Finding rearrangements in nanopore DNA reads with LAST and dnarrange

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Abstract

Long-read DNA sequencing techniques such as nanopore are especially useful for characterizing complex sequence rearrangements, which occur in some genetic diseases and also during evolution. Analyzing the sequence data to understand such rearrangements is not trivial, due to sequencing error, rearrangement intricacy, and abundance of repeated similar sequences in genomes.

The \texttt{last} and \texttt{dnarrange} software packages can resolve complex relationships between DNA sequences, and characterize changes such as gene conversion, processed pseudogene insertion, and chromosome shattering. They can filter out numerous rearrangements shared by controls, e.g. healthy humans versus a patient, to focus on rearrangements unique to the patient. One useful ingredient is \texttt{last-train}, which learns the rates (probabilities) of deletions, insertions, and each kind of base match and mismatch. These probabilities are then used to find the most likely sequence relationships/alignments, which is especially useful for DNA with unusual rates, such as DNA from \textit{Plasmodium falciparum} (malaria) with \(~80\%\, a+t). This is also useful for less-studied species that lack reference genomes, so the DNA reads are compared to a different species’ genome. We also point out that a reference genome with ancestral alleles would be ideal.

Keywords: gene conversion, processed pseudogene, chromothripsis, mutation, probability, alignment, evolution, ancestral

1 Introduction

The LAST software was first made public in 2008: it was intended as a general tool for finding and aligning related regions in gigabase-scale sequence data. It is not really designed for recent long-read data such as nanopore, nevertheless it can be used, and has some unique advantages. The main advantages are:

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Figure 1: Rates (probabilities) of base matches, mismatches, deletions, and insertions between nanopore DNA reads and reference genomes. In each 4×4 matrix, a row corresponds to a genomic base, and a column to a read base.

1. It can learn the rates (probabilities) of deletions, insertions, and each kind of base match and mismatch (Fig. 1), which are due to a combination of sequencing errors and real sequence differences. It then uses these probabilities to determine the most probable alignments (1).

2. It can disentangle complex rearrangements and duplications between sequences, even when confounded by repeats, by finding the most probable split of a sequence into rearranged parts based on the alignment probability of each part (2).

Another useful feature is that LAST can compare DNA to protein sequences, allowing frameshifts, which are frequent in long DNA reads. This has been used for taxonomic and functional binning of metagenomic long reads (3).

Here we shall focus on finding rearrangements from LAST alignments of nanopore DNA reads to a reference genome. The next problem is that we typically find thousands of rearrangements, many of which seem to be artifically-rearranged reads, and many others real-but-benign (4, 5). The artifacts seem to be sporadic, so they can be depleted by sequencing to several-fold coverage of the genome and then discarding reads with unique rearrangements not shared by any other read (4, 5). When seeking rearrangements causing genetic disease, we also use nanopore reads from control individuals without the same disease, and discard benign rearrangements shared with controls. These tasks downstream from LAST are done by dnarrange.

1.1 Examples of match, mismatch, insertion, and deletion rates

Three examples of these rates, for nanopore DNA reads versus reference genomes, are shown in Fig. 1. The human rates are for one set of human nanopore reads, HG02723 (6), versus a reference human genome (hg38). We can see, for example, that a→g substitutions are much more frequent than other substitutions, which is often the case for nanopore sequences (4, 5). Also, the probability (rate) of opening a deletion is much higher than an insertion. The P. falciparum rates are for one set of nanopore reads (7) versus a reference genome: these are interesting because P. falciparum DNA is ∼80% a+t, so the match and mismatch rates are quite different from human. The final example is a set of nanopore reads from the ant Camponotus maculatus (7) versus the closest-available reference genome: Camponotus floridanus. This example has higher substitution rates, as expected between different species.
Figure 2: A made-up example of complex rearrangements during DNA sequence evolution. Each colored block represents a segment of DNA, e.g. a few hundred bp. These segments become rearranged, duplicated, or deleted. The orange blocks labeled n and s are similar sequences from an earlier duplication, i.e. paralogs. Reproduced from ([5]).

1.2 Understanding rearrangements

Various mutational processes can cause almost arbitrarily complex rearrangements, duplications, and deletions of DNA sequence (Fig. 2). For example, chromosomes can shatter into many fragments that rejoin in a different order and orientation, or DNA polymerase can template-switch during DNA replication causing rearrangements, duplications, and deletions. No matter how complex these changes, there is one key point: almost every part of a descendant sequence comes from a unique part of ancestral sequence(s). This is shown in Fig. 2b: we can visually scan the derived sequence from top to bottom and see where each part comes from (red lines).

We might wonder about “insertions”: where does the inserted sequence come from? It must come from somewhere: typically it has been duplicated or moved from elsewhere in the genome. Rarely, an insertion may come from a different genome, such as a virus. There is also non-templated insertion aka “spontaneous generation” of new sequence with no ancestor: this is the only exception to the rule.

Unfortunately we do not have ideal ancestral sequences: in practice we compare DNA reads to a “reference” genome which has separately undergone complex changes. The relationship between two descendant sequences is fundamentally less simple (Fig. 2c), e.g. duplication in both lineages creates a many-to-many relationship, and reference-specific deletion means that parts of
DNA reads correctly align nowhere in the reference, but may align incorrectly to paralogs. Even if we could perfectly detect the correct relationships (red lines in Fig. 2c), it is hard to understand what changes have occurred.

In practice, we assume the reference genome is ancestral, even though it is not. We are interested in rearrangements that cause genetic disease: at least in these cases the reference is likely to represent the ancestral state. If virus insertions are suspected, viral chromosomes can be added as extra chromosomes to the reference. We will find spurious rearrangements because the reference is not perfectly ancestral: the hope is that these will also be found in control data, thus discarded. Another way to detect reference-genome regions with non-ancestral status is by comparing to an outgroup, e.g. an ape genome (4). It would be useful to have a reference genome that contains ancestral alleles (4, 5).

1.3 Simple sequences
DNA is rife with “simple sequences” such as catcatcatcat or aaatattaaaaacaaa. They cause many similarities between sequences that are not correct relationships, because they are not descended from a common ancestor. There are several methods for detecting and “masking” simple sequences, which do not all work equally well (8). LAST uses tantan (8) to detect simple sequences, and converts them to lowercase letters. However, masking can cause problems by hiding correct relationships between sequences, which might cause incorrect relationships to be found instead. So, by default LAST treats lowercase the same as uppercase when finding alignments. Alignments without a significant amount of uppercase-to-uppercase alignment are suspicious, and can optionally be discarded by last-postmask.

2 Methods
2.1 Installation
It may be easiest to install the software from Bioconda (9), or from Debian Med (10). After setting up Bioconda on your computer, this command installs dnarrange and all its dependencies, including LAST:

    conda install dnarrange

At the time of writing, conda seems to have a bug where it may install old versions of software: it may be better to use mamba (https://github.com/mamba-org/mamba). The following discussion applies to LAST version 1296 and dnarrange 1.5.2, and not to older versions.

2.2 Getting the Camponotus rates
Let us first see how to get the ant results in Fig. 1. We need the reference genome sequence in fasta format, which we got by searching “camponotus” at NCBI Genome (https://www.ncbi.nlm.nih.gov/genome/). We renamed this file Cflo_v7.5.fa. The first step is to prepare “index” data-structures for the genome, which enable fast sequence comparison:

    lastdb -P16 -uNEAR antDB Cflo_v7.5.fa

This creates several files whose names start with antDB. The -P16 option makes it faster by running 16 parallel threads, with no effect on the result. Adjust as appropriate for your computer. (As a special case, -P0 uses as many threads as
your computer claims it can run simultaneously, which is a good way to annoy
the other users of a shared server.)

The \texttt{-uNEAR} option changes the seeding scheme. \textsc{last} uses a seed-and-extend
heuristic: it first find “seeds”, simple gapless alignments, then tries to extend
full alignments from the seeds. Although the seeds are gapless, they can allow
some mismatches: the more mismatches they allow, the longer they need to be
to retain specificity. \texttt{-uNEAR} specifies short seeds with few mismatches, which
is appropriate for searching indel-rich nanopore reads against a closely-related
genome. If you omit \texttt{-uNEAR}, it may not make much difference in practice.

Next, we need the DNA reads in fasta or fastq format. We do not use the
extra information in fastq, so fasta is preferable because the files are much
smaller. We can find the rates of substitution etc. between reads and genome
like this, using the original fastq file in this case:

\begin{verbatim}
last-train -P16 -Q0 antDB CmacRNAseq_180528.fastq.gz > ants.train
\end{verbatim}

The \texttt{-Q0} option makes it ignore the fastq quality data. (\textsc{last} can alternatively
use the quality data for more accurate training and alignment. This assumes,
however, that the qualities indicate probability of substitution error, not insertion
or deletion error, and we are not confident this holds for nanopore data.) If you
have fasta instead of fastq, \texttt{-Q0} has no effect and can be omitted.

It’s worth noting that \textsc{last-train} doesn’t actually use all the DNA reads: it
uses a random sample of size one million bases (probably overkill). This means
that giving huge read files to \textsc{last-train} doesn’t make it much slower, except
for the time needed to read the file and get a random sample.

\textsc{last-train} works iteratively: it compares the reads to the genome to find
the rates, then uses these rates to do the comparison more accurately and get
better rates, and so on. It prints data for each iteration (for troubleshooting),
with the final rates at the end. It’s not necessary to look at them: you can just
pass the output file to the next alignment step.

\textbf{2.3 Getting the \textit{Plasmodium falciparum} rates}

The \textit{P. falciparum} rates were found like this:

\begin{verbatim}
lastdb -P16 -uNEAR -R02 plaDB plaFal3D7.fa
last-train -P16 -Q0 plaDB P.falciparum_targeted_seq.fastq > pf.train
\end{verbatim}

The only change is the \texttt{-R02} option, which is recommended for DNA with \textasciitilde80
\texttt{a+t}. This option changes the \texttt{tantan} parameters for defining “simple sequence”,
which is necessary for very \texttt{(a+t)}-rich DNA. If you omit \texttt{-R02}, it might not
matter too much in practice. \textsc{last-train} avoids lowercased simple sequence,
but in this case it hardly affects the result.

\textbf{2.4 Getting the human rates}

The human rates were found like this:

\begin{verbatim}
lastdb -P16 -uRY4 hdb hg38.analysisSet.fa
last-train -P16 -Q0 hdb HG02723_1.fastq.gz > hum.train
\end{verbatim}

The change here is to use \texttt{-uRY4} instead of \texttt{-uNEAR}. This makes \textsc{last} faster and
use less memory, at a cost in sensitivity. The main reason for doing this is that
the fastq file is quite big: 81 gigabases. This is not necessary for \textsc{last-train},
which just uses a sample of the fastq: the point is to speed up the subsequent alignment step.

RY4 makes it use \( \frac{1}{4} \) of the seeds, in a similar way to this: just use seeds starting with \( a \). (It actually uses seeds starting with combinations of purines and pyrimidines \( [7] \), but the idea is the same.) There are faster alternatives using \( \frac{1}{8}, \frac{1}{16}, \) and \( \frac{1}{32} \) of the seeds: RY8, RY16, and RY32.

This last-train run took 23 minutes, most of which was spent decompressing the fastq file to get a random sample from it. It is not necessary to re-run last-train for each dataset, unless there is reason to think the rates may have changed, e.g. due to different versions of sequencing hardware or base-calling software.

2.5 Aligning human DNA reads to a human genome

The next step is to actually align all the reads to the genome. Here is a simple but slightly inefficient way:

```
lastal -P16 -p hum.train hdb HG02723_1.fastq.gz | last-split | gzip > out.maf.gz
```

`lastal` finds and aligns similar regions of the reads and the genome, often aligning the same part of a read to several genome regions. `last-split` cuts the output of `lastal` down to a unique best alignment for each part of each read. This is appropriate if the genome is ancestral to the reads (Fig. [3]). Sometimes, part of a read matches two or more genome regions with almost equal probability: `last-split` chooses the most probable alignment (arbitrarily if exactly equal), and outputs “mismatch probabilities”, the probability that each part of a read is aligned to the wrong place. Finally, `gzip` compresses the output (optional).

The `-P16` option uses 16 threads. This is not perfectly effective, partly because `last-split` and `gzip` are not parallelized, and partly because some threads finish each chunk of data earlier than others so they are not always all running\(^1\). We could shave \(~15\%\) off the total run time by using `gzip -5`, for faster but worse compression, which adds 6% to the output size.

We can get better parallelization by using GNU Parallel \([12]\), which is installed as a `last` dependency by Bioconda. `last` includes `parallel-fasta` and `parallel-fastq`, which run GNU Parallel with options for fasta or fastq, and can be used like this:

```
zcat HG02723_1.fastq.gz |
parallel-fastq -j16 "lastal -p hum.train hdb | last-split | gzip" > out.maf.gz
```

This runs the quoted pipeline 16 times in parallel, and concatenates the results. Fortunately, concatenation of `gzip` outputs is valid `gz` format, so this is ok. To assuage another concern: `lastal` reads the `hdb` files into shared memory, so this memory use does not get multiplied 16 times. The output gets accumulated in large temporary files, whose location can be specified if necessary to ensure enough space. This alignment took 5.5 hours.

2.6 An alternative way using WindowMasker

Our published human studies have mostly not used RY4 (because it’s somewhat new), and instead saved time by “masking” repeats. The main cause of alignment slowness is the abundance of repeats, such as LINEs, SINEs, and simple sequences, so each repeat in a read gets preliminarily aligned to multiple genome locations.

\(^1\)This “chunk” inefficiency might get fixed soon.
We can mitigate this problem by masking repeats. We wish to mask as little as possible in order to maximize sensitivity, but enough to make the run time tolerable. For this aim, WindowMasker (13) seems to work well: it is part of the BLAST package, which can be got from NCBI, Bioconda, or Debian Med. The following commands find repeats in the genome and convert them to lowercase:

```
windowmasker -mk_counts -in hg38.analysisSet.fa > hum.wmstat
windowmasker -ustat hum.wmstat -outfmt fasta -in hg38.analysisSet.fa > hum-wm.fa
```

The next step is to run lastdb like this:

```
lastdb -P16 -uNEAR -R11 -c hwmdb hum-wm.fa
```

The -R11 option retains lowercase from the input and additionally lowers cases simple repeats found by tantan. The -c tells it to “mask” lowercase: this will exclude lowercase from seeds but not from final alignments, and then discard alignments that lack a significant amount of uppercase-to-uppercase alignment (the same as last-postmask).

The subsequent last-train and alignment steps are the same as above. It turns out that lastal -P16 is competitive with GNU Parallel here, because the run time required for each read is more predictable (proportional to read length), so the threads finish each chunk at the same time.

### 2.7 Finding rearrangements with dnarrange

dnarrange gets rearranged DNA reads from the read-to-genome alignments, and performs two optional filtering steps. It discards reads with unique rearrangements not shared by any other read, and discards reads with rearrangements shared by control reads. So we need control DNA reads aligned to the same genome. A few control sets aligned to hg38 are available at https://github.com/mcfrith/dnarrange. We can run dnarrange like this:

```
dnarrange -v out.maf.gz : controls/hg38-* > groups.maf
```

The -v (verbose) option just makes it show progress messages as it’s working (useful for troubleshooting and reassurance that it’s doing something). The “:” separates case from control files. (It’s possible to use more than one case file, which only outputs groups that have reads from all case files.) The output has the alignments of the rearranged reads, in groups where reads that cover the same rearrangement are in the same group. In our studies of human patients, the number of groups per patient declines from thousands without controls to a few dozen after control filtering (5). Filtering is more effective when some controls are from the same family or ethnic group as the case individual.

It’s possible to not use control DNA reads, but then an overwhelming number of rearrangements may be found. They may include dubious rearrangements in regions where the reference genome does not represent the ancestral state, or where the reference is incomplete. For example, if a segment of the reference has been deleted, the corresponding part of a DNA read may get wrongly aligned to a paralog elsewhere in the genome, showing an incorrect rearrangement.

An alternative is to use last-postmask (if repeats were not already masked with lastdb option -c):

```
last-postmask out.maf.gz | dnarrange -v - : controls/hg38-* > groups.maf
```
The “-” has a standard meaning of reading the data that is piped in. As mentioned above, last-postmask discards alignments that are mostly of lower-cased simple sequence. There is doubt that such alignments reflect correct relationships. Nevertheless, such alignments can be informative because they do indicate changes such as expansion or insertion of simple sequence. To analyze tandem repeat changes, we have specialized software tandem-genotypes, described elsewhere in this volume.

2.8 Making dotplot figures of the rearrangements

We can make a dotplot figure of each group in groups.maf like this:

```
last-multiplot -a refGene.txt -a rmsk.txt groups.maf fig-dir
```

This puts the figures in a new directory fig-dir. We have used the -a option to show genes and repeats, using files downloaded from the UCSC Genome Browser (http://genome.ucsc.edu). It’s also possible to show BED, GFF/GTF, or RepeatMasker .out data, or unsequenced gaps in AGP or gap.txt format. Naturally, these dotplots are more useful when there are not too many of them, and making many of them is slow.

Some examples of human rearrangements, in HG02723.1 versus hg38, are shown in the remaining Figures. Fig. 3 shows a classic genetic phenomenon: gene conversion. This is a process where one DNA sequence gets replaced, probably during a DNA repair process, by a copy of a similar sequence. In this case, part of an L1 LINE element in chromosome 20 has been replaced by the homologous part of another L1 element. Here, the converted read parts are aligned to one L1 in chromosome 6, but their mismap probabilities are quite high (not shown), because they have almost equally probable alignments to other L1s. So the specific donor element may not be confidently knowable.

Gene conversion is a poster child for probability-based sequence alignment of the sort done by last-split. A naive alignment method would just align these DNA reads continuously to this L1 in chromosome 20, because the converted sequence is similar to the replaced sequence.

Fig. 4 shows a short-range rearrangement, localized within 5 kb, with a rather small rearranged fragment. This resembles mutations that have been attributed to template switching during DNA replication, which can create numerous complex mutation patterns, and have caused erroneous variant annotations.

Fig. 5 shows inerterion of a processed pseudogene from chromosome 11 into chromosome 13. This is a process where an mRNA molecule, after its introns have been removed, is reverse-transcribed and inserted into the genome (perhaps by a reverse transcriptase enzyme from a retrotransposon).

Finally, Fig. 6 shows a DNA read aligning to chromosome 1, with a gap in the alignment, which might be a deletion in the read or an insertion in the reference. The gap coincides exactly with an L1PA2 element, which is a young LINE retrotransposon. There is no known mechanism for precise excision of a LINE element, whereas LINE insertion is normal, so this is an insertion that occurred in the lineage leading to the reference sequence. This is an example where the reference does not have the ancestral state. Outgroups (other mammal genomes) lack this L1PA2 (Fig. 6).
Figure 3: Alignment of 2 human DNA reads to human genome hg38, showing gene conversion. The read identifiers are shown on the left: the final − or + indicates that the read has been reverse-complemented or not. The figure shows only part of each read. To the left of the vertical black line is 1.5 kb of chromosome 6, to the right is 10 kb of chromosome 20. The vertical stripes show repeat elements in hg38: pink = forward-oriented transposable element, blue = reverse-oriented transposable element, purple = simple repeat.
Figure 4: Alignment of 2 human DNA reads to human genome hg38, showing a short-range rearrangement. The top read is short, and its identifier is omitted.
Figure 5: Alignment of 14 human DNA reads to human genome hg38, showing insertion of a processed pseudogene from chromosome 11 into chromosome 13. The vertical green stripes show exons of USP28, which encodes ubiquitin specific peptidase 28.
Figure 6: Above: alignment of a human DNA read to human genome hg38, showing reference-specific insertion of an L1 LINE retrotransposon in chromosome 1. Below: alignments of other mammal genomes to this part of chromosome 1 (screen shot from http://genome.ucsc.edu).

### 2.9 last-dotplot

last-multiplot (which comes with dnarrange) uses last-dotplot (part of LAST) to draw each figure. It may be useful to run last-dotplot directly. For example, Fig. 4 was made like this:

```
last-dotplot -a refGene.txt -a rmsk.txt -j2 --labels1=2
-j chr4:35117500-35122500 groups.maf fig4.png
```

Option -j2 draws the gray lines joining the aligned segments, --labels1=2 adds the start coordinate and length to the top of the figure, and -1 chr4:35117500-35122500 shows that range of chr4.

### 2.10 Rearrangement types and thresholds

A rearrangement is indicated when two parts of a DNA read align to disjoint places in the genome. dnarrange classifies four kinds of disjointness:

1. Different chromosomes (Fig. 3)

2. Opposite strands of one reference sequence (Fig. 4)

3. Non-colinear alignment of two read parts to the same strand of the same reference sequence. In other words: the alignment of the upstream read part ends at coordinate $X$ in the reference sequence, the alignment of the downstream read part starts at coordinate $Y$, and $Y$ is upstream of $X$.  


4. Two consecutive read parts (i.e. with no aligned part between them in the read) that align colinearly with a big gap in the reference (Fig. 6).

By default, dnarrange ignores “big gaps” < 10 kb, and non-colinearities where Y is less than 1 kb upstream of X. This is because small deletions (colinear gaps) are frequent and arguably not “rearrangements”, and small tandem duplications (non-colinear) are overwhelmingly numerous. You can set the g (gap) and r (reverse jump) thresholds in bp:

dnarrange -v -g1000 -r100 out.maf.gz : controls/hg38-* > groups.maf

We have said that dnarrange discards DNA reads that share rearrangements with control reads, but the truth is a bit more complex. dnarrange prioritizes rearrangement types in this order: inter-chromosome > inter-strand > non-colinear > gap. It discards DNA reads that share rearrangements of the highest-priority type present in the read with control reads.

2.11 Other features of dnarrange

We have covered the most important features of dnarrange, but it can do some further things. dnarrange-merge merges each group of DNA reads into a consensus sequence, using lamassemble. By re-aligning the consensus sequence to the genome, we can perhaps see the rearrangement more clearly and accurately. More simply, dnarrange-merge can just get the rearranged reads without merging them:

dnarrange-merge HG002723_1.fastq.gz groups.maf > some-reads.fastq

This may be useful, because we can re-align just these reads to the genome more slowly and sensitively, for example like this:

lastdb -P16 -uNEAR nearDB hg38.analysisSet.fa

lastal -P16 -m50 -p hum.train nearDB some-reads.fastq | last-split > out2.maf

Here we have used NEAR instead of RY4 for greater sensitivity. We have re-used the last-train result: no need to re-run it. Finally, we added option -m50, which makes it even more slow and sensitive. Higher m values make it increasingly slow and sensitive: the default is 10 (but the -uRY options set the default to 2).

A further issue is that some large and complex rearrangements, e.g. in genetic diseases, are larger even than “long” reads. So each group of reads covers only part of the rearrangement. In order to understand the whole rearrangement, we need to correctly order and orient the groups (i.e. the rearrangement parts). dnarrange includes a method that seeks a most-parsimonious order and orientation, which succeeded in fully characterizing some large and complex rearrangements.

Interestingly, these rearrangements have holistic features, e.g. loss of sequence, that are knowable only from the whole rearrangement and not from the parts. dnarrange can be very slow and memory-consuming. This seems to be because many DNA reads align dubiously to a few genomic hotspots, and dnarrange spends much effort comparing these reads to each other. The solution is to use control reads, which rapidly discard these hotspot reads. This works better if the case and control reads were analysed in the same way, e.g. with/without masking. It also works better if the exact same reference was used. (There are unfortunately different versions of e.g. hg38). Another way to mitigate this problem is to use last-postmask.
It is useful if the DNA reads are long enough to cover a whole rearrangement, but on the other hand long reads have a disadvantage. The problem is that if a read overlaps two rearrangements, and one of them is shared by controls, the whole read may get discarded. This is a limitation of dnarrange, because it is hard to know whether a read covers two small rearrangements or two parts of one large rearrangement. Therefore, a combination of longer and not-so-long reads may be best.

Our previous publications show further interesting rearrangements found by these methods, such as tandem heptuplication, 3’-transduction from a LINE, or chromosomeshattering (4, 5).

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