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DACCOR - Detection, charACterization, and reconstruction of Repetitive regions in bacterial genomes

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ABSTRACT

The reconstruction of genomes using mapping based approaches with short reads experiences difficulties when resolving repetitive regions. These repetitive regions in genomes result in low mapping qualities of the respective reads, which in turn lead to many unresolved bases of the genotypers. Currently, the reconstruction of these regions is often based on modified references in which the repetitive regions are masked. However, for many references such masked genomes are not available or are based on repetitive regions of other genomes. Our idea is to identify repetitive regions in the reference genome de novo. These regions can then be used to reconstruct them separately using short read sequencing data. Afterwards the reconstructed repetitive sequence can be inserted into the reconstructed genome. We present the program DACCOR, which performs these steps automatically. Our results show an increased base pair resolution of the repetitive regions in the reconstruction of Treponema pallidum samples, resulting in fewer unresolved bases.

INTRODUCTION

Modern genome reconstruction often relies on mapping programs such as BWA (Li and Durbin, 2009) to align short reads generated by Next-Generation-Sequencing (NGS) technologies to a known reference genome (Veeramah and Hammer, 2014). The consensus sequence of the aligned reads can then be used to generate the genomic sequence of the newly sequenced sample, assuming that the sample was sequenced with a sufficient coverage depth. This allows for the fast identification of short insertions, deletions, and single-nucleotide polymorphisms (SNPs). The mapping programs typically calculate a score for each aligned read that corresponds to the quality of the alignment (Li et al., 2008). The score quantifies the probability that a read is placed at the correct genomic position. Reads with a low mapping quality can be filtered out to remove reads that might stem from contaminations or were sequenced with low sequencing quality (Smith et al., 2008). Besides bad quality reads, also reads mapping to repetitive regions could yield low quality scores if they cannot be mapped to a unique position. Filtering of reads with low mapping qualities would also include these reads. This filtering is often conducted in the context of ancient DNA (aDNA) (Bos et al., 2016), so that for such samples the repetitive regions of the respective reconstructed genomes are generally affected.

However, repetitive regions play an important role in the genome (Shapiro and von Sternberg, 2005). Hundreds to thousands of such regions are present in prokaryotic and eukaryotic chromosomes (Treangen et al., 2009). The human genome, for example, consists of approximately 50% repetitive regions (Lander et al., 2001). Tandem repeat regions appear to encode outer membrane proteins, which suggests that they help pathogens to adapt to their hosts (Denoeud and Vergnaud, 2004). In the case of the bacterium Treponema pallidum, repetitive sequences in the arp gene are used to distinguish between the subspecies that cause veneral syphilis (Treponema pallidum pallidum), nonveneral yaws (Treponema pallidum pertenue), and bejel (Treponema pallidum endemicum), which is not possible using serological tests (Harper et al., 2008).
New sequencing technologies like the Illumina SLR platform, PacBio, or Oxford Nanopore are able to create long reads that can span most repetitive regions in order to resolve them (Huddleston et al., 2014). However, it is not always possible to apply these technologies to DNA samples. For example, in aDNA projects the average extracted fragment length is approximately 44 to 72 base pairs (Sawyer et al., 2012). Sequencing these fragments with long read technology would not result in any information gain. Additionally, the sequencing of hard to cultivate pathogens, like *Treponema pallidum*, also result in short DNA fragments similar to aDNA (Arora et al., 2016).

In order to better resolve the repetitive regions using short reads, researchers often first mask duplicated and low-complexity regions prior to the read mapping (Frith et al., 2010). For the human reference genome, for example, a masked reference is already available (UCSC, 2014). If no masked genome is available, programs like RepeatMasker (Smitt et al., 1996) can identify these regions and create a masked reference. RepeatMasker uses libraries of known repetitive regions and compares them to the input sequence. While this allows masking of the genome, *de novo* identification of repetitive regions is not possible.

A program that can identify repetitive regions *de novo* is VMatch (Kurtz, 2003). It uses suffix-arrays (Weiner, 1973) to identify the repetitive regions. VMatch has been applied in multiple genome projects for annotation of repetitive regions (e.g. by Lindow and Krogh (2005)), as well as masking tasks (Assuncao et al., 2010).

The general idea of our approach first starts with a *de novo* identification of all repetitive regions in a given reference genome. These identified regions, together with the full reference genome, are then used as separate references to reconstruct them for each individual sample, again using a mapping approach. The reconstructed repetitive regions can then be combined with the reconstruction of the full genome to increase the base-pair resolution of the reconstructed genome for each sample.

**METHODS**

The methodology of DACCOR (see Figure 1) first identifies all repetitive regions in a genome *de novo*. Each of these regions are then reconstructed individually, before they are combined with the rest of the reconstructed genome.

To create an integrated version to identify repetitive regions, our *de novo* approach uses a k-mer based approach, similar to WindowMasker (Morgulis et al., 2005). The workflow to identify these regions can be split up into six steps (see left part of Figure 1). In the first step, the reference genome is divided into its distinct k-mers and all k-mers that occur more than once are stored. In the second step, matching k-mer pairs overlapping at \((k-1)\) positions are combined into \((k+1)\)-mers and stored again. This second step is repeated until all maximal unique repetitive regions are identified. Afterwards low complexity regions, e.g. long regions consisting only of one base, are identified. This step is necessary, because they are identified as two identical repetitive regions directly next to each other and can be combined.
into one region. In the following two steps, repetitive regions with mismatches are identified. Here, a
mismatch-marker representing any unknown base is added to the end of all currently identified maximally
exact repetitive regions. This representation allows us to combine two previously identified repetitive
regions that are separated by these mismatches. Matching pairs of separated regions can then be combined
into repetitive regions with mismatches. In the last step, leading and trailing mismatch markers are
removed. All remaining mismatch markers are replaced with the character N.

DACCOR is not limited to this repeat finding approach. It fully supports the output format of Vmatch,
meaning that the integrated repeat finding approach can be substituted with the result of a previous run
of Vmatch. The regions identified by Vmatch are then extracted and saved in separate fasta files to be
used in the next step.

The repetitive regions that are identified after the six steps are saved to a multi-fasta file. In order
to be able to use each repetitive region directly as reference for a mapping pipeline, it is also possible
to write each identified sequence to a separate file. Additionally a summary of all identified repetitive
regions, as well as a bed file to view the location of the regions in a genome viewer, are created. Users
can specify a minimal length for all reported repeats.

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In order to reconstruct the repetitive region from sequenced fastq files, we use the EAGER pipeline
developed by Peltzer et al. (2016). It can preprocess the raw reads, including adapter clipping and quality
trimming, map them against a given reference, and generate a consensus sequence in the fasta file
format. For the consensus reconstruction, it uses the results of the genotyping results, following GATK
BEST Practice’s guidelines (Van der Auwera et al., 2013).

The de novo identification of repeats in a genome, the automatic separate mapping of NGS data
against all repetitive regions, as well as the subsequent enhanced reconstruction of the genome of NGS
samples has been implemented in DACCOR (Detection charACterization and reConstruction of Repetitive
regions in genomes), a stand-alone program written in Java.

The repeat identification methodology, as described above, is implemented in the identify sub-
program of our program and can be used to identify repetitive regions in a given reference genome de
novo.

The reconstruct subprogram of DACCOR automatically generates EAGER configuration files for
a given reference, its identified repetitive regions, and multiple sequencing samples.

Additionally, the combine subprogram can use the EAGER output of the reconstructed regions, as
well as the EAGER output of the whole genome and combine these reconstructions. Because the origin
of the reconstructed subsequences are known, they can replace the bases in the original reconstruction
generated without specific repeat resolution. For each repetitive region, the respective positions in the
genome are replaced if the original reconstruction resulted in an unknown (N) character.

To be able to automatically assemble a genome with all its repetitive regions, the subprogram
pipeline first identifies all repetitive regions in a given reference. Afterwards the NGS samples
are automatically reconstructed against both the complete reference and each identified repetitive
region individually using the EAGER pipeline. Finally, these individual regions are combined with the
reconstructed genome sequence to increase the resolution in repetitive regions.

The identify subprogram can identify repetitive regions within as well as between different
chromosomes or a bacterial genome and its plasmids. This is done by combining the identified k-mers of
all sequences in a given multi-fasta reference file. To be able to match identified repetitive regions to
the corresponding sequence, a unique offset is added to the indices of the start location of each region.

To evaluate our method, we applied DACCOR to several bacterial genomes of various lengths and
repetitiveness. We first compared the step of repeat identification with VMatch (Kurtz, 2003), allowing
for up to five mismatches in repetitive regions of a minimum length of 101 base-pairs (the length typical
Illumina HiSeq reads). We then applied our proposed reconstruction method to the syphilis samples
published by Arora et al. (2016), Pinto et al. (2016), and Sun et al. (2016) to reconstruct the sequences
of the 16S and 23S rRNA, which are duplicated in the bacterium Treponema pallidium. For this, we
first used DACCOR to identify all repetitive regions with at most five mismatches and a minimal length
Table 1. Comparison of the identified repetitive positions in different bacterial genomes of DACCOR and VMATCH, which was seen as the golden truth to evaluate against. Both programs were run allowing for one mismatch and reporting only regions of at least 101 bp. The k-mer size for DACCOR was set to 17.

|          | T. pallidum | S. flexneri | E. coli | M. leprae |
|----------|-------------|-------------|---------|-----------|
| true positives | 22,382      | 376,669     | 107,456 | 74,406    |
| true negatives | 1,116,478   | 4,211,322   | 4,535,106 | 3,192,194 |
| false positives | 0          | 354         | 463     | 131       |
| false negatives | 773        | 18,857      | 3,307   | 1,472     |
| accuracy (%)  | 99.93       | 99.58       | 99.92   | 99.95     |

of 101 base pairs in the Nichols strain. We also reconstructed the full genomes of the samples using the standard EAGER pipeline. This allowed us to compare our reconstruction of the two genes to the reconstruction generated by the standard method using the full genome as a reference. In order to identify specific variations in either copy of the genes, we searched for heterozygous positions in the individual reconstructions of the extracted sequences that show an allele frequency between 25 and 75%.

Finally we applied the full DACCOR pipeline to reconstruct the whole genomes, including the identified repetitive regions, of two syphilis samples. For this we chose two samples from Arora et al. (2016), one with a moderate coverage (AR1, 7X) and one with a very high coverage (AR2, 157X).

RESULTS

We first evaluated the identification of repetitive regions by comparing these to the repeats identified by VMATCH. This comparison (see Table 1) shows that the results of both programs are almost identical. We considered VMATCH as the “golden truth” and could therefore compute an accuracy for the repetitive regions reported by DACCOR. We achieved a very good accuracy with over 99% in all tested cases with only very few false negatives as well as false positives.

Next we reran DACCOR with a higher sensitivity allowing for up to five mismatches in repetitive regions of lengths at least 101 base pairs. These results of the different bacterial genomes (see Table 2) show that the Shigella genome contains by far the most repetitive regions (1,249 compared to 29 in Nichols, 242 in E. coli, and 190 in M. leprae). It also contains the longest repetitive regions of the four bacteria (5,383 compared to 3,283, 3,141, and 2,578). The average lengths of the repetitive regions are quite similar in all four bacteria (between 570 and 823 base-pairs). The number of repetitive regions that can be identified when allowing for up to five mismatches also varies between the different bacteria. There are 127 of regions containing mismatches in the Shigella genome, whereas there are only 47 in the genome of E. coli, 9 in M. leprae, and 1 in Nichols. Overall 8.3% of the Shigella genome is comprised of repetitive regions. The genomes of E. coli and M. lepra are comprised of 2.4% and 2.3% repetitive regions respectively, and only 2.0% of the Nichols genome is repetitive.

The runtime linearly correlates with the number of identified repetitive bases (see Figure 2). After an initial preprocessing step, DACCOR identifies about 12,000 repetitive bases per minute. The k-mer size

Table 2. Statistics of identified repetitive regions using several bacterial genomes for a k-mer size of 17, at most five mismatches, and a minimum length of 101 base pairs.

|          | T. pallidum | S. flexneri | E. coli | M. leprae |
|----------|-------------|-------------|---------|-----------|
| genome size [bp] | 1,139,633    | 4,607,202   | 4,646,332 | 3,268,203  |
| # repetitive regions | 29          | 1,249       | 242     | 190       |
| different repetitive regions | 14          | 482         | 104     | 76        |
| max length of repetitive regions | 3,283       | 5,383       | 3,141   | 2,578     |
| average length | 823         | 570         | 706     | 643       |
| repetitive regions with mismatches | 1           | 127         | 47      | 9         |
| sum of repetitive bases | 23,892      | 660,261     | 160,897 | 119,871   |
| % of genome repetitive (non overlapping) | 2.0         | 8.3         | 2.4     | 2.3       |
Figure 2. Runtime of DACCOR in relation to the number of identified repetitive bases.

does not influence the runtime (see supplementary Figure 2).

The two longest identified repetitive regions in the Nichols genome correspond to the 16S and 23S rRNAs, respectively. Both operons contain two copies of the gene. We extracted these regions and used

Figure 3. Histogram of resolved bases (as fraction relative to the length of the respective gene) of the 16S rRNA (a) and 23S rRNA (b) in 106 clinical syphilis samples (Arora et al., 2016; Pinto et al., 2016; Sun et al., 2016). The number of resolved base-pairs when mapping against each copy of the gene individually in comparison to the standard mapping approach using both copies has been computed.
Table 3. SNPs in the 16S and 23S rRNAs identified after extracting the repetitive region. *Site-specific* positions refers to positions that appear to be different in the two copies of the respective gene (allele frequency between 25 and 75%).

|                | 16S rRNA | 23S rRNA |
|----------------|----------|----------|
| length of gene | 1495     | 2900     |
| number of variant positions | 36        | 46        |
| number of site-specific positions | 30        | 43        |

them as two independent references for their reconstruction off all samples published by Arora et al. (2016), Pinto et al. (2016), and Sun et al. (2016). We mapped against each copy of the gene individually and count the number of resolved bases in each copy. This number was compared to the number of resolved bases in the two copies of the respective gene when mapping against the whole genome without repeat masking. We then computed the difference of these two numbers and divided by the length of the gene. We could improve the base-pair resolution (see Figure 3) by a median value of 82.7% for the 16S and 87.4% for the 23S rRNA. It shows that the percentage of the resolved base-pairs was at least as high when mapping only against the extracted sequences, compared to the mapping against the whole genome for all analyzed samples. This means that we do not lose resolution when mapping only against these sequences, but in almost all cases gain information up to an improvement of 100% of the sequence of the respective gene.

Using the individual reconstructions of the 16S rRNA and the 23S rRNA gene sequences, we tried to identify SNPs that are specific for only one copy of the respective gene. The results of this analysis identified 36 positions that have a SNP call in at least one of the 106 samples for the 16S rRNA (see Table 3). Of those 36 positions, 30 show evidence for a site-specific SNP in at least one analyzed clinical samples. There are two positions (884 and 888 relative to the start of the 16S rRNA), which show site-specific variance in about 20% of the samples (see supplementary material). In the 23S rRNA there are 46 positions where at least one sample has a SNP. Of these 46, 43 show evidence for site-specific SNPs in at least one sample. Here, one position (2003) shows evidence for site-specificity in 37% of the samples. Additionally, there are nine positions that show site-specificity in at least 15% of the samples.

Finally, we compared the total number of unresolved positions in the samples AR1 and AR2, published by Arora et al. (2016), between DACCOR and EAGER with the full genome as reference (see Table 4).

|                | AR1 EAGER | AR1 DACCOR | AR2 EAGER | AR2 DACCOR |
|----------------|-----------|------------|-----------|------------|
| #N             | 23,348    | 4,473      | 17,549    | 964        |
| %N             | 76.98     | 14.75      | 57.86     | 3.18       |
new draft genome is created by combining the reconstructed repetitive regions with the rest of the genome that has been reconstructed using a standard mapping approach.

Our de novo identification of repeats uses a *k*-mer approach. The choice of *k* is important for the identification of repeat regions. For de novo assembly based on de Bruijn graphs, the optimum depends on the genome length, the coverage, the quality, and the length of the reads (Zerbino and Birney, 2008). In the case of different read lengths, as often observed in aDNA projects, it has been shown that using multiple different *k*-mers improves the assembly (Seitz and Nieselt, 2017). Other approaches choose an optimal *k*-mer size so that the uniqueness is maximized (Gardner and Hall, 2013). However, as we want to identify repetitive regions, we do not want to use a *k*-mer size in order to maximize uniqueness. In our approach the *k*-mer size defines the minimum length of the repetitive regions that can be identified in the first step. Thus, a *k*-mer size as small as possible should be used to be able to identify all putative repeat regions. However, very small *k*-mer sizes lead to an exponential increase in the runtime (see supplementary Figure 2), due to the increasing number of random occurrences of these small *k*-mers that have to be accounted for. Therefore, we propose a minimum size for *k* of 17. On the other hand, the *k*-mer size should also not be longer than half the length of the input reads.

The runtime of **DACCOR** correlates mainly with the repetitiveness of the genome, i.e. with the number of repetitive bases that are identified in the respective genome. When allowing mismatches, a slight increase in runtime is also noted.

To identify all repetitive *k*-mers, all possible *k*-mers are stored in the first step of the identify subprogram of **DACCOR**. Those that are not repetitive are removed after the screening. Nonetheless, this results in high memory usage if the genome contains many repetitive regions. For the example of *Shigella flexneri*, we observed a memory footprint of 8 GB with a *k*-mer size of 17.

The comparison between **DACCOR** and **VMatch** showed that **VMatch** is slightly more sensitive, probably due to its suffix array approach. A possibility to increase the sensitivity of **DACCOR** would be to elongate all identified repetitive regions based on a local alignment.

When mapping short reads from typical NGS data, a number of approaches recommend to use genomes as references whose repetitive regions are masked (Tarailo-Graovac and Chen, 2009). However, this may be problematic, because repetitive regions often overlap and can be quite complex. An example for this is the *arp* gene of *Treponema pallidum* (see Figure 4). It contains several overlapping repetitive regions. It can be seen that the region labeled *repeat_3* is partly repetitive with itself. Thus a masking of one of the repetitive regions would mask most of itself. Additionally, the masking of the second occurrence of *repeat_1* would also mask most of both occurrences of *repeat_3*. When masking the first occurrence of *repeat_1*, the masking of either occurrence of *repeat_2* would also mask part of the unmasked region of *repeat_1*. Thus, masking of repetitive regions could result in either loosing genome information or leaving repetitive regions unmasked. We therefore propose to use the identified repetitive regions as separate references for the mapping, and merge all individually reconstructed regions into a common draft genome.

Since the *de novo* identification of repeats in genomes, using these then as individual references for the mapping and merging all reconstructed genomic regions into a final draft genome, require many different steps, our main goal of **DACCOR** was to present a fully automatic procedure encompassing all these steps. **DACCOR** makes use of **EAGER**, a pipeline for the automatic reconstruction of genomic data sets. For highly identical repeat regions, each copy is stored as individual sequences together with a margin at the 5’ and 3’ end of the region. We propose to set the margin to twice the maximum read length to cover all

![Figure 4. Repetitive regions contained in the *arp* gene in *Treponema pallidum*](image)
reads spanning over the ends of the repeats.

Using DACCOR, we have shown that a higher base-pair resolution, compared to the reconstruction using the standard mapping approach in repeat regions, can be achieved. In the standard approach, reads that can be mapped to different locations result in a mapping quality of zero, which in turn decreases the genotyping quality (McKenna et al., 2010). By mapping to the repeat region only, these reads have a higher mapping quality, a higher genotype quality, and thus result in more resolved positions. However, one has to acknowledge that other unresolved bases, that may be due to lack of coverage or low sequencing quality, can not be resolved with our approach. In the case of the two syphilis samples, we could show that the majority of unresolved bases in repeat regions could be resolved, and that the remaining unresolved bases lie outside of the identified repetitive regions. Furthermore, we have shown that using DACCOR, the identification of SNPs in repeat regions can be improved. This is especially useful for the 23S rRNA, as it is known to play a role in the antibiotics resistance of bacteria (Arora et al., 2016).

In conclusion, we have developed a fully automatic pipeline that first conducts a de novo repeat identification in bacterial genomes and then uses the repeat regions for an enhanced mapping of short read NGS data. Increasing the resolution of a draft genome has an effect on many downstream analyses, such as population genetics or phylogenetic analyses. For future improvements we plan to reduce the runtime and memory usage by adjusting our data structure and by adding more parallelization to some of the compute steps. With this we hope to eventually be able to identify repetitive regions also in large eukaryotic genomes, like the human genome.

SOFTWARE AVAILABILITY

We have developed an automated software pipeline, written in Java, which allows other researchers to use our methodology. This pipeline is available on github:

https://github.com/Integrative-Transcriptomics/Daccor

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