# **An Efficient Index Structure for String Databases** \*

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# **Abstract**

We consider the problem of substring searching in large databases. Typical applications of this problem are genetic data, web data, and event sequences. Since the size of such databases grows exponentially, it becomes impractical to use inmemory algorithms for these problems. In this paper, we propose to map the substrings of the data into an integer space with the help of wavelet coefficients. Later, we index these coefficients using MBRs (Minimum Bounding Rectangles). We define a distance function which is a lower bound to the actual edit distance between strings. We experiment with both nearest neighbor queries and range queries. The results show that our technique prunes significant amount of the database (typically 50-95%), thus reducing both the disk I/O cost and the CPU cost significantly.

# 1 Introduction

String data naturally arises in many real world applications like genetic data, web data and event sequences. There is a frequent need to find similarities between such data sequences. For example, the similarity of two DNA strings from different organisms may correspond to some functional or physical relationship between these organisms. Such similarities may be used to predict diseases, or to design new drugs. Significant breakthroughs have already been achieved in genome research using the analysis of similar genetic strings. Identification of the genetic code of the deadly E.coli bacteria, or genetic clues for fibrodysplasia ossificans progressiva (FOP), a disease that affects muscle and skeleton growth, and the vital proteins for the bone growth, or identification of the genes that hasten the healing

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of some venous ulcers are only a few of the achievements obtained recently.

Another application of substring searching is the identification of similar patterns in large text databases while allowing some amount of typographical errors. This application includes searching a word in a dictionary, or a phrase in a large collection of text. Spell checkers and web searchers are some specific examples of such applications.

Video data can be viewed as an event sequence if some prespecified set of events are detected and stored as a sequence. These events can be voices, faces, objects, or text. Video databases support a wide variety of applications including security cameras, interviews, documentaries, movies, and TV news. Searching similar event subsequences can be used to find related video segments. Some companies like CNN, ABC, CNET, and AltaVista are already encoding and indexing video data. For example, ABC uses a search engine which enables one to search some specific text that appeared in ABC news. A number of universities are also recording lectures and seminars, with the aim of providing online access and search capabilities.

String data applications generally involve very large databases. GenBank [7], a database of nucleotide and protein strings built by National Center for Biotechnology Information (NCBI), is an example of such a database. Figure 1 plots the growth of the size of this database from year 1982 to 2000. The statistics show that the size of GenBank has doubled every 15 months [8]. Similarly, the size of a video database can also increase dramatically: CNN has more than 150 hours of news feed every day, and plans to encode more than 100,000 hours of archived material.

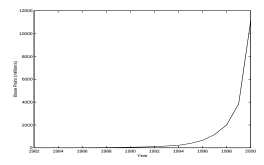


Figure 1: The growth of the NCBI database in recent years.

Most of the string search algorithms proposed so far are in-memory algorithms [5, 6, 11, 17, 20, 21, 22]. That is, these techniques have to scan the whole database for each query. Therefore, these techniques suffer from disk I/Os when the database is too large. In-memory algorithms can become impractical for string databases because the database size grows faster than the available memory capacity, and extensive memory requirements make the search techniques impractical. The size of the index structure for the index based techniques [2, 4, 15, 18] are even larger than the size of the database, and their performance deteriorates for long query patterns. Therefore, efficient external memory algorithms are needed for most string comparison applications of the future.

A string  $s_1$  can be transformed into another string  $s_2$  by using three edit operations, namely insert, delete, and replace, on individual characters of the string  $s_1$ . Figure 2 presents a transformation of the string ACTTAGC to AAT-GATAG using edit operations. The transformation given in Figure 2 consists of 1 replace, 2 insert, and 1 delete operations. The difference between two strings  $s_1$  and  $s_2$  is generally defined as the minimum number of edit operations to transform  $s_1$  to  $s_2$ , called *edit distance* (ED). Let m and n be the lengths of strings  $s_1$  and  $s_2$ , then the edit distance,  $ED(s_1, s_2)$ , and the corresponding edit operations can be determined in O(mn) time and space using dynamic programming [11]. The space complexity can be reduced to  $O(min\{m, n\})$  if only the edit distance is needed (i.e. the corresponding edit operations are not required). Some applications assign different weights to different edit operations or different character pairs [12], leading to a weighted edit distance. The time and space complexity of finding the weighted edit distance is also O(mn) by using dynamic programming.

Figure 2: Transformation of the string ACTTAGC to AAT-GATAG using edit operations.

An alignment of strings  $s_1$  and  $s_2$  is obtained by matching each character of  $s_1$  to a character in  $s_2$  in increasing order. All the unmatched characters in both strings are matched with space. An alignment of strings ACTTAGC and AATGATAG is given in Figure 2. The dashes (i.e., -) in this figure correspond to spaces. Each character pair is assigned a score based on their similarity, and these values are stored in a score matrix. The value of an alignment is defined as the sum of the scores of all of their character pairs. Global alignment (or similarity) of  $s_1$  and  $s_2$  is defined as the the maximum valued alignment of  $s_1$  and  $s_2$ . Finding the similarity of two strings is the dual of finding the distance between them. Local alignment [20] of  $s_1$ and  $s_2$  is defined as the highest valued alignment of all the substrings of  $s_1$  and  $s_2$ . Both global and local alignments can be determined in O(mn) time using dynamic programming.

In this paper, we consider the problem of range queries

and nearest neighbor queries. The typical databases that we work with include very long strings. For example, the string corresponding to chromosome-22 of humans has about 35 million base pairs. (A base pair is one of A,C,G, or T characters corresponding to the four different kinds of nucleic acids.) A k-nearest neighbor returns the k closest substrings from the database to a given query. A range query, on the other hand, returns the substrings that lie within a given distance of the input query.

We propose a wavelet-based method to map the substrings of the database into a multidimensional integer space. The number of dimensions is determined by the alphabet size and the number of wavelet coefficients. We define a notion of distance in this integer space that is a lower bound to the actual edit distance. A sliding window is used to translate a set of contiguous substrings into an MBR (Minimum Bounding Rectangle). Repeating this over all the strings generates an array of MBRs corresponding to one resolution (window size) for the database. We use a hierarchical scheme in which windows of successive coarser grain are used. This generates an approximation to the database at different granularities, and results in a grid of MBRs. The resulting index structure is quite compact and can be stored in memory. Typical size of this index structure ranges between 1-2% of the database size. Range queries and nearest-neighbor queries are first performed using this in-memory index structure using the lower-bound distance. The resulting set of candidate pages are then accessed from the disk to remove false hits (using the actual edit distance).

According to experimental results, our method runs 5 to 45 times faster than existing techniques for nearest neighbor queries of 10 to 200, nearest neighbors and 2 to 12 times faster than existing techniques for range queries.

The rest of the paper is as follows. Section 2 discusses the related work. Section 3 discusses the substring searching problem and defines our index structure and algorithms. Section 4 discusses the experimental results. We end with a brief discussion in Section 5.

# 2 Related Work

The dynamic programming solution to the problem of finding the substrings of a given string s of length n, which are within a distance of  $r=\epsilon \times |q|$  to a query string q of length m, runs in O(mn) time and space. This technique is a variation of the dynamic programming algorithm that finds the edit distance between two strings by generating a distance matrix of size  $m\times n$ . For long data and query strings, this technique becomes infeasible in terms of both time and space. Myers [17] improved the time and space complexity to O(rn) by maintaining only the required part of the distance matrix. However, for large error rates r is O(m), and hence the complexity is still O(mn).

Wu and Manber [22] proposed a technique that uses r binary masks  $M_1$ ,  $M_2$ , ...,  $M_r$  of length m. They scan through the data string s and update these masks for each character in s. After the  $j^{th}$  character is processed, the value of  $M_k[i]$  becomes 1 if the last i characters of s are

within k edit operations to the first i characters of q. If w is the size of a word, the algorithm runs in  $O(n\frac{mr}{w})$  time. The space requirement of this technique is  $O(r\frac{m}{w})$ . The algorithm runs efficiently for small values of m (close to O(nr)), but the performances degrades for large m. Furthermore, the space requirement may become much larger than the data string for large m and r.

In another paper, Myers [18] proposed a technique that preprocesses the data string s and creates an index of size O(n). This technique assumes a lower bound l on the length of the query string (l = log(n) here). All possible strings of length l are mapped to integers using a perfect hashing function. Later, the leftmost points of all the occurrences of these strings in s are stored in separate lists. For a given query q and query radius r, the technique generates the set of strings which are within edit distance of r to q, called *condensed r-neighborhood*. The strings in the condensed r-neighborhood are searched in the index to find the answers to the query. If the query length is larger than l, the technique splits the query string into subqueries, searches each subquery separately and combines the results. Since this technique indexes all possible strings of some prespecified length, we call it a dictionary based technique. The author proves that if the database is created as a result of equi-probable Bernoulli trials, then the technique runs in sublinear time. There are two drawbacks with this technique. First, although the space complexity is O(n), the index size can be 7-9 times larger than the data size. This may cause a drop in performance if the index does not fits in memory. Second, the worst case running time complexity of this technique is very high.

Baeza-Yates and Navarro proposed an NFA-based solution in [5]. They propose an NFA of  $(r+1) \times (m+1)$  states, which accepts s as the input string. The NFA is constructed using the query string. The NFA goes into an accepting state whenever a substring within edit distance of r is processed. The authors propose to use only the required states of the NFA at any time. The expected running time of this technique is  $O(mn\frac{r}{w})$ , where w is the size of a word. The experimental results presented in the paper show that for short queries and small alphabets, this technique performs well. The performance of this technique deteriorates when s is very long (i.e. it does not fit in memory).

Altschul, Gish, Miller, Myers, and Lipman proposed the BLAST technique [3] to find local similarities. BLAST, the most popular string matching tool for biologists, runs in two phases. In the first phase, all the substrings of the query of some prespecified length (typically between 3 and 11) are searched in the database for an exact match. In the second phase, all the matches obtained in the first phase are extended in both directions until the similarity between the two substrings falls below some threshold. This technique keeps a pointer to the starting locations of all possible substrings of the prespecified length in the database to speedup the first phase. Therefore, the space requirement of BLAST is more than the size of the database. Furthermore, BLAST does not find a similar substring to the whole query string, only similarities between the query substrings and the database substrings.

Muthukrishnan and Sahinalp [16] proposed an index structure for approximate nearest neighbor search. This technique uses an index structure based on suffix arrays and a partitioning of the pattern. The resulting index structure is four times the size of the database.

Giladi, Walker, Wang, and Volkmuth [10] considered a heuristics-based solution which runs in O(logn) expected time. This technique splits the data strings into overlapping windows of length l for some prespecified overlap amount of  $\Delta$ . For each such window, they count the number of repetitions of all the possible k-tuples, and store this value in a  $\sigma^k$  dimensional vector, where  $\sigma$  is the alphabet size. Later, these vectors are indexed using a hierarchical binary tree. The authors propose to approximate the similarity between the query string and a substring by using the  $D_1$  distance between these vectors. Experimental results show that this technique runs 25 to 50 times faster than BLAST. The authors also note that this technique can be used as a preprocessing step to speed up any string search program. There are two drawbacks with this method: it allows false drops, and the index size increases exponentially with k.

A special case of the substring matching problem is exact matching (i.e. r=0). One can solve this problem using suffix trees [11] in which all the suffixes of a database string are stored in a tree. However, the size of the suffix tree may be more than ten times larger than the database size. Manber and Myers [15] propose a data structure called suffix arrays to reduce the space requirement for the index structure. However, the space requirement is still more than four times the database size. Ferragina and Manzini [9] proposed a technique to compress the suffix arrays, decreasing the query performance slightly.

# 3 Proposed Solution

String matching problem can be classified in two groups. These are whole matching and substring matching. The simpler case, whole matching, considers the problem of finding the edit distance ED(q, s) between a data string s and a query string q. Substring matching considers all the substrings s[i:j] of s which are close to the query string, where s[i:j] is the substring of s between (and including) the  $i^{th}$  and  $j^{th}$  characters. In this paper, we confine our attention to substring matches.

Given a string database  $S = \{s_1, s_2, ..., s_d\}$  consisting of very long strings, we consider two types of queries:

- Range search seeks all the substrings of S which are within an edit distance of r to a given query string q, where r is the query range. We define  $\epsilon = \frac{r}{|q|}$  as the error rate.
- *k-nearest neighbor search* seeks the *k* closest substrings of *S* to *q*.

There are several challenges in solving the substring matching problem. 1) Finding the edit distance is very costly in terms of both time and space. 2) The strings in the database may be very long. For example, the length of chromosome-22, one of the shortest chromosomes in human genome library, is approximately 35 million base pairs. For a query

string of length ten thousands and an error rate of  $\epsilon=0.01$ , there are billions of possible substrings. Therefore it is infeasible to check all the substrings. 3) The database size for most applications grows exponentially. Therefore, a solution method based on sequentially scanning the database will suffer from extensive disk I/Os. Our approach based on distance approximations and the ensuring hierarchical index structure adresses all the above issues.

The rest of this section is as follows. Section 3.1 defines a lower bound distance for substring searching. Section 3.2 improves this lower bound by using the idea of wavelet transformation. Section 3.3 presents the MRS-index structure based on the aforementioned distance formulations. Sections 3.4 and 3.5 present the algorithms for range queries and nearest neighbor queries.

#### 3.1 A new distance function

We define a transformation, f(s) that maps a string s to a point in a multidimensional integer space as follows:

**Definition 1** Let s be a string from the alphabet  $\Sigma = \{\alpha_1, \alpha_2, ..., \alpha_{\sigma}\}$ . Let  $n_i$  be the number of occurrences of the character  $\alpha_i$  in s for  $1 \le i \le \sigma$ . We define the frequency vector, f(s), of s as:

$$f(s) = [n_1, n_2, ..., n_{\sigma}].$$

For example, if s=TACTTAG is a genetic string (i.e. from alphabet  $\Sigma = \{A, C, G, T\}$ ), then f(s) = [2, 1, 1, 3] (We use alphabetic order in the construction of f(s).). Three important notes follow from this definition. 1) The transformed string f(s) has  $\sigma$  dimensions independent of the length of s. 2) The sum of the entries of f(s) is independent of the contents of s. 3) All the entries of f(s) are nonnegative.

**Lemma 1** Let s be a string from the alphabet  $\Sigma = \{\alpha_1, \alpha_2, ..., \alpha_{\sigma}\}$ . Let  $f(s) = [v_1, v_2, ..., v_{\sigma}]$  be the frequency vector of s, then  $\Sigma_{i=1}^{\sigma} v_i = |s|$ .

As a result of Lemma 1, the transformation of a strings of length n lie on the  $\sigma-1$  dimensional plane that passes through the point [n,0,0,...,0] and is perpendicular to the normal vector [1,1,...,1]. The relationship between the edit operations and the frequency vectors is captured in Theorem 1.

**Theorem 1** Let s be a string from the alphabet  $\Sigma = \{\alpha_1, \alpha_2, ..., \alpha_{\sigma}\}$ . Let  $f(s) = [v_1, v_2, ..., v_{\sigma}]$  be the frequency vector of s. An edit operation on s has one of the following effects on f(s), for  $1 \le i, j \le \sigma$ , and  $i \ne j$ :

1. 
$$v_i := v_i + 1$$

2. 
$$v_i := v_i - 1$$

3. 
$$v_i := v_i + 1$$
 and  $v_i := v_i - 1$ 

#### Proof:

Case 1 corresponds to inserting  $\alpha_i$  in some location of s. Case 2 corresponds to deleting  $\alpha_i$  from s.

Case 3 corresponds to replacing  $\alpha_i$  with  $\alpha_i$  in s.

Theorem 1 shows that a single edit operation on a string results in a limited change in the corresponding integer space. Keeping this fact in mind, we define *neighboring points* as follows:

**Definition 2** Let u and v be integer points in  $\sigma$  dimensional space, then u and v are called neighbors if one of them can be obtained from the other using a single edit operation.

Next we define a new distance function, frequency distance  $(FD_1)$ , for the frequency vectors, which is a lower bound on the edit distance of the corresponding strings. The idea is based on Theorem 1.

**Definition 3** Let u and v be integer points in  $\sigma$  dimensional space. The frequency distance,  $FD_1(u,v)$ , between u and v is defined as the minimum number of steps in order to go from u to v (or equivalently from v to u) by moving to a neighbor point at each step.

Theorem 2 proves that the frequency distance between the frequency vectors of two strings is a lower bound on their edit distance.

**Theorem 2** Let  $s_1$  and  $s_2$  be two strings from the alphabet  $\Sigma = {\alpha_1, \alpha_2, ..., \alpha_{\sigma}}$ , then

$$FD_1(f(s_1), f(s_2)) \le ED(s_1, s_2).$$

**Corollary 1** *Let q and s be two strings from the alphabet*  $\Sigma = \{\alpha_1, \alpha_2, ..., \alpha_{\sigma}\}$ , then

if 
$$r < FD_1(f(q), f(s))$$
 then  $r < ED(q, s)$ .

Given a query string q and a query range r, one can prune a string s without computing ED(q,s) if  $r < FD_1(f(q),f(s))$ . Another important attribute of the frequency distance function is that it is a metric.

Figure 3 presents an efficient algorithm to compute  $FD_1(u,v)$  for  $\sigma$  dimensional integer points u and v. The algorithm computes two values, namely posDistance, and negDistance. PosDistance defines the number of decrement operations that must be applied to u, and similarly negDistance defines the number of increment operations that must be applied to u. Since each edit operation changes the values of posDistance and negDistance by at most one, the larger of the two values equals  $FD_1(u,v)$ .

### 3.2 Improving the lower bound distance

The frequency distance can be improved by storing local frequencies of the characters in the string as well as the global frequencies. For this, we define the wavelet transformation of a string. Assume n is a power of 2 for simplicity in the following development.

/\* u and v are  $\sigma$  dimensional integer points. \*/
Algorithm  $FD_1(u,v)$ 

- posDistance := negDistance := 0
- for i := 1 to  $\sigma$ 
  - if  $u_i > v_i$  then  $posDistance += u_i v_i$
  - else  $negDistance += v_i u_i$
- ullet if posDistance > negDistance then return posDistance else return negDistance

Figure 3: Computation of  $FD_1$ 

**Definition 4** Let  $s = c_1c_2...c_n$  be a string from the alphabet  $\Sigma = \{\alpha_1, \alpha_2, ..., \alpha_{\sigma}\}$ , then  $k^{th}$ -level wavelet transformation,  $\psi_k(s)$ ,  $0 \le k \le log_2n$ , of s is defined as:

$$\psi_k(s) = [v_{k,1}, v_{k,2}, ..., v_{k,\frac{n}{s-k}}], \text{ where } v_{k,i} = [A_{k,i}, B_{k,i}],$$

$$A_{k,i} = \begin{cases} f(c_i) & k = 0\\ A_{k-1,2i} + A_{k-1,2i+1} & 0 < k \le \log_2 n, \end{cases}$$

and

$$B_{k,i} = \left\{ \begin{array}{ll} 0 & k = 0 \\ A_{k-1,2i} - A_{k-1,2i+1} & 0 < k \le \log_2 n \end{array} \right.$$

As an example of the above definition, it is possible to show that  $\psi_3(TCACTTAG)=[[2,2,1,3],[0,2,-1,-1]]$ . The above formulation is similar to the Haar wavelet except for a scaling factor. Some important properties of the wavelet transformation of a string s are as follows. 1) The A coefficients of  $\psi_0$  defines the string. 2) Global frequency vector f(s) as developed in Section 3.1 is simply  $A_{log_2n,0}$  3) Each  $A_{k,i}$  coefficient corresponds to the frequency vector of a substring of s of length  $2^k$ . Formally,  $A_{k,i}=f(s[2^ki:2^k(i+1)-1])$ . 4) Each  $B_{k,i}$  coefficient corresponds to the difference of the frequency vectors of two consecutive substrings of s of length  $2^{k-1}$ . Formally,  $B_{k,i}=f(s[2^ki:2^ki+2^{k-1}-1])-f(s[2^ki+2^{k-1}])$ . Hence,  $B_{log_2n,0}=f(s[0:\frac{n}{2}-1])-f(s[\frac{n}{2}:n-1])$ .

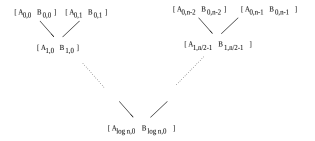


Figure 4: Wavelet decomposition of a string

The hierarchical structure of the wavelet transformation of a string is given in Figure 4. We name  $A_{log_2n,0}$  as the first wavelet coefficient, and  $B_{log_2n,0}$  as the second wavelet

coefficient. If the A and B coefficients of  $\psi_k$  are known, then the A coefficients of  $\psi_{k-1}$  can be computed. The B coefficients of  $\psi_{log_2n-1}$  are called third and fourth wavelet coefficients. In general, if the first wavelet coefficient and all the B coefficient of  $\psi_i$  for  $0 \le i \le log_2n$  are known, then all the A coefficients can be determined.

In Section 3.1, we transformed the strings to their first wavelet coefficients. As the number of wavelet coefficients increases, the accuracy of the lower bound function increases at the cost of a larger index size. This is shown next. We will focus our development on the first two wavelet coefficients; however, the idea can be generalized to any number of coefficients. Hereon, we will use  $\psi(s)$  instead of  $\psi_{log_2|s|}(s)$  for simplicity.

**Theorem 3** Let s be a string from the alphabet  $\Sigma = \{\alpha_1, \alpha_2, ..., \alpha_{\sigma}\}$ . Let  $\psi(s) = [A, B]$  be the first and the second wavelet coefficients of s. Let  $A = [a_1, ..., a_{\sigma}]$  and  $B = [b_1, ..., b_{\sigma}]$ . An edit operation on s has one of the following effects on A and B for  $1 \le i, j \le \sigma$ , and  $i \ne j$ :

1. 
$$a_i := a_i + 1$$
,  $a_j := a_j - 1$ ,  $b_i := b_i + 1$ ,  $b_j := b_j - 1$ .

2. 
$$a_i := a_i + 1$$
,  $a_j := a_j - 1$ ,  $b_i := b_i - 1$ ,  $b_j := b_j + 1$ .

3. 
$$a_i := a_i \pm 1, b_i := b_i \pm 1.$$

4. 
$$a_i := a_i \pm 1$$
,  $b_i := b_i + 1$ ,  $b_i := b_i - 2$ .

5. 
$$a_i := a_i \pm 1, b_i := b_i - 1, b_i := b_i + 2.$$

#### **Proof:**

This can be proven by splitting the string into 2 equal parts and inspecting the effect of the edit operations on these substrings.

The wavelet transform  $\psi(s)$  can be considered as a point in a  $2\sigma$  dimensional integer space. Theorem 3 lists the legal steps that can be used to move from  $\psi(s_i)$  to  $\psi(s_j)$ , where  $s_i$  and  $s_j$  are strings. The transformation of  $s_1$  to  $s_2$  using the edit operations corresponds to a legal path between their wavelet transformations. Therefore, the edit distance between  $s_1$  and  $s_2$  is at least the number of steps in the shortest legal path from  $\psi(s_i)$  to  $\psi(s_j)$ . Lemma 2 defines a lower bound,  $FD_2(\psi(s_i), \psi(s_j))$ , to the number of steps in the shortest legal path in  $2\sigma$  dimensional integer space based on the legal operations given in Theorem 3.

**Lemma 2** Let  $s_1$  and  $s_2$  be strings from the alphabet  $\Sigma = \{\alpha_1, \alpha_2, ..., \alpha_\sigma\}$ . Let  $\psi(s_1) = [A_1, B_1]$  and  $\psi(s_2) = [A_2, B_2]$  be the first and second wavelet coefficients of  $s_1$  and  $s_2$ . Let  $A_1 = [a_{1,1}, ..., a_{1,\sigma}]$ ,  $B_1 = [b_{1,1}, ..., b_{1,\sigma}]$ ,  $A_2 = [a_{2,1}, ..., a_{2,\sigma}]$ , and  $B_2 = [b_{2,1}, ..., b_{2,\sigma}]$ . Let

$$pos = \sum_{i, a_{1,i} > a_{2,i}} (a_{1,i} - a_{2,i}) + \sum_{i, b_{1,i} > b_{2,i}} (b_{1,i} - b_{2,i}),$$

$$neg = \sum_{i,a_{1,i} < a_{2,i}} (a_{2,i} - a_{1,i}) + \sum_{i,b_{1,i} < b_{2,i}} (b_{2,i} - b_{1,i}).$$

Let min be the minimum of pos and neg, then

$$FD_2(\psi(s_i), \psi(s_j)) = \begin{cases} \frac{|pos - neg|}{2}, & \text{if } min < \frac{|pos - neg|}{2} \\ \frac{|pos - neg|}{2} + \frac{min - \frac{|pos - neg|}{2}}{2}, & \text{else} \end{cases}$$

### **Proof:**

Let max be the maximum of pos and neg. Steps 4 and 5 of Theorem 3 results in the largest change to the difference  $\psi(s_i)-\psi(s_j)$  (+3, -1 or -3, +1). Therefore, these steps must be used whenever |pos-neg|>0. If  $min<\frac{|pos-neg|}{2}$ , then these steps can be used at most min times. After that, only Step 3 ( $\pm 2,0$ ) can be used to increment or decrement the remaining value by 2 at each step. Therefore, at least  $\frac{max-3min}{2}$  more steps are needed. This makes the total distance  $min+\frac{max-3min}{2}=\frac{|pos-neg|}{2}$ .

A similar resulting can be used for the second part. If

A similar reasoning can be used for the second part. If  $\min > \frac{|pos-neg|}{2}$ , then steps 4 and 5 can be used at most  $\frac{|pos-neg|}{2}$  times. This makes pos=neg, so Step 1 and Step 2 (+2, -2 or -2, +2) can be used for the rest. Therefore, at least  $\frac{|pos-neg|}{2} + \frac{min-\frac{|pos-neg|}{2}}{2}$  steps are needed to move from  $\psi(s_i)$  to  $\psi(s_j)$ .

Lemma 2 constructs a lower bound to the edit distance using the first and second wavelet coefficients at the same time. However,  $FD_1(f(s_1), f(s_2))$  is not necessarily less than  $FD_2(\psi(s_1), \psi(s_2))$ . We define the maximum frequency distance (FD) between  $s_1$  and  $s_2$  as

$$FD(s_1, s_2) = max\{FD_1(f(s_1), f(s_2)), FD_2(\psi(s_1), \psi(s_2))\}.$$

#### 3.3 The MRS index structure

Let  $S=\{s_1,s_2,...,s_d\}$  be a database consisting of potentially long strings from alphabet  $\Sigma=\{\alpha_1,\alpha_2,...,\alpha_\sigma\}$ . Let  $w_1=2^a$  be the length of the shortest possible query string. Our index structure stores a grid of trees  $T_{i,j}$ , where i ranges from a to a+l-1, and j ranges from 1 to d. The parameter l represents the number of resolution levels available in the index structure. Tree  $T_{i,j}$  is the index structure for the  $j^{th}$  string corresponding to window size  $2^i$ . Figure 5 shows a layout of this index structure.

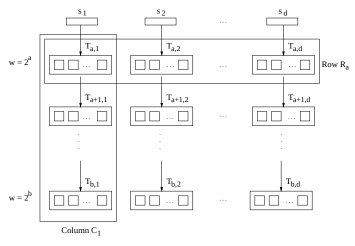


Figure 5: Layout of the MRS-index structure

In order to obtain  $T_{i,j}$ , we slide a window of length  $2^i$  on string  $s_j$ , starting from the leftmost point of  $s_j$ . For each possible placement of the window, we compute the

wavelet transformation of the corresponding substring of  $s_i$ , and store the first two wavelet coefficients. Note that each substring corresponds to a point in the  $2\sigma$  dimensional integer space. We begin with the initial substring and find the minimum box, called Minimum Bounding Rectangle (MBR), that covers the wavelet coefficients of this substring. This box is later extended to cover the transformations of the first c substrings, where c is the box capacity. (We will later discuss the impact of the value of c on the efficiency of the index structure.) After the first c substrings are transformed, a new MBR is created to cover the next c substrings. This process continues until all substrings are transformed. Note that, we only store the lower and higher end points of the MBRs along with the starting locations of the first substring contained in that MBR. Since the index structure stores frequencies at different resolutions, we call this index structure the Multi Resolution String (MRS) Index Structure.

The  $i^{th}$  row of the MRS index structure is represented by  $R_i$ , where  $R_i = \{T_{i,1}, ..., T_{i,d}\}$  corresponds to the set of all trees at resolution  $2^i$ . Similarly, the  $j^{th}$  column of the MRS index structure is represented by  $C_j$ , where  $C_j = \{T_{a,j}, ..., T_{a+l-1,j}\}$  corresponds to the set of all trees for the  $j^{th}$  string in the database.

Let q be a query string of length  $2^i$ , where  $a \le i \le a+l-1$ . Given an MBR B, we define  $FD(q,B) = min_{s \in B}FD(q,s)$ . We observe the following:

- 1. if  $r \leq FD(q, B)$  then  $r \leq FD(q, s)$  for all  $s \in B$ .
- 2. As the box capacity c increases, the box volume increases. As a result, the query-box distance FD(q,B) decreases, and the performance of the index structure deteriorates.
- 3. For a fixed value of c, FD(q, B) decreases as the window size (w) decreases. This can be explained as follows. Recall that the sum of the entries of the frequency vectors of substrings of length w is constant (i.e. w). Furthermore, frequency vectors contain a fixed number of dimensions (i.e.  $\sigma$ ), and each dimension has a nonnegative value. Hence, there are  $C(w + \sigma - 1, w)$  possible frequency vectors for substrings of length w. Consequently, with decreasing w, the set of frequency vectors in the MBR constitutes a higher percentage of the set of all possible frequency vectors. As a result, the probability that f(q)is contained in the MBR increases, and FD(q, B) decreases. We verified this for our datasets by computing the average volume of an MBR for different window sizes. This is plotted in Figure 6. According to this figure, the average box volume increases exponentially as w decreases.
- 4. The wavelet coefficients of the substrings obtained by sliding the window by a single character are very close to each other. Therefore, the set of wavelet coefficients in an MBR are generally highly clustered.

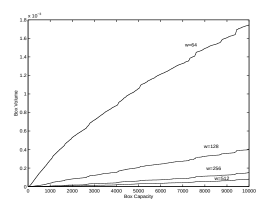


Figure 6: The average volume of MBRs for various box capacities and window sizes.

## 3.4 Range Queries

Our search technique partitions a given query string of arbitrary length into a number of subqueries at various resolutions available in our index structure. Later, it performs a partial range query for each of these subqueries on the corresponding row of the index structure. This is called a partial range query, because it only computes distance of the wavelet transform of the query substring to the MBRs, not the distance of the query string to the substrings contained in the MBRs.

Given any query q of length  $k2^a$  and a range  $\epsilon$ , there is a unique partitioning  $^l$ ,  $q=q_1q_2...q_t$ , with  $|q_i|=2^{c_i}$  and  $a \leq c_1 < ... < c_i \leq c_{i+1} \leq ...c_t \leq a+l-1$ . This partitioning technique chooses the longest possible suffix of q, such that its length is equal to one of the resolutions available in the index, as the last query substring. Later, it recursively partitions the rest of the strings to find the other query substrings.

We first perform a search using  $q_1$  on row  $R_{c_1}$  of the index structure. As a result of this search, we obtain a set of MBRs that lie within a distance of  $r = \epsilon \times |q|$  from  $q_1$ . Using the distances to these MBRs, we refine the value of r for each MBR and make a second query using  $q_2$  on row  $R_{c_2}$  and the new value of r. This process continues for the remaining rows  $R_{c_3}$  ...  $R_{c_t}$ .

The relationship between an MBR and the substrings of the string that forms this MBR is captured in the following lemma.

**Lemma 3** Let s be a string and B be an MBR that covers the wavelet transforms of all the substrings of length w in s. Let q be a query string of length w. Let d be the minimum edit distance between q and all substrings of s, then

$$FD(q, B) \leq d$$
.

Figure 7 presents the complete search algorithm. Step 1 partitions the query q into separate pieces corresponding to a subset of the rows  $R_{c_1}$ ,  $R_{c_2}$ , ...,  $R_{c_t}$  of the

 $Algorithm RANGE\_SEARCH(q,r)$ 

- 1. Partition the query q into t parts as  $q_1, q_2, ..., q_t$  such that  $|q_i| = 2^{c_i}$  and  $a \le c_1 < c_2 < ... < c_i \le c_{i+1} \le ... \le c_t \le b$ . Let  $T_{c_1,j}$  contain  $M_j$  boxes.
- 2. For j:=1 to d /\* process each sequence \*/ 3. For k:=1 to  $M_j$  /\* process each MBR \*/
  - (a) distance[k] := r
  - (b) For i := 1 to t
    - i. Perform range query on MBRs corresponding to  $q_i$  using radius distance[k]. Let  $Res_{c_i,j}$  be the resulting set of MBRs whose distances to  $q_i$  are less than distance[k].
    - ii.  $distance[k] := max_{B \in Res_{c_i,j}} \{ distance[k] FD(q_i, B) \}$
- 4. Read disk pages corresponding to  $Res_{c...i}$
- 5. Perform postprocessing to eliminate false retrievals.

Figure 7: Range search algorithm.

index structure. In Step 2, every database sequence is searched independently. Then for every sequence, we process its MBRs independently (Step 3). The distance vector for a given MBR is initialized to r (Step 3a). Successive rows of the index structure for the sequence are then used to refine this distance vector (Step 3b). When all rows have been searched, the disk pages corresponding to the last result set are read (Step 4). Finally, postprocessing is carried out to eliminate false retrievals (Step 5). Note that any of the distance computation techniques available [2, 3, 4, 5, 6, 11, 15, 17, 18, 20, 21, 22] can be used in the postprocessing step.

As a consequence of Theorem 2 and Lemma 3 we have the following theorem.

**Theorem 4** The MRS index structure does not incur any false drops.

We note the following about the search algorithm. 1) For each MBR, the refinement of radius is carried out independently, and proceeds from top to bottom. 2) For each substring, no disk reads are done until the termination of the for loop in Step 3b. Furthermore, the target pages are read in the same order as their location on disk. As a result, the average cost of a page access is much less than the cost of a random page access.

# 3.5 Nearest neighbor queries

A k-nearest neighbor (k-NN) query seeks the k closest substrings from the database to a query string s. We perform a k-NN query in two phases. In the first phase, the k closest MBRs in the index structure are determined by an inmemory search on the index structure. Once the k closest MBRs are determined, the algorithm reads the substrings contained in these MBRs, and finds the k<sup>th</sup> smallest edit

<sup>&</sup>lt;sup>1</sup> If the length of the query string is not a multiple of the minimum window size, then the longest prefix of the query whose length is a multiple of the minimum window size can be used.

Algorithm  $k - NN\_SEARCH(q, k)$ 

- 1. Phase 1
  - $\mathcal{B}$  := The set of k closest MBRs to the query q.
  - $r_1 := k^{th}$  smallest edit distance to strings in  $\mathcal{B}$ .
- 2. Phase 2
  - $RANGE\_SEARCH(q, r_1)$
  - Return the k closest strings in the answer set.

Figure 8: *k*-nearest neighbor algorithm

distance of these substrings to the query string. We represent this distance by  $r_1$ . Note that, generally a small percentage of the database is processed at this stage. In the second phase, we perform a range query using  $r_1$  as the query radius. Figure 8 presents the complete k-NN search algorithm.

It is guaranteed that the k nearest neighbors are retrieved in the second phase. This can be explained as follows. The edit distance of the query to the actual  $k^{th}$  nearest neighbor (say r) is at most  $r_1$ . Let B be an MBR that contains the wavelet coefficients of at least one of the substrings in the actual answer set of the k-NN query, then  $FD(q,B) \leq r$ . Hence,  $FD(q,B) \leq r_1$ , and all the MBRs that contain answer strings are retrieved in the second phase.

One can retrieve any number of MBRs in the first phase as long as they contain more than k substrings. For example, the substrings in the closest MBR is sufficient to prove correctness of the algorithm if the box capacity is at least k. The postprocessing and disk read cost of the first phase decreases if fewer MBRs are retrieved in the first phase. However, as the number of MBRs retrieved in the first phase decreases, the radius of the range query in the second phase increases. Hence, it causes more pages reads and postprocessing in the second phase.

Korn, Sidiropoulos, Faloutsos, Siegel, and Protopapas [14] propose a similar k-NN search. The authors propose a technique in which k closest points are obtained in the first phase using an approximate distance function. The actual distance to these points is computed, and a range query with the greatest actual distance is performed in the second phase. Seidl and Kriegel [19] propose an optimal iterative k-nearest neighbor search technique. They iterate over both the feature and the object spaces to ensure that no unnecessary objects are accessed. Our algorithm is closer in spirit to the former algorithm except that, we work with MBRs instead of data points in the first phase.

# 4 Experimental results

We used four *homo sapiens* chromosome strings in our experiments taken from [1]. These are chromosome 2 (chr02), chromosome 18 (chr18), chromosome 21 (chr12), and chromosome 22 (chr22). These chromosome strings are composed of the alphabet  $\Sigma = \{A, C, G, T, N\}$ . The letter "N" stands for *not known*. We treat the letter N as a different letter, resulting in an alphabet size of 5. The

lengths of these strings and the number of occurrences of each letter are presented in Figure 9. The chr18 dataset contains 4M characters, and all the other datasets contain more than 31M characters. The numbers of A, C, G, and T characters in the the chr22 dataset are about equal, while the other datasets contain more A/T than C/G. The percentage of the unknown characters for the chr22 dataset is 3%, while it is less than 0.007% for the other datasets.

We implemented single wavelet coefficient and two wavelet coefficient versions of the MRS-index structure for window sizes  $w = \{128, 256, 512, 1024\}$ . We compared the performance of our technique to the NFA based method proposed by Baeza-Yates and Navarro [5]. This is a recent technique for range queries. This technique sequentially reads a data string and feeds it to an NFA constructed using the query string and the query radius. The NFA goes into the accepting state whenever an answer substring is processed. Our measure of cost is based on the number of disk pages accessed, and the number of string comparisons made later using the accessed disk pages. Since the number of string comparisons needed for a given disk page is constant over all techniques, the total cost is proportional to the number of disk pages read. We assume that the page size is 1K in our experiments.

The rest of this section is as follows. Section 4.1 discusses the effect of box capacity on the performance of queries. Section 4.2 discusses the effect of window size on the performance of queries. Sections 4.3 and 4.4 discusses our results for NN and range queries.

## 4.1 Effect of box capacity

The first experiment compares the performance of the MRS-index for different box capacities. In this experiment, we perform 100 arbitrary length nearest neighbor queries for box capacities 100, 500, 1000, 2000, 3000, and 4000 for k = 10 on the chr18 dataset. The length of the queries varies between 512 and 10000. Figure 10 plots the cost of the MRS-index technique and the NFA technique for 10-NN queries. The cost of the MRS-index increases as the box capacity increases. This effect of increasing box capacity on the performance was observed earlier in Section 3.3 (Observation 2). The cost of the MRS-index is much lower than the NFA technique for all these box capacities. The MRS-index runs up to 120 times faster than the NFA technique when the box capacity is 500, and up to 4 when the box capacity is 4000. Although using 2-wavelet coefficient slightly improves the performance for the same box capacity, the size of the index structure is doubled. If we use the same amount of memory, the single coefficient version performs better.

#### 4.2 Effect of window size

The second experiment reports the impact of the window size (i.e. resolution) on the performance of the MRS-index structure. In this experiment, we use only one row of the index structure. We ran 100 arbitrary length k-NN queries for window sizes 128, 256, 512, 1024 for k=10 on the chr18 dataset. The queries are randomly selected from the

Dataset	A	C	G	T	N	Total
chr18	1300930	823103	819387	1282027	30	4225477
chr21	10032226	6921020	6908202	9962534	174	33824156
chr22	8751963	8002860	8000421	8721658	1076929	34553831
chr02	9394422	6251334	6299566	9565401	2266	31512989

Figure 9: Frequency of symbols in the datasets.

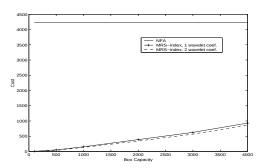


Figure 10: The cost of our method with one and two wavelet coefficients and the NFA technique for different box capacities for 10-NN queries on the chr18 dataset.

chr18 dataset. Figures 11 plots the cost of the MRS-index structure and the NFA technique. The MRS-index structure outperforms the NFA technique for all the window sizes. Furthermore, the performance of the MRS index structure itself improves as the window size increases. The effect of increasing window size on the performance was observed earlier in Section 3.3 (Observation 3).

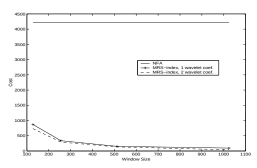


Figure 11: The cost of our method with one and two wavelet coefficients and the NFA technique for different window lengths for 10-NN queries on the chr18 dataset.

#### 4.3 Nearest neighbor queries

The third experiment considers k-NN queries on the chr21 dataset for 9 different values of k from 10 to 500 when the box capacity is 1000. In this experiment, we ran 100 random NN queries of arbitrary lengths in the range 512 to 10000 for each value of k. The queries are generated from the same dataset. Figure 12 presents the cost of the MRS-index structure and the NFA technique. The experimental results for the other datasets are similar. The MRS-index structure outperforms the NFA technique for all values of k. Although the performance of the MRS-index structure

drops for large values of k, it still performs better than the NFA technique. We achieved speedups up to 45 for 10 nearest neighbors. The speedup for 200 nearest neighbors is 3. As the number of nearest neighbors increases, the performance of the MRS-index structure approaches to that of the NFA technique. This is because an increasingly large number of MBRs are accessed in Phase 1 of the k-nearest neighbor algorithm.

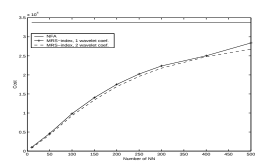


Figure 12: The cost of our method with one and two wavelet coefficients and the NFA technique for different number of nearest neighbors on the chr21 dataset.

#### 4.4 Range queries

We considered range queries in the fourth experiment set for  $\epsilon = 0.01$ , 0.025, 0.05, 0.075, and 0.1 on the chr22 dataset. We performed 100 arbitrary length queries for each error rate. The box capacity is fixed to 1000 in this experiment. Figure 13 presents the cost of the MRS-index structure and the NFA technique. In this experiment the query strings are selected from the chr18 dataset. The MRS-index structure performed up to 12 times faster than the NFA technique. The performance of the MRS-index structure improved when the queries are selected from different data strings. This is because the DNA strings have a high self similarity. Therefore, if the query string is chosen from the data string itself, the likelihood of having more number of matches within a specified range increases. The performance of the MRS index structure deteriorates as the error rate increases. This is because the size of the candidate set increases as the error rate increases.

#### 5 Discussion

In this paper, we considered the problem of searching similar strings in a database consisting of very long strings. The distance between the strings is defined as the minimum number of edit operations (insert, delete, and replace)

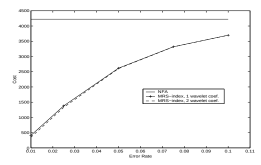


Figure 13: The cost of our method with one and two wavelet coefficients and the NFA technique for range queries of different ranges on the chr18 dataset. The query strings are chosen from the chr22 dataset.

to transform one string to the other.

We transformed the strings to an integer space by mapping them to their frequency vectors. Later, we generalized this idea to map the strings to their local frequencies for different resolutions by using a wavelet transform. We presented an efficient algorithm to find a lower bound on the distance between the wavelet coefficients of the strings.

We adapted the MR-index [13] to cluster the wavelet coefficients of the string. We called this index structure the MRS-index structure. This index structure slides a window on the data string for different resolutions. For each resolution, the index structure clusters the wavelet coefficients of a fixed number (box capacity) of consecutive substrings in an MBR. The MRS-index structure is dynamic, and allows arbitrary length queries.

We presented algorithms for both range queries and nearest neighbor queries. The range query algorithm splits the query into subqueries of available resolutions and performs successive range reduction for each subquery without using the data strings. The k-nearest neighbor algorithm runs in two phases. In the first phase, the distance of the query to the MBRs are computed, and the distance to the  $k^{th}$  closest subsequence in the k closest MBRs is used as the radius for a range query in the second phase. This technique can be used as a preprocessing to speed up any string search technique including BLAST by pruning large amounts of the data strings.

According to our experimental results with four different human chromosomes, our method runs up to 45 times faster than the NFA technique for 10 nearest neighbor queries. The MRS-index structure works efficiently for up to 200 nearest neighbors. Similarly, the MRS-index structure runs efficiently for range queries if the error rate is less than 0.1. The MRS-index structure performs up to 12 times better than other techniques for range queries with error rate 0.01.

In the future, we would like to model other distance measures in which different character pairs have different costs, or where the gaps are affine. Finding high local similarities within the substrings of the query string to the data strings is also an interesting problem. Comparison of the quality of our results with existing software packages is also planned for future.

## References

- [1] www.ncbi.nlm.nih.gov.
- [2] S. Altschul and B. W. Erickson. Optimal sequence alignment using affine gap costs. J. Molecular Biology, 1986.
- [3] S. Altschul and W. Gish. Basic local alignment search tool. J. Molecular Biology, 1990.
- [4] R. A. Baeza-Yates and G. Navarro. A practical index for text retrieval allowing errors. In *CLEI*, volume 1, pages 273–282, November 1997.
- [5] R. A. Baeza-Yates and G. Navarro. Faster approximate string matching. *Algorithmica*, 23(2):127–158, 1999.
- [6] R. A. Baeza-Yates and C. H. Perleberg. Fast and practical approximate string matching. In *Combinatorial Pattern Matching, Third Annual Symposium*, pages 185–192, 1992.
- [7] D.A. Benson, M.S. Boguski, D.J. Lipman, J. Ostell, and B.F. Ouellette. Genban. *Nucleic Acids Research*, 26(1):1–7, 1998.
- [8] D.A. Benson, I. Karsch-Mizrachi, D.J. Lipman, J. Ostell, B.A. Rapp, and D.L. Wheeler. Genbank. *Nucleic Acids Research*, January 2000.
- [9] P. Ferragina and G. Manzini. An experimental study of an opportunistic index. In SODA, pages 269–278, Washington, DC, 2001.
- [10] E. Giladi, M. G. Walker, J. Z. Wang, and W. Volkmuth. Sst: An algorithm for searching sequence databases in time proportional to the logarithm of the database size. In *RECOMB*, Japan, 2000.
- [11] D. Gusfield. Algorithms on Strings, Trees, and Sequences: Computer Science and Computational Biology. Cambridge University Press, 1 edition, January 1997.
- [12] S. Henikoff and J. G. Henikoff. Amino acid substitution matrices from protein blocks. In PNAS, pages 10915–10919, 1989.
- [13] T. Kahveci and A. Singh. Variable length queries for time series data. In *ICDE*, Heidelberg, Germany, 2001.
- [14] F. Korn, N. Sidiropoulos, C. Faloutsos, E. Siegel, and Z. Protopapas. Fast nearest neighbor search in medical databases. In VLDB, pages 215–226, India, 1996.
- [15] U. Manber and E. Myers. Suffix arrays: A new method for on-line string searches. SIAM Journal on Computing, 22(5):935–948, 1993.
- [16] S. Muthukrishnan and S. C. Sahinalp. Approximate nearest neighbors and sequence comparison with block operations. In STOC, Portland, Or, 2000.
- [17] E. Myers. An O(ND) difference algorithm and its variations. Algorithmica, pages 251–266, 1986.
- [18] E. Myers. A sublinear algorithm for approximate keyword matching. Algorithmica, pages 345–374, 1994.
- [19] T. Seidl and H.P. Kriegel. Optimal multi-step k-nearest neighbor search. In SIGMOD, 1998.
- [20] T.F. Smith and M.S. Waterman. Identification of common molecular subsequences. *Journal of Molecular Biology*, March 1981.
- [21] S. Uliel, A. Fliess, A. Amir, and R. Unger. A simple algorithm for detecting circular permutations in proteins. *Bioinformatics*, 15(11):930–936, 1999.
- [22] S. Wu and U. Manber. Fast text searching allowing errors. Communications of the ACM, 35(10):83–91, 1992.