MASTER THESIS

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Variant calling using local reference-helped assemblies

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Title: Variant calling using local reference-helped assemblies

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Abstract:

Despite active development during past years, the task of sequencing a genome still remains a challenge. Our current technologies are not able to read the whole genome in one piece. Instead, we shatter the target genome into a large amounts of small pieces that are then sequenced separately. The process of assembling these small pieces together, in order to obtain sequence of the whole genome, is painful and resource-consuming. Multiple algorithms to solve the assembly problem were developed.

This thesis presents yet another assembly algorithm, based on the usage of de Bruijn graphs, and focusing on sequencing short genome regions. The algorithm is compared to well-known solutions in the field.

Keywords: high-throughput sequencing variant calling local assembly de Bruijn graphs
Here, I would like to thank Mgr. Petr Danček, Ph.D. for supervising and helping me with the thesis in all other ways, and RNDr. Martin Kruliš, Ph.D. for allowing me to use a high-performance computer to speed up computations required by the thesis.
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1. Introduction

Despite technological advances in the past fifteen years, genome sequencing (determining the DNA sequence of an organism) remains a challenging task. With the current technology it is not possible to the DNA sequence in one piece from the beginning to the end. Instead, relatively short fragments must be sequenced individually and then painstakingly assembled into the complete sequence. This is a very laborious and expensive process, complicated by the fact that large proportion of most genomes consists of repetitive sequences. Therefore, instead of the pure de novo approach of assembling the complete sequence from scratch, most of the sequencing done today merely aims to determine differences between the sequenced organism and a complete, gold standard reference genome obtained previously by other means. This approach is known as resequencing and is, comparatively, a much simpler task.

The most widely used sequencing platform today is Illumina and the algorithms developed in this work assume Illumina data on input. The platform takes advantage of massive parallelism, where an ensemble of DNA molecules is fragmented into very short pieces (usually ~500 bp) and then sequenced in parallel from both ends. In one sequencing run, more than $10^9$ of short sequences are obtained simultaneously. These sequences are called reads. Due to technical limitations, reads are only ~150 bp long, but when they are mapped to the reference genome, many will overlap and cover the whole genome, thus enabling to determine the differences between the sequenced organism, such as single nucleotide polymorphisms (SNPs) or short insertions and deletions (indels). Collectively, these differences are referred to as variants and the process of determining the variants is known as variant calling.

In order to distinguish between random sequencing errors and real variants, we need sufficient coverage (the average number of reads mapped to a position of the genome). Thus for variant calling we need first to confidently place the reads to the correct location of the genome (the problem known as mapping), correctly determine the exact sequence of matches, mismatches, insertions and deletions at the location (alignment) and finally apply a statistical model to tell apart true variation from random sequencing errors, mapping errors and alignment errors (variant calling).

Single nucleotide variants are easier to call, indels are more problematic. This is partly because reads containing indels can be often aligned in multiple ways, leading to ambiguous alignments and false calls, especially in difficult parts of the genome of low complexity or high repeat content.

1.1 Basic Terms

For the rest of this thesis, a genome is viewed as a string of character, each represents a nucleotide, at certain position. Since DNA molecules consist of four types of nucleotides, only four characters, A, C, G or T, appear in the string. The four nucleotides have similar chemical structure. They are composed of a sugar molecule and a phosphate group, which are identical for all four of them, and a unique nitrogenous base. Hence the term base is often used when referring to a
nucleotide at a specific position. Each of the bases pairs with one other, A with T and C with G, therefore knowing the sequence of one strand of a DNA double helix determines the sequence of the complementary strand. Hence the terms base pair and base are used interchangeably.

The algorithm developed and discussed in this work expect a large bunch of short reads on input. It expects that the sequenced organism and the reference genome are similar enough to, so an assumption that reads are correctly mapped can be made. Its task is to resolve local differences arising from incorrect read alignments. This is a subproblem of the assembly one, called microassembly.

Some assembly algorithm transform each read into a sequence of k-mers, sub-strings of equal length, usually named k. The read sequence of length l is decomposed into \( l - k + 1 \) k-mers \( k_0, ..., k_{l-k} \) of length k. If we denote bases in the read as \( b_0, ..., b_{l-1} \), the k-mer \( k_i \) covers bases from \( b_i \) to \( b_{i+k-1} \). Such definition implies that adjacent k-mers overlap by \( k - 1 \) bases.

An example of transforming the sequence TACTGGCC into k-mers of length 3 is illustrated on Figure 1.1. The sequence has 8 bases in length and 6 k-mers are created from it. The read sequence, as in all other figures in this work, is marked red, contents of individual k-mers is in yellow.

Figure 1.1: Sequence transformation into k-mers

### 1.2 The Model

The simplest mathematical assembly model, assumes that we are assembling genome string of length \( g \), given a set of \( n \) reads of length \( l \) (\( l \ll g \)), each potentially transformed into a sequence of \( l - k + 1 \) k-mers. The model assumes that reads are sampled from the genome string uniformly and randomly and that they are error-free. Since the probability of sampling a base at certain genome position is very low for single sampling event, and the number of sampling events is large, genome coverage (i.e. the number of reads covering a position) follows a
Poisson distribution. In other words, the probability that base at certain position is covered by \( k \) reads is

\[
\frac{c^k}{k!} \times e^{-c}
\]

c is a base coverage depth, also known as sequencing depth, and can be computed simply as a total number of bases within the input read set divided by the length of the genome.

\[
c = \frac{n \times l}{g}
\]

Very similar relations apply for k-mer coverage depth, only the formula for the k-mer coverage depth needs to be changed to \( \frac{n \times (l-k+1)}{g-k+1} \). These details brings answers to at least two important questions. Primo, how many reads need to be created to (statistically) cover the whole genome string, and, secundo, how to determine whether a given read set is free of sequencing errors?

The percentage of genome not covered by any read from the set is equal to \( P(k = 0) = e^{-c} \). Multiplying it by the genome size \( g \) gives us the expected number of bases with zero coverage. So, to cover the whole genome, this number must be lower than 1 which places condition of \( c > l \times n \). For example, to cover the whole human genome \( (g = 3 \times 10^9) \), the read coverage depth must be at least 22.

Answer to the second question is implied by the facts that k-mer coverage of the genome follows the Poisson distribution. In an ideal case, the k-mer frequency distribution function of an error-free read set would follow the probability mass function of the Poisson distribution, meaning that frequencies of most of the k-mers are near the k-mer average depth. However, when we introduce possibility of sequencing errors, some k-mers would be sampled less often and some become even unique. The k-mer frequency distribution of such a read set does not follow the Poisson distribution. The aspect of sequencing errors and their means of their correction are described in great detail in Chapter 3.

### 1.3 The Real World

In contrast to the idealized model described in the previous section (1.2), read sequencing data contain sequencing errors. In other words, some of the reads contain incorrectly interpreted bases. To help with identifying such bases, each base of a read has its base quality, a probability that the given base is incorrect. Base quality (BQ) is usually represented as a Phred score \( Q \) defined as

\[
P(\text{base is wrong}) = 10^{-\frac{Q}{10}}
\]

Phred score has a natural interpretation, for example \( Q = 40 \) indicates one error in 10000 bases, \( Q = 30 \) one error in 1000 bases and so on (Figure 1.2 map the score values to probabilities). When stored in a text format, such as SAM or FASTQ, base qualities are encoded as ANSI characters. Because the first 32 ASCII characters are not printable, and the space character is coded by 32, the base quality values are incremented by 33. When loading the reads from such a text format, this needs to be taken in account.
Figure 1.2: Mapping Phred score values to probabilities. Be aware of the logarithmic scale on the $y$ axis.

Mapped reads are accompanied by their position (POS) in the reference sequence, by additional alignment details and a mapping quality (MQ), which is a measure of mapping confidence expressed as a Phred score. Although the position information may be wrong, it may help us in cases when we are interested in correcting only a part of the genome and would like to filter out reads that do not map confidently.

The alignment details are given in form of a CIGAR string which describes how the individual bases of the read map to the reference. The string is formatted as a set of numbers defining the number of bases, and a character code describing the alignment alignment. The most common operations include:

- **Match (M)**. Alignment match. That can represent both sequence match and mismatch.
- **Mismatch (X)**. The sequence mismatch. In practice, this code is seldom used since M may code both matches and mismatches.
- **Insertion (I)**. The sequence is inserted to the reference at a given position. Then, the read continues to follow the reference at this position plus one unless alignment operation following the insertion tells otherwise.
- **Deletion (D)**. The read sequence skips the corresponding part of the reference.
- **Hard-clip (H)**. The original read was longer, but was trimmed in downstream analysis.
• **Soft-clip (S).** This part of the read could not be aligned well and was excluded from the alignment.

For example, assume a reference sequence CAGGTGTCTGA and the read GGT-GAATCTA aligned as follows:

```
1 2 3 4 5 6 7 8 9 A B
Reference: C A G G T G T C T G A
Read: G G T G A A T C T A
```

The read was mapped to the reference position 3 and the CIGAR string of the alignment is 4M2I3M1D1M.

Genomes of diploid organisms, introduce another problem not covered by the simple mathematical model covered earlier. Since such organism owns two sets of chromosomes (one from each of its parents), both sets are present within the read set obtained by chopping the genome to short reads. That implies that two genome strings are present within, and need to be reconstructed, from the input read set.

### 1.4 Major Assembly Algorithm Classes

Currently, there are two widely used classes of algorithms: overlap-layout-consensus (OLC) and de-bruijn-graph (DBG) [1]. Algorithms belonging to the former base their idea on construction of a graph recording overlapping reads and extracting the alternate sequences from it. Main idea behind the DBG approach is to decompose all reads into k-mers that are then used to construct a de Bruijn graph which, after some optimizations, is subject to derivations of alternate sequences.

#### 1.4.1 De Bruijn Graphs

De Bruijn graphs (DBG) were originally invented as an answer to the superstring problem (finding a shortest circular string containing all possible substrings of length \(k\) (k-mers) over a given alphabet) [2]. For a given \(k\), de Bruijn graph is constructed by creating vertices representing all strings of length \(k - 1\), one per string. Two vertices are connected with a directed edge if there exist a k-mer such that the source and destination vertices represent its prefix and suffix of length \(k - 1\) respectively. The k-mer labels the edge.

Formally, let \(L_k\) be a language containing all words of length \(k\) over an alphabet \(\sigma\), then de Bruijn graph \(B(V, E)\) is defined as

\[
V = \{ v_w \mid w \in L_{k-1} \} \quad (1.1)
\]

\[
E = \{ (v_u, v_w) \mid x \in L_k, \text{its prefix and suffix \textit{ew}} \} \quad (1.2)
\]

\[
(1.3)
\]

The shortest superstring is found by concatenating all \(k - 1\)-mers represented by vertices on any Eulerian cycle of the corresponding de Bruijn graph. Since a polynomial-time algorithm for finding an Eulerian cycle is known, the superstring problem can be effectively solved.
Application to Genome Assembly

Similarly, de Bruijn graphs can be used for genome assembly, especially if the genome question is circular. Assume that, for a fixed $k$, all $k$-mers of the target genome are known. Then a DBG can be constructed in two different ways:

- **K-mers are represented by edges.** Each edge is labeled by a $k$-mer in the same way as in the graph used for solving the superstring problem. Each edge connects vertices representing (k-1)-mers forming prefix and suffix of its $k$-mer. The genome string can be reconstructed by choosing the right one from all possible Eulerian cycles. This approach is presented and discussed in [2].

- **K-mers are represented by vertices.** Each $k$-mer is represented as a single vertex. Edges reflect the $k$-mer order within reads. To recover the genome, one needs to find a Hamiltonian path of the graph, which is one of NP-complete problems. Among others, HaploCall [3] used by GATK takes advantage of this way of de Bruijn graph usage.

**HaploCall**

Rather than assembling whole genome at once, HaploCall starts with detection of regions that are likely to contain variants[4]. A score is computed for every genome position, reflecting the probability of variant occurrence. Regions are formed around positions classified as interesting. As subset of the input reads is assigned to each of the regions; reads are selected based on their mapping position. Each region is then processed separately and independently of others.

The active region processing phase starts by decomposing the reference sequence covering the region into $k$-mers and using them as vertices in a newly constructed de Bruijn graph. Edges connect vertices representing $k$-mers adjacent by their position within the reference. For each edge, a weight value is initialized to zero.

When the reference is transformed into the graph, a similar process happen with each of the input reads assigned to the active region. The read is decomposed into $k$-mers that are inserted into the graph in the same manner as the reference $k$-mers. Again, edges follow the order of $k$-mers within reads. New vertices are created only for $k$-mers not seen in the reference, similar case applies to new edges — if an edge denoting adjacent position of two $k$-mers already exists, its weight is increased by one. Otherwise, a new edge with weight initialized to 1 is inserted into the graph. The weight values actually count number of reads covering individual edges.

After inserting all the reads into the graph (the HaploCall documentation refers to the step as Threading reads through the graph), it is time to simplify the graph structure a little bit. The main concern here is to remove graph sections created due to sequencing errors present in the input reads. Such sections are identified by their low read coverage and removed. Other structure refinements are performed, including removal of paths not leading to the reference end.

In the next stage, most probable sequences are extracted from the resulting graph. Each sequence is represented by a path leading from the starting $k$-mer of the reference to ending one and its probability score is computed as the product
of so-called transition probabilities of the path edges. Transition probability of
an edge is computed as its weight divided by the sum of weights of all
edges sharing the same source vertex. By default, 128 most probable paths are
selected.

The Smith-Waterman algorithm is used to align each of the selected sequences
to the reference. The alignment is retrieved in form of a CIGAR string that
indicates places of possible variants, such as SNPs and indels. Indel positions are
normalized (left aligned).

The CIGAR strings actually contain a super set of the final set of called
variants. To filter out false positives, other solutions are employed (for example,
the HaploCall documentation mentions UnifiedGenotyper for GATK).

AbySS

AbySS\cite{5} is another representative of assembly algorithms using de Bruijn graphs.
Unlike HaploCall, it does not use the reference sequence for assembly. One of
its interesting features is a support for running on multi-computer environment
with fast network connection. Most of the computation is massively parallelized
and the MPI (Message Passing Interface) is used for communication between
individual computing units.

AbySS starts by decomposing all input reads to k-mers and constructing a de
Bruijn graph. Except being distributed among nodes of the computing cluster,
it treats k-mers and their reverse-complements as equals. Both are represented
by a single vertex and when one is inserted, the other also starts its existence.

Structure of the distributed de Bruijn is then optimized mainly by remov-
ing short branches with dead ends that usually results from sequencing errors.
Then, linear sequences of vertices without branching are merged into contigs,
representing longer sequences of the assembled genome.

The paired-end information is then used to put individual contigs together.
A link is created within two contigs if the number of paired-end reads joining the
contigs exceed a given threshold. The final step lies in determining a path going
through all contigs linked together.

Although utilizing the de Bruijn graph approach, AbySS actually shares a
lot with the overlap-layout-consensus class of assembly algorithms. The main
difference, at a high level of abstraction, is the usage of k-mers, while the OL-
C algorithms construct their graphs from whole reads and do not perform the
decomposition step.

1.4.2 Overlap-Layout-Consensus

Algorithms taking advantage of this approach usually do not decompose reads
into k-mer sequences. Reads are considered to be the smallest units. As the name
suggests, the work is done in three steps.

Main purpose of the overlap phase is, not surprisingly, to compute overlaps
between all possible pairs of the input reads. Since number of short reads within
high coverage data sets may go to millions, the process of overlap computation
may take a lot of processor and memory resources. Overlapping reads are connect-
ed into longer sequences called contigs. Actually, a graph is being constructed,
its vertices represents individual reads (or contigs) and edges represent overlaps.
The definition of read overlapping is not strict. Two reads are considered overlapping (and thus, are connected by an edge) if they overlap at least by $t$ bases, allowing several mismatches. The constant $t$ is a parameter of the algorithm and is called a cutoff.

During the layout phase, a read-pairing information is used to put disconnected parts of the read overlapping graph together and resolving structures created by sequencing errors. The consensus phase is responsible for deriving candidate sequences for variant calling from the graph. In an ideal case, the candidate sequences would be the Hamiltonian paths of the graph.

1.5 Goals of this Work

Main goal of this work is to develop an algorithm capable of variant calling on high-coverage data sets, comparable to, or more precise than methods currently employed. The approach used by HaplotypeCaller (HaploCall) should be used as an inspiration which implies the newly developed algorithm would take advantage of the reference sequence and would belong to the class of algorithms utilizing de Bruijn graphs.

The new algorithm should accept the input data set in widely used formats, such as SAM for the read set and FASTA for the reference sequence, and output the called variants in the VCF format\[6\]. Its results need to be compared to at least two other assembly and variant calling tools. GATK (HaploCall) \[3\], as a representative of the DBG class and reference-aided methods, and Fermikit \[7\], utilizing the OLC concept.
2. The Algorithm

Our algorithm takes a reference sequence and a set of reads as inputs, and outputs a VCF file containing all detected variants. Both inputs are read into memory during the initialization phase, there are no memory-saving optimizations employed. In other words, no index files are used for the input data.

The variant calling is done on region basis. The reference sequence is divided into regions of constant length (2000 bases by default), sometimes also referred as active regions and with 25% overlap. Reads are assigned to individual regions according to their mapping position. All regions are called independently and in parallel. To detect the variants, the following steps are performed:

- the reference sequence covering the active region is transformed into a De Bruijn-like graph,
- reads assigned to the active region are integrated into the graph,
- the graph structure is pruned and optimized,
- variants are extracted,
- genotypes and phasing are determined,

The k-mer size used to build the de Bruijn graph belongs to parameters of the algorithm (Section 2.2). If the graph is too complex after the structural optimizations, it is discarded and the algorithm makes another attempt using higher value for k-mer size. This behavior is very similar to the HaploCall algorithm.

2.1 Input, Output and Preprocessing

The reference sequence is expected to be in the FASTA format and starting at position 1. In a typical scenario, the whole chromosome is provided as an input. The FASTA file may contain multiple distinct reference sequences (covering multiple chromosomes). Reference sequences are processed one at a time which means that at most one sequence is present in memory at any moment. Only characters A, C, G and T are considered valid nucleotides. Other characters, such as N used to mask low-complexity regions, are treated as invalid and reference regions filled with them are not subject to variant calling.

The read set needs to be stored in a text file reflecting the SAM format. Presence of header lines is not required, all information is deduced from the reads. Since the algorithm does not perform any error correction on its own, the input reads must be corrected beforehand. The error correction is supported as a separate command of the tool implementing our algorithm. The correction method was adopted from the Fermi-lite project and Chapter 3 describes it in detail.

After the whole SAM file is read into memory and parsed, reads considered useless for the purpose of variant calling are removed from the set. Contents of the FLAGS column of the SAM file serves as a main filter, since it is used to detect the following types of undesirable reads:
• **unmapped**, recognized by the bit 2 set to 1,

• **secondary alignments**, detected by the bit 8 set to 1,

• **duplicates**, bit 10 is set to 1,

• **supplementary**, bit 11 is 1.

Also, reads with the mapping position (POS) set to zero and with mapping quality (MAPQ) lower than certain threshold, defined to 10 by default, are removed from the set. Read’s mapping quality is also used to update individual base qualities. Each base quality \( q \) is updated according to the formula

\[
q = \min(q, MAPQ)
\]

Several other SAM fields play a role in read preprocessing. The **CIGAR** string is used to detect and remove soft-clipped bases. The **QNAME** values are used to detect paired end reads, **RNAME** gives the name of the input reference sequence (chromosome).

The SAM file format is described in great detail in [9].

When all the reads are preprocessed this way, they are assigned to respective active regions based on the mapping position. Then, the main algorithm comes into play.

### 2.2 Algorithm Parameters

Behavior of the algorithm and produced results can be controlled by several parameters:

• **Initial k-mer size**. Defines the initial k-mer size used to build the de Bruijn-like graph. If the resulting graph is too complex, another attempt is made with higher k-mer size. By default, this parameter is set to 21.

• **Minimum position quality**. Only reads which mapping quality (MAPQ) is greater than this parameter are used. The default value is 10.

• **Global threshold**. Defines a threshold value used globally through the algorithm. Edges of the de Bruijn graph with read coverage lower than the threshold are automatically removed. Similar approach is used when computing genotypes and phasing information; variant connection covered by less reads than the global threshold are ignored. The default value of the parameter is 4.

• **Region length**. Length of active regions. Set to 2000 bases by default.

• **Low quality variant threshold**. If the number of reads supporting a certain variant is not above this value, the variant is considered as a low quality variant. Such variants are subjects to a binomial test that decides whether they will be written to the resulting VCF. By default, this parameter is set to 3.
• **Binomial threshold.** Defines a threshold for accepting low quality variants. The binomial test gives a probability how a given low quality variant is reliable. If this probability is above the threshold (set to 1% by default), the variant is reported in the resulting VCF.

### 2.3 K-mer Representation

Our algorithm works with k-mers only through an interface consisting from several functions and macros. The k-mer implementation is used as a black box which allows us to make several implementations and then choose the best fitting one without any consequences on the rest of the algorithm, except those implied by the interface.

We currently use two implementations representing k-mers in different ways, shown on Figure 2.1. For performance reasons, the implementations may be switched only in compile time.

The upper part of the figure shows the representation of the **debugging** k-mer. The main idea behind this type of k-mer is to make the content easily human-readable, which is excellent for debugging purposes. The k-mer sequence is stored in a character array, each element represents one base. Except its context number, the purpose of which is explained in the next section, each debug k-mer also remembers its size. That value is used for debugging purposes only, mainly to recognize attempts to pass wrong k-mer size argument to one of the k-mer interface routines. Debug k-mers also have an advantage of potentially unlimited maximum size. It is clear that debug k-mers are not a good option in terms of performance, especially for bigger $k$.

**Compact,** k-mers in contrast, are more optimized for performance and quick append and prepend operations (moving the k-mer sequence forward or backward). Their design is greatly inspired by the k-mer implementation of the Fermi-lite project. The k-mer substring is stored in three 64-bit integers. Each base is represented by three bits, each stored separately in the three integer fields. To read a $i^{th}$ base (starting from zero), one needs to combine $(k - i - 1)^{th}$ bits of the integers and then translate the result by rules described in Table 2.1. The table also describes the meaning of the eight possible values.

Structure of a compact k-mer is displayed in lower part of Figure 2.1. Its advantages are a fixed size of 28 bytes regardless of $k$ and performance (on the other hand, space occupied by a debugging k-mer depends on used $k$ value and is not known at the compilation time). However, limiting the maximum k-mer size to 63 bases this way may impose a problem in case of assembling repetitive regions.

All k-mer implementations usable by our algorithm must reserve some space for storing a **k-mer context number.** The number is used to differentiate even between k-mers that contain the same substring of length $k$. That implies that a k-mer, according to this new definition, is unique only if it differs in both the substring and the context number from all other k-mers. K-mers that differ only in the context number are sometimes referred as **equal by (sub)string** or **equal by sequence.** The context number proves to be useful when fighting repeats within the reference sequence (described in Section 2.4).
Table 2.1: Base representations in debugging and compact k-mers

<table>
<thead>
<tr>
<th>Compact value</th>
<th>Debugging value</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>A</td>
<td>The base A</td>
</tr>
<tr>
<td>1</td>
<td>C</td>
<td>The base C</td>
</tr>
<tr>
<td>2</td>
<td>G</td>
<td>The base G</td>
</tr>
<tr>
<td>3</td>
<td>T</td>
<td>The base T</td>
</tr>
<tr>
<td>4</td>
<td>B</td>
<td>Denotes beginning and end of the active region.</td>
</tr>
<tr>
<td>5</td>
<td>H</td>
<td>Used to represent k-mers of helper vertices.</td>
</tr>
<tr>
<td>6</td>
<td>D</td>
<td>Used to represent k-mers of helper vertices.</td>
</tr>
<tr>
<td>7</td>
<td>N</td>
<td>-</td>
</tr>
</tbody>
</table>

The whole chapter uses the term k-mer in this sense described above, unless explicitly specified otherwise.

The Fermi-lite project uses two bits to represent each base in the k-mer. Our algorithm can be modified to do the same, since the helper vertices actually do not need k-mers and exist mostly for the sake of code simplicity (no special handling regarding helper vertices is required). Similarly, there is no real reason for marking beginning and end of the active region by a special character, this was only useful for debugging.

2.4 Reference Transformation

The first step of the algorithm lies in transforming a reference sequence covering the selected active region into a de Bruijn-like graph. The idea behind this step is very similar to other assembly algorithms based on these graph types.

The reference is decomposed into k-mers, each overlaps with the adjacent ones by \( k - 1 \) bases. K-mers representing the same sequence are differentiated by their context number, so each k-mer derived from the reference is unique. Two extra k-mers, denoting the beginning and the end of the active region are added to the set. Then, each k-mer is represented by a single vertex in the graph, and edges are defined by the order of the k-mers within the reference.
Formally speaking, with the active region of length $l$ represented as a string $b_1 \ldots b_l$, k-mers $k_0, \ldots, k_{l-k+2}$ are derived from the region as follows:

- $k_0 = (Bb_1 \ldots b_{k-1}, 0)$
- $k_1 = (b_1 \ldots b_k, 0)$
- $\ldots$
- $k_i = (b_ib_{i+k-1}, c_i), 2 \leq i \leq l - k + 1$
- $\ldots$
- $k_{l-k+2} = (b_{l-k+2} \ldots b_lB)0$

$c_i$ represents the context number of the k-mer $k_i$. The number is set to zero for k-mers unique within the active region. On the other hand, let’s assume that k-mers $k_{i_1}, \ldots, k_{i_n}, i_0 < \ldots < i_n$, contain the same string. Their context numbers are defined as

$$c_{ij} = j,$$

$k_0$ and $k_{l-k+2}$ are special k-mers added to the set in order to show the start and end of the active region within the graph. $B$ is a virtual base that ensures these k-mers are unique. The bases must not appear anywhere else within the active region. All k-mers created in this step and all vertices created from them are also called as reference k-mers and reference vertices. Similarly, k-mers and vertices created during the read integration phase, are sometimes referred as read k-mers and read vertices.

Each k-mer $k_i$ is transformed into a single vertex $v_i$. Edges follow the order of the k-mers in the active region. In other words, the edge set of the graph is

$$E = \{(v_i, v_{i+1})\} \quad 0 \leq i \leq l - k + 1$$

Figure 2.2 displays a graph created by transforming the active region ATCTGTATATATG with k-mer size of 5. The algorithm creates the following k-mers:

- $k_0 = (BATCT, 0)$
- $k_1 = (ATCTG, 0)$
- $k_2 = (TCTGT, 0)$
- $k_3 = (CTGTA, 0)$
- $k_4 = (TGTAT, 0)$
- $k_5 = (GTATA, 0)$
- $k_6 = (TATAT, 0)$
- $k_7 = (ATATG, 0)$
- $k_8 = (TATGB, 0)$
- $k_9 = (ATATG, 0)$
- $k_{10} = (TATGB, 0)$

The figure follows a coloring scheme preserved in de Bruijn graphs across the whole thesis — green color for reference, red for reads, yellow and blue for marking start and end of an active region.
As can be seen, there are two k-mers representing sequence TATAT, namely $k_6$ and $k_8$. Because of their distinct context numbers, they are represented as separate vertices. Introduction of the context numbers removed a loop from the graph. The loop can be observed on Figure 2.3 that shows a standard de Bruijn graph constructed from the same active region. K-mers $k_6$ and $k_8$ are represented by the same vertex. In order to recover the sequence, it is required to know how many times the loop was actually used during the transformation step.

![Figure 2.2: Graph resulting from the transformation of ATCTGTATATATG sequence](image)

Although this solves the problem of cycles for now, at least for now, things become more complicated in the next step of the algorithm which inserts individual reads into the graph.

### 2.5 Adding Reads

The basic idea behind this stage is fairly simple and similar to the approach used in the previous case. The read being added is decomposed into k-mers. If the k-mer is not present in the graph already, a new vertex is created. Again, vertices representing adjacent k-mers in the read are connected by edges.
Figure 2.3: Transformation of the ATCTGTATATATG sequence to a standard de Bruijn graph (with no k-mer context numbers)

Figure 2.4 shows a graph created by transforming a region of ACCGTGGTAAT and adding the read ACCGTAGTAAT to the resulting graph. K-mer size is set to 5. The read is divided into the following k-mers:

\[
\begin{align*}
    k_0 &= (ACCGT, 0) \\
    k_1 &= (CCGTA, 0) \\
    k_2 &= (CGTAG, 0) \\
    k_3 &= (GTAGT, 0) \\
    k_4 &= (TAGTA, 0) \\
    k_5 &= (AGTAA, 0) \\
    k_6 &= (GTAAT, 0)
\end{align*}
\]

In this example, the k-mers \( k_0 \) and \( k_6 \) were already present in the reference graph but new vertices has to be created for the rest.

Finally, edges are added (if necessary) to show the k-mer order within the read. When talking about classical de Bruijn graphs, edges do not need to be
expressed explicitly since they always connect adjacent k-mers. In case of our modification to de Bruijn graph, we chose to make the k-mer connection explicit. Two reasons led us to such decision:

- our definition adds context number to k-mers and we wish to connect only certain k-mers adjacent by their strings,
- we need to make connections between non-adjacent k-mers, e.g. shortcuts representing paths with no branches, or fulfilling purposes discussed mainly in Section 2.6.

Keeping graph edges explicitly has also its drawbacks, especially those related to performance. Classical de Bruijn graphs can be represented only by a set of their vertices, since the edges can be deduced on demand.

The figure also suggests how to retrieve the alternate sequence introduced by the read — by going through edges supported by the read and concatenating the last bases from the k-mers present on the path. K-mers covering start and end of the reference region are the only exceptions; all except the B is used from the former, nothing from the latter.

Figure 2.4 reflects an ideal state, meaning that all places, where the read differs from the reference, are covered by distinct k-mers, and the distance between each two of them is greater than $k$. If these conditions are met, each single $n$-base long difference (SNP, insertion or deletion) adds at most $n + k - 1$ new vertices to the graph. Each difference then creates only two linear paths, one covering the reference, the other for the alternate allele. Such structures are called bubbles and are easy to work with.

However, it may happen that some of the k-mers covering a difference collide by sequence with either k-mers of the reference, or k-mers of other reads covering a totally different place of the active region. To minimize such problematic cases, additional graph transformations need to be made after all the reads are integrated into the graph.

Unfortunately, the basic idea does not work in our case. Introduction of the k-mer context numbers prevented loops in the graph derived purely from the reference. But since multiple k-mers representing the same sequence may exist, it is not always possible to easily determine which of the vertices should be assigned to individual k-mers of the read. For example, if looking at the graph from Figure 2.2, it is not clear which vertex (or vertices) should be assigned to a k-mer with TATAT sequence.

We decided to solve the issue by transforming the basic idea into the following steps:

- decompose the read into sequence of k-mers (a so-called short variant optimization, described later, may be applied),
- to each k-mer assign a set of vertices with the same k-mer sequence (differences in context numbers are permitted),
- from each set, select one vertex to represent the k-mer of the read (the selection process is described later in this section),
- connect all read vertices in a way that respects the order of the k-mers in the read.
2.5.1 Transforming the Read into K-mers

Let’s define a read of length $n$ as a sequence $r_1 \ldots r_n$ of bases. If the short variant optimization, described in the next paragraph, is not applied, the read is decomposed into individual k-mers $k_1, \ldots, k_{n-k+1}$ in the same way as for the reference case, except that no extra k-mers to denote read start and end are created. The k-mers look as follows:

$$k_1 = (r_1 \ldots r_k, 0)$$

$$\ldots$$

$$k_{n-k+1} = (r_{n-k+1} \ldots r_n, 0)$$

Then, the step described in Subsection 2.5.2 is applied.

As described above, in an ideal case, a $n$-base long difference from the reference produces $n + k - 1$ k-mers different from all reference k-mers. To reduce the probability that some of the new k-mers actually collide with either the reference, or another read, the short variant optimization may be applied. The optimization reduces the number of k-mers representing a $n$-base long difference to:
• $n$ for an insertion,
• zero for a deletion,
• 1 for a SNP.

The optimization assumes that when recovering a sequence from the graph, only the last base of each k-mer, with the exception of the starting one, is used. So, only k-mers covering the difference by their last base need to be added; reference k-mers may be used for the rest in case the difference is followed by a reasonable number of bases equal to the reference.

To make the k-mers covering a $n$ base long difference by their last bases really unique, the first $n$ bases of the first k-mer is changed to $D$, a virtual base used for helper purposes only. Rest of the k-mers is obtained by appending the bases of the difference to the first one. Keep in mind that $n$ is always smaller than the k-mer size, since we are talking about the short variant optimization.

Figure 2.5 shows how the graph is optimized for a read containing SNP. The reference and read sequences are taken from Figure 2.4. Since the difference has 1 base in length, only one k-mer (DCTGA_0) is used to represent it. The k-mer is followed by reference k-mers. As can be seen, their last base are equal to one of k-mers from Figure 2.4.

The short variant optimization is applied for k-mer $k_i$ if the following holds:

- There is only one reference vertex with a k-mer equal to $k_{i-1}$ by sequence. Let’s assume this is vertex $v_{j-1}$.
- There is at most one vertex for $k_i$ that either is a result of a read addition, or is a reference one but do not immediately follows the vertex $v_{j-1}$ in the reference.

Such conditions are met when the read differs from the reference at base $r_{i+k-1}$. To determine whether the difference is only a short one, the Smith-Watterman algorithm is applied. If this is the case, the action depends on the difference type:

- for $n$-base long deletion, $k_i$ is defined as $v_{j+n-1}$,
- for insertion of length $n$, $k_i$, ..., $k_{i+n-1}$ are defined as with no short variant optimization and $k_{i+n}$ is set to $v_j$,
- for SNP, $k_i$ is left as such and $k_{i+1}$ is defined as $v_{j+1}$

### 2.5.2 Assigning Sets of Vertices to K-mers

The process of assigning vertex sets to individual k-mers derived from the read in the previous step is quite straightforward — a set assigned to a certain k-mer $x$ contains exactly the vertices sharing the same k-mer string. The k-mer context number is not taken into account. If a k-mer is not represented by any vertex of the current graph (thus, the k-mer would receive an empty set), a new vertex is created to represent it. With a classical de Bruijn graph, all sets would contain exactly one vertex. With k-mer context numbers, there can be multiple vertices per k-mer. This complicates the task of integrating reads into the graph.
Figure 2.5: Short Variant optimization
because the graph may contain multiple paths representing a single read (by using
different vertices with k-mers equal by sequence).

Previous steps of the algorithm, described above, impose the following condi-
tions on the assigned vertex sets:

- each set contains either read, or reference vertices, but not both,
- if a set contains read vertices, its size is always one,
- sets containing reference vertices do not have such restriction,
- each two sets are either distinct (their intersection is an empty set) or equal.

The second and third condition holds because k-mer context numbers are used to
differentiate reference k-mers but not the read ones. The fourth one is implied by
the fact that each set contains all existing vertices with k-mers equal by sequence
to the read k-mer being processed. So, if two sets have a non-empty intersection,
they actually refer to the same read k-mer (k-mer context numbers are not used
when processing read k-mers).

In formal terms, a set $M_i$ is assigned to a k-mer $k_i$ where

$$M_i = \{v_{ij} | v_{ij} \in V(G), kmer(v_{ij}) \text{ equals to } k_i \text{ by sequence}\}$$

When a set is assigned to each k-mer, it is time to integrate the read into the
graph in the form of a path, starting in the vertex representing $k_1$ and ending in
the vertex covering $k_{n-k+1}$. Since $M_i$ sets may contain more than one vertex, it
is necessary to select vertices to form a path best fitting to the read. To derive a
good path, we decided to assume the following:

- they should follow the reference sequence in the forward direction,
- The probability of skipping large number of reference vertices (long dele-
tions) is low,
- multiple reads cover one place, sharing appropriate parts of their paths.

These requirements cannot be enforced too strictly as de Bruijn graphs are not
very suitable for coping with repeats of length $k$ or more. The case of a difference
containing a copy of reference at least $k$ bases in length might be enough to break
the first assumption. The second assumption permits exceptions by definition.
The third forms a base for most of the genome assembly algorithms.

In order to choose the correct vertices from the $M_i$ sets, we decided to reduce
it to a shortest-path problem on a helper graph the structure of which is defined
by the sets and their contents as explained in 2.5.3.

2.5.3 The Helper Graph

The helper graph is an oriented layered graph. Each layer consists of all vertices
contained in one reference $M_i$ set. The order of the layers respect the order of
$M_i$ sets. Sets consisting of read vertices are not part of the helper graph. Only
adjacent layers are connected by edges, their orientation reflects the order of the
sets. Each subgraph consisting of two adjacent layers is a full bipartite graph.
The structure of the helper graph does not take equality of $M_i$ into account. In other words, when $M_i = M_j$ for $i \neq j$, both sets are represented within the helper graph as individual layers, even if they refer to the same vertices of the (main, non-helper) graph.

Formally speaking, let $M_i = \{v_i^1, ..., v_i^n\}$ and let $i$ index the reference sets only. Then the helper graph $G_h$ can be defined as follows:

$$G_h = (V_h, E_h)$$

$$V_h = \bigcup_i M_i$$

$$E_h = \{(u, v) | u \in M_i, v \in M_{i+1}\}$$

By finding the shortest path leading from a vertex in the first layer to one in the last layer, we perform the process of selection of vertices representing the read in the main graph. The shortest path depends on weights of edges connecting the adjacent layers. In general, the weighting function follows these rules:

- the weight is increased by a **missing edge penalty** if there is a missing edge on the path from $u$ to $v$ in the main graph,
- the weight is increased by a **reference backward penalty** if reference position of $u$ is greater or equal to the reference position of $v$,
- the weight is increased by a **reference forward penalty** if reference position of $u$ is far less than reference position of $v$.

The rules actually indicate why $M_i$ sets covering read vertices are not parts of the helper graph – since their vertices maintain no reference position, only the missing edge penalties would apply and that can be included within missing edge penalties of the reference vertices only.

For an example of a helper graph, let’s have a reference sequence ACTATACTA and a read ACTAGACTA. The left part of Figure 2.6 shows the main graph just after adding vertices for the reference and the read k-mers with short variant optimization applied. The resulting helper graph is shown on the right part of the figure.

Six k-mers are derived from the read which means that vertex sets $M_0, ..., M_5$ are assigned to them. Since the second k-mer is represented by a read vertex, the $M_1$ set is not included as a layer of the helper graph. Other sets contain reference vertices, so they form individual layers. Adjacent layers are then connected. Edges with applied penalties (only the reference backward penalty in this case) are depicted red. The black edges show the shortest path.

The shortest path select the first vertex from $M_0$ and the second one from $M_5$ to represent the read within the main graph (since other sets contain only one vertex, the selection process is trivial there). The resulting path in the graph can be used to correctly recover the sequence covered by the read.

As Figure 2.7 indicates, both graphs look a little bit differently when the short variant optimization is not applied. The main graph contains more read vertices which reduces the number of layers in the helper graph. Although the graphs are different, the sequence covered by the rad is the same and can be correctly recovered again.
2.6 Graph Structure Optimization

When all reads are integrated into the de Bruijn-like graph, it is time to optimize its structure in order to get rid of unpopulated paths, usually created by read errors, and resolve some other issues caused mostly by repetitive regions inside either the reference or the reads.

2.6.1 Connecting Bubbles

Figure 2.6 demonstrates one class of the structural problems. A subset of reference sequence was transformed into vertices 1, 2, 3, 4, 5 and 6. A set of reads is represented by a “path” 2, 3, 4, 5, R1, R2, 1, 2, 3, R3, 6. The left part of the figure shows how such a graph would look like without any optimizations. It is clear that recovering the correct sequence would not be trivial.

However, since we know that the path leads from R2 to R3 (through 1, 2,
3), we can theoretically replace edges (R2, 1) and (3, R3) by a special edge (R2, R3), as the right part of the figure suggests. An information about the sequence covered by vertices 1, 2 and 3 needs to be recorded within the new edge. Recovering the correct sequence from the right part of the figure does not impose a problem since it is just a simple bubble.

A more general, and a very typical, situation is shown on Figure 2.9. The subgraph contains a subset of the reference (vertices 1, ..., n+1) and two bubbles; one ending by \( R_1 \) and connected to 2, another starting at \( R_2 \) and leading from \( n \). If edges \( I_1 \) (the input edge) and \( O_1 \) (the output edge) share reasonable amount of reads (\(|\text{reads}(I_1) \cap \text{reads}(O_1)| > \text{threshold}\)), the subgraph may also be interpreted as that the reads contain a sequence of length \( n - 1 \) that is also present in the reference. In that case, it is wise to connect the vertices \( R_1 \) and \( R_2 \) directly the same way as on Figure 2.8 by bypassing the reference part. The edge maintaining the direct link is marked as \( C_1 \) (the connecting edge). Reads shared by the input and the output edges are moved to the connecting one.

If \( C_1 \) is created, the read set intersection is also used to decide whether the edges \( I_1 \) and \( O_1 \) should be removed. The input edge is deleted if does not share enough reads with the next reference edge (meaning there are no valid sequences leading through both these edges). Similarly, the output edge is deleted in case it does not share enough reads with the last reference edge (no valid paths goes through the edges).

Figure 2.10 depicts probably the most general case; multiple reads share the
same sequence of $n$ bases ($R_1, \ldots, R_n$). There is $k$ input and $l$ output edges.
To determine the association between individual input and output edges, the intersection of covering reads is used again and connecting edges are created if necessary. More precisely, the following rules apply:

- If $i^{th}$ input and $j^{th}$ output edges share reasonable amount of reads ($|\text{reads}(I_i) \cap \text{reads}(O_j)| > \text{threshold}$), a connecting edge $C_{i,j}$ is created and the shared reads are moved to it. The new edge starts in $\text{source}(I_i)$ and ends in $\text{dest}(O_j)$.
- $I_i$ and $O_i$ are removed in case their read coverage drops below threshold as a result of moving it to the newly created $C_i$ edges.

### 2.6.2 Helper Vertices

The bubble connection optimization works well when the bubbles being connected contain at least one read vertex. If this is not the case, however, the effect of
replacing input and output edges with connecting edges leads to a destruction of the sequences recorded within the graph.

As an example, consider a case illustrated by the left part of Figure 2.11. The relevant part of the reference runs from vertex 1 to 7 and the read coverage supports a path of $3 \rightarrow 4 \rightarrow 5 \rightarrow 6 \rightarrow 1 \rightarrow 2 \rightarrow 4 \rightarrow 5 \rightarrow 7$. Applying steps described in this section results in creation of connecting edges $6 \rightarrow 4$ and $2 \rightarrow 7$ and removal of the edges $6 \rightarrow 1$, $2 \rightarrow 4$ and $5 \rightarrow 7$. Such a graph cannot be used to recover the alternate sequence.

As this problem arises only when the bubbles are formed entirely by edges connecting reference vertices (the short variant optimization produces such cases for deletions), the countermeasure is quite straightforward; the solution is to insert special placeholder vertices with an empty sequence. We use a conservative approach for their insertion which means they divide the following types of edges:

- output degree of their source vertex is greater than one,
- input degree of their destination is greater than one.

The right part of Figure 2.11 indicates where the helper vertices would be inserted. They divide edges $6 \rightarrow 1$, $2 \rightarrow 4$ and $5 \rightarrow 7$. If the bubble connection is applied now, it leads to creation of edges $H1 \rightarrow H2$ and $H2 \rightarrow H3$ and deletion of $H1 \rightarrow 2$, $3 \rightarrow H2$, $H2 \rightarrow 4$ and $5 \rightarrow H3$. Thanks to the optimization, the alternate sequence can be now recovered ($3 \rightarrow 4 \rightarrow 5 \rightarrow 6 \rightarrow H1 \rightarrow H2 \rightarrow H3 \rightarrow 7$).

### 2.7 Variant Calling

The graph structure optimization phase ends by removing nodes and edges with insufficient read coverage. Then, alternate sequences covered by the reads are extracted. Because the intra species variability of biological sequences is relatively small, it is practical to work with a list of differences (variants), rather than whole sequences. The algorithm identifies parts of the sequence shared with the
Variants are extracted by detecting certain subgraphs. When such a subgraph is discovered, the variant sequence is determined and integrated back into the graph by replacing its reference edges by a single variant edge. Read edges unique to the variant are also removed from the subgraph which simplifies its structure. The process of variant detection stops when no suitable subgraphs are found.

Figures 2.12 and 2.13 show four types of subgraphs used for variant extraction and demonstrate how they are modified in the process. The reference part of the graph must always start in vertex 1, end in \( n \) and all inner nodes must have one input and one output edge. Only edges of reference or variant type may be present in the reference part. In all cases, this path is replaced with a variant edge, shown as blue, connecting directly 1 and \( n \). All edges that are removed as a result of the extraction are plotted as discontinuous lines.

**Simple Bubble**

Read edges and vertices form a linear path leading from 1 to \( n \). Since the edges are not part of any other variant, they all are removed and the whole subgraph degenerates only to two reference vertices connected by a variant edge (Figure 2.13a).
Figure 2.12: Variant detection cases; bubble with outputs (a), bubble with inputs (b)

**Bubble with Inputs**

Figure 2.12b shows a subgraph where the read node has more than one input edge. In such case, only the sequence of read edges leading from the 1 vertex to the first vertex with more inputs than one is removed. Other read edges may take part also in other variants.

**Bubble with Outputs**

This subgraph may be viewed as an opposite to the bubble with inputs. All read vertices in the subgraph are allowed to have more outputs than one, but only one input. Only the last sequence of edges, starting in the last read vertex with output greater than one and ending in the $n$ vertex is removed, since it may participate only in one variant (Figure 2.12a).

**Diamond**

The most complicated case is shown under in Figure 2.13b. The sequence of read vertices may contain one with output degree greater than one followed (not necessarily directly) by one with unrestricted input degree. Only the edges covering one part of the supposed diamond are present in one variant only and hence are removed.

### 2.8 Variant Graph and Variant Filtering

When variants are extracted from the de Bruijn-like graph, they need to be filtered and their genotype and phasing computed. For this, a *variant graph* is built and
the task is transformed into a graph coloring problem.

However, not all variants reach this stage. If a variant extracted in the previous step is considered as a low quality variant (number of supporting reads is not above the low quality variant threshold), a binomial test is applied to check the variant reliability. Variants with low read support may be caused by sequencing errors and such need to be eliminated. The binomial test computes a probability that the variant is not false positive by the formula

\[
P(\text{variant is reliable}) = 2 \cdot I_{0.5}(r, a + 1)
\]

where \(r\) is a number of reads supporting the reference part of the variant, \(a\) counts the read support for the alternate part and \(I\) is the regularized incomplete beta function.

If the probability falls below the binomial threshold, the variant is treated as a false positive and is removed.

The variant graph represent each variant by two vertices; one for its reference and one for its alternate sequence. The final task is to color each vertex by one of these colors:

- **Blue.** The variant part is used by the first sequence.
- **Red.** The variant part is used by the second sequence.
- **Purple.** Both sequences go through the variant part.

If a vertex is colored purple, the vertex representing the other part of the variant is not required (no sequence goes through it) and is removed from the graph. The deletion usually happens only to the vertices representing the reference parts, since removing a vertex of the alternate path means that the variant was filtered out.

Before coloring, graph vertices are connected by several types of bidirectional edges that place various conditions on the color of their sources and destinations.
- **Variant edges** connect vertices representing parts of one variant. Their source and destination must be colored differently.

- **Read edges** connect variant parts covered by the same subset of reads with size greater than a threshold, such vertices need to be colored by the same color, with exception of purple. If one of the vertices is purple, an arbitrary color may be assigned to the other.

- **Pair edge** put together variant parts that are covered by paired-end reads. Since paired-end reads indicate that the variants covered by them belong to the same alternate sequence, the coloring restrictions are the same as for the read edges. Two vertices are connected by a paired edge if the share a sufficient number of paired-end reads.

The graph usually has a form of many components. Variants in one component share the same phasing. All the information used to create read and pair edges remain stored in the graph vertices and can be used to detect false positives among the variants, possibly together with a detection method based on searching for certain types of subgraphs.

An example of a colored component of the variant graph can be seen on Figure 2.14. The component puts together two variants (at reference positions 534324 and 534447) and a residuum after a variant evaluated as a false positive (position 534547). Blue edges connect parts of each variant together, black edges report connections based on paired-end information and parts sharing some amounts of reads are connected by red. Numbers on the edges specify number of reads participating in the connection.

The coloring clearly indicates that both variants are present only in one of two alternate sequences (blue one). Since the variant at 534547 is considered as false positive, both alternate sequences are directed through its reference part and its alternate part is removed from the graph. Keeping residues after fake variants may makes sense because other variants may connect through them. It can also be observed that the phasing of the two true variants is enforced by the paired-end information, not by reads covering both variants directly.

When the graph is colored, the genotype and phasing information are known. The variants are written to the resulting VCF file.
Figure 2.14: An example of a variant graph component
3. Read Error Correction

Read data used as an input to many assembly algorithms contain plenty of errors, such as wrongly read bases. To make the data usable for assembly, an error correction step is required. However, it does not remove all the errors and assembly algorithms must cope with that fact, especially when dealing with read ends.

Currently, two different approaches are used to correct read errors, and both are based on transforming individual reads into series of k-mers. One is based on detecting errors as low covered edges (or paths) in a de Bruijn graph, the other works with a k-mer frequency distribution. During development of our algorithm presented in this thesis, we made several attempts to implement an error correction algorithm based on de Bruijn graphs. Since we use these graphs also during assembly performing error corrections on them seemed to be a natural choice. Although they definitely helped to improve the quality of input reads, all our attempts performed worse compared to the k-mer frequency distribution approach.

In the end, we decided to adopt the error correction algorithm used by the Fermi-lite library [8] and based on k-mer frequency distribution. This chapter describes both approaches.

3.1 De Bruijn Graphs

This method transforms the input set of reads into a de Bruijn graph in a way very similar to one used by our assembly algorithm. Although implementation details may differ, the basic idea is the same: each read mapped to certain active region is divided into a sequence of k-mers, each k-mer serves as a vertex and the edges follow the k-mer order within the sequence. Reference sequence, covering the active region, may also be included in the graph.

The basic assumption is that errors produce unique k-mers and thus also nodes and edges with low coverage. Low-covered edges with source vertices that have output degree greater than one are especially interesting. A change of even a single base can divert a read path through edges with higher read coverage. The locality of the change depends on the k-mer size used.

A simple example demonstrating the main idea behind the method is displayed in Figure 3.1. Many reads share a sequence of TTGCGCTAA. However, there is also a single read that contains one error — TTGCACTAA. The de Bruijn graph shown on the figure uses k-mers 4 bases long. Combination of both sequences produces a standard bubble.

If the bubble was supported by reasonable amount of reads, it would be treated as a SNP. However, because there is only one supporting read, it may be reasonable to consider its divergence from other reads as an error, and to correct it, so the read path would follow more populated edges. When the correction is done, the resulting graph becomes linear, as shown on the right side in Figure 3.1.
3.2 K-mer Frequency Distribution

The method is based on the assumption that k-mer frequency distribution of an error-free read set has certain statistical properties. Specifically, it follows the Poisson distribution centered around the average k-mer coverage depth. Figure 3.2 shows the frequency distribution for an error-free read set and for a read set with error rate 1%.

As can be seen in the figure, errors lead to enrichment of unique and rare k-mers. The idea behind the correction algorithms based on this method is to transform the low-frequency k-mers into those with frequencies in the right place.

This approach is also used by the Fermi-lite software and is covered in the next section.

3.3 The Fermi-lite Approach

Fermi-lite is a standalone C library as well as a command-line tool for assembling Illumina short reads in regions from 100 bp to 10 million bp in size. It is largely
a light-weight in-memory version of Fermikit without generating any intermediate files. Results of the assembly are not produced in the VCF format, but as a graph. Read error corrections are not the main goal of the project, although this step is definitely required for a successful assembly.

We have successfully extracted the error correction algorithm from the project. The implementation should work well on multiprocessor systems and trades performance over memory consumption. The algorithm proceeds in the following steps:

- **Preprocessing.** The input read sequences are divided into k-mers, k-mer frequencies are calculated.

- **Error correction.** The problem is reduced into a shortest path graph problem and is solved by a modified version of Dijkstra’s algorithm.

- **Unique k-mer filtering.** Unique k-mers introduced during the error correction phase are removed from the read sequences.

Description of the error-correction algorithm is based on analysis of Fermi-lite’s source code, Since the source, especially the analyzed part, lacks comments and other characteristics typical for easy-to-read codes, some minor aspects of the algorithm still remain unclear.

### 3.3.1 Data Preprocessing

The main goal of the preprocessing phase is to compute frequencies for all k-mers found in the input read set. The frequencies are computed by inserting the k-mers into a k-mer table. During this phase, several terms related to k-mers and their occurrences are introduced:

- A k-mer occurrence is defined as **high quality** if quality of all bases covered by it is greater than certain threshold (set to 20 by default). If not all bases satisfy this condition, the occurrence is considered as **low quality**.

- A k-mer is considered **solid** if its frequency is greater than certain threshold.

- A k-mer is considered **unique** if it its frequency is zero.

- A k-mer is referred as **absent** if it its frequency is is below certain threshold.

K-mers are implemented as four 64-bit integers (Figure 3.3). Each base is represented by two bits. This limits the maximum k-mer size to 64 but allows effective implementations of standard k-mer operations. The first 64-bit integer stores least significant bits of the k-mer bases, the second contains their most significant bits. The bases are stored in an opposite order — the first base resides in bits 1, the second to k – 2 and so on. The second two integers store the same content, but with different order — the first base is stored in bits 0, the second in bits 1 etc. Such a k-mer form vastes memory; only two 64-bit integers are required to hold all the necessary information. The error correction algorithm often takes advantage of the two-integer representation.

Such a representation allows quick appends or individual base changes. To append a base into the first two integers, they just need to be shifted by one to
the left, ORed with the new base, and ANDed with $2^k - 1$ to set the unused bits to zero.

Since the k-mer representation reserves only 2 bits for one base, it is used to represent k-mers containing at most four types of bases which seems to be OK since the genome also consists from four types of bases. However, a virtual base N may appear within a read sequence, denoting places that the sequencing technology was unable to read. Fermi-lite solves the issue by ignoring all k-mers containing any number of Ns.

Figure 3.3: K-mer representation used by the Fermi-lite project

The k-mer table is actually a set of $2^{l_{pre}}$ khash tables. When a k-mer is being inserted or looked up, $l_{pre}$ bits of its data are used to select the table and the rest serves as an input to the hash function. $l_{pre} = 20$ by default. This representation of the k-mer table increases overall memory consumption, but has great impact on its performance in parallel environments.

The table uses 14 bits to track occurrences of each k-mer. Lower 8 bits count low quality occurrences, higher 6 bits are used by high quality ones. The counting stops on values of 255 and 63, no integer overflow happens. The table actually stores their hashes and occurrences, rather than full k-mers. Hence, distinct k-mers may be treated as one, since the table may use up to $l_{pre}$ of k-mer content to choose the right khash table, and up to 50 bits may be stored as a key. That means, key collisions may be avoided if the k-mer size drops below 35 (the table actually has a different hash function for k-mers size below 33).

After the insertion stage, a k-mer frequency distribution is computed individually for low quality and high quality occurrences. The most common frequency is named mode and is used during the second and third phase of the algorithm.

All phases of the algorithm are partially driven by its parameters. Table 3.1 provides a short description of them.
3.3.2 Error Correction

The error correction is performed separately for each read sequence. The correction problem is transformed into a shortest-path search in a layered oriented graph. Vertices represent individual k-mers, edges connect adjacent ones and their weights reflect the cost of transforming one k-mer to another. The weight function is defined by formula

\[ w(k_i) = w_{\text{absent,high}} \cdot nhq(k_i) + w_{\text{absent}} \cdot nlq(k_i) + w_{\text{ec}} \cdot ec(k_i) + w_{\text{ec,high}} \cdot bq(k_i) \]

\[ nhq(k_i) = \begin{cases} 0, & \text{k}_i \text{ is a high quality k-mer} \\ 1, & \text{otherwise} \end{cases} \]

\[ nlq(k_i) = \begin{cases} 0, & \text{k}_i \text{ is a low quality k-mer} \\ 1, & \text{otherwise} \end{cases} \]

\[ ec(k_i) = \begin{cases} 0, & \text{the edge ending in } k_i \text{ does not introduce an error correction} \\ 1, & \text{otherwise} \end{cases} \]

\[ bq(k_i) = \begin{cases} 0, & \text{the last base of } k_i \text{ has quality above q} \\ 1, & \text{otherwise} \end{cases} \]

Dijkstra’s search algorithm is used and its main loop can be decomposed into the following steps:

- Retrieve the vertex with lowest price, and its k-mer from the heap.
- by separately appending A, C, G, and T to the k-mer, touch the adjacent vertices on the next layer and compute the cost of their connections. Without appropriate pruning, this step would produce exponential amount of new vertices. Thus, error correction resulting into non-solid k-mers are not tracked. Additional criteria, such as proximity of a corrected high-quality k-mer, or clustering of corrected bases, are applied, however, this part of the code still remains unclear.
- Insert the newly created vertices into the heap.

Each path from the starting vertex to a vertex in the last layer represents one possible corrected part of the read sequence. Four such paths are computed. The path computation stops if a gap greater than max_path_diff is detected in their costs.

The error correction algorithm described above is applied to both the read sequence and its reverse complement, which is reverse-complemented again after the correction is finished. Thus, two potentially error-free sequences are generated for one read. The resulting corrected sequence is their combination — if one of the source sequences contains the N base at certain position, content of the second one is used instead. This approach reduces number of Ns in the corrected read set.
3.3.3 Unique k-mer Filtering

The error correction phase may produce unique k-mers which, as Figure 3.2 indicates, are not desirable. The Fermi-lite library attempts to get rid of such k-mers. Each corrected read sequence is processed separately (and in parallel with others).

At first, the longest k-mer sequence covered by non-unique k-mers is found. Denote its length, in k-mers, as $n$ and the read sequence length as $l$. Then, the read sequence between the read start and the first base covered by the found k-mer sequence is removed from the read. If the read is covered only by non-unique k-mers, nothing is removed, since the k-mer sequence covers the whole read. However, if the following condition holds:

$$\frac{n + k - 1}{l} < \text{min\_trim\_frac}$$

the read is removed from the read set. This case includes also zero-length k-mer sequences that appear when the read contains unique k-mers only.
Table 3.1: Parameters of the Fermi-lite algorithm

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Default value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k$</td>
<td>-</td>
<td>K-mer size. By default, this value is set to the base-two logarithm of the total number of bases.</td>
</tr>
<tr>
<td>$q$</td>
<td>20</td>
<td>Base quality threshold used to recognize high quality content from low quality one. $P(error) = 10^{-\frac{q}{10}}$</td>
</tr>
<tr>
<td>min_cov</td>
<td>4</td>
<td>Minimum frequency for solid k-mers.</td>
</tr>
<tr>
<td>win_multi_ec</td>
<td>10</td>
<td>Takes part in pruning the graph created during the error correction phase. The exact meaning, however, remains unclear.</td>
</tr>
<tr>
<td>$l_{pre}$</td>
<td>20</td>
<td>Number of khash tables in the k-mer table, defined as $2^{l_{pre}}$</td>
</tr>
<tr>
<td>min_trim_frac</td>
<td>0.8</td>
<td>Defines how long the sequence of solid k-mers must be in order not to remove the read during unique k-mer filtering.</td>
</tr>
<tr>
<td>$w_{ec}$</td>
<td>1</td>
<td>Participates in the weighing function used in the error correction step.</td>
</tr>
<tr>
<td>$w_{ec\text{.high}}$</td>
<td>7</td>
<td>Participates in the weighing function used in the error correction step.</td>
</tr>
<tr>
<td>$w_{absent}$</td>
<td>3</td>
<td>Participates in the weighing function used in the error correction step.</td>
</tr>
<tr>
<td>$w_{absent\text{.high}}$</td>
<td>1</td>
<td>Participates in the weighing function used in the error correction step.</td>
</tr>
<tr>
<td>max_path_diff</td>
<td>15</td>
<td>Defines a maximum cost difference between paths found by the modified Dijkstra’s algorithm. If this value is reached, no additional paths are examined.</td>
</tr>
</tbody>
</table>
4. Results

When developing a new algorithm, an important part is testing and evaluation against existing solutions. This chapter describes this stage, informing about a data set used to debug and improve our solution, and the method of comparison with other solutions, such as Fermikit and GATK.

4.1 Test Data Set

chr1:1-40,000,000 region of the high-coverage data from the 1000 Genome Project. In addition to the input reads [10], also variants called by Fermikit and GATK are available in form of VCF files [11]. The VCF files were used to compare the accuracy of the algorithm. In order to be able to make meaningful comparisons and to avoid remapping of all reads, we used the same version of the reference genome which was used also by the 1000 Genome Project (GRCh37 [12]).

The test read set consists of 12,475,011 reads with length of 151 bases. Figure 4.1 shows k-mer frequency distribution of the set with k-mer size of 21 bases. The shape of the graph, when compared to Figure 3.2 suggests that the set indeed contains read errors, therefore the error correction step was applied.

As Table 4.1 indicates, the error correction process removed and shortened a significant proportion of the reads. Approximately 21% of the input reads was subject to repairs. Figure 4.2 shows a distribution of the number of repaired bases per read, not including effects of read trimming.

Figure 4.3 shows the k-mer frequency distribution of the corrected read set.
Table 4.1: Statistics related to error correction of the test data set

<table>
<thead>
<tr>
<th>Category</th>
<th>Value</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total reads</td>
<td>12,475,011</td>
<td>-</td>
</tr>
<tr>
<td>Removed</td>
<td>64,653</td>
<td>0.52% of all reads.</td>
</tr>
<tr>
<td>Shortened</td>
<td>944</td>
<td>0.0076% of all reads</td>
</tr>
<tr>
<td>Total bases</td>
<td>1,880,123,991</td>
<td>-</td>
</tr>
<tr>
<td>Bases repaired</td>
<td>5,098,764</td>
<td>0.27% of all bases</td>
</tr>
</tbody>
</table>

Figure 4.2: Distribution of a number of repaired bases in a single read

Although still not perfect, the distribution of corrected reads resembles the theoretical distribution of an ideal error-free set more than the distribution of raw reads.

As described in Section 2.1, not all input reads, even from the corrected set, can be processed by our algorithm. Table 4.2 summarizes numbers of reads discarded for various reasons. The preprocessing phase removed nearly one fifth of the corrected data set (18.99%). Most of the reads were removed due to being possible duplicates (87.65%). Quite a large portion of reads were not accepted because of their low mapping quality (12.97%). Also, about 3% of all the reads were shortened in order to remove soft-clipped regions.

4.2 Quality Evaluation

In order to evaluate the algorithm, the generated VCF files were compared to those generated by the following variant calling toolchains:

- **GATK.** These VCFs were taken as reference points, since the method used
by GATK should be very similar to our algorithm. Hence, results of all algorithms were compared against them.

- **Fermikit.** Unlike GATK and our algorithm, Fermikit’s assembly algorithm is based on the OLC concept.

- **Fermikit on regions.** This is a combination of the OLC approach used by Fermikit and the short region one adopted by our algorithm (and also by GATK). The fermikit assembly algorithm was run on exactly the same regions as our algorithm. The aim of this test case was to force the Fermikit to minimize differences of outputs generated by DBG and OLC algorithms.

- **samtools mpileup, bcftools call.** A traditional variant caller implemented by SAMtools and BCFtools packages [13]. Results generated by these algorithms were compared to those of GATK using

Table 4.2: Categories of reads present within the corrected test data set

<table>
<thead>
<tr>
<th>Name</th>
<th>Value</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total reads</td>
<td>12,410,475</td>
<td>-</td>
</tr>
<tr>
<td>Bad reads</td>
<td>2,357,002</td>
<td>18.99% of all reads</td>
</tr>
<tr>
<td>Low MAPQ</td>
<td>305,588</td>
<td>12.97% of bad reads</td>
</tr>
<tr>
<td>Unmapped</td>
<td>5,020</td>
<td>0.21% of bad reads</td>
</tr>
<tr>
<td>Supplementary</td>
<td>33,621</td>
<td>1.43% of bad reads</td>
</tr>
<tr>
<td>Duplicate</td>
<td>2,065,795</td>
<td>87.65% of bad reads</td>
</tr>
<tr>
<td>Soft-clipped</td>
<td>305,209</td>
<td>3.04% of accepted reads</td>
</tr>
</tbody>
</table>

Figure 4.3: K-mer frequency distribution of the corrected read set
the tool \textit{rtgeval}. \textit{Rtgeval} is a wrapper for RTG’s \textit{vcfeval}, a sophisticated open source variant comparison tool developed by Realtime Genomics. It simplifies the use of \textit{vcfeval} and potentially helps to get consistent results given VCFs produced by different variant callers [14].

\textit{Rtgeval} accepts two VCF files on input: a test set and a truth set. The truth set is a VCF file generated by a reference algorithm, in our case GATK. The test set VCF is produced by the algorithm to be tested. The evaluation is done separately for SNPs and indels and each variant is sorted into one of three categories:

- **True positive (TP).** The variant is present in both sets.
- **False negative (FN).** It is present in the truth set only.
- **False positive (FP).** It can be found only in the test set.

\textit{Rtgeval} can compare VCF files in three different modes: positional, allelic and genotypic. The positional mode is intuitive; the tool is determining whether the same variants are present at approximately the same position inside both test and truth set. If the positions difference does not exceed 10 bases, the variants are considered true positives. Otherwise, either false negative, or false positive is reported. In allelic mode, the tool focuses on biallelic variants and evaluates whether they are correctly detected by both tested algorithms. The genotyping mode compares genotype and phasing information of the variants.

### 4.2.1 Positional

The comparison in positional mode is shown in Table 4.3. It is clear that variants called by OLC and DBG classes of algorithms indeed are not the same. The percentages were counted from the total number of variants in GATK’s VCF file, separately for SNPs and indels.

Fermikit, as a representative of the OLC class, performs well when looking at numbers of false positives (below 1% in case of SNPs, ~2.3% indels), which is, however, traded off by quite big percentage of undetected variants (about 10% for SNPs and 28% indels). Numbers of false negatives are reduced when the algorithm is run on the region basis, although false positives roughly triple. Such situation seems to be typical also for other algorithms or different parameter settings of one algorithm; when the number of false negatives drops, false positives rise and vice versa.

Our algorithm was run with default settings described in Section 2.2. The value of 4 for the global threshold is, compared for example to HaploCall that uses 2, high. It was chosen to fight high numbers of false positives. The default settings actually removes the binomial test from the variant filtering process since no low quality variants actually emerge. As discussed later, the binomial test, although useful in some cases, did not prove to be a world-saver.

When talking about SNPs, results produced by the SAMTools and BCFTools tandem form quite the opposite to those of the Fermikit run at the whole genome. Less than 3% of SNPs found by GATK were left undiscovered, however, however, the number of false positives increased. It is true that our algorithm did not
Table 4.3: Positional comparison of results generated by our algorithm

<table>
<thead>
<tr>
<th>/</th>
<th>Fermikit</th>
<th>Fermikit (regions)</th>
<th>mpileup</th>
<th>Our algorithm</th>
</tr>
</thead>
<tbody>
<tr>
<td>SNP TP</td>
<td>45,241 (89.3%)</td>
<td>47,630 (94%)</td>
<td>49,201 (97.1%)</td>
<td>48,172 (95%)</td>
</tr>
<tr>
<td>SNP FN</td>
<td>5,432 (10.7%)</td>
<td>3,043 (6%)</td>
<td>1,472 (2.9%)</td>
<td>2,501 (5%)</td>
</tr>
<tr>
<td>SNP FP</td>
<td>385 (0.76%)</td>
<td>1,441 (2.84%)</td>
<td>2,090 (4.12%)</td>
<td>816 (1.61%)</td>
</tr>
<tr>
<td>INDEL TP</td>
<td>7,853 (71.6%)</td>
<td>9,602 (87.55%)</td>
<td>8,707 (79.39%)</td>
<td>9,476 (86.4%)</td>
</tr>
<tr>
<td>INDEL FN</td>
<td>3,114 (28.4%)</td>
<td>1,365 (13.45%)</td>
<td>2,260 (20.61%)</td>
<td>1,491 (13.6%)</td>
</tr>
<tr>
<td>INDEL FP</td>
<td>250 (2.28%)</td>
<td>835 (7.61%)</td>
<td>1,412 (12.95%)</td>
<td>1,255 (11.4%)</td>
</tr>
</tbody>
</table>

achieved nearly the same number of false negatives even when run with low global threshold causing loads of false positives.

A deeper analysis of the results obtained by running our algorithm with lower values of the global threshold (2 and 3 to be precise) and different initial k-mer sizes suggests that the variant filtering step needs additional work to be done. It is true that information stored inside the variant graph can be used to get hints what variants should be filtered out. Variants with the reference and alternate sequences covered by the same reads, or paired-end reads seem to be, at first glance, good candidates for false positives. Unfortunately, these information need not to be treated as hard facts. Simply classifying variants with such properties ad false positives did not bring expected results. Furthermore, coloring restrictions of the variant graph do not permit to deduce genotypes for such cases.

Furthermore, the analysis suggested that many of the undetected GATK variants were removed because of their low read coverage. The low number of supporting reads was not, however, caused by a sequencing error, but by a lower read coverage of the whole active region. Hence, introducing a binomial test to resolve such cases seemed to be a good idea. Effects of the test addition were observed more closely on our algorithm run with the initial k-mer size set to 31, and with global threshold of 2. Figure 4.4 shows number of TPs and FPs filtered out when the binomial test had been applied to all called variants. Figure 4.5 is a result of applying the test only to variants with less than 4 supporting reads (the low quality variant threshold was set to 3).

The test run produced a total of 59062 variants shared with GATK’s VCF, and 4738 false positives. As Figure 4.4 illustrates, even setting a value of the binomial threshold to 1% removes about 2100 of true positives. Although this is roughly only 3%, we do not consider the case as a good trade-off even with regard to percentage of removed false positives (nearly 50%).

When applied only to the variants with low read support, the binomial test gives much more sense. Setting the binomial threshold to 1% removes only 411 true positives but has great effect on sorting out the false variants. Unfortunately, we do not consider the effect as great enough, since running our algorithm with lower global threshold produces more false positives than the binomial test removes.
Figure 4.4: Numbers of TPs and FPS removed by the binomial test applied to all called variants

Figure 4.5: Numbers of TPs and FPS removed by the binomial test applied to variants with less than 4 supporting reads
4.2.2 Genotyping

We were unable to use the Rtgeval tool in the genotypic mode, causes of its failure are unknown. Error reporting mechanisms of the tool are not too advanced because the failure is reported as an exception caused by opening a non-existent file. All input VCF files were validated.

Nevertheless, results would not be better than the positional case. Our algorithm fails to color about 5% of the variant graph. Genotyping information of uncolored variants is incorrect by design. Also, we are filling the phasing information, based on paired-end information and read coverage intersections of individual variants, much more aggressively than other solutions. For this reason, the test data set may prove improper for genotypic evaluation.
5. Conclusion and Future Work

We have developed an algorithm for reference-aided local genome assembly. We have compared it against well-known solutions, such as GATK, Fermikit or the traditional caller implemented in SAMTools. Although the comparison did not ended badly, there definitely is a space for improvements. Our implementation, stored on a DVD-ROM attached to this work, should be viewed as a prototype, not a solution ready for production. The implementation is not as fast as it should be, since no extra performance optimizations were made.

The algorithm should also be tested with more data sets, not just only at a 40 MB region of the first human chromosome. Additional testing may prevent over-tuning to certain test cases. One of the test cases should include variants with more phasing information. Our algorithm deduces the phasing information much more aggressively than others.

Probably the weakest part of the algorithm is the variant filtering step which should recognize and eliminate fake variants. More sophisticated methods than a binomial test and read coverage should be applied. For example, the algorithm does not utilize base qualities because our attempts to do so did not improve the results in any significant manner. Because of an absence of a good variant filtering mechanism, a constant quality value is assigned to all called variants since we are not able to reliably tell how much a certain variant may be fake. The binomial test seemed to be a reasonable solution, however, it succeeded only on variants with low read coverage, and even then its successes were limited.

The variant graph constructed during computation of genotype and phasing information, also may have a great potential in recognizing fake variants. Unfortunately, all attempts to do so failed so far.
References


[10] 100 Genomes Project: chr1 Illumina high coverage Read set ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/technical/pilot2_high_cov_GRCh37_bams/data/NA12878/alignment/ 20 Jul 2017


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