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# A New String Matching Algorithm for Searching Biological Sequences

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### **Abstract**

String matching algorithms play a key role in many computer science problems, and in the implementation of computer software. This problem has received, and continues to receive a great deal of attention due to various applications in text manipulation, information retrieval, speech recognition, image and signal processing and computational biology. In this study, we propose a new algorithm called the Odd and Even algorithm (OE). OE combines an enhanced preprocessing phase from the Berry Ravindran algorithm with our proposed new searching phase procedure. This variety of searching order allows our proposed algorithm to reduce the number of comparison characters and enhances the searching response time. Experimental results show that OE algorithm offers a smaller number of comparisons and offers improved elapsed searching time when compared to other well-known algorithms for searching any length of alphabets and patterns. The proposed algorithm is applicable to searching protein sequence databases as well as any other string searching applications.

### 1. Introduction

Protein data can be found in many different forms such as sequences data, structure data, microarray data, and image data. Proteins are fundamental to the structure and the function of all living cells and viruses. Protein compounds are made of 20 different amino acids arranged in a linear chain. They are "complex organic compounds that consist of amino acids joined by peptide bonds" [1]. Computational biology and chemistry that use computational methods handle large amount of data. Protein sequence technologies have produced many extremely large sets of biological data which need faster techniques to process them.

The Swiss-Prot database is one of the main protein sequence databases containing descriptions of protein functions, domain structures, post-translational modifications and variants with a low level of redundancy and a high level of integration with other databases [2].

String-matching algorithms aim to find all occurrences of a given pattern  $P=p_1p_2...p_m$  in a text  $T=t_1t_2...t_n$ . They work as follows: they first align the left ends of the

pattern and the text, then compare text characters with pattern characters and after a mismatch between the pattern and the text or a whole match between them they shift the pattern to the right. This procedure is repeated until the right end of the pattern reaches the right end of the text

Most string-matching algorithms consist of a preprocessing phase and a searching phase to search for the pattern in the given text. The preprocessing phase analyses the characters in the pattern in order to use this information to determine the pattern shift in case of a mismatch or a whole match, with the aim of reducing the total number of character comparisons, while the searching phase defines the order of comparison of characters in each attempt between the pattern and the text. The main aim in algorithm development is to decrease the searching phase during each attempt and to increase the shifting value of the pattern.

String matching algorithms can be classified into seven categories according to the preprocessing function in the algorithm [3]. The first category, e.g. the Brute Force algorithm (BF) [4], shifts the pattern only one position at each attempt. The second category, which includes the Boyer-Moore algorithm (BM) [5]-[7] and the Fast Search algorithm (FS) [8], uses two preprocessing functions. The third category, a good example of which is the Boyer Moore Horspool algorithm (BMH) [9]-[11], uses one preprocessing function based on the rightmost character in the current window. The fourth category, e.g. the Quick Search algorithm (QS) [12], uses one preprocessing function based on the character next to the current window. The fifth category, such as the Berry-Ravindran algorithm (BR) [13], uses one preprocessing function based on the next two characters to the current window. The sixth category, e.g. the Karp-Rabin algorithm (KR) [14] and the Zhu Takaoka algorithm (ZT) [15], uses a preprocessing hashing function. The final category uses hybrid algorithms and includes the SSABS [16], TVSBS [17], ZTMBH [18], BRFS [19], BRBMH [20] and the BRQS [3] algorithms.

The paper is organized as follows: section II includes a survey of the main string-matching algorithms. Section III describes the proposed algorithm and its two main phases. Section IV provides a working example. Section V includes the experimental results with an evaluation of our new algorithm comparing it to other common string-

matching algorithms. Finally the conclusion is presented in section VI.

# 3. Proposed algorithm

### 3.1. Preprocessing phase

In this phase, the proposed algorithm uses our enhanced brBc preprocessing function by counting the shifting values for each character in the pattern and storing them in the one-dimensional brBc array [20]. Fig. 1 shows the pseudo code for the pre-processing phase.

#### /\*Pre-Processing Phase\*/

FOR i=0 TO m-2
SET brBcShiftArray[i] TO m-i

---- --

END FOR

 $IF \ t[end]+1 = p[end] \ THEN$ 

SET shiftvalue to 1

 $ELSE\ IF\ t[end]+2=p[start]\ THEN$ 

 $COMPUTE\ shift value\ AS\ m+1$ 

ELSE

COMPUTE shiftValue AS m+2

END IF

Fig. 1 OE Pre-processing Phase

## 3.2. Searching phase

After implementing several algorithms, we found out that the best order in the searching phase is to compare the pattern and the text window characters from right to left.

Our proposed algorithm searches the pattern from right to left with new order. It starts with the last character of the text window and the pattern, and after a match, it moves backward to compare the odd index positions of pattern and text window characters. If all these characters match, it will return and compare whole even index pattern and text window characters. In case of a mismatch or whole match during the comparison in odd or even positions it uses our enhanced brBc preprocessing function to shift the pattern. Fig. 2 shows the pseudo code for the searching process phase.

## 4. Working example

A sample file has been taken from the Swiss-Prot database which consists of 8740 proteins [23]. The following example illustrates our proposed algorithm:

#### /\*Searching Phase\*/

WHILE odd>=t[start] AND p[odd]=t[odd]

DECREMENT odd - 2

**ENDWHILE** 

IF odd > t[start] THEN

WHILE even>=t[start] AND p[even]=t[even]

DECREMENT even – 2

**ENDWHILE** 

ELSE

SET notMatch TO true

 $SET\ textPortion\ TO(\ t[end]\ +1)+\ (\ t[end]\ +2)$ 

CALL brBcShiftArray WITH textPortion

END IF

 $IF\ whole Match = true\ THEN$ 

CALL brBcShiftArray WITH textPortion

END IF

END WHILE

### /\*Searching in the brBcShiftArray\*/

FOR i=0 TO m

IF p[i] = textPortion THEN

SET shiftValue TO brBcShiftValue

END IF

END FOR

Fig. 2 OE Searching Phase

Given:

Pattern(p)="LAVKLATAIVLA", length (m) =12

 $\label{eq:text} Text(n) = \text{``KRFDSLYKQILAMGIFSIANQHIVLAVK} \\ LATAIVLATHTSPVVPVTTPGTKPDLNASFVSANAE \\ \text{''}, length(n) = 64$ 

### 4.1. Preprocessing phase

The shift values for the pattern characters are calculated according to Fig. 1. Table 1 shows the brBc one-dimensional array for the pattern characters.

### 4.2. Searching phase

The searching phase in this example is implemented according to Fig. 2. The following tables illustrate the searching phase for the given pattern (p) in the sample text (t).

LA	AV	VK	KL	LA	AT	TA	ΑI	IV	VL	LA
12	11	10	9	8	7	6	5	4	3	2

Table 1: The preprocessing phase

**4.2.1. Attempt 1:** in this attempt, Table 2 shows that  $t_0 - t_{11}$  is the current text that is compared with the pattern  $p_0 - p_{11}$ . The  $t_{11}$  comparison with  $p_{11}$  has matched, so the algorithm will move backward to the next odd index which compares  $t_9$  to  $p_9$  which causes a mismatch. The

pattern will be shifted to the right according to the precounted shifting value for the next two characters of the current window which are t<sub>12</sub> and t<sub>13</sub> (MG) and in this attempt will shift by 14 positions.

0	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17
K	R	F	D	S	L	Y	K	Q	I	L	A	M	G	Ι	F		
									2		1						
L	Α	V	K	L	Α	T	A	I	V	L	A						

Table 2: Attempt 1 in searching phase

**4.2.2. Attempt 2:** in this attempt, Table 3 shows that  $t_{14} - t_{25}$  is the current text which is compared with the  $p_0 - p_{11}$ . The  $t_{25}$  comparison with  $p_{11}$  has matched, so the algorithm will move backward to the next odd indices which are  $t_{23}$  and  $p_9$  which match. The next comparison

is between  $t_{21}$  and  $p_7$  which causes a mismatch. The pattern will be shifted to the right according to the precounted shifting value for the next two characters of the current window which are  $t_{26}$  and  $t_{27}$  (VK) giving a shift of 10 positions.

14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	29	30	31
I	F	S	I	A	N	Q	Н	I	V	F	Α	V	K	L	A		
							3		2		1						
L	Α	V	K	L	Α	T	Α	I	V	L	A						

Table 3: Attempt 2 in searching phase

**4.2.3. Attempt 3:** in this attempt, Table 4 shows that  $t_{14} - t_{25}$  is the current text that is compared with the  $p_0 - p_{11}$ . The first comparison between  $t_{35}$  and  $p_{11}$  produces a match, so the algorithm will move backward to the next odd indexes which are  $t_{33}$  and  $p_9$  which produces a match again. The next comparison is between  $t_{31}$  and  $p_7$  which also produces a match. Then the same procedure is repeated until the all the odd indices match. It will

then go back to the first even indexes (from the right) which they are  $t_{34}$  and  $p_8$ . This produces a match also and it will proceed to move back to compare further even indices. After a whole match between pattern and text it shifts the pattern to the right according to the precounted shifting value for the next two characters to the current window which are  $t_{36}$  and  $t_{37}$  (TH). In this attempt it will be 14 positions.

24	25	26	27	28	29	30	31	32	33	34	35	36	37	38	39	40	41
L	Α	V	K	L	Α	T	A	I	V	L	A	T	Н	T	S		
12	6	11	5	10	4	9	3	8	2	7	1						
L	Α	V	K	L	Α	T	A	I	V	L	Α						

Table 4: Attempt 3 in searching phase

**4.2.4. Attempt 4:** in this attempt, Table 5 shows that  $t_{38} - t_{49}$  is the current text that is compared with the  $p_0 - p_{11}$ . The comparison of  $t_{49}$  with  $p_{11}$  causes a mismatch. The pattern will be shifted to the right according to the pre-counted shifting value for the two characters next to

the current window which are  $t_{50}$  and  $t_{51}$  (KP) and in this attempt it will be 14 positions. But in this case the algorithm will cancel the pattern shifting since the length of the remaining text is 13 which is less than the pattern length.

38	39	40	41	42	43	44	45	46	47	48	49	50	51	52	53	54	55
T	S	P	V	V	P	V	T	T	P	G	T	K	P	D	L		
											1						
L	Α	V	K	L	Α	T	Α	I	V	L	A						

Table 5: Attempt 4 in searching phase

## 6. Experimental results

To evaluate our new algorithm, we implemented it based on the code in Fig.1 and Fig.2. Additionally, we implemented six other algorithms; two of them are our previous algorithms using the enhanced brBc preprocessing phase; two developed in 2008 using the original BR as the fastest algorithms to search the proteins; one of them using the rightmost character in the current window and the last one using only one character next to the current text window.

The performance of the algorithm proposed in this research is evaluated using the number of comparison between the pattern and the text and the elapsed time of searching.

A sample file has been taken from the Swiss-Prot database which consists of 8740 proteins to test the efficiency of our algorithm compared to other algorithms. Table 6 below shows the number of comparison and Fig. 3 below shows the average elapsed time (s.) for searching different length of patterns in the protein sample file.

Pattern Length	OE	BRQS	BRMH	BRFS	TVSBS	QS	ВМН
32	95384	95498	95595	96356	95682	172936	161089
64	50973	51171	51202	52101	51258	133723	113597
128	26985	27099	27180	27388	27214	87426	59229
256	10012	10040	10058	11925	10075	45394	38005
512	2950	2978	2987	3186	2997	16120	8502
1024	1233	1235	1239	1282	1243	2647	2186

Table 6: Number of comparison

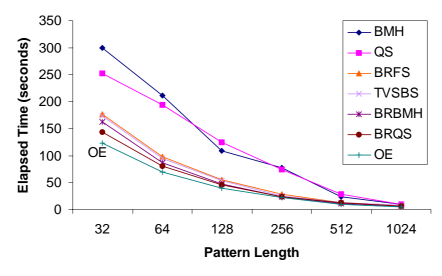


Fig. 3 average elapsed time (s.) for searching different length of patterns

Table 6 and Fig. 3 show that the number of comparisons and the elapsed searching time between the pattern and the text using our proposed algorithm is better in all cases than other algorithms.

### 7. Conclusion

In this paper, we have presented a new algorithm. The OE algorithm is a fast string matching algorithm. It combines our enhanced preprocessing phase from the Berry Ravindran algorithm with our new searching phase procedure. Experimental results show that our algorithm uses fewer comparisons to perform searches and has a shorter elapsed searching time. Our proposed algorithm is therefore suitable for searching the protein sequences in the Swiss-Prot database as well as in any other string searching applications.

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