5 BLAST and Aho-Corasick

This exposition was prepared by Clemens Gröpl, based on versions by Daniel Huson and Knut Reinert. It is based on the following sources, which are all recommended reading:


5.1 BLAST, FASTA, ...

Pairwise alignment is used to detect homologies between different protein or DNA sequences, either as global or local alignments.

This can be solved using dynamic programming in time proportional to the product of the lengths of the two sequences being compared.

However, this is too slow for searching current databases and in practice algorithms are used that run much faster, at the expense of possibly missing some significant hits due to the heuristics employed.

Such algorithms are usually on seed and extend approaches in which first small exact matches are found, which are then extended to obtain long inexact ones.

5.2 Some BLAST terminology

BLAST, the Basic Local Alignment Search Tool, is perhaps the most widely used bioinformatics tool ever written. It is an alignment heuristic that determines "local alignments" between a query q and a database d.

A segment is simply a substring s of q or d.

A segment-pair \((s, t)\) consists of two segments, one in q and one in d, of the same length.

\[
\begin{align*}
V & A L L A R \\
P & A M M A R \\
\end{align*}
\]

We think of s and t as being aligned without gaps and score this alignment using a substitution score matrix, e.g. BLOSUM or PAM.

The alignment score for \((s, t)\) is denoted by \(\sigma(s, t)\).

A locally maximal segment pair (LMSP) is any segment pair \((s, t)\) whose score cannot be improved by shortening or extending the segment pair.

Given a cutoff score C, a segment pair \((s, t)\) is called a high-scoring segment pair (HSP), if it is locally maximal and \(\sigma(s, t) \geq C\).
Given a cutoff score, the goal of BLAST is to compute all high-scoring segment pairs.

### 5.3 BLAST algorithm for protein sequences

A *word* is simply a short substring.

BLAST computes high-scoring segments pairs using short words as *seeds*. Exact matches of words are searched using a special algorithm. Then the short matches are *extended* to both sides, leading to locally maximal segment pairs.

All this is fine-tuned by a couple of parameters: the *word size* $w$, the *similarity threshold* $T$ used when generating the list of short words to be searched for, and a minimum match score $C$ which applies to HSPs.

The BLAST algorithm for *protein* sequences operates as follows:

1. The list of all words of length $w$ that have similarity $\geq T$ to some word in the query sequence $q$ is generated.
2. The database sequence $d$ is scanned for all *exact matches* $t$ of words $s$ in the list.
3. Each such *seed* $(s, t)$ is *extended* until its score $\sigma(s, t)$ falls a certain distance below the best score found for shorter extensions. All extensions are reported that have score $\geq C$.

In practice, $w$ is around 3 for proteins.

**Example.**

Using words of length 2:

```
Initial Words
ql
ln
nf
fs
sa
ag
gw
```

Expanding the initial word list:

<table>
<thead>
<tr>
<th>Initial Words</th>
<th>Expanded List</th>
</tr>
</thead>
<tbody>
<tr>
<td>ql</td>
<td>ql, qm, hl, zl</td>
</tr>
<tr>
<td>ln</td>
<td>ln, lb</td>
</tr>
<tr>
<td>nf</td>
<td>nf, af, ny, df, qf, ef, gf, hf, kf, sf, tf, bf, zf</td>
</tr>
<tr>
<td>fs</td>
<td>fs, fa, fn, fd, fg, fp, ft, fb, ys</td>
</tr>
<tr>
<td>sa</td>
<td>(nothing scores 8 or higher)</td>
</tr>
<tr>
<td>ag</td>
<td>ag</td>
</tr>
<tr>
<td>gw</td>
<td>gw, aw, rw, nw, dw, qw, ew, hw, hw, iw, kw, mw, pw, sw, tw, vw, bw, zw, xw</td>
</tr>
</tbody>
</table>
(1) For the query find the list of high scoring words of length $w$.

For each word from the query sequence find the list of words that will score at least $T$ when scored using a pairscore matrix (e.g. PAM 250). For typical parameters there are around 50 words per residue of the query.

(2) Compare the word list to the database and identify exact matches.

With a careful implementation, the list of all words of length $w$ that have similarity $\geq T$ to some word in the query sequence $q$ can be produced in time proportional to the number of words in the list.

The similar words are immediately placed in a keyword tree and then, for each word in the tree, all exact locations of these words in the database $d$ are detected in time linear to the length of $d$, using a variation of the Aho-Corasick algorithm for multiple exact string matching.

As BLAST does not allow indels, also the hit extension is very fast.

Note that the use of seeds of length $w$ and the termination of extensions with fading scores are both steps that speed up the algorithm, but also imply that BLAST is not guaranteed to find all HSPs.
5.4 BLAST algorithm for DNA sequences

For DNA sequences, BLAST operates as follows:

- The list of all words of length \( w \) in the query sequence \( a \) is generated.
- The database \( d \) is scanned for all hits of words in this list. Blast uses a two-bit encoding for DNA. This saves space and also search time, as four bases are encoded per byte.

In practice, \( K \) is around 12 for DNA.

5.5 The BLAST family

There are a number of different variants of the BLAST program:

- BLASTN: compares a DNA query sequence to a DNA sequence database,
- BLASTP: compares a protein query sequence to a protein sequence database,
- TBLASTN: compares a protein query sequence to a DNA sequence database (6 frames translation),
- BLASTX: compares a DNA query sequence (6 frames translation) to a protein sequence database, and
- TBLASTX: compares a DNA query sequence (6 frames translation) to a DNA sequence database (6 frames translation).

BLAST is constantly being developed further and many internet services is available, e.g.,
http://blast.wustl.edu/

5.6 Multiple string matching

The task at hand is to find all occurrences of a given set of \( r \) patterns \( P = \{p^1, \ldots, p^r\} \) in a text \( T = t_1, \ldots, t_n \). Each \( p^i \) is a string \( p^i = p^i_1, \ldots, p^i_{m_i} \). Usually \( n \) is much bigger than \( m_i \).

Idea of the Aho-Corasick algorithm:

The Aho-Corasick algorithm belongs to the class of \textit{prefix-based} approaches. We assume that we have read the text up to position \( i \) and that we know the length of the longest suffix of \( t_1, \ldots, t_i \) that is also a prefix of some pattern \( p^i \in P \).

The Aho-Corasick algorithm maintains a data structure, called the \textit{Aho-Corasick automaton}, to keep track of the longest prefix of some pattern that is also a suffix of the text window.

The AC automaton is built on top of another data structure called trie, so we explain this one first.
5.7 The trie data structure

A trie is a compact representation of a set of strings; in our case, the set $P$. It is a rooted, directed tree. The path label of a node $v$ is the string $L(v)$ read when traversing the trie from the root to the node $v$. A node is a terminal node if its path label is within the set of strings to be represented by the trie. Each $p' \in P$ corresponds to a terminal node.

Example: $P = \{\text{annual}, \text{annually}, \text{announce}\}$

In the following pseudocode, $\delta(current, p'_j)$ denotes the successor state reached from state $current$ by the outgoing edge that is labeled with $p'_j$. The symbol $\theta$ means “undefined”.

The set of terminal states is denoted $F$ (for “final”). (In the code each state has a set $F$ of indices indicating that it is terminal for the respective strings.)

The size of the trie and the running time of its basic operations depend on the implementation of the transition function $\delta$.

- We can provide each node with a table of size $|\Sigma|$. This yields access in $O(1)$, but may still be worse in practice because the high space consumption will lead to many processor (L1) cache misses.
- We can use a sorted array of size $\sum_{p \in P} |p|$ and store only the existing edges. The edge list is accessed using binary search which takes $O(\log |\Sigma|)$ time per transition.
- In practice, a hash table will be the best compromise for most applications. It uses $O(\sum_{p \in P} |p|)$ space and usually $O(1)$ time per access (but $O(\sum_{p \in P} |p|)$ time in the worst case).
5.8 The Aho-Coracick automaton

The Aho-Coracick automaton augments the trie for $P$ with a supply function. Except for the root it holds that for each node $x$, the supply link $S(x)$ points to a node $y$ such that $L(y)$ is the longest proper suffix of $L(x)$ that is represented by a trie node (that means of some string in the input set). $S(x)$ can be the root. The root represents the empty string, i.e., $L(root) = \epsilon$.

The supply links can be computed in $O(\sum_{k}|p_{k}|)$ time while constructing the trie. They are used to perform safe shifts in the Aho-Coracick algorithm.

Here is an example. The patterns are

$$P = \{aab, ababc, abc, bab\}$$

Let’s conduct a multiple string matching on

$$T = abaababcabbb$$

You can see that it really pays off not to search individually for the four strings.

```plaintext
/\ n[0] = length of pat[0]  \\
/\ n[1] = number of patterns  \\
final = empty set
FOR a = 1 TO n[1]  \\
  i = root
  FOR j = 1 TO n[0]  \\
    IF pat[j] eq pat[0][j] = NULL  \\
      index = pat[j][j]
    ELSE
      index = -1
    ENDIF
    /\ visualization point /
    push(index, i)
    final = union(final, i)
  ENDFOR
  /\ visualization point /
  parent(index) = root
  /\ visualization point /
ENDFOR
```

Code to create trie

```plaintext
h(root) = fail
ENDWHILE
IF e == fail
  h(e) = g(root, e)
  /\ visualization point /
ELSE
  h(e) = g(e)
  IF g(e) eq final
    final = union(final, e)
  ENDIF
  /\ visualization point /
ENDEWHILE
```

Code to add supply links
A “translation table” for the animation

<table>
<thead>
<tr>
<th>Animation code</th>
<th>Our code</th>
</tr>
</thead>
<tbody>
<tr>
<td>$q'$ (green)</td>
<td>current</td>
</tr>
<tr>
<td>$q$ (orange)</td>
<td>parent</td>
</tr>
<tr>
<td>$h$</td>
<td>$S$</td>
</tr>
<tr>
<td>$r$ (pink)</td>
<td>down</td>
</tr>
<tr>
<td>fail</td>
<td>$\theta$</td>
</tr>
<tr>
<td>$c$ (cyan)</td>
<td>$\sigma$</td>
</tr>
</tbody>
</table>

Matches: abaababcbabcbb
         aab
         ababc
         abc abc
         bab bab

Note that when the match for ababc is found, we have to report an occurrence of abc as well. One pattern can be a suffix of another.

The discussion how we just built the example together with the pseudocode follows.

First the trie is constructed, then it is augmented with the supply links by a breadth-first search (= BFS traversal.) See the pseudocode below.

Inductively can assume that we have already computed the supply links of all states before state current in BFS order. Now consider the parent parent of current in the trie and assume that it leads to current via the symbol $\sigma$. That means, $\delta($parent, $\sigma$) = current. The supply link for parent has already been computed. It points to $S($parent$)$.

Let $v = L($current$)$ and let $u$ be the longest proper suffix of $v$ that labels a path in the trie, or let $u = \epsilon$ (the empty string). The string $v$ has the form $v'\sigma$, for some string $v'$. In fact, we have $v' = L($parent$)$.

Thus, if $u$ is non-empty, it must have the form $u = u'\sigma$, for some string $u'$, and in that case $u'$ is a proper suffix of $v'$ that is the label of a path in the trie. In fact, we have $u' = L($parent$)$.

Now, if $S($parent$)$ does have an outgoing transition labeled by $\sigma$ to a state $h$, i.e., $\delta($parent, $\sigma$) = $h$, then $u'\sigma = L($parent$)\sigma$ is the longest suffix of $v$ that is the label of a path, namely $u'\sigma = L(h)$. Hence $S($current$)$ has to be set to $h$.

If $S($parent$)$ does not have an outgoing transition labeled by $\sigma$, then we consider the second-longest proper suffix of $v'$ that is the label of a path. That is, we consider $S($parent$)$, and if that fails, then $S($parent$)$, and so on.
We repeat that until either we find a non-empty suffix of $u$ that has an outgoing transition labeled with $\sigma$ or we reach the initial state.

1. Build $AC(P = p_1, \ldots, p_r)$;
2. $ACtrie = Trie(P)$; $\delta$ is transition function, root initial state
3. $S(init) = \emptyset$;
4. for current in BFS order do
   5. $parent = par(current)$;
   6. $\sigma$ = label of transition from parent to current;
   7. $down = S(parent)$;
   8. while $down \neq \emptyset$ and $\delta(down, \sigma) = \emptyset$ do
      9. $down = S(down)$;
   10. od
   11. if $down \neq \emptyset$ then
      12. $S(current) = \delta(down, \sigma)$;
      13. if $S(current) = terminal$ then
         14. mark current as terminal;
         15. $F(current) = F(current) \cup F(S(current))$;
      16. fi
   17. else
      18. $S(current) = root$;
   19. fi
   20. od

Searching in the Aho-Corasick automaton is straightforward and very similar to the building phase. While scanning the text, we walk through the automaton. Whenever we enter a terminal state we report the set of strings as matching. If we cannot walk on, we follow supply links to find a safe shift.

5.9 Running time

We start with the search phase, since the time bound is a bit easier to obtain for it than for the preprocessing.

Lemma. The search phase for the Aho-Corasick algorithm (just reporting whether there exists a match or not) takes $O(|T|)$ time.

Proof. We need to bound the number of transitions within the AC automaton. Let $\sigma$ be the next letter in $T$. If the transition to $\delta(current, \sigma)$ is defined, then $L(current)$ increases by 1. Otherwise, $L(current)$ will become strictly smaller. Since $0 \leq |L(current)|$, it follows that, during the whole search over $T$, at most $|T|$ times a supply link is being used – we cannot walk more often upward than downward in the automaton.

Lemma. The preprocessing for the Aho-Corasick algorithm takes $O\left(\sum_{p \in P} |p|\right)$ time.

Proof. The argument is the similar to the search argument. After the construction of the trie, we conduct a BFS search. Both clearly need time $O\left(\sum_{p \in P} |p|\right)$.

During the construction of the supply links, we process a total of $\sum_{p \in P} |p|$ characters. For each processed character we spend constant work except for following already inserted supply links. So in total, we cannot follow more supply links than there are characters in the patterns.

As a final comment, one can implement the set union in line (15) by a simple concatenation of lists in constant time, since the sets are disjoint.
5.10 ‘Advanced’ Aho-Corasick algorithm

We can further speed up the search phase the AC algorithm if we are willing to spend more time for an extended preprocessing.

Note that it is possible to precompute all transitions implied by the supply links in advance. Essentially, the formula is

$$\delta(\text{current}, \sigma) = \delta(S(\text{current}), \sigma)$$

for all $\sigma$ where $\delta$ would otherwise be undefined.

The extended preprocessing pays off for relatively small sets of patterns and for small alphabets. However, we need more space to represent $\delta$.

A compromise is to compute the additional edges “on the fly”, when they are “discovered” for the first time. This technique is also known as path compression. We can simply add the compressed transitions to the hash table representing $\delta$. Then the next time we will not need to follow supply links, but will be directly forwarded to the destination. This strategy was implemented in the first version of the well-known unix application grep.
6 Fast filtering algorithms

This exposition is based on the following sources, which are all recommended reading:

2. Burkhardt et al.: q-gram Based Database Searching Using a Suffix Array (QUASAR), RECOMB 99

We will present the hierarchical filtering approach of Navarro and Baeza-Yates and the simple QUASAR idea.

6.1 Filtering algorithms

The idea behind filtering algorithms is that it might be easier to check that a text position does not match a pattern string than to verify that it does.

Filtering algorithms filter out portions of the text that cannot possibly contain a match, and, at the same time, find positions that can possibly match.

These potential match positions then need to be verified with another algorithm like for example the bit-parallel algorithm of Myers (BPM).

Filtering algorithms are very sensitive to the error level $\alpha := k/m$ since this normally affects the amount of text that can be discarded from further consideration. ($m =$ pattern length, $k =$ errors.)

If most of the text has to be verified, the additional filtering steps are an overhead compared to the strategy of just verifying the pattern in the first place.

On the other hand, if large portions of the text can be discarded quickly, then the filtering results in a faster search.

Filtering algorithms can improve the average-case performance (sometimes dramatically), but not the worst-case performance.

6.2 The pigeonhole principle

The idea behind the presented filtering algorithm is very easy. Assume that we want to find all occurrences of a pattern $P = p_1, \ldots, p_m$ in a text $T = t_1, \ldots, t_n$ that have an edit distance of at most $k$.

If we divide the pattern into $k + 1$ pieces $P = p^1, \ldots, p^{k+1}$, then at least one of the pattern pieces has to match without error.

There is a more general version of this principle first formalized by Myers in 1994:

Lemma 1. Let $\text{Occ}$ match $P$ with $k$ errors, $P = p^1, \ldots, p^j$ be a concatenation of subpatterns, and $a_1, \ldots, a_j$ be nonnegative integers such that $A = \sum_{i=1}^j a_i$. Then, for some $i \in 1, \ldots, j$, $\text{Occ}$ includes a substring that matches $p^i$ with $\lfloor a_i k/A \rfloor$ errors.
Proof: Exercise.

So the basic procedure is:

1. **Divide**: Divide the pattern into \( k + 1 \) pieces of approximately the same length.

2. **Search**: Search all the pieces simultaneously with a multi-pattern string matching algorithm. According to the above lemma, each possible occurrence will match at least one of the pattern pieces.

3. **Verify**: For each found pattern piece, check the neighborhood with a verification algorithm that is able to detect an occurrence of the whole pattern with edit distance at most \( k \). Since we allow indels, if \( p_1 \ldots p_{i_2} \) matches the text \( t_{j-(i_1-1)-k} \ldots t_{j+(m-i_2)+k} \), then the verification has to consider the text area \( t_{j-(i_1-1)-k} \ldots t_{j+(m-i_2)+k} \), which is of length \( m + 2k \).

### 6.3 An example

Say we want to find the pattern `annual` in the texts

\[ t_1 = \text{any annealing} \] and \[ t_2 = \text{an unusual example with numerous verifications} \]

with at most 2 errors.

1. **Divide**: We divide the pattern `annual` into \( p^1 = an \), \( p^2 = nu \), and \( p^3 = al \). One of these subpattern has to match with 0 errors.

2. **Search**: We search for all subpatterns:

   1: searching for `an`: in \( t_1 \): find positions 1, 5
   in \( t_2 \): find position 1
   2: searching for `nu`: in \( t_1 \): find no positions
   in \( t_2 \): find positions 5, 25
   3: searching for `al`: in \( t_1 \): find position 9
   in \( t_2 \): find position 9

3. **Verification**: We have to verify 3 positions in \( t_1 \) and 4 positions in \( t_2 \), to find 3 occurrences at positions (indexed by the last character) 9, 10, 11 in \( t_1 \) and none in \( t_2 \).

### 6.4 Hierarchical verification

The toy example makes clear that many verifications can be triggered that are unsuccessfull and that many subpatterns can trigger the same verification. Repeated verfications can be avoided by carefully sorting the occurrences of the pattern (exercise).

It was shown by Baeza-Yates and Navarro that the running time is dominated by the multipattern search for error levels \( \alpha = k/m \) below \( 1/(3 \log_{|\Sigma|} m) \). In this region, the search cost is about \( O(kn \log_{|\Sigma|} m) \). For higher error levels, the cost for verifications starts to dominate, and the filter efficiency deteriorates abruptly.

Baeza-Yates and Navarro introduced the idea of hierarchical verification to reduce the verification costs, which we will explain next. Then we will work out more details of the three steps.
Navarro and Baeza-Yates use Lemma ?? for a hierarchical verification. The idea is that, since the verification cost is high, we pay too much for verifying the whole pattern each time a small piece matches. We could possibly reject the occurrence with a cheaper test for a shorter pattern.

So, instead of directly dividing the pattern into \( k + 1 \) pieces, we do it hierarchically. We split the pattern first in two pieces and search for each piece with \( \lfloor k/2 \rfloor \) errors, following Lemma ?? . The halves are then recursively split and searched until the error rate reaches zero, i.e., we can search for exact matches.

With hierarchical verification the area of applicability of the filtering algorithm grows to \( \alpha < 1/\log_{|\Sigma|} m \), an error level three times as high as for the naive partitioning and verification. In practice, the filtering algorithm pays off for \( \alpha < 1/3 \) for medium long patterns.

**Example.** Say we want to find the pattern \( P = \text{aaabbbcccccddd} \) in the text \( T = \text{xxxbbbxxxxxx} \) with at most \( k = 3 \) differences. The pattern is split into four pieces \( p^1 = \text{aaa}, p^2 = \text{bbb}, p^3 = \text{ccc}, p^4 = \text{ddd} \). We search with \( k = 0 \) errors in level 2 and find bbb.

```
level 0 aaabbbcccccddd with k=3 errors
  /  \
level 1 aaabbb cccddd with k=1 errors
    /  \
level 2 aaa bbb ccc ddd with k=0 errors
```

Now instead of verifying the complete pattern in the complete text (at level 0) with \( k = 3 \) errors, we only have to check a slightly bigger pattern (aaabbb) at level 1 with one error. This is much cheaper. In this example we can decide that the occurrence bbb cannot be extended to a match.

```
level 0 aaabbbcccccddd with k=3 errors
  /  \
level 1 AAABBB cccddd with k=1 errors
    /  \
level 2 aaa BBB ccc ddd with k=0 errors
```

### 6.5 The PEX algorithm

**Divide:** Split pattern into \( k + 1 \) pieces, such that each piece has equal probability of occurring in the text. If no other information is available, the uniform distribution is assumed and hence the pattern is divided in pieces of equal length.

**Build Tree:** Build a tree of the pattern for the hierarchical verification. If \( k + 1 \) is not a power of 2, we try to keep the binary tree as balanced as possible.

Each node has two members \( \text{from} \) and \( \text{to} \) indicating the first and the last position of the pattern piece represented by it. The member \( \text{err} \) holds the number of allowed errors. A pointer \( \text{myParent} \) leads to its parent in the tree. (There are no child pointers, since we traverse the tree only from the leaves to the root.) An internal variable \( \text{left} \) holds the number of pattern pieces in the left subtree. \( \text{idx} \) is the next leaf index to assign. \( \text{plen} \) is the length of a pattern piece.

Algorithm CreateTree generates a hierarchical verification tree for a single pattern. (Lines ?? and ?? are justified by Lemma ??.)
CreateTree

Create new node node

Search:

Example:

Search phase of algorithm PEX
We search for annual in annual\_CPM\_anniversary. We constructed the tree for annual. A multi-pattern search algorithm finds: (1,1), (12,1), (3,2), (5,3). (Note that leaf i corresponds to pattern $p^{i+1}$). For each of these positions we do the hierarchical verification:

**Initialization for (1,1);**
n=n0; in=1; n=n3; cand=true;
While loop;
  a) $p1=1-(1-1)-1=0$; $p2=1+(4-1)+1=5$;  
    verify pattern annu in text annua with 1 error => found !  
  b) $p1=1-(1-1)-2=-1$; $p2=1+(6-1)+2=8$;  
    verify pattern annual in text annual\_C => found !  
  c) report end positions (6,7,8)

**Initialization for (3,2);**
n=n1; in=3; n=n3; cand=true;
While loop;
  a) $p1=3-(3-1)-1=0$; $p2=3+(4-3)+1=5$;  
    verify pattern annu in text annua with 1 error => found !  
  b) $p1=3-(3-1)-2=-1$; $p2=3+(6-3)+2=8$;  
    verify pattern annual in text annual\_C => found !  
  c) report end positions (6,7,8)

**Initialization for (12,1);**
n=n0; in=1; n=n3; cand=true;
While loop;
  a) $p1=12-(1-1)-1=11$; $p2=12+(4-1)+1=16$;  
    verify pattern annu in text _anniv with 1 error => found !  
  b) $p1=12-(1-1)-2=10$; $p2=12+(6-1)+2=19$;  
    verify pattern annual in text M\_annivers => NOT found !

### 6.6 Quasar

QUASAR, or "Q-gram Alignment based on Suffix ARrays", is a filtering approach. QUASAR finds all local approximate matches of a query sequence S in a database $D = \{d,...\}$. The verification is performed by other means.

**Definition.** A sequence d is locally similar to S, if there exists at least one pair $(S_{i,i+w-1}, d')$ of substrings such that:

1. $S_{i,i+w-1}$ is a substring of length w and $d'$ is a substring of $D$, and
2. the substrings $d'$ and $S_{i,i+w-1}$ have edit distance at most k.

We call this the approximate matching problem with k differences and window length w.

For simplicity, we assume that the database consists of only one sequence, i.e. $D = \{d\}$. 
6.7 The q-gram lemma

A short subsequence of length \( q \) is called a \( q \)-gram. In the following we start by considering the first \( w \) letters of \( S \). The algorithm uses the following lemma:

**Lemma 2.** Let \( P \) and \( S \) be strings of length \( w \) with at most \( k \) differences. Then \( P \) and \( S \) share at least \( w + 1 - (k + 1)q \) common \( q \)-grams.

In our case, this means:

**Lemma 3.** Let an occurrence of \( S_{1,w} \) with at most \( k \) differences end at position \( j \) in \( D \). Then at least \( w + 1 - (k + 1)q \) of the \( q \)-grams in \( S_{1,w} \) occur in the substring \( D_{j-w+1,j} \).

**Proof:** Exercise. . .

That means that as a necessary condition for an approximate match, at least \( t = w + 1 - (k + 1)q \) of the \( q \)-grams contained in \( S_{1,w} \) occur in a substring of \( D \) with length \( w \). For example the strings \( ACAGCTTA \) and \( ACACCTTA \) have \( 8 + 1 - (1 + 1)3 = 3 \) common 3-grams, namely \( ACA \), \( CTT \) and \( TTA \).

6.8 q-gram index

The algorithm builds in a first step an indexing structure as follows:

1. Build a suffix array \( A \) over \( D \).
2. Given \( q \), compute for all possible \( | \Sigma | \) \( q \)-grams the start position of the hitlist. This allows to lookup a \( q \)-gram in constant time.
3. If another \( q \) is specified, \( A \) is used to recompute the above table.
6.9 Counting q-grams

Now we have to find all approximate matches between $S_{1,w}$ and $D$, that means we have to find all substrings in $D$ that share at least $t$ $q$-grams with $S_{1,w}$. The algorithm proceeds in the following basic steps on which we will elaborate:

1. Define two arrays of non-overlapping blocks of size $b \geq 2w$. The first array is shifted by $b/2$ against the other.
2. Process all $q$-grams in $S_{1,w}$ and increment the counters of the corresponding blocks.
3. All blocks containing approximate matches will have a counter of at least $t$. (The reverse is not true).
4. Shift the search window by one. Now we consider $S_{2,w+1}$.

6.10 Blocking

Since we want to count the $q$-grams that are in common between the query and the database, we use counters. Ideally we would use a counter of size $w$ for each substring of this size. Since this uses too much memory, we build larger, non-overlapping blocks. While this decreases the memory usage, it also decreases the specificity.

Since the blocks are not overlapping we might miss $q$-grams that cross the block boundary. As a remedy, we use a second, shifted array of blocks.

6.11 Window Shifting

We started the search for approximate matches of window length $w$ with the first $w$-mer in $S$, namely $S_{1,w}$. In order to determine the approximate matches for the next window $S_{2,w+1}$, we only have to discard the old $q$-gram $S_{1,q}$ and consider the new $q$-gram $S_{w-q+2,w+1}$.

To do that we decrement the counters of all blocks that contain $S_{1,q}$ that have not reached the threshold $t$. However, if the counter has already reached $t$ it stays at this value to indicate a match for the extension phase.

For the new block we use the precomputed index and the suffix array to find the occurrences of the new $q$-gram and increment the corresponding block counters (at most two).

6.12 Alignment

After having computed the list of blocks, QUASAR uses BLAST to actually search the blocks. Here are some results from the initial implementation. QUASAR was run with $w = 50$, $q = 11$, and $t$ such that windows with at most 6% differences are found. Reasonable values for the block size are 512 to 4096.
A database in BLAST format is built in main memory which is then passed to the BLAST search engine. The construction of this database requires a significant amount of time and introduces unnecessary overhead.

### 6.13 Summary

- Filtering algorithms prevent a large portion of the text from being looked at.
- The larger $\alpha = \frac{k}{m}$, the less efficient filtering algorithms become.
- Filtering algorithms based on the pigeonhole principle need an exact, multi-pattern search algorithm and a verification capable approximate string matching algorithm.
- The PEX algorithm starts verification from short exact matches and considers longer and longer substrings of the pattern as the verification proceeds upward in the tree.

<table>
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<th>DB size</th>
<th>query</th>
<th>id. res.</th>
<th>filtr. ratio</th>
<th>QUASAR s</th>
<th>BLAST s</th>
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</thead>
<tbody>
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<td>73.5 Mb</td>
<td>368</td>
<td>91.4%</td>
<td>0.24%</td>
<td>0.123</td>
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<tr>
<td>280 Mb</td>
<td>393</td>
<td>97.1%</td>
<td>0.17%</td>
<td>0.38</td>
<td>13.27</td>
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</tbody>
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