined repetitions. For example, the word $(000100)^n$ of length $6n$ is a run of 1-mismatch tandem repeats of period 3, which contains $(2n - 1)$ 1-mismatch globally-defined repetitions. Below is another example.

Example 2.6 *The following run of 1-mismatch tandem repeats of period 4 contains 8 1-mismatch globally-defined repetitions, shown below in positions*

aligned with their occurrences.

```

0000 1000 1100 1110 1111 1111 0111 0011 0001 0000
0000 1000 1
  000 1000 1100 11
    00 1100 1110 111
      0 1110 1111 1111
        1111 1111 0111 0
          111 0111 0011 00
            11 0011 0001 000
              1 0001 0000

```

It is easily seen that the run can be iterated and therefore this gives another example of a family of runs containing a linear number of globally-defined repetitions.

In general, the following observation holds.

Lemma 2.7 *Let $w[1 : n]$ be a run of K -mismatch tandem repeats of period p and let s be the number of mismatches $w[i] \neq w[i + p]$, $1 \leq i, i + p \leq n$ (equivalently, $s = h(w[1..n - p], w[p + 1..n])$). Then w contains $s - K + 1$ globally-defined repetitions.*

Note that both definitions can be criticized as for their relevance to practical situations. An obvious property of runs, as shown in Example 2.6, is that the repeated pattern can change completely along the run regardless the value of K . For example, *aaa aba abb abb bbb* is a run of 1-mismatch tandem repeats of period 3, although 3-letter patterns *aaa* and *bbb* have nothing in common. On the other hand, globally-defined repetitions put a global limit on the number of mismatches and therefore may not capture some repetitions that one would possibly like to consider as such, in particular repetitions of big exponent where the total number of mismatches can exceed K while the relative number of mismatches remains low. However, these two structures are of primary importance as they provide respectively the weakest and strongest notions of repetitions with K mismatches, and therefore “embrace” all practically relevant repetitions. In what follows we propose efficient algorithms to find both those types of repetitions.

3 Finding K -mismatch globally-defined repetitions

In this section we describe how to find, in a given word w , all maximal K -mismatch globally-defined repetitions occurring in w (K is a given constant). Our algorithm extends, on the one hand, the one for exact maximal repetitions [Mai89, KK99a] and on the other hand, generalizes the one of [LS93] (see also [Gus97]) by using a special factorization of the word to speed-up the algorithm.

To proceed, we need more definitions. Consider a globally-defined repetition $r = w[i..j]$ of period p in a word $w[1 : n]$. $w[i..i + p - 1]$ is called the *left root* of r and $w[j - p + 1..j]$ its *right root*. r is said to *contain* the character $w[l]$ iff $i \leq l \leq j$, and is said to *touch* $w[l]$ iff r contains $w[l]$, or contains one of characters $w[l - 1]$, $w[l + 1]$.

We assume we fixed a minimal bound p_0 for the period of repetitions we are looking for. For example, p_0 can be taken to be $K + 1$ having in mind that if a period $p \leq K$ is allowed, a tandem repeat of length $2p$ with no common characters between the left and the right root would fall into the definition. This is purely pragmatic assumption which does not affect the method nor the complexity bounds.

Our first basic technique is described by the following auxiliary problem: Given a word $w[1 : n]$ and a distinguished character $w[l]$, $l \in [2..n - 1]$, we wish to find all K -mismatch globally-defined repetitions in w which touch $w[l]$. We distinguish two disjoint classes of repetitions according to whether their right root starts to the left or to the right to $w[l]$. We concentrate on repetitions of the first class, those of the second class are found similarly.

For each $p \in [p_0..l - 1]$, and for all $k \in [0..K]$, we compute the following functions :

$$LP_k(p) = \max\{j | h(w[l - p..l - p + j - 1], w[l..l + j - 1]) \leq k\}, \quad (1)$$

$$LS_k(p) = \max\{j | h(w[l - p - j..l - p - 1], w[l - j..l - 1]) \leq k\}. \quad (2)$$

Informally, $LP_k(p)$ is the length of the longest subword in w starting at position $l - p$ and equal, within k mismatches, to a subword starting at l . Similarly, $LS_k(p)$ is the length of the longest subword ending at position $l - p - 1$ equal, within k mismatches, to a subword ending at position $l - 1$. These functions

are variants of *longest common extension functions* [LS93, Gus97] and can be computed in time $O(nK)$ using suffix trees combined with the lowest common ancestor computation in a tree. We refer to [Gus97] for a detailed description of the method.

Consider now a K -mismatch globally-defined repetition r of period p which has its right root starting to the left to $w[l]$. Note that character $w[l - p]$ is contained in r , and that r is uniquely defined by the number of mismatches $w[i - p] \neq w[i]$, $i \geq l$, contained in r . Let k be the number of those mismatches. Then

$$LP_k(p) + LS_{K-k}(p) \geq p. \quad (3)$$

Conversly, (3) can be used to detect a repetition. The following theorem holds (see [LS93, Gus97]), which is a generalization of the corresponding result of [ML84, Mai89].

Theorem 3.1 *Let $w[1 : n]$ be a word and $w[l]$, $1 < l < n$, a distinguished character. There exists a K -mismatch globally-defined repetition of period p which contains $w[l]$, and has its right period starting to the left to $w[l]$, iff for some $k \in [0..K]$,*

$$LP_k(p) \leq p, \quad (4)$$

and inequation (3) holds. In this case, this repetition starts at position $l - p - LS_{K-k}(p)$ and ends at position $l + LP_k(p) - 1$.

Inequation 4 ensures that the right root starts to the left of $w[l]$.

Theorem 3.1 provides an $O(nK)$ algorithm for finding all considered globally-defined repetitions: compute longest extension function (1) (2) (this takes time $O(nK)$) and then check inequations (3), (4) for all $p \in [p_0..l - 1]$ and all $k \in [0..K]$ (this takes time $O(nK)$ too). Each time the inequations are verified, a new repetition is identified. Finding repetitions with the right root starting to the right to $w[l]$ is a symmetric problem, which is solved within the same time bound.

The algorithm solving the auxiliary problem described above will be referred to as Algorithm 1. Its pseudo-code is shown below.

The second important tool is Lempel-Ziv factorization used in the well-known compression method. Let w be a word and assume that the last symbol of w does not occur elsewhere. In this paper, we need two variants of

Algorithm 1 Computing all K -mismatch globally-defined repetitions repetitions in w touching a distinguished character

Input: word $w[1 : n]$, position l , $1 < l < n$

Output: all K -mismatch globally-defined repetitions in w which touch $w[l]$
 {Find those repetitions which have their right root starting to the left to $w[l]$ }

```

1: for all  $p \in [p_0..l - 1]$ ,  $k \in [0..K]$ , compute longest common extension
   functions  $LP_k(p)$ ,  $LS_k(p)$  defined as in (1), (2)
2: for  $p = p_0$  to  $\min\{n - l + 1, n/2\}$  do
3:   for  $k = 0$  to  $K$  do
4:     if  $LP_k(p) + LS_{K-k}(p) \geq p$  and  $LP_k(p) \leq p$  then
5:       output a  $K$ -mismatch globally-defined repetition starting at posi-
         tion  $l - p - LS_{K-k}(p)$  and ending at position  $l + LP_k(p) - 1$ 
   {Similarly, find those repetitions which have their right root starting to
    the right to  $w[l]$ }
```

the Lempel-Ziv factorization, that we call *with copy overlap* and *without copy overlap*².

Definition 3.2 *The Lempel-Ziv factorization of w with copy overlap (respectively without copy overlap) is the factorization $w = f_1 f_2 \dots f_m$, where f_i 's are defined inductively as follows:*

- $f_1 = w[1]$,
- for $i \geq 2$, f_i is the shortest word occurring in w immediately after $f_1 f_2 \dots f_{i-1}$ which does not occur in $f_1 f_2 \dots f_i$ other than in prefix (respectively, does not occur in $f_1 f_2 \dots f_{i-1}$).

As an example, the Lempel-Ziv factorization with copy overlap of the word $aabbabababbcb$ is $a|ab|ba|bababb|bc$; the factorization without copy overlap is

²The s -factorization used in [Mai89, KK99a] is a minor modification of the Lempel-Ziv factorization with copy overlap. The difference is that the s -factorization considers the longest factor occurring earlier, while the Lempel-Ziv factorization considers the shortest factor which does not occur earlier (see [Gus97] for a related discussion). In this paper, we use the Lempel-Ziv factorization which suits better to our purposes.

$a|ab|ba|bab|abbb|c$. Both variants of Lempel-Ziv factorization can be computed in linear time [RPE81, Gus97]. If $w = f_1 f_2 \dots f_m$ is the Lempel-Ziv factorization, we call f_i 's *Lempel-Ziv factors* or simply *factors* of w . The last character of f_i will be called the *head* of f_i .

We are now ready to describe the algorithm for finding all K -match globally-defined repetitions. Consider the Lempel-Ziv factorization of w *with copy overlap*. The algorithm consists of three stages. The key to the **first stage** is the following lemmas.

Lemma 3.3 *The right root of a K -mismatch globally-defined repetition cannot contain as subword $K + 1$ consecutive Lempel-Ziv factors.*

Proof: Each factor contained in the right root contains a character mismatching the one located one period to the left. Indeed, if it does not contain a mismatch, it has an exact copy occurring earlier, which contradicts to the definition of factorization. As the right root contains at most K mismatches, it cannot contain $K + 1$ or more factors. \square

We divide w into consecutive *blocks* of $K + 2$ Lempel-Ziv factors. Let $w = B_1 \dots B_m$ be the partition of w into such blocks. The last character of B_i will be called the *head character* of this block. At the first stage, we find, for each block B_i , those repetitions which touch the head character of B_i but does not touch that of B_{i+1} . First, concentrate on those of such repetitions with the right root starting before the head character of B_i .

Lemma 3.4 *Assume a K -mismatch globally-defined repetition r touches the head character of B_i but not that of B_{i+1} . Then $|r| < 2|B_i B_{i+1}|$.*

Proof: Lemma 3.3 implies that the right root of r cannot start before the first character of B_i . Therefore, the period of r is bounded by $|B_i B_{i+1}|$. On the other hand, by the argument of the proof of Lemma 3.3, r cannot extend by more than a period to the left of B_i . Therefore, the total length of r is bounded by $2|B_i B_{i+1}|$. \square

Lemma 3.4 allows us to apply Algorithm 1 : Consider the word $w_i = v B_i B_{i+1}$, where v is the suffix of $B_1 \dots B_{i-1}$ of length $|B_i B_{i+1}|$. Then find, using Algorithm 1, all repetitions in w_i touching the head character of B_i and discard

those which touch the head character of B_{i+1} . The resulting complexity is $O(K(|B_i| + |B_{i+1}|))$.

After processing all blocks, we find all repetitions touching block head characters. Observe that repetitions resulting from processing different blocks are distinct. Summing up over all blocks, the resulting complexity of the first stage is $O(nK)$. The repetitions which remain to be found are those which lie entirely within a block – this is done at the next two stages.

At the **second stage** we find all repetitions inside each block B_i which touch factor heads other than the block head (=last character of the block). For each B_i , we proceed by simple binary division approach:

- (i) divide current block of factors $B = f_i f_{i+1} \dots f_{i+m}$ into two sub-blocks $B' = f_i \dots f_{\lfloor m/2 \rfloor}$ and $B'' = f_{\lfloor m/2 \rfloor + 1} \dots f_{i+m}$,
- (ii) using Algorithm 1, find the repetitions in B which touch the head character of $f_{\lfloor m/2 \rfloor}$, but discard those which touch the head character of f_{i+m} or contain the first character of f_i ,
- (iii) process recursively B' and B'' .

The above algorithm has $\lceil \log K \rceil$ levels of recursion, and since at each step the word is split into disjoint sub-blocks, the whole complexity of the second stage is $O(nK \log K)$.

Finally, at the **third stage**, it remains to find the repetitions which occur entirely inside each Lempel-Ziv factor, namely which don't contain its first character and don't touch its head character. By definition of factorization with copy overlap (Definition 3.2), each factor without its head character has another (possibly overlapping) occurrence to the left. Therefore, each of these repetitions has another occurrence to the left too. Using this observation, these repetitions can be found using the same technique as the one of [KK99a]: When constructing the Lempel-Ziv factorization we keep for each factor w a pointer to a copy of w to the left. Then processing factors from left to right, recover repetitions inside the factor from its pointed copy. We refer to [KK99a] for algorithmic details. The complexity of this stage is $O(n + S)$, where S is the number of repetitions found.

The following theorem summarizes this section.

Theorem 3.5 *All K -mismatch globally-defined repetitions can be found in time $O(nK \log K + S)$ where n is the word length and S is the number of repetitions found.*

The algorithm of finding all K -mismatch globally-defined repetitions, referred to as Algorithm 2, is given below.

Algorithm 2 Computing all K -mismatch globally-defined repetitions in w

Input: word $w[1 : n]$

Output: all K -mismatch globally-defined repetitions in w

- 1: Compute the Lempel-Ziv factorization with copy overlap $w = f_1 \dots f_m$
 - 2: Partition the factorization into blocks of $K + 2$ consecutive factors; let $w = B_1 \dots B_{m'}$ be the decomposition of w into such blocks
 - {first stage}
 - 3: **for** each block B_i **do**
 - 4: find, using Lemma 3.4 and Algorithm 1, globally-defined repetitions which touch the head character of B_i but not that of B_{i+1}
 - {second stage}
 - 5: **for** each block B_i **do**
 - 6: starting from B_i apply the following recursive procedure
 - 7: divide the current block $B = f_i f_{i+1} \dots f_{i+m}$ into two sub-blocks $B' = f_i \dots f_{\lfloor m/2 \rfloor}$ and $B'' = f_{\lfloor m/2 \rfloor + 1} \dots f_{i+m}$
 - 8: find, using Algorithm 1, globally-defined repetitions in B which touch the last character of $f_{\lfloor m/2 \rfloor}$, but discard those which touch the head character of f_{i+m} or contain the first character of f_i
 - 9: process recursively B' and B''
 - {third stage}
 - 10: **for** each Lempel-Ziv factor f_j **do**
 - 11: retrieve all globally-defined repetitions in f_j which don't contain its first character and don't touch its last character, from its left copy (see [KK99a] for details)
-

4 Finding runs of K -mismatch tandem repeats

In this section we describe an algorithm for finding all runs of K -mismatch tandem repeats in a word.

The general structure of the algorithm is the same as for globally-defined repetitions (Algorithm 2) – it has the three stages playing similar roles. At the first and second stages, the key difference is the type of objects we are looking for : instead of computing globally-defined repetitions we now compute *subruns* of K -mismatch tandem repeats. Formally, a subrun is a run of K -mismatch tandem repeats, which is not necessarily maximal. At each point of the first and second stage when we search for repetitions touching some head character $w[l]$, we now compute subruns of those K -mismatch tandem repeats which touch $w[l]$. This can be seen as outputting by Algorithm 1 only the part of the globally-defined repetition falling to the interval $l - 2p..l + 2p$. The modified Algorithm 1, referred to as Algorithm 3, is given below.

Algorithm 3 Computing subruns of K -mismatch tandem repeats in w touching a distinguished character

Input: word $w[1 : n]$, position l , $1 < l < n$

Output: all subruns of K -mismatch tandem repeats in w which touch $w[l]$
 {Find those tandem repeats which have their right root starting to the left to $w[l]$ }

- 1: for all $p \in [p_0..l - 1]$, $k \in [0..K]$, compute longest common extension functions $LP_k(p)$, $LS_k(p)$ defined as in (1), (2)
- 2: **for** $p = p_0$ to $\min\{n - l + 1, n/2\}$ **do**
- 3: **for** $k = 0$ to K **do**
- 4: **if** $LP_k(p) + LS_{K-k}(p) \geq p$ and $LP_k(p) \leq p$ **then**
- 5: create a subrun of K -mismatch tandem repeats ending at positions $start(p, k) = \max\{l + p - LS_{K-k}(p) - 1, l - 1\}$ through $end(p, k) = \min\{l + LP_k(p) - 1, l + p - 1\}$
- 6: **if** $k > 0$ and $end(p, k) \leq end(p, k - 1) + 1$ **then**
- 7: merge this subrun with the subrun computed for the previous value of k

{Similarly, find those tandem repeats which have their right root starting to the right to $w[l]$ and thus end in the interval $l + p..l + 2p$ }

The major additional difficulty in computing runs is *assembling* subruns into runs. To perform the assembling, we need to store subruns in an additional data structure and to carefully manage merging of subruns into bigger runs. We have to ensure that the number of subruns we come up with and the work spent on processing them do not increase the resulting complexity bound.

The assembling occurs already at the level of Algorithm 3, as subruns found for different values of k (for-loop at line 3) may overlap or immediately follow each other, in which case we join them into a bigger subrun (lines 6-7). Similarly, subruns of tandem repeats with the right root starting to the right to $w[l]$ (case non-shown in Algorithm 3) may have to be joined with subruns found by instructions 1-7 of Algorithm 3). We leave out the details of how this is done.

Below we describe the three stages of the algorithm in more details. We identify a subrun with the interval of *end positions* of the tandem repeats it contains.

For the input word w , we compute the Lempel-Ziv factorization *without copy overlap* and divide it into blocks $B_1 \dots B_{m'}$, each containing $K + 2$ consecutive Lempel-Ziv factors. **At the first stage**, we find subruns of all those tandem repeats which touch block head characters. For each block B_i , we find the tandem repeats which touch the head character of B_i but not that of B_{i+1} . Let l_i be the position of the head character of B_i . Then the subruns of period p , found at this step, belong to the interval $[l_i - 1, \min\{l_i + 2p, l_{i+1} - 2\}]$. We call this interval the *explored interval* for $w[l_i]$ and p . The subruns found at this step can be seen as subintervals of this explored interval. These subruns are stored into a double-linked list in increasing order of positions. (We leave it to the reader to check out that such a list can be easily computed by Algorithm 3 by making at each step a constant amount of extra work.) If the explored interval for $w[l_{i-1}]$ and p ends at position $l_i - 2$, it is merged with the explored interval for $w[l_i]$, thus forming a bigger explored interval. Accordingly, the lists of subruns associated with these intervals are merged into a single list. All additional operations take constant time, and the resulting complexity of the first stage is $O(nK)$.

The second stage is modified in a similar way. Recall that at each call of Algorithm 3 we are searching for repetitions occurring between some factor head, say $w[l']$, another factor head $w[l'']$, and touching some factor head $w[l]$

($l' < l < l''$). Assuming that recursive calls are executed in preorder (see the description of the second stage in the previous section), no factor head between $w[l']$ and $w[l'']$ has been processed yet. In this case, the explored interval is $[\max\{l' + 2p + 1, l - 1\}.. \min\{l + 2p, l'' - 2\}]$, and we may have to merge it either with the previous explored interval, or with the next one, or both. The complexity of the second stage stays $O(nK \log K)$.

After accomplishing the first and second stages, we have, *for each period* p , a set of non-intersecting explored intervals. Each interval is associated with a sequence of successive head characters $w[l_i], w[l_{i+1}], \dots, w[l_m]$ such that $l_{j+1} - l_j \leq 2p + 2$ for $j \in [i..m - 1]$, and the interval itself is $[l_i - 1..l_m + 2p]$. In particular, the interval is associated to $w[l_i]$ and $w[l_m]$ - the first and the last head characters of this sequence. Those subruns of tandem repeats which have been actually found within this interval, are stored in a double-linked list associated to the interval.

At **the third stage**, we have to find subruns of those tandem repeats which lie entirely inside Lempel-Ziv factors. For each period, potential occurrences of these subruns correspond precisely to the gaps between explored intervals. Thus, the third stage can be also seen as closing up the gaps between explored intervals for this period.

As in the previous section, the key observation here is the fact that Lempel-Ziv factors without their head character have a copy to the left (here required to be non-overlapping), and the idea is again to process w from left to right and to retrieve the subruns inside each factor from its copy. However, the situation here is different in comparison to globally-defined repetitions: we may have to “cut out” a chain of subruns belonging to the factor copy from a longer list and then to “fit” it into the gap between two explored intervals. The “cutting out” may entail splitting subruns which span over the borders of the factor copy, and “fitting into” may entail merging those subruns with subruns from the neighboring explored intervals. Below we sketch the algorithm for the third stage, which copes with these difficulties. Algorithm 4 given below provides a detailed description of the third stage.

During the computation of the Lempel-Ziv factorization, for each Lempel-Ziv factor $f_i = va$ we choose a copy of v occurring earlier and point from the end position of this copy to the head character a of f_i . It may happen that one position has to have several pointers, in which case we organize them in

Algorithm 4 Third stage of the algorithm of finding runs of K -mismatch tandem repeats

Input: word $w[1 : n]$; lists of subruns found at the first and second stages

Output: all runs of K -mismatch tandem repeats in w

$\{activerun(p)\}$ will be maintained to be the last considered run of period p
 $\{startingruns(i)\}$ will be maintained to be the list of runs starting at i

```

1: for each position  $i \in [1..n]$  do
2:   for each run  $r$   $startingruns(i)$  do
3:     let  $p$  be the period of  $r$ 
4:      $activerun(p) := r$ 
5:     if  $r$  is not the last in its list then
6:       let  $i'$  be the first position of the next run in the list
7:       add the next run to  $startingruns(i')$ 
8:   for each factor copy ending at position  $i$  do
9:     let  $w[j..i]$  be this copy and  $f_m$  the corresponding factor
10:    for each period  $p \leq (j - i + 1)/2$  do
11:      if  $activerun(p)$  contains tandem repeats inside  $w[j..i]$  then
12:        link/merge this subrun of tandem repeats to/with the first subrun
        of the explored interval associated with the head symbol of  $f_m$ 
13:         $currentrun :=$  the predecessor of  $activerun(p)$  in its list
14:        while  $currentrun$  contains tandem repeats inside  $w[j..i]$  do
15:          link the subrun of those tandem repeats to the previously
          copied subrun in  $f_m$ 
16:        link/merge the last processed subrun to/with the last subrun of
        the explored interval associated with the head symbol of  $f_{m-1}$ 
17:      else
18:        link the last subrun of the explored interval associated with the
        head symbol of  $f_{m-1}$  with the first subrun of the explored interval
        associated with the head symbol of  $f_m$ 
19:      if  $activerun(p)$  is the last subrun of the explored interval associated
        with the head symbol of  $f_{m-1}$  then
20:        let  $i'$  be the first position of the next run in the list
21:        add the next run (if any) to  $startingruns(i')$ 
22:        close up the gap corresponding to  $f_m$  by merging intervals associated
        to the head symbol of  $f_m$  and the head symbol of  $f_{m-1}$ 

```

a list. We traverse w from left to right and maintain for the current position the last runs (of all possible periods) which start before this character. To this purpose, we also maintain the following invariant: at the moment we arrive at a position, we know the list of all subruns which start at this position. This information is collected according to the following general rule: for each subrun starting at the current position, we assign the starting position of the next subrun in the list (instructions 2-7 in Algorithm 4). Of course, there may be no next subrun if the current subrun is the last one in the explored interval. In this case, the starting position of the subrun following the current subrun will be set at the moment we fill the gap after this explored interval (instructions 19-21).

When we arrive at the end position of a copy of a Lempel-Ziv factor, we need to copy into the factor all the subruns which this copy contains. Therefore, we scan *backwards* the subruns contained in the copy and copy them to the factor (instructions 11-18). Copying the subruns closes up two explored intervals into one interval, and links together two lists of subruns, possibly inserting a new list of runs in between. Copying subruns in the backward direction is important for the correction of the algorithm – this guarantees that no subruns are missed. It is also for these reason that we need the copy to be non-overlapping with the factor.

After the whole word has been traversed, no more gaps between explored intervals exist anymore. This means that for each period, we have a list of subruns with this period occurring in the word, which are actually the searched runs. The complexity of the third stage is $O(n + S)$, where S is the number of resulting runs. Putting together the three stages, we obtain the main result of this section.

Theorem 4.1 *All runs of K -mismatch tandem repeats can be found in time $O(nK \log K + S)$ where n is the word length and S is the number of runs found.*

Once all runs have been found, we can easily output all tandem repeats. We then have the following result improving the result of [LS93].

Corollary 4.2 *All K -mismatch tandem repeats can be found in time $O(nK \log K + S)$ where n is the word length and S is the number of tandem repeats found.*

5 Other definitions of approximate repetitions

K -mismatch globally-defined repetitions limit by K the *total* number of mismatches, and therefore provide the strongest notion of approximate repetition. On the other hand, runs of K -mismatch tandem repeats provide the weakest notion of repetition with K mismatches, as they impose only the minimal requirement that every tandem repeat in a such repetition contains no more than K mismatches. For practical applications, such as genome analysis, it might be interesting to consider intermediate definitions with respect to the two “extreme” cases. In this section we introduce two such types of repetitions and point out very briefly some of their properties. A more detailed analysis, together with illustrating examples, is to appear in the extended version of this paper.

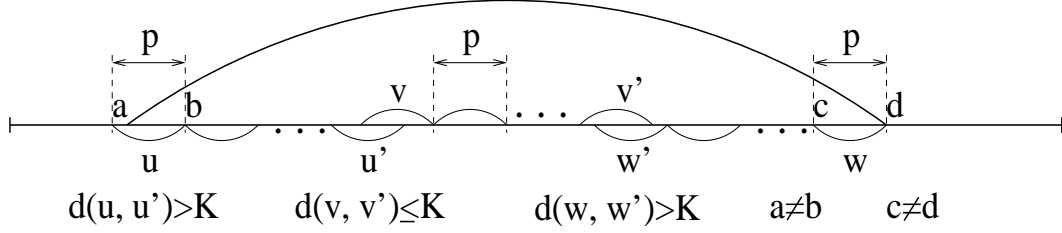
One natural way to loosen the definition of globally-defined repetitions is to limit by K the Hamming distance between two subwords of length p or less (p period), located within any distance which is a multiple of p .

Definition 5.1 *A word $r[1 : n]$ is called a K -mismatch uniform repetition of period p , $p \leq n/2$ iff for every two subwords $r[i..i+j-1]$, $r[i+kp..i+kp+j-1]$ of r such that k is any integer and $1 \leq j \leq p$, we have $h(r[i..i+j-1], r[i+kp..i+kp+j-1]) \leq K$.*

For example, *cbcabbbcaab* is a 1-mismatch uniform repetition of period 4, but *abcaabcaabc* is not since *ab* at position 1 is at Hamming distance 2 from *bc* at position 9. In general, K -mismatch uniform repetitions is a weaker notion than K -mismatch globally-defined repetitions: any word from $(abcadc)^+$ is a 1-mismatch uniform repetition of period 3, whereas none of these words is a 1-mismatch globally-defined repetition.

The following technical remark concerning uniform repetitions is important. When we consider *maximal* uniform repetition in a word we have to add an additional condition: $w[i..j]$ is a maximal K -mismatch uniform repetition of period p in w if neither $w[i-1..j]$ nor $w[i..j+1]$ is a K -mismatch uniform repetition of period p and the following inequalities hold:

$$w[i-1] \neq w[i-1+p], \quad w[j+1-p] \neq w[j+1] \quad (5)$$

Figure 4: K -mismatch uniform repetition

In contrast to both notions of repetition considered before, inequalities (5) don't hold automatically for uniform repetitions and should be added explicitly in order to ensure, in particular, that every K -mismatch uniform repetition in a word is the union of K -mismatch globally-defined repetitions containing in it. The following example illustrates the situation.

Example 5.2 Consider the following word of length $4p$. It contains two maximal K -mismatch globally-defined repetitions of period p starting at position 1 and $p+1$ and shown below in regular font. Both of these repetitions are also maximal K -mismatch uniform repetitions. However, if we don't require inequalities (5) to hold, we obtain a series of $K-1$ “superfluous” K -mismatch uniform repetitions shown in *italic*.

$$\begin{array}{ccccccc}
 \overbrace{0 \dots 00 \dots 0}^p & \overbrace{0 \dots 0 \dots 01}^{p-1} & \overbrace{0 \dots 01 \dots 1}^{p-K} & \overbrace{0 \dots 01 \dots 1}^K & \overbrace{0 \dots 01 \dots 111}^{p-K-1} & \overbrace{0 \dots 01 \dots 111}^{K+1} & \\
 0 \dots 00 \dots 0 & 0 \dots 0 \dots 01 & 0 \dots 01 \dots 1 & 0 \dots 01 \dots 1 & 0 \dots 0 & & \\
 & \overbrace{0 \dots 0}^{K-1} & 0 \dots 0 \dots 01 & 0 \dots 01 \dots 1 & 0 \dots 01 & & \\
 & & & \dots & & & \\
 & & & & & \overbrace{0 \dots 01 \dots 1}^{K-1} & \\
 & & & & & 0 \dots 01 \dots 1 & 0 \dots 01 \dots 111
 \end{array}$$

Figure 4 gives an illustration to the definition of maximal K -mismatch uniform repetition. The relationship of uniform repetitions to globally-defined repetitions and runs are summarized in the following lemma.

Lemma 5.3 (i) *Any K -mismatch globally-defined repetition can be extended to a (possibly not unique) K -mismatch uniform repetition. Any K -mismatch uniform repetition is the union of K -mismatch globally-defined repetitions it contains.*

(ii) *Any K -mismatch uniform repetition can be extended to a unique run of K -mismatch tandem repeats. A run of K -mismatch tandem repeats is the union of K -mismatch uniform repetition it contains.*

Another possible definition is obtained when one thinks about an approximate repetition as an “exact repetition with no more than K replacement errors per period”. This viewpoint is somewhat similar to the one of [SM98], where a repetition is defined through a *consensus* such that each repeated motif is within a specified distance from the consensus.

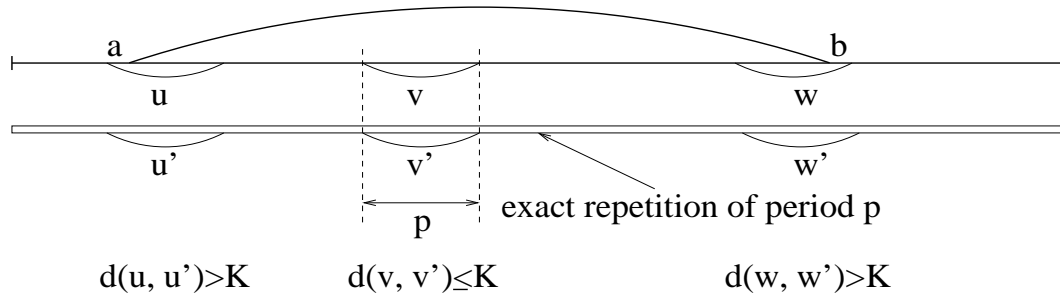
Definition 5.4 *Word $r[1 : n]$ is a K -mismatch consensus repetition of period p , $p \leq n/2$, iff there exists an exact repetition $v[1 : n]$ of period p such that for any subword $r[i..j]$ of r such that $j - i \leq p$, we have $h(r[i..j], v[i..j]) \leq K$.*

Example 5.5 *Consider the word from Example 2.6. The 1-mismatch consensus repetitions it contains are shown together with a possible consensus for each of them shown below in italic.*

```

0000 1000 1100 1110 1111 1111 0111 0011 0001 0000
0000 1000 1100 11
1000 1000 1000 10
000 1000 1100 1110 111
100 1100 1100 1100 110
00 1100 1110 1111 1111
10 1110 1110 1110 1110
0 1110 1111 1111 0111 0
1 1111 1111 1111 1111 1
1111 1111 0111 0011 00
0111 0111 0111 0111 01
111 0111 0011 0001 000
011 0011 0011 0011 001
11 0011 0001 0000
01 0001 0001 0001

```

Figure 5: K -mismatch consensus repetition

The notion of K -mismatch consensus repetition is illustrated on Figure 5. Similar to uniform repetitions, consensus repetitions provide an intermediate structure between globally-defined repetitions and runs:

Lemma 5.6 *Assume K is even.*

- (i) *Any K -mismatch globally-defined repetition can be extended to a (possibly not unique) $K/2$ -mismatch consensus repetition. Any $K/2$ -mismatch consensus repetition is the union of K -mismatch globally-defined repetitions it contains.*
- (ii) *Any $K/2$ -mismatch consensus repetition can be extended to a unique run of K -mismatch tandem repeats. A run of K -mismatch tandem repeats is the union of $K/2$ -mismatch consensus repetition it contains.*

Concerning the relationship between uniform and consensus repetitions, it is easily seen that a $K/2$ -mismatch consensus repetition can be extended to a K -mismatch uniform repetition. Any K -mismatch uniform repetition can be, in turn, extended to a K -mismatch consensus repetition. More subtle relationship between these two notion require an additional analysis. Designing an efficient algorithm for finding these types of repetitions remains also an open question.

Relations between different notions of repetition, studied in this paper, is summarized on Figure 6. A solid arrow denotes the extension relation (see Figure 3 and remark after Example 2.5), and a flashed arrow means just that

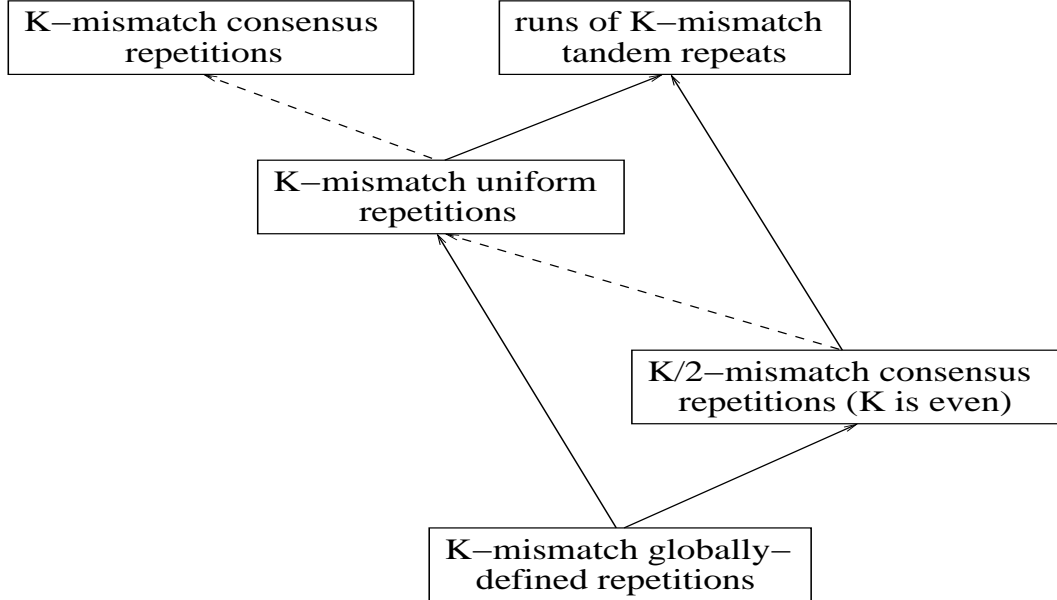


Figure 6: Relations between different notions of repetitions

the “source structure” can be extended to a “target structure”, but not necessarily that a target structure is the union of the source structures contained in it.

6 Concluding remarks

We proposed $O(nK \log K + S)$ algorithms for finding K -mismatch globally-defined repetitions and runs of K -mismatch tandem repeats (S the output size). Note that if K is considered constant, we have $O(n + S)$ algorithms for finding each of these structures. This is an interesting result, which has been long time unknown even for the exact case.

The algorithms presented in this paper are now being implemented within the `mreps` software³. Currently, `mreps` implements the algorithm of finding

³<http://www.loria.fr/~kucherov/SOFTWARE/MREPS/index.html>

exact maximal repetitions [KK99a]. Some interesting experiments have been done by applying `mreps` to genomic sequences [GK00].

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