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Indexing Sequences for Mapping Reads with a Single Mismatch

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Abstract. To be written.

1 Introduction

In the classical string matching problem we are given a text of length $n$ and a pattern of length $m$ and we need to answer the query whether the pattern exists in the text (existence query) as a factor or substring and further to provide the start (or equivalently end) positions of the corresponding occurrences (occurrence query). Using the famous KMP algorithm, due to Knuth, Morris and Pratt [32], the classical string matching problem can be solved in optimal $O(n + m)$ time. For a review on KMP algorithm as well as various other (exact) string matching algorithms the readers are referred to [10].

In many application settings, a number of patterns are searched in a particular text which gives rise to the indexing version of the problem. In this version, the text is given beforehand for preprocessing with a goal to construct an index data structure. Subsequently, a number of patterns are queried against the text. Clearly, this indexing variant can also be solved using KMP algorithm; however, if we have $k$ patterns (each of length $m$), the running time will be $O(k(n + m))$. Since all the queries are done against a single text, it is only natural to construct an index data structure for it once, preferably in $O(n)$ time and then answer the subsequent queries in $O(m)$ time each, which gives a total of $O(km + n)$ running time. And, indeed there exist efficient data structures (e.g., suffix trees [45, 38, 17, 46] and suffix arrays [33, 31, 30, 2]) that can be constructed in $O(n)$ time and are capable of answering an existence query in $O(m)$ time and an occurrence query in $O(m + |Occ|)$ time, where Occ is the set of output.

The string matching problem has tremendous application in different branches of science including, but not limited to, Computational Molecular Biology, Bioinformatics, Computer Vision, Information retrieval, Computational Musicology, Data Mining, Network Security etc. However, in many, if not most, practical settings, instead of the exact matching some approximate matching schemes become more relevant and useful. The requirement of such approximate matching schemes is more prominent in Computational Molecular Biology and Bioinformatics as discussed below. Notably, in the context of Computational Biology, the string matching problem is essential for mapping reads (i.e., patterns) to a reference biological sequence (i.e., a text).
While most of the biology labs are using dedicated high-throughput equipments to produce large DNA sequences on a daily basis, the need for automatic annotation and content analysis is greater everyday. Unfortunately, even with the tremendous advancement of the current state of the art technology, the quality of the automatically obtained sequences is sometimes questionable. The reason for this can be attributed to, among others, the intrinsic limitations of the equipments and the natural polymorphism that can be observed between individual samples (e.g., a Simple Nucleotide Polymorphism, that is a unique mutation, can either stop the translation of a mRNA into a protein sequence, or create a binding site for a protein complex that will prevent the complete formation of the functional protein [9, 34]). Analysing such uncertain sequences is therefore much more complicated than the traditional problem. This gives the computer scientists, in particular, the stringology researchers the challenge to solve the string matching problem where in the given text or pattern some positions may be uncertain in some sense. To capture this phenomenon of uncertainty, the idea of mismatches and of gaps was introduced. Additionally, a popular and useful framework of don’t care pattern matching was introduced under this approximate matching scheme, where a pattern and/or text may contain don’t care characters that match with any character in the underlying alphabet.

The use of don’t care paradigm to capture the approximate pattern matching scenario mentioned above is not new. It was introduced and solved efficiently using convolutional methods in [20]. Slightly tighter solutions have been presented in [11, 13, 28, 29]. There exist some solutions avoiding the convolution method as well [40, 39]. A number of solutions exists in the literature that consider the problem of text indexing with don’t cares [12, 35, 43, 44]. Notably, in the literature, the don’t cares are also referred to as wildcards. Some work has also been done on a generalised model of the don’t care paradigm known as the degenerate or indeterminate string model in the literature [41, 25, 42]; in this model, some positions of a string may contain more than one letters and don’t care is essentially modelled as a position that contain all the letters of the alphabet. Another popular approximate model in the literature is where some $k \geq 0$ mismatches are allowed while doing the pattern matching. There exists a number of results on this problem [16, 6, 7, 3, 23, 22, 36]. However, from indexing point of view the results are only few [5, 12]. Notably, the $k$ mismatch model can be represented/captured in the don’t care model by assuming $k$ don’t care characters at all possible permutations of the positions in the pattern (and/or the text).

In this paper, we focus on the indexed version of the pattern matching problem with restricted number of mismatches and/or gaps. In particular, we are interested in the pattern matching problem when at most one mismatch or gap is allowed. Here gap refers to consecutive mismatches. To the best of our knowledge the only work that deals with this problem directly is the work of Amir et al. [5]. We will give a brief review of the work of [5] and compare their results with ours in a subsequent section.

The contribution of this current paper is twofold. First we attack a restricted version of the problem in hand where we assume that the size of the pattern
is fixed, i.e., we are given the size of the pattern to be queried against our data structure beforehand. As discussed below, this particular version of the problem has strong motivation from Computational Biology and Bioinformatics, especially in the context of the Next Generation Sequencing. In the sequel we consider the general version of the problem where this restriction is lifted. The solution we propose makes use of suffix arrays and range search data structures borrowed from Computational Geometry literature. In particular, in order to present an efficient data structure for our problem, we utilise a reduction from our problem at hand to the range search problem in geometry. As will be further discussed in later sections, similar reduction has also be employed in [5] to solve this particular problem and in different other papers (e.g., [15, 27, 26]) to solve some other interesting problems. Note however that the reduction itself is not enough to get a good solution. As will be clear later, we need to do some non-trivial work based on some useful observations and lemmas to achieve an efficient running time for the queries.

The motivation of our work comes both from the field of Stringology as well as from that of Computational Biology and Bioinformatics. Firstly, a solution to the approximate pattern matching with a single mismatch would be useful as a theoretical advancement in the context of the general variant where \( k \) mismatches are considered. Secondly, approximate pattern matching problem with at most a single mismatch is a useful problem on its own right. This is specially true in Computational Molecular Biology, where with the advent of new state of the art technologies, the chance of experimental mistakes has become much lower than it was before. Also, the existence of close mismatches called SNPs is rare. Thirdly, because of the recent high throughput sequencing technologies we are particularly interested to provide a fast solution to the restricted version of the problem where the pattern size is fixed. In the so-called Next Generation Sequencing (NGS) technologies, millions of short reads are generated. Usually, the first region of these reads, called the seeds, are almost error-free. The seed region is followed by a region having very small possibilities of error and hence from mapping point of view, only a single mismatch or a gap (i.e., consecutive mismatches) are expected. Now, in NGS, the generated sequences, which are usually several millions in number, are of the same size; this size depends on the technologies used (e.g., Illumina etc.). As a result, the fixed-pattern length version of the problem is of particular interest in this particular scenario.

The rest of the paper is organised as follows. After the definitions and problem setting in Section 2, we give in Section 3 the general idea of the solution. It serves as a guideline for the development of algorithms. Section 4 solves a simpler problem, while Section 5 describes a solution for the complete problem. Experiments? and conclusion follow.

2 Preliminaries

A string is a sequence of zero or more symbols from an alphabet \( \Sigma \). A string \( \mathcal{T} \) of length \( n \) is denoted by \( \mathcal{T}[1 \ldots n] = T_1T_2\ldots T_n \), where \( T_i \in \Sigma \) for \( 1 \leq i \leq n \).
The length of $\mathcal{T}$ is denoted by $|\mathcal{T}| = n$. The string $\bar{T}$ denotes the reverse of the string $T$, i.e., $\bar{T} = T_n T_{n-1} \ldots T_1$.

A string $w$ is a factor of $\mathcal{T}$ if $\mathcal{T} = uwv$ for $u, v \in \Sigma^*$; in this case, the string $w$ occurs at position $|u| + 1$ in $\mathcal{T}$. The factor $w$ is denoted by $|\mathcal{T}|u + 1 \ldots |u| + |w|$. A $k$-factor is a factor of length $k$. A prefix (or suffix) of $\mathcal{T}$ is a factor $\mathcal{T}[x \ldots y]$ such that $x = 1$ ($y = n$), $1 \leq y \leq n$ ($1 \leq x \leq n$). We define the $i$th prefix to be the prefix ending at position $i$ i.e. $\mathcal{T}[i \ldots i]$, $1 \leq i \leq n$. Dually, the $i$th suffix is the suffix starting at position $i$ i.e. $\mathcal{T}[i \ldots n]$, $1 \leq i \leq n$.

The Hamming distance between two strings of equal length is the number of positions at which the corresponding letters are different. More formally, the Hamming distance between $u, v \in \Sigma^*$ is said to match $v$ (or equivalently $v$ is said to match $u$) with at most $k$ mismatches if $\delta(u, v) \leq k$. Essentially, the exact match is characterised by a zero Hamming distance.

Given a text $\mathcal{T}$ of length $n$ and a pattern $\mathcal{P}$ of length $m$ such that $m \leq n$, $\mathcal{P}$ is said to occur in $\mathcal{T}$ at position $i$ (i.e., exact match) if and only if $\mathcal{P} = \mathcal{T}[i \ldots i + m - 1]$. The position $i$ is said to be an occurrence of $\mathcal{P}$ in $\mathcal{T}$. We denote by Occ$^{\mathcal{T}}_{\mathcal{P}}$ the set of occurrences of $\mathcal{P}$ in $\mathcal{T}$. As an extension, $\mathcal{P}$ is said to occur in $\mathcal{T}$ at position $i$ with at most $k$ mismatches if and only if $\delta(\mathcal{P}, \mathcal{T}[i \ldots i + m - 1]) \leq k$. We use Occ$^{\mathcal{T}}_{\mathcal{P}, \leq k}$ to denote the set of positions where $\mathcal{P}$ matches $\mathcal{T}$ with at most $k$ mismatches. Similarly, we use Occ$^{\mathcal{T}}_{\mathcal{P}, = k}$ to denote the set of positions where $\mathcal{P}$ matches $\mathcal{T}$ with exactly $k$ mismatches. Clearly, our interest is in calculating Occ$^{\mathcal{T}}_{\mathcal{P}, \leq 1}$.

In traditional full-text indexing problems one of the basic data structures used is the suffix array data structure. Others are suffix trees and suffix automata. In our indexing problem we make use of this suffix array data structure. A complete description of a suffix array is beyond the scope of this paper, and can be found in [33, 31, 30, 2] or in any textbook on stringology (e.g., [14, 24]). Here we give a very concise definition of suffix array. The suffix array SA$^\mathcal{T}[1 \ldots n]$ of a text $\mathcal{T}$ is an array of integers $j \in \{1 \ldots n\}$ such that SA$^\mathcal{T}[i] = j$ if, and only if, $\mathcal{T}[j \ldots n]$ is the $i$-th suffix of $\mathcal{T}$ in (ascending) lexicographic order. Suffix arrays were first introduced in [37], where an $O(n \log n)$ construction algorithm and $O(m + \log n + |\mathrm{Occ}^\mathcal{T}_{\mathcal{P}}|)$ query time were presented. Later, linear time construction algorithms for space efficient suffix arrays were presented [33, 31, 30]. The query time is also improved to optimal $O(m + |\mathrm{Occ}^\mathcal{T}_{\mathcal{P}}|)$ in [1] with the help of another array essentially storing the lengths of longest common prefixes between lexicographically consecutive suffixes. We recall that, the result of a query for a pattern $\mathcal{P}$ on a suffix array SA$^\mathcal{T}$ of $\mathcal{T}$, is given in the form of an interval $[s \ldots e]$ such that Occ$^{\mathcal{T}}_{\mathcal{P}} = \{SA^\mathcal{T}[s], SA^\mathcal{T}[s + 1], \ldots, SA^\mathcal{T}[e]\}$. In this case, the interval $[s \ldots e]$ is denoted by Int$^{\mathcal{T}}_{\mathcal{P}}$.

3 The Underlying Idea for a Solution

In this section, we discuss how we can efficiently compute Occ$^{\mathcal{T}}_{\mathcal{P}, \leq 1}$, given $\mathcal{T}$ and $\mathcal{P}$. In other words, we will in fact discuss a solution of our problem albeit not
from a indexing point of view. In the sequel we will be using the underlying idea to construct the index data structures of our interest. Notably, the same idea was employed in [5] to provide an indexing solution for the problem. We will discuss the problems of the solution in [5] and highlight the differences between their solution and the solution presented in this paper in a later section.

Suppose we know the position of the mismatch and assume that the position is \( j \). In other words, we suppose that the mismatch is only allowed with \( P_j \). Let us use \( \text{Occ}\_P^\leq_j \) to denote the corresponding set of occurrences. Then, we may proceed as follows. We compute \( \text{Occ}\_P^\leq_1 \) and \( \text{Occ}\_P^\leq_{j+1} \). Then, clearly, our desired set of occurrences \( \text{Occ}\_P^\leq_j \) can be computed as follows:

\[
\text{Occ}\_P^\leq_j = \{ i \mid i \in \text{Occ}\_P^\leq_1 \text{ and } (i + j) \in \text{Occ}\_P^\leq_{j+1} \}.
\]

Now to lift the restriction we can simply run a loop on all possible values of \( j \). This simple idea works perfectly and the steps are formally presented in Algorithm 1. However, transforming this idea to handle the indexing version is a non-trivial task.

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**Algorithm 1** Computing \( \text{Occ}\_P^\leq_1 \)

1: Set \( \text{Occ}\_P^\leq_1 = \emptyset \)
2: for \( j \in [1..m] \) do
3: \( \text{Compute } \text{Occ}\_P^\leq_1 \)
4: for \( i \in \text{Occ}\_P^\leq_1 \) do
5: \( i = i + j \)
6: end for
7: \( \text{Compute } \text{Occ}\_P^\leq_{j+1} \)
8: Set \( \text{Occ}\_P^\leq_j = \text{Occ}\_P^\leq_1 \cap \text{Occ}\_P^\leq_{j+1} \)
9: Set \( \text{Occ}\_P^\leq_1 = \text{Occ}\_P^\leq_1 \cup \text{Occ}\_P^\leq_j \)
10: end for
11: return \( \text{Occ}\_P^\leq_1 \)

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**4 The Index Data Structure**

In this section, we focus on constructing an index data structure for our problem. We however first consider a restricted variant of the original problem. In particular, for the time being, we assume that we are given the pattern-length \( m \) which we use during the index data structure construction. In other words, the index constructed will be \( m \)-specific, i.e., it will be able to handle patterns of length \( m \) only. Recall from Section 1 that this particular variant of the problem is of particular interest especially in the context of Next Generation Sequencing. The general version of the problem will be handled in a later section.
We basically extend the idea of Algorithm 1. We maintain two suffix array data structures $SA_T$ and $SA_{←T}$. We use $SA_T$ to find the occurrences of $P[1..j-1]$. We can find the occurrences of $P[j+1..m]$ using $SA_{←T}$ as well. But we need to take a different approach because we have to “align” the occurrences of $P[1..j-1]$ (Step 4) with the occurrences of $P[j+1..m]$ so that we can compute $Occ_{P[j] \leq 1}$ by intersecting (Step 8) them just as is done in Algorithm 1. However it is not as straightforward as Algorithm 1 because our aim is to maintain an index rather than finding a match for a particular pattern. We use the following trick, which has been applied to solve this and different other problems in the literature before (see for example [5, 15, 27]).

Our approach is as follows. We use the suffix array of the reverse string of $T$, i.e. $SA_{←T}$, to find the occurrences of $P[j+1..m]$. By doing so, in fact, we get the end positions of the occurrences of $P[j+1..m]$ in $T$. However we still have to do a bit more “shifting” for the intersection to work because from $SA_{←T}$, we get the start positions of the occurrences of $P[1..j-1]$. This is where the information of a fixed pattern-length, i.e., $m$ comes handy. To achieve the “shifting” effect automatically, we appropriately rename the entries of $SA_{←T}$ as follows. For all $1 \leq i \leq n$, if originally $SA_{←T}[i] = j$, we assign, $SA_{←T}[i] = n - (j + m - 1) + 1 = n - j - m - 2$. It is easy to see that this will effectively transform the position of each of the occurrences of $P[j+1..m]$ to the appropriate position that will facilitate the intersection.

Now, it remains to show how we can perform the intersection (Step 8) efficiently in the context of indexing. Clearly we now have two arrays, namely, $SA_T[1..n]$ and $SA_{ ←T}[1..n]$. And we also have two intervals, namely, $Int_T[1..j-1]$ and $Int_{←T}[j+1..m]$. Of course, the second interval will give us slightly different positions based on our requirement. And, now our problem is to find the intersection of the positions within these two intervals. This problem is a known problem in geometry and is called the Range Set Intersection Problem.

**Problem 1.** [Problem RSI] Let $V[1..n]$ and $W[1..n]$ be two permutations of $[1..n]$. Preprocess $V$ and $W$ to answer the following form of queries.

Query: Find the intersection of the elements of $V[i..j]$ and $W[k..\ell]$, $1 \leq i \leq j \leq n, 1 \leq k \leq \ell \leq n$.

In order to solve the above problem we reduce it to the well-studied Range Search Problem on a Grid.

**Problem 2.** [Problem RSG] Let $A$ be a set of $n$ points on the grid $[0..U] \times [0..U]$. Preprocess $A$ to answer the following form of queries.

Query: Given a query rectangle $q \equiv (a,b) \times (c,d)$ find the set of points of $A$ contained in $q$.

We can see that Problem RSI is just a different formulation of the Problem RSG. This can be realised as follows. We set $U = n$. Since $V$ and $W$ in Problem RSI are permutations of $[1..n]$, every number in $[1..n]$ appears precisely once in each of them. We define the coordinates of every number $i \in [1..n]$ to be $(x,y)$,
where \( V[x] = W[y] = i \). Thus we get the \( n \) points on the grid \([0 \ldots n] \times [0 \ldots n] \), i.e., the array \( A \) of Problem RSG. The query rectangle \( q \) is deduced from the two intervals \([i \ldots j] \) and \([k \ldots \ell] \) as follows: \( q \equiv (i, k) \times (j, \ell) \). It is easy to verify that the above reduction is correct and hence we can solve Problem RSI using the solution of Problem RSG. So, this completes our description for constructing the index data structure for our problem. Algorithm 2 formally states the steps to build our data structure.

**Algorithm 2** Algorithm to build the index data structure for the fixed-length pattern case

1: Build a suffix array \( SA_T \) of \( T \).
2: Build a suffix array \( SA_{\overleftarrow{T}} \) of \( \overleftarrow{T} \).
3: for \( i = 1 \) to \( n \) do
4:   Set \( j = SA_{\overleftarrow{T}}[i] \)
5:   Set \( SA_{\overleftarrow{T}}[i] = n - j - m - 2 \)
6: end for
7: for \( i = 1 \) to \( n \) do
8:   Set \( A[i] = \epsilon \)
9: end for
10: for \( i = 1 \) to \( n \) do
11:   if there exists \((x, y)\) such that \( SA_T[x] = SA_{\overleftarrow{T}}[y] = i \) then
12:     \( A[i] = (x, y) \)
13:   end if
14: end for
15: Preprocess \( A \) for Range Search on the grid \([0 \ldots n] \times [0 \ldots n] \).

4.1 Analysis

Let us analyse the running time of Algorithm 2, i.e., the data structure construction. The computational effort spent for Steps 1, 2 and 3 is \( O(n) \). In Step 10, we construct the set \( A \) of points in the grid \([0 \ldots n] \times [0 \ldots n] \) on which we will apply the range search. This step can also be done in \( O(n) \) as follows. We construct \( SA_{\overleftarrow{T}}^{-1} \) such that \( SA_{\overleftarrow{T}}^{-1}[SA_T[i]] = i \). Similarly, we can construct \( SA_T^{-1} \). Then, it is easy to construct \( A \) in \( O(n) \). A detail is that in our case there may exist \( i, 1 \leq i \leq n \) such that \( SA_{\overleftarrow{T}}^{-1}[i] = i \) for all \( 1 \leq j \leq n \). This is because \( SA_{\overleftarrow{T}}^{-1} \) is a permutation of \([-m + 2 \ldots n - m + 1] \) instead of \([1 \ldots n] \). Now, it is easy to observe that any \( i \in SA_{\overleftarrow{T}}^{-1} \) such that \( i > n \) or \( i < 1 \) is irrelevant in the context of our search. So we ignore any such \( i \in SA_{\overleftarrow{T}}^{-1} \) while creating the set \( A \). After \( A \) is constructed we perform Step 15. There has been significant research work on Problem RSG. For example, we can use the data structure of Alstrup et al. [4]. This data structure can answer the query of Problem RSG in \( O(\log \log n + k) \) time where \( k \) is the number of points contained in the query rectangle \( q \). The data structure requires
We simply compute \( \text{Occ}_T \) is the set of those points. Then it is easy to verify that \( \text{Occ}_T \) have changed the entries of all intervals first and store them with a little book-keeping to implement Step 5 for the points in \( A \).

Note carefully that here we slightly abuse the notation \( \text{Int}_T \) to facilitate the desired shift. Now, we find all the points in \( A \) that are inside the rectangle \( q \equiv (s_1^\ell, s_r^\ell) \times (e_1^\ell, e_r^\ell) \). Let \( B_j^\ell \) is the set of those points. Then it is easy to verify that \( \text{Occ}_T \) of 1 is \( B_j^\ell \).

Now, we can easily lift the restriction that \( j \) is the position of the mismatch. We simply, compute \( \text{Occ}_T \) for all values of \( j \) and compute \( \text{Occ}_T \) = \( \bigcup_{j=1}^{m-1} B_j^\ell \). The steps are formally presented in the form of Algorithm 3.

### Algorithm 3 Algorithm for Query Processing

1. \( B = \emptyset \)
2. for \( j = 1 \) to \( m-1 \) do
   3. Compute \( \text{Int}^T_{[1..j-1]} \equiv [s_1^\ell \ldots e_\ell^\ell] \) using \( \text{SA}_T \).
   4. Compute \( \text{Int}^T_{[j+1..m]} \equiv [s_r^\ell \ldots e_r^\ell] \) using \( \text{SA}_T \).
   5. Set \( B_j^\ell = \{(x, y) \mid (x, y) \in A \text{ and } (x, y) \text{ is contained in } q \equiv (s_1^\ell, s_r^\ell) \times (e_1^\ell, e_r^\ell)\} \)
   6. \( B = B \cup B_j^\ell \)
3. end for
8. return \( \text{Occ}_T \) of 1 = \( B \)

The running time of the query processing is deduced as follows. Step 3 and Step 4 can be done in \( O(m) \) time. And Step 5, i.e., range search query can be done in \( O(\log \log n + K) \) time, where \( K \) is the number of points in the query rectangle. Now, there is a loop at Step 2. So, a straightforward analysis of Algorithm 3 leads us to a running time of \( O(m^2 + m \log \log n + K) \). Here assume that \( K \) is the size of the output returned by the algorithm. We will focus on \( K \) shortly. However, we can do far better than \( O(m^2 + m \log \log n + K) \) as follows. Note that we can compute \( \text{Int}^T_{[1..j-1]} \) incrementally as we increment \( j \) from 1 to \( m-1 \). This means we can get all the intervals from \( \text{SA}_T \), namely, \( [s_1^\ell \ldots e_\ell^\ell], 1 \leq j < m \) spending \( O(m) \) time in total. Similarly we can get all the intervals from \( \text{SA}_T \), namely, \( [s_r^\ell \ldots e_r^\ell], 1 \leq j < m \) spending \( O(m) \) time in total. So, we can compute all the intervals first and store them with a little book-keeping to implement Step 5 for all \( j \), \( 1 \leq j \leq m - 1 \), afterwards. This will give a much better running time of \( O(m \log \log n + K) \).
4.3 Discussion on $\mathcal{K}$

Finally, a brief discussion on the value of $\mathcal{K}$ is in order. Since we are computing the occurrences with at least 1 mismatch, the occurrences of exact matches will also be reported. And in our algorithm when we compute $\text{Occ}_T^{|m| \leq 1}$ for a particular value of $j$, it will also contain the exact occurrences. In other words, for all values of $j$, we have $\text{Occ}_T^j \subseteq \text{Occ}_T^{|m| \leq 1}$. So, every exact occurrence will be reported $m-1$ times. To make the value of $\mathcal{K}$ optimal we need to employ some further tricks. First we need the following easy lemma.

**Lemma 1.** Suppose we are given a suffix array $SA_T[1 \ldots n]$ of a text $T[1 \ldots n]$ and a pattern $P[1 \ldots m]$. Suppose $\text{Int}_T[P[1 \ldots j-1]] \equiv [s] \ldots [e]$ and $\text{Int}_T[P] \equiv [s \ldots e]$. If $1 \leq j < m$, we must have $s|P \leq s \leq e|P$.

Clearly, Lemma 1 tells us that the range identifying the occurrences of a pattern must be contained in or equal to the range identifying the occurrences of a prefix of that pattern. How do we use the lemma to ensure an optimal value of $\mathcal{K}$? We modify our query algorithm as follows. We only need to modify the part that uses $SA_T$ and do some more work while we compute $B_j$. At the beginning (before the loop at Step 2 of Algorithm 3), we compute $\text{Int}_T[P] \equiv [s \ldots e]$ using $SA_T$. Now, suppose that we have computed $\text{Int}_T[P[1 \ldots j-1]] \equiv [s|P \ldots e|P]$ using $SA_T$. By Lemma 1, we know that $s|P \leq s \leq e|P$. So, we divide the range into (at most) two ranges $[s|P \ldots s-1]$ and $[s+1 \ldots e|P]$. Then the computation of $B_j$ is modified as follows: $B|P = \{(x, y) \mid (x, y) \in A \text{ and } (x, y) \text{ is contained in } q_1 \equiv (s|P, s|P) \times (s-1, e|P) \text{ or in } q_2 \equiv (e+1, e|P) \times (e|P, e|P)\}$. In other words, what we are doing is that the interval of each of the prefixes $P[1 \ldots j], 1 \leq j < m$, of $P$ is divided into at most two sub-intervals so as to remove the exact occurrences of $P$. Hence finally, we need to include the exact occurrences of $P$ before returning the set. In other words, instead of returning $B = \bigcup_{i=1}^{\ell} B|P$, we need to return $B \cup \text{Occ}_T^P$. Clearly, now the output size $\mathcal{K}$ will be optimal.

Finally, what will be the query time of the augmented query algorithm? Clearly, now for each of the prefixes $P[1 \ldots j], 1 \leq j \leq m-1$, we would have at most two ranges (against one range of $P[j+1 \ldots m], 1 \leq j \leq m-1$). So, we need to make at most $2m$ range search queries in total keeping the total query time asymptotically the same as before.

4.4 Exactly one mismatch

In some practical application, especially in Bioinformatics, the occurrences with exactly one mismatch is sought. In other words, in such cases, the exact occurrences are to be excluded, i.e., we are to compute $\text{Occ}_T^{|m| = 1}$ instead of $\text{Occ}_T^{|m| \leq 1}$. Clearly this can be achieved without any change in the query time. In this case, we simply need to return $B = \bigcup_{i=1}^{\ell} B|P$ in the end in our augmented algorithm (without including $\text{Occ}_T^P$).
5 General Case

In this section we relax the assumption that the pattern length $m$ is given as part of the input. So, we cannot take advantage of using $m$ during the data structure construction as we did in Step 3 of Algorithm 2. For this case, we use the same idea but the shifting need be done in a different way so that we do not require the knowledge of $m$. This in the sequel requires a different query processing algorithm as will be discussed shortly. Below, we first discuss the index construction and then focus on to the query processing algorithm.

5.1 Index Construction

As has been mentioned previously, we will use the same basic idea and hence will depend on Algorithm 1. Let us suppose that we know the position of the mismatch and assume that the position is $j$. Similar to the previous approach we will maintain two suffix arrays $SA_T$ and $SA_{\overline{T}}$. However, we now reverse the role of the two suffix arrays as follows. We use $SA_T$ to find the occurrences of $P[j+1..m]$ (rather than $P[1..j−1]$ as we did previously). We use the suffix array of the reverse string of $T$, i.e. $SA_{\overline{T}}$, to find the occurrences of $\overline{T}[1..j−1]$.

By doing so, in fact, we get the end positions of the occurrences of $P[1..j−1]$ in $T$. Now, note that we have the end positions of $P[j+1..m]$ in $T$ and the start positions of $P[1..j−1]$ in $T$. So, to get the desired alignment all we need to do is to consider the mismatch position and relabel accordingly. Since now the alignment is done at the start position of $P[j+1..m]$ (instead of the start position of $P[1..j−1]$), we do not need the knowledge of $m$. It is easy to verify that, now, each $i \in SA_{\overline{T}}$ needs to be relabelled as $(n+1)−i+1+1 = n−i+3$ to get the desired alignment. The rest of the steps are identical to Algorithm 2. Clearly, the running time of the index construction remains the same as before.

5.2 Query Processing

The query processing algorithm is built on the same principle as before. Let us suppose that we know the position of the mismatch and assume that the position is $j$. Then the query can be answered as follows. We first compute $Int_{P[j+1..m]} = [s_r|j..e_r|j]$ using $SA_T$. Then we compute $Int_{P[1..j−1]} = [s_l|j..e_l|j]$ using $SA_{\overline{T}}$. Like before, here we slightly abuse the notation $Int_{P[1..j−1]}$ because we have changed the entries of $SA_{\overline{T}}$ to facilitate the desired shift. Now, we find all the points in $A$ that are inside the rectangle $q \equiv (s_l|j..e_l|j) \times (s_r|j..e_r|j)$.

Let $B|j$ is the set of those points. Now, note that we have done the alignment at the start position of $P[j+1..m]$. So, we need to undo this shift to get the actual start position of the desired occurrence of the pattern $P$. So we compute, $B'|j = \{(SA_T[x]−1−(j−1)) \mid (x, y) \in B|j\}$. It is easy to verify that $Occ_{P}|_{\leq 1} = B'|j$. 

Now, we can easily lift the restriction that $j$ is the position of the mismatch. We simply, compute $\text{Occ}^T_{P|j \leq 1}$, i.e., $B^\prime|j$ and $B^\prime'|j$ for all values of $j$ and compute $\text{Occ}^T_{P|j \leq 1} = \bigcup_{j=1}^{m-1} B^\prime|j$.

5.3 Analysis of the Query Processing Algorithm

The analyses of the two algorithms, namely, the query processing algorithm presented in Section 4.2 and that presented in the previous section (Section 5.2) are similar but with a distinct difference that makes the running time of the latter worse. For the query processing of the fixed $m$ case, we computed $\text{Int}^T_{P[1..j-1]}$ incrementally as we increment $j$ from 1 to $m$ and got all the intervals from $SA_T$, namely, $[s_d|j \ldots e_d|j], 1 \leq j < m$ spending $O(m)$ time in total. Similarly we computed all the intervals from $SA_{\leftarrow P}$, namely, $[s_r|j \ldots e_r|j], 1 \leq j < m$ spending $O(m)$ time in total. For the general case unfortunately, we cannot apply the above trick readily. This is because, now, we need to compute $\text{Int}^T_{P[j+1..m]}$ incrementally as we increment $j$ from 1 to $m-1$. This means that, unlike the previous case, where we were looking for intervals for prefixes of increasing sizes, now we have to compute intervals for suffixes of decreasing sizes. The same applies for the computation of intervals for $P[1..j-1], 1 \leq j \leq m-1$ using $SA_T$. As a result the query time becomes $O(m^2 + m \log \log n + K)$.

5.4 Improved Query Time

Clearly the bottleneck is the computation of the appropriate intervals for $P[j+1..m]$ and $P[1..j-1]$ for different values of $j$. To achieve the same query time of $O(m \log \log n + K)$ we need to be able to compute the intervals for the suffixes of decreasing sizes incrementally as we did for the prefixes of increasing sizes. This would ensure a $O(m)$ running time for the computation of the appropriate intervals and thereby keeping the desired running time intact. To achieve this improvement we resort to the famous backward search technique of Ferragina and Manzini [18] using their data structure popularly known as the FM-Index. In particular, we use the following result which is the heart of the Backward Search Algorithm (BSA) of [18] using an FM-index.

**Lemma 2.** [Ferragina, Manzini [18]] Assume $[s..e] = \text{Int}^T_P$ is already computed. Then for any character $c$, the interval $[s'..e'] = \text{Int}^T_{P|c}$ can be computed in $O(1)$ time.

It is clear that with Lemma 2 at our disposal, we can compute the appropriate intervals for suffixes of decreasing sizes of a pattern $P$ spending $O(|P|)$ time in total. So, using BSA of [18, 19] we are now able to compute the appropriate intervals for $P[j+1..m]$ and $P[1..j-1]$ for different values of $j$ spending a total time of $O(m)$. Therefore, the query time improves to $O(m \log \log n + K)$, where $K$ is the output size.
5.5 Data Structure Construction Time with FM-Index

In the previous section we have used BSA (Lemma 2) to achieve the desired running time for the query. However, this means that we would need to include the FM-index in our index data structure. Therefore, a further analysis of the data structure construction time is in order. The FM-Index and hence the BSA, make clever use of the so-called Rank query. Suppose we have a string $X$, $c \in \Sigma$ and $f$ is an integer. Then, $\text{Rank}_X(c, f)$ is defined to be the number of occurrences of $c$ (i.e., rank) in the prefix $X[1..f]$. FM-Index uses a wavelet tree [21] to facilitate constant time Rank queries. It also requires an auxiliary array $C$ such that $C[c]$ stores the total number of occurrences of all $c' \leq c$, where ‘$\leq$’ here means lexicographically smaller than or equal to. Finally, FM-Index and BSA utilise the famous Burrows-Wheeler transformation (BWT) technique [8]. A complete description of BWT is out of the scope of and not required for our discussion. We just give a brief definition of BWT in relation to suffix array as follows. The BWT encoding of the string $T$ is another string $B_T$ as defined below. Assume that the $i$th element of the suffix array of $T$ is $SA_T[i]$. Then $B_T[i] = T[SA_T[i] - 1]$ where $T[0] = T[n]$. The computation of $B_T$ can be done in $O(n)$ time. FM-Index needs to preprocess $B_T$ for constant-time rank queries. Since wavelet trees can be constructed in linear time, this can also be achieved in $O(n)$ time. The auxiliary array $C$ can also be prepared in $O(n)$ time. So overall the data structure construction time remains dominated by the range search data structure construction.

5.6 Optimality of $\mathcal{K}$

The problem with the optimality of $\mathcal{K}$ as discussed in Section 4.3 applies for the current algorithm as well. Unfortunately, The method employed in Section 4.3 to make $\mathcal{K}$ optimal cannot be used now. The technique employed in Section 4.3 was based on Lemma 1, which basically states that the interval provided by a suffix array for a pattern always lies within the interval of any of its prefixes. But this relation does not hold for suffixes. Therefore, to compute $\mathcal{K}$ we cannot use Lemma 1. Nevertheless, we apply another technique to achieve our goal as described below. We need the following lemma.

Lemma 3. Suppose we are given a suffix array $SA_T[1..n]$ of a text $T[1..n]$ and a pattern $P[1..m]$. Suppose $\text{Int}_{P[1..m]}^{-} = [s_1..e_1]$ and $\text{Int}_{P[1..m]}^{+} = [s_0..e_0]$, where $1 \leq j < m$. Then the followings hold true:

1. The size of the interval $\text{Int}_{P[j+1..m]}^{-}$ is greater than or equal to the size of the interval $\text{Int}_{P[j..m]}^{+}$, i.e., $e_1 - s_1 + 1 \geq e_0 - s_0 + 1$.
2. We have $SA_T[s_0] + 1 \equiv SA_T[p]$ for some $s_1 \leq p \leq e_1$.
3. Suppose, $SA_T[s_0] + 1 = SA_T[p]$. Then, $SA_T[s_0 + i] + 1 = SA_T[p + i]$ for all $1 \leq i \leq e_1 - s_1$.

Now let us discuss how we can obtain the optimal $\mathcal{K}$ as follows. The basic idea is similar as before. We want to divide an interval into at most two subintervals
such that the subinterval responsible for the exact occurrences of the complete pattern can be excluded when we do the intersection. Now, suppose we have computed \( \text{Int}_T^{P[j+1\ldots m]} \) and \( \text{Int}_T^{P[j\ldots m]} \). By Lemmas 3(2) and 3(3) we know that the occurrences of \( P[j+1\ldots m] \) in some sense 'contain' the occurrences of \( [j\ldots m] \). This is because \( P[j\ldots m] = P[j+1\ldots m] \).

Now, let us go back to our computation assuming that we know the position of the mismatch and assume that the position is \( j \). So, we need to do the intersection between \( \text{Int}_T^{P[j+1\ldots m]} \equiv [s_r^j \ldots e_r^j] \) (computed using \( SA_T \)) and \( \text{Int}_T^{P[j\ldots j-1]} \equiv [s_{i[l]} \ldots e_{i[l]}] \) (computed using \( SA_T \)). Note carefully that, here \( P[j] \) corresponds to the mismatch position. So, if from the interval \( \text{Int}_T^{P[j+1\ldots m]} \) we can remove the positions which follows the occurrence of \( P[j] \) in \( T \) we are done. Assuming that we have \( \text{Int}_T^{P[j\ldots m]} \) at our hand, we can easily identify those positions using Lemma 3 as follows. Following the hypothesis of Lemma 3, let us assume that \( \text{Int}_T^{P[j+1\ldots m]} \equiv [s_1 \ldots e_1] \) and \( \text{Int}_T^{P[j\ldots m]} \equiv [s_0 \ldots e_0] \). By Lemma 3(2), we have an \( s_1 \leq p \leq e_1 \) such that \( SA_T[s_1^j] + 1 = SA_T[p] \). We identify this \( p \). How can we identify \( p \) efficiently? To do this efficiently, we slightly augment our index data structure by maintaining an auxiliary array \( \text{Next}[1\ldots n] \) of length \( n \) as follows. We store \( \text{Next}[i] = j \) if and only if \( SA_T[i] + 1 = SA_T[j] \). Clearly this will facilitate \( O(1) \) time identification of the index \( p \). Also, note that the computation of \( \text{Next}[1\ldots n] \) can be done in linear time during the index data structure construction.

Once \( p \) is identified, we compute \( q = p + e_0 - s_0 \). Now, we have two subintervals, namely, \([s_1 \ldots p - 1] \) and \([q + 1 \ldots e_1] \). It is not very difficult to realise that these two intervals, namely, \([s_1 \ldots p - 1] \) and \([q + 1 \ldots e_1] \) give us the positions where \( P[j+1\ldots m] \) occurs in \( T \) but is not preceded by an occurrence of \( P[j] \); this is exactly what we desired. So, now we finish off by modifying the computation of \( B_j \) appropriately. To show the modification of the computation of \( B_j \) we need to keep the notational convention followed so far. Note that the position of \( P[j+1\ldots m] \) is to the right with respect to the fixed mismatch position \( j \). So, to keep the notational symmetry we will now rename \( s_1 \) and \( e_1 \) as \( s_r^j \) and \( e_r^j \) respectively. Now the modified computation of \( B_j \) is shown below.

\[
B_j^\beta = \{ (x, y) \mid (x, y) \in A \text{ and } (x, y) \text{ is contained in } q_1 \equiv (s_r^j, x_l^j) \times (e_r^j, p - 1) \text{ or in } q_1 \equiv (s_r^j, q + 1) \times (e_r^j, e_r^j) \}
\]

Finally, we need to include the exact occurrences of \( P \) before returning the final output set. In other words, instead of returning \( B = \bigcup_{i=1}^n B_i^\beta \), we need to return \( B \cup \text{Occ}_P^T \). Clearly, now the output size \( K \) will be optimal. By similar argument as presented in Section 4.3, the query time remains asymptotically the same, i.e., \( O(m \log \log n + K) \).

6 Discussion on the solution of [5]

As has been mentioned above, the underlying idea used in our solution approach is identical to that of the solution proposed by Amir et al. [5]. In this section we present a detailed discussion between the two solutions and identify the shortcomings of the solution of [5]. In what follows, we will refer to the data structure
of [5] as $DS_{AKLLLR}$ (using the first letters of the authors surnames). And we will use $DS_{Fixed}$ and $DS_{Gen}$ to refer to our data structure for the fixed $m$ version and the general version, respectively. In both cases, we will use the names to refer to the corresponding algorithms as well.

$DS_{AKLLLR}$ consists of two suffix trees, one for $T$ and one for $\overline{T}$. The role of these two suffix trees is exactly the same as the role of the two suffix arrays in our data structure $DS_{Fixed}$. Recall that in $DS_{Gen}$ the roles of the two suffix arrays (FM-Index, to be precise) are in fact reversed. The range search data structure used by $DS_{AKLLLR}$ is similar but not identical to the one used by $DS_{Fixed}$ and $DS_{Gen}$, as will be clear shortly. So, overall, the data structure construction is almost similar in both algorithms. The query algorithm however is drastically different as discussed below.

$DS_{AKLLLR}$ query algorithm requires that the suffix trees are constructed using the Weiner construction [46]. According to Weiner’s algorithm, given the text $T$ of length $n$, the suffix $T[n \ldots n]$ is first considered, followed by $T[n-1 \ldots n]$, then $T[n-2 \ldots n]$ and so on. Now similar to $DS_{Fixed}$, $DS_{AKLLLR}$ needs to compute $Int_{P[1 \ldots j-1]} \equiv [s_{\ell}^j \ldots e_{\ell}^j]$ using $ST_T$ and $Int_{P[j+1 \ldots m]} \equiv [s_r^j \ldots e_r^j]$ using $ST_{\overline{T}}$. So with the suffix tree $ST_T$ of $T$ at its disposal, it proceeds as follows. $DS_{AKLLLR}$ ‘feeds’ the pattern $P$ to the suffix tree in some sense. In particular, it continue from $ST_T$ and builds a suffix tree of $P_1P_2 \ldots P_m\#T$, where $\# \notin \Sigma$. While doing this extended construction, $DS_{AKLLLR}$ cleverly keeps track of the locus positions for each suffix of $P$. And hence they can easily get the range $Int_{P[1 \ldots j-1]} \equiv [s_{\ell}^j \ldots e_{\ell}^j]$. After the construction ends, they ‘undo’ this to keep the suffix tree $ST_T$ as before. Identical operation on $ST_{\overline{T}}$ gives the range $Int_{P[j+1 \ldots m]} \equiv [s_r^j \ldots e_r^j]$. Subsequently, all is needed is the application of an appropriate range search query on the range search data structure which is a part of $DS_{AKLLLR}$.

Now, there are two important points that need be carefully noted as highlighted below.

- First, the discussion, in Section 4.3 and later again in Section 5.6, on the output size $K$ applies to $DS_{AKLLLR}$ as well. To make $K$ optimal $DS_{AKLLLR}$ employs a different technique than ours. In particular, it resorts to a higher dimensional range search query. We omit the details because this is not relevant. But the point is that, this technique required the data structure of 3D range search and 3D query. This makes both the data structure construction time and query time (slightly) inferior to the time of ours.

- Secondly, and more importantly, the claim that the computation of $Int_{P[1 \ldots j-1]} (Int_{P[j+1 \ldots m]}$) by ‘feeding’ $P$ ($\overline{P}$) to $ST_T$ ($ST_{\overline{T}}$) requires asymptotically $O(m)$ is somewhat flawed as follows. The linear time of the Weiner construction of suffix tree (and in fact all other linear time construction, e.g., [45]) depends on an amortised analysis. Hence, while we can certainly say that the construction of a suffix tree for the string $P\#T$ can be done in $O(|P| + |T|)$ time, given a suffix tree for $T$ we cannot always claim that extending it for $P\#T$ can be done in $O(|P|)$ time.
7 Experimental Results

To be written.

8 Conclusion

To be written.

References


