KPop: Unleash the full power of your k-mers!

KPop is an assembly-free and scalable method for the comparative analysis of microbial genomes and environmental samples. It is based on full k-mer spectra and dataset-specific transformations; it allows to accurately compare hundreds of thousands of assembled, or thousands of unassembled microbial genomes or sequenced samples, in a matter of hours. It provides excellent resolution across a very large number of use cases and applications. More details can be found in our bioRxiv preprint.

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0. Installation

There are several possible ways of installing the software on your machine: through conda; by downloading pre-compiled binaries (Linux and MacOS x86_64 only); or manually.



🚺 Note that the only operating systems we officially support are Linux and MacOS. 🛕



Both OCaml and R are highly portable and you might be able to manually compile/install everything successfully on other platforms (for instance, Windows). Please let us know if you succeed or if you encounter some unexpected behaviour. However, please note that in general we are unable to provide installation-related support or troubleshooting on specific hardware/software combinations.

0.1. Conda channel



Coming soon!



0.2. Pre-compiled binaries

You can download pre-compiled binaries for Linux and MacOS x86 64 from our releases. After doing so, just copy or move them to a directory which is accessible from your PATH.

For instance, supposing that you've downloaded programs to directory ~/.local/bin/, in order to make them accessible from everywhere you'll have to execute a command such as

export PATH=~/.local/bin:\$PATH

or add it to one of your login scripts (such as ~/.bashrc or similar for bash).

0.3. Manual install

Alternatively, you can install KPop manually by cloning and compiling its sources. You'll need an up-to-date distribution of the OCaml compiler and the Dune package manager for that. Both can be installed through OPAM, the official OCaml distribution system. Once you have a working OPAM distribution you'll also have a working OCaml compiler, and Dune can be installed with the command

```
opam install dune
```

if it is not already present. Make sure that you install OCaml version 4.12 or later.

Cloning should be done with the option --recursive, as in

```
git clone --recursive git@github.com:PaoloRibeca/KPop.git
```

so as to make sure that all the subrepositories get correctly pulled.

Then go to the directory into which you have downloaded the latest KPOP sources, and type

```
./BUILD
```

That should generate all the executables you'll need (as of this writing, KPopCount, KPopCounterDB, KPopTwist_, KPopTwist_, KPopTwistDB) in the directory ./build . Copy them to some favourite location in your PATH, for instance ~/.local/bin.

For the time being, due to the presence of some legacy code, you'll also need to install some R packages (and possibly R itself) by using your favourite R package manager. Those packages are:

```
data.table ca
```

1. Quick start

Download file clusters-small.fasta from the directory test. Then run the following commands (throughout this documentation we'll assume that you're using a fairly up-to-date version of bash):

```
export K=5
date

for CLASS in C1 C2 C3 C4 C5 C6 C7 C8 C9 C10; do cat clusters-small.fasta | awk -v CLASS=$CLASS '{nr=(NR-1)%4; ok=
(nr==0?$0~("-"CLASS"$"):nr==1&&ok); if (ok) print}' | KPopCount -k $K -L -f /dev/stdin | KPopCountDB -k /dev/stdin -R "~." -A $CLASS -L $CLASS -N -D --table-transform none -t /dev/stdout; done | KPopCountDB -k /dev/stdin -o
Classes.$K -v
KPopTwist -i Classes.$K -v
cat clusters-small.fasta | awk -v K="$K" '{nr=(NR-1)%4; if (nr==2) split($0,s,"[>-]"); if (nr==3) print ">"s[2]"-
"s[3]"\n"$0}' | KPopCount -k $K -L -f /dev/stdin | KPopTwistDB -i T Classes.$K -k /dev/stdin -o t /dev/stdout |
KPopTwistDB -i T Classes.$K -i t Classes.$K -s /dev/stdin Test_prediction.$K -v
echo -n ">>> Misclassified sequences: "; cat Test_prediction.$K.KPopSummary.txt | awk -F '\t' 'BEGIN{0FS="\t"}
{$1=gensub("-","\"\t\"",1,$1); print}' | awk -F '\t' '{if ($2!=$7) print}' | wc -1
```

That should produce an output such as this one:

```
Wed 14 Feb 16:46:55 GMT 2024
This is KPopCountDB version 38 [07-Feb-2024]
 compiled against: BiOCamLib version 242 [23-Jan-2024];
                   KPop version 368 [07-Feb-2024]
(c) 2020-2024 Paolo Ribeca <paolo.ribeca@gmail.com>
(Dune_exe_KPopCountDB.KMerDB.add_files.(fun)): [1/1] File '/dev/stdin': Read 10 spectra on 5127 lines.
(Dune_exe_KPopCountDB.KMerDB.to_binary): Outputting DB to file 'Classes.5.KPopCounter'... done.
This is KPopTwist version 17 [02-Jan-2024]
 compiled against: BiOCamLib version 242 [23-Jan-2024];
                  KPop version 368 [07-Feb-2024]
(c) 2022-2024 Paolo Ribeca <paolo.ribeca@gmail.com>
Wed 14 Feb 2024 16:47:19 GMT: [1/13] Exporting table...
Wed 14 Feb 2024 16:47:19 GMT: [2/13] Splitting table...
Wed 14 Feb 2024 16:47:19 GMT: [3/13] Reading table...
Wed 14 Feb 2024 16:47:19 GMT: [4/13] Resampling table...
Wed 14 Feb 2024 16:47:19 GMT: [5/13] Transforming table...
Wed 14 Feb 2024 16:47:19 GMT: [6/13] Writing twisted...
Wed 14 Feb 2024 16:47:19 GMT: [7/13] Writing inertia...
Wed 14 Feb 2024 16:47:19 GMT: [8/13] Normalizing table...
Wed 14 Feb 2024 16:47:19 GMT: [9/13] Transposing table...
Wed 14 Feb 2024 16:47:19 GMT: [10/13] Writing twister...
Wed 14 Feb 2024 16:47:19 GMT: [11/13] Encoding twisted...
Wed 14 Feb 2024 16:47:19 GMT: [12/13] Encoding twister...
Wed 14 Feb 2024 16:47:19 GMT: [13/13] Cleaning up...
Wed 14 Feb 2024 16:47:19 GMT: All done.
This is KPopTwistDB version 27 [17-Jan-2024]
compiled against: BiOCamLib version 242 [23-Jan-2024];
                   KPop version 368 [07-Feb-2024]
(c) 2022-2024 Paolo Ribeca <paolo.ribeca@gmail.com>
(Dune_exe__KPopTwistDB.Twister.of_binary): Reading DB from file 'Classes.5.KPopTwister'... done.
(KPop__Matrix.of_binary): Reading DB from file 'Classes.5.KPopTwisted'... done.
(KPop__Matrix.of_binary): Reading DB from file '/dev/stdin'... done.
(KPop__Matrix.Base.get_normalizations): Done 500/500 rows.
(KPop__Matrix.Base.get_normalizations): Done 10/10 rows.
(KPop__Matrix.summarize_rowwise): Writing distance digest to file 'Test_prediction.5.KPopSummary.txt': done 500/500
>>> Misclassified sequences: 0
Wed 14 Feb 16:47:41 GMT 2024
```

This is an example of κ_{POP} -based classifier. The input FASTA file clusters-small.fasta contains 1000 sequences having names such as S2-C1, meaning "sequence 2 belonging to class 1". There are 10 different classes. We will see how the process works in more detail in the following sections, but, to summarise:

- 1. Sequences with an odd index are taken to be part of the training set, sequences with an even index are considered part of the test set
- 2. For each class c1, c2, ... c10, if the variable CLASS contains the name of the class, the command

```
cat clusters-small.fasta | awk -v CLASS=$CLASS '{nr=(NR-1)%4; ok=(nr==0?$0~("-"CLASS"$"):nr==1&&ok); if (ok) print}' | KPopCount -k $K -L -f /dev/stdin | KPopCountDB -k /dev/stdin -R "~." -A $CLASS -L $CLASS -N -D --table-transform none -t /dev/stdout
```

runs KPopCount (with k=5) on each training sequence belonging to CLASS; the results, which are text files each one containing a list of k-mers with their respective frequencies, are concatenated and sent to KPopCountDB through a pipe — if you wanted to look into the format, you could do so with the command

```
cat clusters-small.fasta | awk -v CLASS=CLASS '\{nr=(NR-1)\%4; ok=(nr==0?\$0~("-"CLASS"\$"):nr==1\&ok); if (ok) print}' | KPopCount -k $K -L -f /dev/stdin | less
```

After that, KPopCountDB collects all spectra for each class into a temporary database, replaces them with a combination of the input spectra, names the combined spectrum \$class, and re-outputs it in the same format used before for the spectra produced by KPopCount

3. The command

```
KPopCountDB -k /dev/stdin -o Classes.$K -v
```

receives the 10 spectra, one per class, and outputs them to a binary database called Classes.5 (actually that corresponds to a file, which gets automatically named Classes.5. KPopCounter)

4. The command

```
KPopTwist -i Classes.$K -v
```

"twists" spectra to a reduced-dimensionality space, storing the results in binary form (actually that corresponds to two files, which are automatically named Classes.5.KPopTwister and Classes.5.KPopTwisted). Both the twisted spectra and the "twister" — i.e., the operator that can be used to convert the original spectra to twisted space — are stored and can be reused later on

5. The command

```
cat clusters-small.fasta | awk -v K="$K" '{nr=(NR-1)%4; if (nr==2) split($0,s,"[>-]"); if (nr==3) print ">"s[2]"-
"s[3]"\n"$0}' | KPopCount -k $K -L -f /dev/stdin | KPopTwistDB -i T Classes.$K -k /dev/stdin -o t /dev/stdout |
KPopTwistDB -i T Classes.$K -i t Classes.$K -s /dev/stdin Test_prediction.$K -v
```

selects test sequences, runs each of them separately through KPopCount to produce a spectrum, and concatenates and pipes all the spectra thus generated to KPopTwistDB, which twists them according to the twister generated at the previous stage (named Classes.5). The results are output to a summary text file, which gets automatically named Test_prediction.5.KPopSummary.txt. The file contains information about the two closest classes for each sequence

6. Finally, the command

```
echo -n ">>> Misclassified sequences: "; cat Test_prediction.$K.KPopSummary.txt | awk -F '\t' 'BEGIN{0FS="\t"} {$1=gensub("-","\"\t\"",1,$1); print}' | awk -F '\t' '{if ($2!=$7) print}' | wc -l
```

parses the results in Test_prediction.5.KPopSummary.txt and computes the number of misclassified sequences.

Congratulations! You have now moved your first steps in the fascinating world of KPop.

1.2. Visualisation of Quick Start results

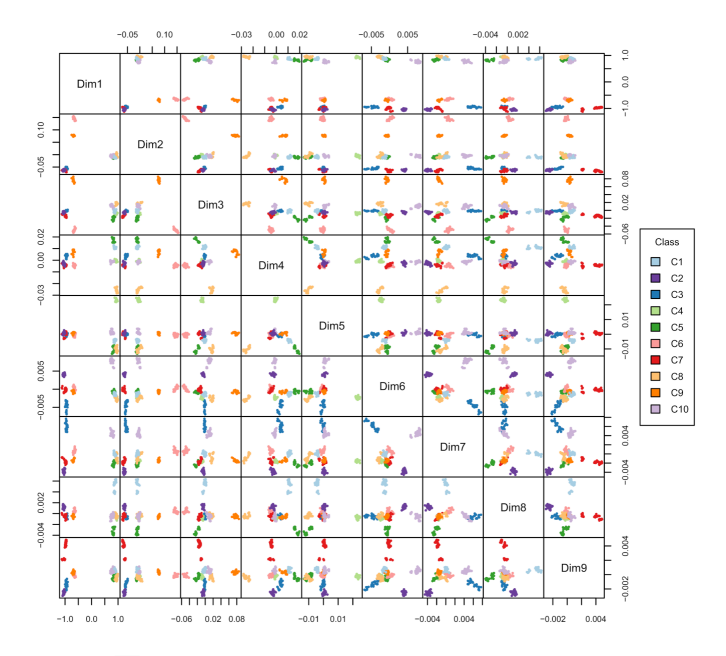
Now we can visualise the embeddings (a.k.a. twisted spectra, or multidimensional vectors in KPOP space) generated by KPOP for each test sequence as a scatterplot matrix for all possible 2-dimensional projections of the KPOP space. This normally wouldn't be recommended as there will likely be a large number of dimensions, but in this example we only have 9.

To do this, we firstly create a new_data.5.KPopTwisted.txt using KPopTwistDB with only the test sequences:

```
cat clusters-small.fasta | awk -v K="$K" '{nr=(NR-1)%4; if (nr==2) split($0,s,"[>-]"); if (nr==3) print ">"s[2]"-
"s[3]"\n"$0}' | KPopCount -k $K -L -f /dev/stdin | KPopTwistDB -i T Classes.$K -k /dev/stdin -o t /dev/stdout -v |
KPopTwistDB -i t /dev/stdin -0 t Test.$K
```

We then visualise the results as a scatterplot matrix using R:

▼ Script to visualise Quick Start results



One can see how KPop is able to embed sequences (i.e., to map them to an abstract multi-dimensional space) in such a way that they cluster according to their original colouring — which represents, in this case, the ground truth. As the clusters are well-defined and have neat boundaries along a suitable number of dimensions, as shown by the projections, that results in a good (actually perfect) classifier for this example if the cluster/class centroids are considered as reference points and the closest centroid is used to predict the class each sequence belongs to. However, as KPOP is able to explicitly turn sequences into vectors, any kind of more complex classifier based on machine learning or AI techniques (for instance *k*-NN, decision trees, random forests or others) can also be used. We provide some examples later on.

As mentioned previously, this plot is a great way to visually check that test sequences cluster in the correct class — however, it wouldn't be as useful for a dataset with 1000 dimensions!

2. Frequently asked questions

There are no binaries available for my OS, what do I do?

KPop is almost entirely implemented in OCaml, an industry-strength functional statically-typed programming language that offers remarkable concision, symbolic power, robustness, compiled speed, and portability. Due mostly to historical reasons, a small part of KPop is still written in R, although we hope to eventually migrate everything to OCaml.

Both OCaml and R are highly portable and you might be able to manually compile/install everything successfully on other platforms. See the Installation section below.

How does KPop compare with minimiser-based methods?

KPop is much more sensitive than minimiser-based methods, allowing out-of-the-box accurate comparison of assembled or unassembled long sequences irrespective of whether they differ by single nucleotides or by large portions. This is usually not true for minimiser-based methods, irrespective of the hashing scheme they use, as they only produce a much coarser comparison. A number of benchmarks and a detailed explanation of the differences between the two approaches can be found in our bioRxiv preprint.

Crucially, while minimiser-based methods are geared towards providing distances between sequences, KPOP explicitly produces *embeddings*, i.e., it is able to turn sequences into points of a low-dimensionality latent space. Having an explicit latent space is important for many reasons — it helps explainability; it also makes it possible to perform direct clustering and use vector DBs or libraries to store and search embedded sequences.

Can I run it on my laptop?

Depending on the problem at hand, KPop analysis can require a relatively large amount of resources, memory in particular. That is a conscious design choice — one could not leverage the power of full *k*-mer spectra otherwise. If you need to explore datasets with millions of sequences, we recommend that you do so on a robustly sized HPC node. However, not everything is expensive — for instance, and similar to what happens with other big-data frameworks, while the "training" phase needed to create KPop -based classifiers is typically resource-intensive, the classifiers themselves can typically be run on lower-end machines.

All KPop programs are parallelised and will automatically use as many CPUs as are available on your machine, in order to reduce as much as possible the wallclock time taken by your tasks.

How do I choose k?

As explained in our bioRxiv preprint, the choice of k can be tricky, and you should always make sure you try different results and select the lowest value of k for which results become stable. One empirical criterion that might help you in most cases is to choose \$\$ k := { \mathrm{argmax}k \displaystyle \mathrm{min}s \frac{\mathrm{max}h C^k{h s}}{\text{ft}(\prod{h} C^k{h s} \right)^{1/{n_h}}} }\$\$ \$\$ where \$C^k_{h s}\$ is a non-zero count for k-mer \$h\$ in sample \$\$\$ and there are \$n_h\$ non-zero k-mers. I.e., you would select the \$k\$ maximising the minimum across samples of the ratio between the maximum non-zero count and the (harmonic) mean of the non-zero counts for each sample.

3. Overview of commands

KPop is implemented as a suite of different programs, which can be combined into complex workflows in a modular fashion. They are:

- KPopCount . It implements extraction of *k*-mer spectra from files containing sequences or sequencing reads (in FASTA format; and both single- and paired-end FASTQ format)
- **KPopCountDB**. It implements collection of *k*-mer spectra into binary databases, allowing the user to export the resulting objects as either binary files or text tables. Spectra can be transformed before export
- KPOPTWiST. It implements the unsupervised generation of coordinate transformations (or "twisters") from databases of *k*-mer spectra. Each transformation is optimised for the database at hand, and turns ("twists") *k*-mer spectra into numerical vectors of a typically very much reduced dimensionality. The transformation can be stored as a binary object for future use
- **KPOPTWISTOB** . It implements a number of operations on twisted spectra. It can: use an existing twister to twist *k*-mer spectra; generate databases of twisted spectra and output/input them as binary files or text tables; compute and summarise distances between twisted spectra; and more.

3.1. General design

While KPopCount and KPopTwist implement one-shot operations (computing spectra from sequences and generating a twister, respectively), KPopCountDB and KPopTwistDB need to be versatile and perform a number of complex tasks, ranging from householding ones (management of spectra and twisted spectra, respectively) to complex operations that are an essential component of larger workflows.

In response to such needs, KPopCountDB and KPopTwistDB are able to execute a series of actions. Actions are specified each one as a different command line option, and performed in order of specification — i.e., saying something like KPopCountDB -D -N is not the same as saying KPopCountDB -N -D.

In addition, both KPopCountDB and KPopTwistDB have *registers*, i.e., slots to/from which the contents of external files and the results of computations can be loaded/saved. You can see registers as program variables, each one having a specific type. For instance, the KPop programs know that only twisters can be loaded to/saved from twister registers, and will enforce that by

requiring and checking that your external file has the correct extension (.KPopTwister) and format when doing I/O to/from a twister register.

KPopCountDB has two registers:

- a *database* register, holding a set of *k*-mer spectra (i.e., a set of *k*-mers and their associated frequencies). There is one spectrum per sample, and additional metadata in the form of strings or numbers can be associated to the samples (see our bioRxiv preprint for more information). Due to space optimisation reasons, modifications of the database happen in-place
- a selection register, holding a set of sample labels. Labels can be specified on the command line or read from the database present in the database register. Collective operations (selection, deletion, combination, and so on) can be performed on the samples present in the database register according to the set of sample names specified by the selection register, which acts as a guide to enable complex operations on the database. For instance, one might extract from the database all the sample names matching some regular expression and put them in the selection register, add some more sample names to the selection register, and finally delete from the database all the samples whose names do not appear in the selection register.

KPopTwistDB has four registers:

- a *twister* register holding a twister, i.e., an object able to transform *k*-mer spectra to "twisted" embeddings/vectors in an abstract KPop space (note that the transformation is dataset-dependent and generated by the user when runnning KPopTwist on a set of spectra, as explained in our bioRxiv preprint and the next section)
- a twisted register holding "twisted" vectors, i.e., KPop embeddings of biological sequences into an abstract KPop space
- a *metrics* register holding a metric, i.e., a vector of numbers specifying the relevance of each one of the dimensions of the KPOP space (see our bioRxiv preprint for more details)
- a *distance* register holding a distance matrix between couples of samples. The matrix can be computed between the vectors present in the twisted register and themselves, or between the vectors present in the twisted register and those present in an external file containing twisted vectors.

Options are provided to write content from/to these registers, both in binary and textual form — the latter can be used, for instance, to import/export data from/to R . In addition to input/output to/from registers, both KPopCountDB and KPopTwistDB are able to read and process external files containing k-mer spectra in the format produced by KPopCount — this ensures that new spectra/twisted spectra can be added to their respective databases. A figure showing some of the available data transformations and formats can be found in the next section.

In general, registers provide "memory" (or a "state" in computer science parlance) to KPopCountDB and KPopTwistDB — by using them and the rich set of command line options we implemented, one can easily combine KPop programs and build complex workflows performing sophisticated tasks (see the section on examples below).

3.2. Building workflows

The following figure summarises the main typical data transformations that are possible and the file formats used to encode inputs and outputs in each case:

KPopTwist KPopCount Twister (binary, .KPopTwister) k-mer spectra DB equences (FASTA) k-mer spectra (binary, KPopTwist Reads (FASTQ) (tab-separated) .KPopCounter) Twisted spectra .KPopTwisted) **KPopCountDB KPopTwistDB** k-mer spectra (tab-separated) k-mer spectra (tab-separated) k-mer spectra DB Twisted spectra **KPopCountDB** (binary, (binary, KPopCounter) KPopTwisted) Twister Sample metadata (binary, (tab-separated) .KPopTwister) Twisted spectra k-mer spectra (tab-separated) .KPopTwisted) k-mer spectra DB Distance matrix KPopCountDB KPopTwistDB (binary. (binary, .KPopDMatrix) .KPopCounter) (binary, (binary, .KPopCounter) .KPopTwisted) Distance matrix Distance summary k-mer spectra DB k-mer spectra DB (binary, KPopTwistDB KPopCountDB (tab-separated, (tab-separated) .KPopDMatrix) .KPopSummary.txt) .KPopCounter)

There are many more possible ways of using KPopCountDB and KPopTwistDB. For instance, with KPopCountDB one can compute distances between untwisted *k*-mer spectra; with KPopTwistDB one can accumulate twisted *k*-mer spectra or distances into existing databases, and convert binary files from/to plain-text tab-separated tables. In the latter case, as explained, automatic naming rules are applied depending on the type of the register being used — for example, a command such as

```
KPopTwistDB ... -o t Classes
```

would result in the production of a binary file named Classes. KPOpTwisted containing twisted k-mer spectra, while the command

```
KPopTwistDB ... -0 t Classes
```

will produce the same results but output them to a tab-separated tabular file named <code>classes.KPopTwisted.txt</code> . In general, in the style of BLAST indices, you do not have to worry about formats or extensions as long as you consistently use the same prefix (in this case, <code>Classes</code>) to indicate logically related files.

Note that as a rule, sequence/sample names, as well as metadata field names and field values, cannot contain double quote '"' or tabulation '\t' characters.

4. Command line syntax

4.1. KPopCount

This is the list of command line options available for the program (KPopCount). You can visualise the list by typing

KPopCount -h

in your terminal. You will see a header containing information about the version:

```
This is KPopCount version 14 [18-Mar-2024]
compiled against: BiOCamLib version 248 [18-Mar-2024];
KPop version 417 [05-Mar-2024]
(c) 2017-2024 Paolo Ribeca <paolo.ribeca@gmail.com>
```

```
KPopCount -1 <output_vector_label>|-L [OPTIONS]
```

Algorithmic parameters

Option	Argument(s)	Effect	Note(s)
-k -Kk-mer- sizek-mer- length	k_mer_length	<i>k</i> -mer length (must be positive, and <= 30 for DNA or <= 12 for protein)	default= <mark>12</mark>
-M max- results- size	positive_integer	maximum number of <i>k</i> -mer hashes to be kept in memory at any given time. If more are present, the ones corresponding to the lowest cardinality will be removed from memory and printed out, and there will be repeated hashes in the output	<u>default=<mark>16777216</mark></u>

Input/Output

Option	Argument(s)	Effect	Note(s)
-C content	DNA-ss DNA-single- stranded DNA-ds DNA- double-stranded protein	how file contents should be interpreted. When content is DNA-ss or protein, the sequence is hashed; when content is DNA-ds, both sequence and reverse complement are hashed. DNA-ss prevents automatic matching of reverse-complemented sequences; use it only when comparing a set of single, homogeneus sequences	<u>default=</u> DNA- <u>ds</u>
-f fasta	fasta_file_name	FASTA input file containing sequences. You can specify more than one FASTA input, but not FASTA and FASTQ inputs at the same time. Contents are expected to be homogeneous across inputs	
-s single-end	fastq_file_name	FASTQ input file containing single-end sequencing reads You can specify more than one FASTQ input, but not FASTQ and FASTA inputs at the same time. Contents are expected to be homogeneous across inputs	
-p paired-end	fastq_file_name1 fastq_file_name2	FASTQ input files containing paired-end sequencing reads You can specify more than one FASTQ input, but not FASTQ and FASTA inputs at the same time. Contents are expected to be homogeneous across inputs	
-1 label	output_vector_label	label to be given to the <i>k</i> -mer spectrum in the output file. It must not contain double quote '"' characters. Either option -1 or option -L is mandatory	
-L one- spectrum-per- sequence		output one spectrum per input sequence, using the sequence name as label. Sequence names must not contain double quote '"' characters. Either option -1 or option -L is mandatory	
-o output	output_file_name	name of generated output file	<u>default=<mark>stdout</mark></u>

Miscellaneous

Option	Argument(s)	Effect	Note(s)
-v verbose		set verbose execution	default= <mark>false</mark>
-V version		print version and exit	
-h help		print syntax and exit	

4.2. KPopCountDB

This is the list of command line options available for the program KPopCountDB . You can visualise the list by typing

```
KPopCountDB -h
```

in your terminal. You will see a header containing information about the version:

KPopCountDB [ACTIONS]

 $\mbox{\bf Actions.}$ They are executed delayed and in order of specification.

Actions on the database register:

Option	Argument(s)	Effect	Note(s)
-e empty		put an empty database into the register	
-i input	binary_file_prefix	load into the register the database present in the specified file (which must have extension .KPopCounter)	
-m metadata add- metadata	metadata_table_file_name	add to the database present in the register metadata from the specified file. Metadata field names and values must not contain double quote characters	
-kkmersadd-kmersadd-kmer- files	k-mer_table_file_name[, , k- mer_table_file_name]	add to the database present in the register <i>k</i> -mers from the specified files	
summary		print a summary of the database present in the register	
-o output	binary_file_prefix	dump the database present in the register to the specified file (which will be given extension .KPopCounter)	
distance distance- function	<pre>euclidean minkowski(non_negative_float)</pre>	set the function to be used when computing distances. The parameter for minkowski() is the power	default= <mark>eucl</mark>
distance- normalize normalize- distances distance- normalization	true false	whether spectra should be normalized prior to computing distances	
-ddistancescompute- distancescompute- spectral- distances	REGEXP_SELECTOR REGEXP_SELECTOR binary_file_prefix	where REGEXP_SELECTOR:= metadata_field ~ regexp[,, metadata_field ~ regexp] and regexps are defined according to https://ocaml.org/api/Str.html: select two sets of spectra from the register and compute and output distances between all possible pairs (metadata fields must match the regexps specified in the selector; an empty metadata field makes the regexp match labels. The result will have extension .KPopDMatrix)	
table- output-row- names	true false	whether to output row names for the database present in the register when writing it as a tab- separated file	default= <mark>true</mark>
table- output-col- names	true false	whether to output column names for the database present in the register when writing it as a tabseparated file	default= <mark>true</mark>
table- output- metadata	true false	whether to output metadata for the database present in the register when writing it as a tab-separated file	default= <mark>fals</mark>
table- transpose	true false	whether to transpose the database present in the register before writing it as a tab-separated file (if true: rows are spectrum names, columns [metadata	default= <mark>fals</mark>

Option	Argument(s)	Effect	Note(s)
		and] <i>k</i> -mer names; if <code>false</code> : rows are [metadata and] <i>k</i> -mer names, columns spectrum names)	
table- threshold	non_negative_integer	set to zero all counts that are less than this threshold before transforming and outputting them	default= <mark>1</mark>
table-power	non_negative_float	raise counts to this power before transforming and outputting them. A power of 0 when the pseudocounts method is used performs a logarithmic transformation	default= <mark>1.</mark>
table- transform table- transformation	none normalize pseudocounts clr	transformation to apply to table elements before outputting them	default= <mark>normali.</mark>
table- output-zero- rows	true false	whether to output rows whose elements are all zero when writing the database as a tab-separated file	default= <u>false</u>
table- precision	positive_integer	set the number of precision digits to be used when outputting counts	default= <mark>15</mark>
-t table	file_prefix	write the database present in the register as a tab- separated file (rows are <i>k</i> -mer names, columns are spectrum names; the file will be given extension .KPopCounter.txt)	

Actions involving the selection register:

Option	Argument(s)	Effect	Note(s)
-L labels selection- from-labels	spectrum_label[, , spectrum_label]	put into the selection register the specified labels	
-R regexps selection- from-regexps	metadata_field ~ regexp[, , metadata_field ~ regexp]	put into the selection register the labels of the spectra whose metadata fields match the specified regexps and where regexps are defined according to https://ocaml.org/api/Str.html. An empty metadata field makes the regexp match labels	
selection- combination- criterion combination- criterion	mean median	set the criterion used to combine the <i>k</i> -mer frequencies of selected spectra. To avoid rounding issues, each <i>k</i> -mer frequency is also rescaled by the largest normalization across spectra (mean averages frequencies across spectra; median computes the median across spectra)	default= <mark>mean</mark>
-Aadd- combined- selectionselection- combine-and- add	spectrum_label	combine the spectra whose labels are in the selection register and add the result (or replace it if a spectrum named spectrum_label already exists) to the database present in the database register	
-D delete selection- delete		drop the spectra whose labels are in the selection register from the database present in the register	
-N selection- negate		negate the labels that are present in the selection register	
-P selection- print		print the labels that are present in the selection register	
-C selection- clear		purge the selection register	
-F selection- to-table- filter		filter out spectra whose labels are present in the selection register when writing the database as a tab-separated file	

Miscellaneous options. They are set immediately

Option	Argument(s)	Effect	Note(s)
-T threads	computing_threads	number of concurrent computing threads to be spawned (default automatically detected from your configuration)	default= <mark>nproc</mark>
-v verbose		set verbose execution	default= <mark>false</mark>
-V version		print version and exit	
-h help		print syntax and exit	

4.3. KPopTwist

This is the list of command line options available for the program KPOPTWist . You can visualise the list by typing

```
KPopTwist -h
```

in your terminal. You will see a header containing information about the version:

```
This is KPopTwist version 20 [29-Feb-2024]
compiled against: BiOCamLib version 245 [14-Feb-2024];
KPop version 411 [01-Mar-2024]
(c) 2022-2024 Paolo Ribeca <paolo.ribeca@gmail.com>
```

followed by detailed information. The general form the command can be used is:

```
KPopTwist -i|--input <input_table_prefix> [OPTIONS]
```

Algorithmic parameters

Option	Argument(s)	Effect	Note(s)
-f -F -s -S fraction sampling sampling-fraction	fractional_float	fraction of the <i>k</i> -mers to be considered and resampled before twisting	default= <mark>1.</mark>
threshold- counts	non_negative_integer	set to zero all counts that are less than this threshold before transforming them. A fractional threshold between 0. and 1. is taken as a relative one with respect to the sum of all counts in the spectrum	default= <mark>1</mark>
power	non_negative_float	raise counts to this power before transforming them. A power of 0 when the pseudocounts method is used performs a logarithmic transformation	default= <mark>1.</mark>
transform transformation	binary power pseudocounts clr	transformation to apply to table elements	default= <mark>normalize</mark>
-n normalize normalize- counts	true false	whether to normalize spectra after transformation and before twisting	default= <mark>true</mark>
threshold- kmers	non_negative_integer	compute the sum of all transformed (and possibly normalized) counts for each <i>k</i> -mer, and eliminate <i>k</i> -mers such that the corresponding sum is less than the largest sum rescaled by this threshold. This filters out <i>k</i> -mers having low frequencies across all spectra	default= <mark>0.</mark>

Input/Output

Option	Argument(s)	Effect	Note(s)
-i input	binary_file_prefix	load the specified <i>k</i> -mer database in the register and twist it. File extension is automatically determined (will be .KPopCounter). The prefix is then re-used for output (and the output files will be given extensions .KPopTwister and .KPopTwisted)	(mandatory)
-o output	binary_file_prefix	use this prefix when dumping generated twister and twisted sequences. File extensions are automatically determined (will be .KPopTwister and .KPopTwisted	(mandatory)

Miscellaneous

Option	Argument(s)	Effect	Note(s)
-T threads	computing_threads	number of concurrent computing threads to be spawned (default automatically detected from your configuration)	default= <mark>nproc</mark>
keep- temporaries		keep temporary files rather than deleting them in the end	default= <mark>false</mark>
-v verbose		set verbose execution	default= <mark>false</mark>
-V version		print version and exit	
-h help		print syntax and exit	

4.4. KPopTwistDB

This is the list of command line options available for the program KPopTwistDB . You can visualise the list by typing

```
KPopTwistDB -h
```

in your terminal. You will see a header containing information about the version:

followed by detailed information. The general form the command can be used is:

```
KPopTwistDB [ACTIONS]
```

Actions. They are executed delayed and in order of specification.

Option	Argument(s)	Effect	Note(s)
-e empty	T t d	load an empty database into the specified register (T =twister; t =twisted; d =distance)	
-i input	T t d <i>binary_file_prefix</i>	load the specified binary database into the specified register (T =twister; t =twisted; d =distance). File extension is automatically determined depending on database type (will be: .KPopTwister; .KPopTwisted; or .KPopDMatrix, respectively)	
-I Input	T t d table_file_prefix	load the specified tabular database(s) into the specified register (T =twister; t =twisted; d =distance). File extension is automatically determined depending on database type (will be: .KPopTwister.txt and .KPopInertia.txt; .KPopTwisted.txt; or .KPopDMatrix.txt , respectively)	
-a add	t d binary_file_prefix	add the contents of the specified binary database to the specified register (t = twisted; d = distance). File extension is automatically determined depending on database type (will be: .KPopTwisted; or .KPopDMatrix, respectively)	
-A Add	t d table_file_prefix	add the contents of the specified tabular database to the specified register (t = twisted; d = distance). File extension is automatically determined depending on database type (will be: .KPopTwisted.txt; or .KPopDMatrix.txt, respectively)	
-kkmersadd-kmersadd-kmer- files	k-mer_table_file_name[, , k- mer_table_file_name]	twist k -mers from the specified files through the transformation present in the twister register, and add the results to the database present in the twisted register	
distancedistance- functionset- distanceset- distance- function	euclidean cosine minkowski(non_negative_float)	set the function to be used when computing distances. The parameter for minkowski is the power. Note that euclidean is the same as minkowski(2), and cosine is the same as (euclidean ^2)/2	<u>default=<mark>euclidea</mark>.</u>
distance- normalization set- distance- normalization	true false	set whether twisted vectors should be normalized prior to computing distances	default= <mark>true</mark>
-mmetricmetric- functionset-metricset-metric- function	flat powers(POWERS_PARAMETERS)	where POWERS_PARAMETERS := non_negative_float , fractional_float , non_negative_float : set the metric function to be used when computing distances. Parameters are: internal power; fractional accumulative threshold; external power.	default= <mark>powers(1</mark>
-d distances compute- distances	twisted_binary_file_prefix	compute distances between all the vectors present in the twisted register and all the vectors present in the specified twisted binary file (which must have extension .KPopTwisted) using the metric provided by	

Option	Argument(s)	Effect	Note(s)
compute- twisted- distances		the twister present in the twister register. The result will be placed in the distance register	
-o output	Т t d binary_file_prefix	dump the database present in the specified register (T =twister; t =twisted; d =distance) to the specified binary file. File extension is automatically assigned depending on database type (will be: .KPopTwister; .KPopTwisted; or .KPopDMatrix, respectively)	
precisionset- precisionset-table- precision	positive_integer	set the number of precision digits to be used when outputting numbers	default= <mark>15</mark>
-O Output	Т t d m table_file_prefix	dump the database present in the specified register (T = twister; t = twisted; d = distance; m = metric) to the specified tabular file(s). File extension is automatically assigned depending on database type (will be: .KPopTwister.txt and .KPopInertia.txt; .KPopTwisted.txt; .KPopDMatrix.txt; or .KPopMetrics.txt, respectively)	
-Kkeep-at- mostset-keep- at-mostsummary- keep-at-most	positive_integer all	set the maximum number of closest target sequences to be kept when summarizing distances. Note that more might be printed anyway in case of ties	default= <mark>2</mark>
-scompute- and- summarize- distancescompute- and- summarize- twisted- distances	twisted_binary_file_prefix summary_file_prefix	compute distances between all the vectors present in the twisted register and all the vectors present in the specified twisted binary file (which must have extension .KPopTwisted) using the metric provided by the twister present in the twister register; summarize them and write the result to the specified tabular file. File extension is automatically assigned (will be .KPopSummary.txt)	
-Ssummarize- distancessummarize- twisted- distances	summary_file_prefix	summarize the distances present in the distance register and write the result to the specified tabular file. File extension is automatically assigned (will be .KPopSummary.txt)	

Miscellaneous options. They are set immediately.

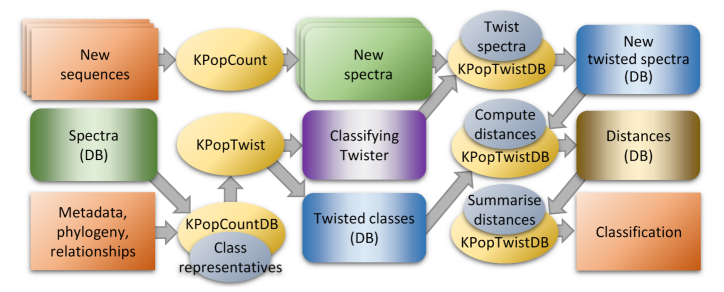
Option	Argument(s)	Effect	Note(s)
-T threads	computing_threads	number of concurrent computing threads to be spawned (default automatically detected from your configuration)	default= <mark>nproc</mark>
-v verbose		set verbose execution	default= <mark>false</mark>
-V version		print version and exit	
-h help		print syntax and exit	

5. Examples

By using the programs just described, it is possible to implement a number of interesting high-throughput workflows. We illustrate some examples here - for a more general description, please refer to our bioRxiv preprint.

5.1. Sequence classification

One can implement a sequence classifier, starting from either a set of genomic sequences or a deep-sequencing dataset containing multiple samples, by using the following strategy:



Shortly, one first generates a collection of spectra that describe the "classes" understood by the classifier; from the classes a "twister", i.e. a specialised transformation, is derived; the twister is then used to transform new sequences; and distances are computed and summarised between the twisted spectra for the classes and the twisted spectra for the sequences in order to understand what is the class closest to each sequence (possibly with some additional statistical refinement).

Note that the classifier should be generated according to the data type of the input, i.e., you should not use a classifier trained on genomic sequences to process NGS samples or vice-versa. Doing so might occasionally work if contaminations are low and the sequencing bias is reasonably flat along the sequence, but in order to get consistent results significant post-processing might be needed.

5.1.1. Classifier for simulated *M.tuberculosis* sequencing reads

As described in our bioRxiv preprint, we simulated sequencing reads for 1,000 *M.tuberculosis* genomes. The data thus generated is large (~127 GB) and hence we are not making it directly available for download. However, the scripts used to (re-)generate it can be found in the test directory of this repository.

5.1.1.1. Data preparation

In the following, we assume that the input files derived from the simulation have been organised into directories relative to your current location, and their placement reflects the set (training/test) and sequence cluster each sample belongs to.

So, we'll have two directories,

```
Train/
Test/
```

and each directory will contain subdirectories, one for each sequence cluster:

```
Train/1
Train/2
...
Train/10
```

For instance, directory Train/8 will contain input files for all the samples used as training data for sequence class 8, and command

```
ls Train/8
```

will return

```
683_1.fastq 691_1.fastq 699_1.fastq 707_1.fastq 715_1.fastq 723_1.fastq 731_1.fastq 739_1.fastq 747_1.fastq 755_1.fastq 763_1.fastq 771_1.fastq 779_1.fastq 683_2.fastq 691_2.fastq 699_2.fastq 707_2.fastq 715_2.fastq 723_2.fastq 731_2.fastq 739_2.fastq 747_2.fastq 755_2.fastq 763_2.fastq 771_2.fastq 779_2.fastq 685_1.fastq 693_1.fastq 701_1.fastq 709_1.fastq 717_1.fastq 725_1.fastq 733_1.fastq 741_1.fastq 749_1.fastq 757_1.fastq 765_1.fastq 773_1.fastq 781_1.fastq 685_2.fastq 693_2.fastq 701_2.fastq 709_2.fastq 717_2.fastq 725_2.fastq 733_2.fastq 741_2.fastq 741_2.fastq 725_2.fastq 765_2.fastq 701_2.fastq 781_2.fastq 687_1.fastq 687_1.fastq 703_1.fastq 741_1.fastq 741_1.
```

A similar structure will have been put in place for test data under the Test/ directory, with files equally split under Train/ and Test/ for cross-validation purposes (however, different choices would be possible). This conventional arrangement will be assumed in the rest of this document for all the examples involving KPop -based classifiers.

5.1.1.2. Data analysis

In order to analyse sequences in parallel fashion and decrease waiting times, we first prepare a short bash script named process_classes, as follows:

```
#!/usr/bin/env bash

while read DIR; do
   CLASS=$(echo "$DIR" | awk '{l=split(gensub("[/]+$","",1),s,"/"); print s[1]}')
   echo "Processing class '${CLASS}'..." > /dev/stderr
   ls "$DIR"/*_1.fastq |
        Parallel --lines-per-block 1 -- awk '{l=split($0,s,"/"); system("KPopCount -k 12 -l
"gensub("_1.fastq$","",1,s[1])" -p "$0" "gensub("_1.fastq$","_2.fastq",1))}' |
        KPopCountDB -k /dev/stdin -R "~." -A "$CLASS" -P -L "$CLASS" -N -P -D --summary --table-transform none -t
/dev/stdout 2> /dev/null
done
```

The program Parallel can be obtained from the BiOCamLib repository, on which the implementation of KPop depends.

The script takes as input a list of directories. For each directory, the script generates the 10-mer spectra for all the files contained in that directory, combines the spectra into an in-memory KPOPCOUNT database, generates their linear combination, discards the database, and writes the combination to standard output.

So, for instance,

```
echo Train/8 | ./process_classes
```

would process all files present in directory Train/8.

Note that the script is implicitly parallelised, in that both Parallel and KPopCountDB will automatically check for the number of available processors, and start an adequate number of computing threads to take full advantage of them. KPopCount is not by itself parallel, but multiple copies of it will be invoked through Parallel.

At this point, in order to perform the "training" phase, we need to issue the commands

```
ls -d Train/*/ | Parallel --lines-per-block 1 -- ./process_classes | KPopCountDB -k /dev/stdin -o Classes KPopTwist -i Classes
```

The first command will generate one combined, representative spectrum for each class in the training set, and subsequently, thanks to KPopCountDB, combine the spectra for all the representatives into a database having prefix Classes and full name Classes. KPopCounter.

The second command will "twist" the database, i.e., generate a dataset-specific transformation that turns *k*-mer spectra of the size chosen when generating them (10 in this case) into vectors embedded in a reduced-dimensionality space. The transformation is stored in the file Classes.KPopTwister. In addition, KPopTwist also produces a file containing the positions of the class representatives in twisted space (Classes.KPopTwisted). Such files will be reused in what follows.

Once that has been done, the command

```
ls Test/*/*_1.fastq | Parallel --lines-per-block 1 -- awk '{l=split($0,s,"/"); system("KPopCount -k 12 -l "gensub("_1.fastq$","",1,s[l])" -p "$0" "gensub("_1.fastq$","_2.fastq",1))}' | KPopTwistDB -i T Classes -k /dev/stdin -o t Test -v
```

will generate separate *k*-mer spectra for each file in the test set, and twist them according to the "classifying" transformation stored in file Classes.KPopTwister. The results will be stored in an additional file, Test.KPopTwisted.

All the files generated so far are binary — their content is laid out according to the way OCaml data structures are stored in memory.

Should you be curious to see what the content looks like in human-readable format, you can always use KPopTwistDB to convert the file (or any other file generated by KPopTwistDB) to a tab-separate textual table as those you can import into, or export from, R. For instance, the command

```
KPopTwistDB -i t Test -0 t Test
```

will load a binary file (hence option <code>-i</code>) of the "twisted" type (hence option <code>-i</code> t) with prefix <code>Test</code> (hence option <code>-i</code> t <code>Test</code>), i.e., according to the automatic naming conventions enforced by <code>KPop</code>, it will load file <code>Test.KPopTwisted</code> into the "twisted" register of <code>KPopTwistDB</code>. After that, the command will output the content of the "twisted" register in tabular form (hence option <code>-0</code> t) to a file with prefix <code>Test</code>, i.e., according to <code>KPop</code> 's automatic naming conventions, to file <code>Test.KPopTwisted.txt</code>. You can see how such naming conventions free you from thinking about files extensions, and avoid name clashes between files having the same prefix but a different content.

The file Test.KPopTwisted.txt will contain a header

```
"" "Dim1" "Dim2" "Dim3" "Dim4" "Dim5" "Dim6" "Dim7" "Dim8" "Dim9"
```

followed by rows such as

```
"121" 0.461489766036905 0.568113255163704 -0.882699288338637 -0.0272667586657692 0.0581302393092316 -0.0189776114363531 0.00161058296863769 0.0225299502172142 -0.0278201311947722
```

expressing the coordinates of twisted sequences or samples (in this case, sample 121) in twisted space.

We now have several files in our directory, namely

```
Train/
Test/
Classes.KPopCounter
Classes.KPopTwister
Classes.KPopTwisted
Test.KPopTwisted
Test.KPopTwisted.txt
```

What is left to do in order to classify the sequences in our test set is to compute the distance in twisted space between each sequence (rows of file Test.KPopTwisted) and the representative of each equivalence class of the training set (rows of file Classes.KPopTwisted); the classification for each given sequence will be the closest equivalence class (provided that the closest and second closest match are separated by some reasonable margin). Computing all pairwise distances in twisted space between sequences and classes also requires transformation values from Classes.KPopTwister, and is accomplished by the command

```
KPopTwistDB -i t Classes -i T Classes -d Test -0 d Test-vs-Classes -o d Test-vs-Classes
```

which loads the twisted file Classes.KPopTwisted, computes pairwise distances with the contents of Test.KPopTwisted — the results will be placed in the "distance" register of KPopTwistDB —, and writes results into a binary file Test-vs-Classes.KPopDMatrix and a tabular file Test-vs-Classes.KPopDMatrix.txt (we write to a text here for illustration). File Test-vs-Classes.KPopDMatrix.txt will contain a header

```
"" "10" "1" "2" "3" "4" "5" "6" "7" "8" "9"
```

i.e., the list of the training classes, followed by rows of corresponding distances, such as

```
"121" 3.298234166382 3.33901585931896 1.85388698190156 3.35586331250312 3.35677089495605 3.35069768834933 3.31043491848486 3.30792669036762 3.32894004017435 3.31694544810727
```

In this case, for instance, sequence 121 has distance in twisted space of ~1.9 from class 2, while the distance from all other classes is >3.3. That classifies the sequence as belonging to class 2 (which is correct according to the truth table of the simulation).

In fact, an automated way of summarising distances and finding the \$n\$ closest ones is implemented in the option -s of KPopTwistDB, and that is what one would probably use in real life. The command

```
KPopTwistDB -i T Classes -i t Classes -s Test Test-vs-Classes
```

will produce a file Test-vs-Classes.KPopSummary.txt made of tab-separated lines, one per sequence, such as

```
"121" 3.18187160005451 0.467075454492746 3.32894004017435 0.0217576481749768 "2" 1.85388698190156 -2.84319076367473 "10" 3.298234166382 0.249130124925674
```

The first 5 fields have a fixed meaning, being:

- 1. Sequence name
- 2. Mean of distances from all classes for the sequence being considered
- 3. Standard deviation of distances from all classes for the sequence being considered
- 4. Median of distances from all classes for the sequence being considered
- 5. MAD of distances from all classes for the sequence being considered.

Those are followed by groups of 3 fields (how many groups depends on the parameters given when the summary was created), each field being:

- 1. Class name
- 2. Distance from this class for the sequence being considered
- 3. *z*-score of the distance from this class for the sequence being considered. The groups list the classes closest to the sequence (2 by default). They are sorted by increasing distance.

For instance, in the example line shown above the summary examines sequence 121, revealing that its mean distance from classes is 3.18; the closest class is 2, at a distance of 1.85 (corresponding to a z-score of -2.84) while the second closest class is 10,

at a distance of 3.30 (corresponding to a *z*-score of 0.25). This line confirms in a statistically more sound way what we had glimpsed by eye above from the complete list of distances.

Alternatively, Test-vs-Classes.KPopSummary.txt can be generated using Test-vs-Classes.KPopDMatrix.txt with the KPopTwistDB option -i d, as follows

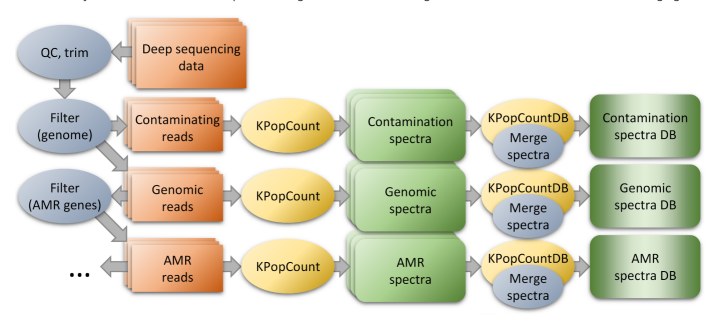
```
KPopTwistDB -i T Classes -i d Test-vs-Classes -S Test-vs-Classes
```

5.1.2. Classifier for deep-sequencing *M.tuberculosis* samples

This example requires a large sequencing dataset available from the Short Read Archive (~1300 samples) to be downloaded and processed. As before, the data is too large to be stored for direct download; however, the lists of SRA identifiers used as a starting point for this experiment can be found in the test directory of this repository.

5.1.2.1. Data preparation

As described in our bioRxiv preprint, one of the advantages offered by KPop is its versatility. In particular, by pre-processing reads one can easily have the method focus on particular regions or features of the genome of interest, as shown in the following figure:



For this example, we are mainly interested in performing a pre-processing step to discard contaminations — i.e., we want to only keep sequencing reads that are likely to originate from *Mycobacterium*. In order to do so, we process each set of sequencing reads through the following script (here we assume to be operating on sample ERR275184 , which comes as the two paired-end FASTQ files ERR275184_1.fastq.gz and ERR275184_2.fastq.gz):

```
# First, trim
trim_galore -j 8 --paired ERR275184_1.fastq.gz ERR275184_2.fastq.gz
# Second, filter
 FASTools \ e \ -p \ <(zcat \ ERR275184\_1\_val\_1.fq.gz) \ <(zcat \ ERR275184\_2\_val\_2.fq.gz) \ | \ FASTools \ -S \ | \ awk \ -F \ '\t' \ 'function \ | \ FASTools \ -S \ | \ awk \ -F \ '\t' \ 'function \ | \ FASTools \ -S \ | \ Awk \ -F \ '\t' \ ' function \ | \ FASTools \ -S \ | \ Awk \ -F \ '\t' \ ' function \ | \ FASTools \ -S \ | \ Awk \ -F \ '\t' \ ' function \ | \ FASTools \ -S \ | \ Awk \ -F \ '\t' \ ' function \ | \ FASTools \ -S \ | \ Awk \ -F \ '\t' \ ' function \ | \ FASTools \ -S \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk \ -F \ '\t' \ ' function \ | \ Awk 
round(x)\{return (x\%1>=0.5?x+(1-x\%1):x-x\%1)\} BEGIN\{MIN\_LEN=50; SEGM\_LEN=25\} \{1=length(\$2); if (1>=MIN\_LEN)\}
 {n=round(1/SEGM\_LEN)}; segm\_len=round(1/n); n=round(1/segm\_len); split($1,s,""); for (i=1;i<=n;++i) {lo=1+segm\_len*}
 (i-1); hi=(i=-n?1:segm\_len*i); print "@"(i==1?s[1]"~"$2"~"$3:s[1])" \n"substr($2,lo,hi-lo+1)" \n+\n"substr($3,lo,hi-lo+1)" \n+\n
function process_current(){if (current!="") print hits"\t"current} {l=split($1,s,"~"); if (l==3) {process_current();
 current = gensub("^-"," t","g",\$1); \ hits=0 \} \ if \ (\$4^-"([0:|0:0:)[1-9]") \ ++hits \} \ END\{process\_current()\}' \ | \ awk \ -F \ 't' \ -v' \ | \ -F \ 't' \ -V' \ | \ -F \ 't' \ | \
FIRST=ERR275184_1.fastq -v SECOND=ERR275184_2.fastq 'BEGIN{MIN_HITS=3; current=""; found=0} function
process\_current() \{ if (current! = ""\&\&hits > HIN\_HITS\&\& found = 2) \ \{ print first > FIRST; \ print second > SECOND \} \} \ \{ if (current! = ""\&\&hits > HIN\_HITS\&\& found = 2) \} \} \} 
 second="@"$2"\n"$3"\n+\n"$4}} END{process_current()}'
# Third, merge
flash -m 20 -M 500 ERR275184_1.fastq ERR275184_2.fastq -o ERR275184
# Fourth, count k-mers
KPopCount -l ERR275184 -s ERR275184.extendedFrags.fastq -p ERR275184.notCombined_1.fastq
ERR275184.notCombined_2.fastq > ERR275184.k12.txt
```

The script requires several additional components, namely:

• Trim Galore and dependencies. It is run at the very beginning of the script to do quality and adapter trimming on the reads

- FASTools . It is a Swiss-knife tool to manipulate FASTA/FASTQ files. It can be obtained from the BiOCamLib repository. on which the implementation of KPOP depends
- The GEM mapper. It is a high-performance and very accurate aligner with an expressive native output format that we exploit in our script to quickly select matches
- FLASH. It is a fast program to merge partially overlapping FASTQ paired-end reads. While our approach is not very sensitive to their presence, merging them helps keeping the overall *k*-mer normalisation more even, and it is hence considered good practice.

The main idea behind the script is to split each read into small segments (SEGM_LEN, i.e., 25 nt in the code above), map each chunk into a *Mycobacterium* "pangenome", and keep the pair only if a sufficient number of segments (MIN_HITS, i.e., 3 in the code above) align to the pangenome. The sequences we used for the pangenome are listed in the test directory of this repository and can be downloaded from the SRA — we'll assume that they have all been concatenated into file Mycobacterium.CompleteGenomes.fasta. Once that has been done, an index for the pangenome can be generated with the command

```
gem3-indexer -i Mycobacterium.CompleteGenomes.fasta -o Mycobacterium.CompleteGenomes
```

That will generate the file Mycobacterium.CompleteGenomes.gem appearing in the script.

After each read has been split into chunks and aligned to the pangenome, the input of the first awk command executed after gem3-mapper will have FASTQ record in tabular format, with the number of mapping chunks coming before it:

and the second awk commands counts the sum of aligned segments and decides whether or not to keep the pair.

However, equivalent techniques that discard extraneous reads using different approaches or programs would also be acceptable.

5.1.2.2. Data analysis

Once reads have been pre-processed and their *k*-mer spectrum generated for each sample, data analysis proceeds along the lines of the simulated *M.tuberculosis* example above — we assume that the spectra have been placed in a Train and Test directory and separated into subdirectory according to their class, as per the convention previously described. We then generate representative spectra for the classes; twist them; and use the resulting transformation to twist the test sequences, processing each directory in parallel. The results are then collected in the database Test.KPopTwisted.

```
ls -d Train/*/ | awk '{print substr(gensub("Train/","",1),1,length($0)-7)}' | Parallel -l 1 -t 4 -- awk '{CLASS=$0; system("cat Train/"CLASS"/*.txt | KPopCountDB -k /dev/stdin -R \"~.\" -A "CLASS" -L "CLASS" -N -D -v --table-transform none -t "CLASS)}' cat M_*.KPopCounter.txt | KPopCountDB -k /dev/stdin -o Classes -v KPopTwist -i Classes -v ls -d Test/M_*/ | Parallel -l 1 -t 4 -- awk '{system("cat "$0"*.kl2.txt | awk -F \047\\t\047\if ($1==\"\") print $0\"\\001"substr($0,6,length($0)-6)"\"; else print}\047 | KPopTwistDB -i T Classes -k /dev/stdin -o t "substr($0,1,length($0)-1)" -v")}' KPopTwistDB $(ls Test/*.KPopTwisted | awk '{split($0,s,"[.]"); printf " -a t "s[1]}') -o t Test
```

Note that in this example we annotate the name of each test sequence with its class, by adding to it a non-printable character $\$ followed by the class ($\$ is fine with non-printable characters in labels, provided that they are not $\$ $\$ $\$ We'll do the same below for the training sequences below, so as to be able to count the majority class when we use a $\$ $\$ $\$ A-NN approach.

As detailed in our bioRxiv preprint, for this example we explored three different classifiers. Although they all provide very good performance, these alternative approaches are easy to implement using KPop tools and illustrate the versatility of our framework.

This approach works exactly as in the simulated *M.tuberculosis* example above:

```
KPopTwistDB -i T Classes -i t Test -d Classes -o d Test-vs-Classes -s Test Test-vs-Classes -v
cat Test-vs-Classes.KPopSummary.txt | awk '{print gensub("\001","\"\t\"","g")}' > RESULTS-2C.txt
```

Note the use of the distance capability provided by KPOPTWistDB, as in the previous example, which produces results with the format explained above. We also post-process such results to split the name of the test sequence and its annotated class into two separate columns — as mentioned before, we had joined them through a non-printable character \001 that we now turn into \t.

k-NN approach

In this case, we implement the classifier by using a *k*-NN approach, i.e. by performing a majority call among the *k* nearest neighbours found for each test sequence - the result is the class which is most represented in the *k* closest data points belonging to the training set. In order to be able to do so, we need to separately compute the twisted spectrum of each training sequence — we also collect them into a database named Train:

```
ls -d Train/M_*/ | Parallel -l 1 -t 4 -- awk '{system("cat "$0"*.k12.txt | awk -F \047\\t\047 \047{if ($1==\"\") print $0\"\\001"substr($0,7,length($0)-7)"\"; else print}\047 | KPopTwistDB -i T Classes -k /dev/stdin -o t "substr($0,1,length($0)-1)" -v")}'
KPopTwistDB $(ls Train/*.KPopTwisted | awk '{split($0,s,"[.]"); printf " -a t "s[1]}') -o t Train
```

We then use the distance capability of KPOpTwist as before. However, in this case we alter the default to output information about 5 nearest neighbours rather than 2:

```
KPopTwistDB $(ls Test/*.KPopTwisted | awk '{split($0,s,"[.]"); printf " -a t "s[1]}') -o t Test -d Train -o d Test-
vs-Train --keep-at-most 5 -s Test Test-vs-Train -v
```

This produces the usual digest of the distance matrix containing, in this case, the 5 training sequences that are closest to the test sequence (we adopt a 5-NN classification approach in this example, but of course the number of neighbours could be chosen to be different). At this point we are almost done, but we still have to parse the digest in order to compute how many of the nearest neighbours belong to the most frequent class, which translates into a slightly more complex script:

```
cat Test-vs-Train.KPopSummary.txt | awk '{print gensub("\001","\"\t\"","g")}' | awk '{delete c; delete o; n=0; for (i=8;i<=NF;i+=4) {++n; if (!($i in c)) o[length(c)+1]=$i; ++c[$i]} max=0; max_class=0; l=length(o); for (i=1;i<=l;++i) {if (c[o[i]]>max) {max=c[o[i]]; max_class=o[i]}} print 0\t"max_class=\t"(max/n)}' > RESULTS-kNN.txt
```

Notably, we solve ties by selecting the class that appears first, i.e. which has a sequence that is closer to the test sequence.

Random forest approach

This is a more sophisticated approach that uses random forests to implement the predictor. Note that this is possible because KPOP is capable of producing an explicit embedding of the sequences as a vector in twisted space. As we are going to use an external R library to implement the predictor, we first have to export to a tabular text file the coordinates of the sequences in twisted space — which we can do with KPOPTwist — and then, once more, de-mangle the class information into a separate column:

```
KPopTwistDB -i t Train -0 t Train -i t Test -0 t Test
cat Train.KPopTwisted.txt | awk '{print gensub("\001","\"\t\"","g")}' | awk -F '\t' 'BEGIN{0FS="\t"} {if (NR==1)
{$1="\"Lineage\""}; print}' > Train.Truth.KPopTwisted.txt
cat Test.KPopTwisted.txt | awk '{print gensub("\001","\"\t\"","g")}' | awk -F '\t' 'BEGIN{0FS="\t"} {if (NR==1)
{$1="\"Lineage\""} else $2="\"\""; print}' > Test.Truth.KPopTwisted.txt
```

The files Train.Truth.KPopTwisted.txt and Test.Truth.KPopTwisted.txt will now contain the coordinates of train and test sequences in twisted space, respectively. We then run the following R commands:

```
library(randomForest)
train<-read.table("Train.Truth.KPopTwisted.txt")
test<-read.table("Test.Truth.KPopTwisted.txt")
rf<-randomForest(Lineage~.,data=train,mtry=4)
write.table(as.data.frame(predict(rf,test)),"RF.txt",sep="\t")</pre>
```

which will produce a file RF.txt with the preliminary results of the predictor.

And finally, this command re-annotates the sequences with their original class for comparison purposes:

```
cat RF.txt | awk -F '\t' 'BEGIN{while (getline < "NGS-TB-Test.txt") t["\""$1"\""]="\""$2"\"" {if (NR>1) print $1"\t"\{1\}" > RESULTS-RF.txt
```

generating a final result RESULTS-RF.txt . The file NGS-TB-Test.txt contains the original annotation for each test sequence, and is available from the test directory of this repository.

It should be noted that the structure of RESULTS-RF.txt is simpler than that of the final files produced by the previous methods. It contains lines such as

```
"ERR046839"
              "M_tuberculosis_L4"
                                    "M_tuberculosis_L4"
              "M_tuberculosis_L5"
"ERR046933"
                                     "M_tuberculosis_L5"
"ERR072050" "M_tuberculosis_L4"
                                    "M_tuberculosis_L4"
"ERR1023314" "M_tuberculosis_L4" "M_tuberculosis_L4"
"ERR1023338" "M_tuberculosis_L3"
                                    "M_tuberculosis_L3"
"ERR1023449"
              "M_tuberculosis_L2"
                                    "M_tuberculosis_L2"
"ERR1034887" "M_tuberculosis_L4"
                                    "M tuberculosis L4"
```

each one reporting samples name, original class, and classification generated by random forest — there is no additional statistical information reported in the default output generated by randomForest .

5.1.3. Classifier for SARS-CoV-2 sequences (Hyena)

This is a rather more complex example, that showcases many of the good qualities of KPop (mainly its being high-throughput and accurate). It's also not for the faint of heart, in that it requires large amounts of disk space and computing time — at the time of this writing, the file containing all COVID-19 sequences made available on GISAID has an uncompressed size of ~303 GB, and counting. (And we know that technically we should call them SARS-CoV-2 sequences, but in this section we'll sloppily say COVID-19 instead.) In fact, most of the needed preprocessing does not really have much to do with KPop, but we work out the complete example anyway for clarity's sake. Feel free to skip it if you are only interested in the KPop, part.

5.1.3.1. Data preparation

▼ Preparing input data for the SARS-CoV-2 classifier

We assume as a starting point that you have downloaded from GISAID and decompressed in your current directory the sequences.fasta file containing all the sequences available until that moment (note that the file is *not* publicly available — you'll have to obtain access to GISAID if you want to be able to download it). We'll also assume you have downloaded to your local directory the file lineages.csv containing the Pangolin designations of COVID-19 lineages. There are some ~1600 lineages — i.e., classes to recognise — that are currently in use.

So you'll have two files in your directory to start with,

```
sequences.fasta
lineages.csv
```

As a first step, we must extract the actual COVID-19 sequences that we need in order to train and test our classifier - the file lineages.csv does not contain the actual sequences, just their classification (which is assumed to be the ground truth). In order to do so, we run the following script:

```
CHUNK_BLOCKS=125; BLOCK_SIZE=$(( 1024 * 1024 )); rm -rf Split; mkdir Split; echo sequences.fasta | awk -v
 CHUNK_BLOCKS="$CHUNK_BLOCKS" -v BLOCK_SIZE="$BLOCK_SIZE" 'END{get_size="ls -1 \047"$0"\047 | awk \047{print $5}\047"
  |& getline size; close(get_size); blocks=int((size+BLOCK_SIZE)/BLOCK_SIZE);
 \verb|chunks=int((blocks+CHUNK_BLOCKS)/CHUNK_BLOCKS)|; for (i=0;i<chunks;++i) | print $0"\t"i*CHUNK_BLOCKS}|' | Parallel -l 1 | l 
  -t $(( $(nproc) * 3 )) -- awk -F '\t' -v BLOCK_SIZE="$BLOCK_SIZE" '{get_chunk="dd bs="BLOCK_SIZE" if=\047"$1"\047
skip="$2" 2> /dev/null"; overhang=""; line=""; while (get_chunk |& getline line) {if (line==""||line~"^>") break;}
overhang = overhang \ line "\n" \} \ print $1"\t"($2*BLOCK\_SIZE) + length(overhang); \ close(get\_chunk) \}' \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t' \ -v \ | \ awk \ -F \ '\t'
 INPUT = sequences. fasta" + (if (NR>1) print $1"+t"old"+t"($2-old); old=$2$ END{get\_size="ls -l \047"INPUT"\047 | awk fold=$1.5 | awk fold=$
  \047{print $5}\047" |& getline size; close(get_size); print $1"\t"old"\t"(size-old)}' | Parallel -l 1 -- awk -F '\t'
  -v BLOCK_SIZE="$BLOCK_SIZE" 'function remove_spaces(name,
                                                                                                                                                                                                                                                                                                                                                                                                                                 s){split(name,s,"/"); return gensub("[
    _]","","g",s[1])"/"s[2]"/"s[3]} BEGIN{nr=0; while (getline < "lineages.csv") {++nr; if (nr>1) {split($0,s,",");
t[remove\_spaces(gensub("^BHR/","Bahrain/",1,s[1]))] = s[2]\}\} \ function \ output\_sequence() \{ ++ counts[offset"/"class]; \} \} 
 print ">"name" \\ n"seq > "Split/"offset"/"class".fasta"; return} {offset=$2; system("mkdir -p Split/"offset); return} {offset=$2; system("mkdir -p Split/"offset"); return} {offset=$2; system("mkdir -p Split/"
 \label{lock_size} $$ get_chunk="dd bs="BLOCK_SIZE" if=\047"$1"\047 iflag=\"skip_bytes,count_bytes\" skip="offset" count="$3" 2> $$ for the count="$3" 2> $$ for the count="$3" 2> $$ for the count="$4" f
 \label{lem:condition} $$ \dev/null"; first=0; while (get\_chunk | \& getline) {if ($0^"^>") {if (first&&active) output\_sequence(); first=1; } $$ $$ \dev/null"; first=0; while (get\_chunk | \& getline) {if ($0^"^>") {if (first&active) output\_sequence(); first=1; } $$ $$ \dev/null"; first=0; while (get\_chunk | \& getline) {if ($0^"^>") {if (first&active) output\_sequence(); first=1; } $$ $$ \dev/null"; first=0; while (get\_chunk | \& getline) {if ($0^"^>") {if (first&active) output\_sequence(); first=1; } $$ $$ \dev/null"; first=0; while (get\_chunk | \& getline) {if ($0^"^", $) {if (first&active) output\_sequence(); first=1; } $$ \dev/null"; first=0; while (get\_chunk | \& getline) {if ($0^", $) {if (first&active) output\_sequence(); first=1; } $$ \dev/null"; first=0; while (get\_chunk | \& getline) {if ($0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", $0^{", 
 active=0; split(substr($0,10),s,"[|]"); res=remove_spaces(s[1]); if (res in t) {active=1; name=res; class=t[res];
seq=""\} else \{if (active) \ seq=seq \$0\}\} \ if (active) \ output\_sequence()\} \ END\{for (i \ in \ counts) \ print \ i"\t"counts[i]\}' \ else \{if (active) \ seq=seq \$0\}\} \ if (active) \ output\_sequence()\} \ END\{for (i \ in \ counts) \ print \ i"\t"counts[i]\}' \ else \{if (active) \ seq=seq \$0\}\} \ if (active) \ output\_sequence()\} \ END\{for (i \ in \ counts) \ print \ i"\t"counts[i]\}' \ else \{if (active) \ seq=seq \$0\}\} \ if (active) \ output\_sequence()\} \ END\{for (i \ in \ counts) \ print \ i"\t"counts[i]\}' \ else \{if (active) \ seq=seq \$0\}\} \ if (active) \ output\_sequence()\} \ else \{if (active) \ seq=seq \$0\}\} \ if (active) \ output\_sequence()\} \ else \{if (active) \ seq=seq \$0\}\} \ if (active) \ output\_sequence()\} \ else \{if (active) \ seq=seq \$0\}\} \ if (active) \ seq=seq \$0\}
 \mid awk -F '\t' '{split($1,s,"/"); offset=s[1]; class=s[2]; system("cat \"Split/"$1".fasta\" >>
\"Split/"class".fasta\"; rm \"Split/"$1".fasta\"; rmdir --ignore-fail-on-non-empty Split/"offset);
counts[class]+=$2} END{for (i in counts) print i"\t"counts[i]}' > Stats.txt
```

Although apparently very complex, the script is just performing a parallelised scan of the big file, in order to find the sequences specified in lineages.csv and write them as files — one per class — in the directory Split. In particular, the script is made of three parts, each one performing a specific task.

Indexing

```
CHUNK BLOCKS=125
BLOCK_SIZE=$(( 1024 * 1024 ))
rm -rf Split
mkdir Split
echo sequences.fasta |
awk -v CHUNK_BLOCKS="$CHUNK_BLOCKS" -v BLOCK_SIZE="$BLOCK_SIZE"
    get_size="ls -l \047"$0"\047 | awk \047{print $5}\047" |& getline size
    close(get size)
    blocks=int((size+BLOCK_SIZE)/BLOCK_SIZE)
    \verb|chunks=int((blocks+CHUNK_BLOCKS)/CHUNK_BLOCKS)|\\
    for (i=0;i<chunks;++i)</pre>
      print $0"\t"i*CHUNK_BLOCKS}
' | Parallel -l 1 -t $(( $(nproc) * 3 )) -- awk -F '\t' -v BLOCK_SIZE="$BLOCK_SIZE" '
    get\_chunk="dd bs="BLOCK\_SIZE" if=\\047"$1"\\047 skip="$2" 2> /dev/null"
    overhang=""
    line=""
    while (get_chunk |& getline line) {
      if (line==""||line~"^>")
        break:
      overhang=overhang line"\n"
    print $1"\t"($2*BLOCK_SIZE)+length(overhang)
    close(get_chunk)
' | awk -F '\t' -v INPUT="sequences.fasta" '
    if (NR>1)
      print $1"\t"old"\t"($2-old);
    old=$2
 END {
    \label{lem:get_size} get\_size="ls -l \047"INPUT" \047 | awk \047{print $5} \047" | \& getline size
    close(get_size)
    print $1"\t"old"\t"(size-old)
 }
```

After creating an empty directory Split, this script logically partitions the input file, sequences.fasta, into blocks of 125 MB, or \$125 * 1024 *1024 \$ B. As the sequences have variable length, however, before doing anything meaningful we have to locate the boundaries between blocks by locating where the first record after the logical boundary between each two blocks effectively starts. Notably, the script uses the Unix command dd in order to stream the contents of the file starting from the logical boundaries between blocks rather than from the beginning — the operation is performed in parallel for many couples of blocks thanks to the

use of Parallel. The result is an index file in tabular form, with each line describing a block as a range, expressed as starting and ending position in bytes with respect to the beginning of the file. It should be noted that this technique is general and goes beyond this specific example — one could apply this script to any FASTA file, and even modify it to index other formats by suitably modifying the regular expression

```
line~"^>"
```

which located the beginning of the next FASTA record.

Parallel scan

```
Parallel -l 1 -- awk -F '\t' -v BLOCK_SIZE="$BLOCK_SIZE" '
 function remove_spaces(name,
                                 s) {
   split(name,s,"/")
   return gensub("[ _]","","g",s[1])"/"s[2]"/"s[3]
 }
 BEGIN {
    nr=0
   while (getline < "lineages.csv") {</pre>
     ++nr;
     if (nr>1) {
       split($0,s,",")
        t[remove_spaces(gensub("^BHR/", "Bahrain/", 1, s[1]))]=s[2]
   }
  }
  function output_sequence() {
   ++counts[offset"/"class]
    print ">"name"\n"seq > "Split/"offset"/"class".fasta"
    return
 }
  {
    offset=$2
    system("mkdir -p Split/"offset)
    get_chunk="dd bs="BLOCK_SIZE" if=\047"$1"\047 iflag=\"skip_bytes,count_bytes\" skip="offset" count="$3" 2>
/dev/null"
    first=0
    while (get_chunk |& getline) {
     if ($0~"^>") {
       if (first&&active)
         output_sequence();
        first=1
        active=0
        split(substr($0,10),s,"[|]")
        res=remove spaces(s[1])
        if (res in t) {
         active=1
          name=res
          class=t[res]
          seq=""
       3
     } else {
        if (active)
          seq=seq $0
     }
    if (active)
      output_sequence()
  }
 END {
    for (i in counts)
     print i"\t"counts[i]
 }
```

This is the core of the method, with several "worker" processes that are spawned and executed at the same time by Parallel. Each one operates on one of the chunks identified by the previous part, streaming it with dd and identifying the class each sequence belongs to by comparing it with the information contained in lineages.csv (some additional manipulations are needed in order to cope with several inconsistencies between the way metadata is represented in sequences.fasta and lineages.csv, but they are not really relevant to the general workflow).

Each worker outputs its results in a different directory named <code>Split/<offset></code>, where <code><offset></code> is the position within <code>sequences.fasta</code> of the beginning of the chunk processed by the worker. The sequences seen by each worker are split into different sub-directories of <code>Split/<offset></code>, named <code>Split/<offset>/<lineage></code>, according to their class. Each worker also outputs a summary made of a series of records of the form

```
<offset> / eage> Tab <frequency>
```

that is subsequently fed to the next stage.

Sequential collection

```
awk -F '\t' '
{
    split($1,s,"/")
    offset=s[1]
    class=s[2]
    system("cat \"Split/"$1".fasta\" >> \"Split/"class".fasta\"; rm \"Split/"$1".fasta\"; rmdir --ignore-fail-on-
non-empty Split/"offset)
    counts[class]+=$2
}
END{
    for (i in counts)
        print i"\t"counts[i]
}
' > Stats.txt
```

Implementing our usual convention, this last part accumulates the worker-specific results, which had been stored by each worker into a separate <code>split/<offset></code> directory, into global results that have the same structure (one sub-directory per lineage/class) but live in the <code>split</code> directory. Worker-specific files and temporary directories are then deleted. Finally, the script outputs a file named <code>stats.txt</code> containing the number of sequences successfully found and written for each lineage.

Unfortunately this step cannot be easily parallelised, as a concurrent write to the same file by several processes would result in corruption of the file — so it turns out to be the bottleneck. We could implement some locking/mutexing mechanism, but, after all, we have to prepare data only once. So we sit down and wait a couple hours.

By the end of the script, the Split directory is populated by files names as COVID-19 lineages, such as

```
AY.4.2.fasta
BA.1.15.1.fasta
```

Each file contains (almost) all the sequences used as the ground truth for that lineage. Note that the procedure we used does not retrieve all sequences because some of the ones used to train Pangolin are not in GISAID. In particular, the following command:

```
 awk -F '\t' 'BEGIN\{nr=0; while (getline < "lineages.csv") \{++nr; if (nr>1&&$0-",") \{split($0,s,","); found[s[2]]; \\ ++lineages[s[2]]\}\} while (getline < "Stats.txt") {found[$1]; stats[$1]=$2} for (i in found) {print i"\t"(i in lineages?lineages[i]:0)"\t"(i in stats?stats[i]:0)}}' | awk -F '\t' '{if (($3<0.9*$2&&$3<20)||$2<10&&$3!=$2) print}'
```

first generates a table containing the number of sequences in each category both in the Pangolin set and returned by our procedure,

```
BA.1.13.1
           1536
                   1536
           1289
AY.4.2.1
                 1288
AY.4.2.2
           3241
                  3234
AY.4.2.3
           1237
                  1235
AY.4.2.4
           114
                  114
B.30 50
            50
     212
            166
B.31
B.1.36.10
            111
                   110
```

and then it only keeps categories for which our procedure has produced too much change or an insufficient number of sequences. Further inspection shows that the only really probelmatic categories are

```
B.60 74 2
AY.3.4 186 5
```

for which most of the sequences are unavailable in GISAID (and B.1.617, which only has 1 sequence in lineages.csv). We might find workarounds for that, but, as that would involve the downloading of more large datasets, we keep things simple and just exclude B.60 and AY.3.4 from the rest of the analysis.

At this point we just have to split each of the files containing lineage-specific sequences into training and test set, as usual. The following command does it:

```
rm -rf Train Test; mkdir Train Test; for FILE in Split/*.fasta; do BASE=$(basename $FILE); echo $BASE; done |
Parallel -l 1 -- awk 'function output_seq(){if (name!="") print name"\n"seq >> (counts%2==1?"Train":"Test")"/"$0}
{input="Split/"$0; counts=0; while (getline line < input) if (line~"^>") {output_seq(); ++counts; name=line} else
seq=line; output_seq()}'
```

suitably populating the subdirectories Test and Train. As in most of the examples so far, we use Parallel to perform the splitting on many files in parallel and hence reduce the overall wallclock time taken by the command.

5.1.3.2. Data analysis

Once the preparation stage has been completed, the analysis proceeds exactly as in the M.tuberculosis example above.

First, we compute the spectra for the class representatives using KPopCount and KPopCountDB:

```
ls Train/*.fasta | Parallel -l 1 -- awk '{l=split($0,s,"/"); class=substr(s[l],1,length(s[l])-6); system("KPopCount -k 10 -l "class" -f Train/"class".fasta")}' | KPopCountDB -k /dev/stdin -o Classes
```

This produces a Classes. KPopCounter file which is ~2.0 GB. The command

```
KPopCountDB -i Classes --summary
```

confirms that indeed this database contains the spectra for all classes:

```
[Vector labels (1636)]: 'A.11' 'A.12' 'A.15' 'A.16' 'A.17' 'A.18' 'A.19' 'A.1' 'A.21' 'A.22' 'A.22' 'A.23.1' 'A.2.3' 'A.23' 'A.2.4' 'A.24' 'A.25.1' 'A.2.5.2' 'A.2.5.3' 'A.2.5' 'A.25' 'A.26' 'A.27' 'A.28' 'A.29' 'A.2' 'A.30' 'A.3' 'A.4' 'A.5' 'A.6' 'A.7' 'A.8' 'A.8
```

According to the usual procedure, we then twist the class representatives with KPopTwist:

```
KPopTwist -i Classes -v
```

and we use the resulting classifying twister to twist and accumulate the test sequences into file Test.KPopTwisted with KPopTwistDB:

```
cat Test/*.fasta | awk 'BEGIN{ok=1} {if ($0~"^>") {ok=!($0 in t); t[$0]} if (ok) print}' | Parallel -l 2 -- awk '{if (NR==1) {job="KPopCount -k 10 -l \""substr($0,2)"\" -f /dev/stdin"; print $0 | \& job} else {print $0 | \& job; close(job, "to"); while (job |& getline) print $0}' | KPopTwistDB -i T Classes -k /dev/stdin -o t Test -v
```

Note that right at the beginning of the script, with the part

```
awk 'BEGIN{ok=1} {if ($0^{-"}>") {ok=!($0 in t); t[$0]} if (ok) print}'
```

we are removing repeated sequences (yes, apparently there are repeated sequences in GISAID ;) as KPOPTWistDB is unhappy with them and will bail out if it finds some in the input.

Also, as there are \sim 650K sequences to be classified, doing so will take long (\sim 14h on the node I'm using for these tests). So, you might wish to further parallelise the process, as I did, by splitting the input into smaller files and processing them separately on different nodes. After that, you can merge together all the pieces with a command such as

```
KPopTwistDB -a t Test.aa -a t Test.ab ... -o t Test -v
```

The final size of the file Test . KPopTwisted containing all the ~650K twisted COVID-19 sequences in the test set is ~8.4 GB.

At this point, the analysis proceeds exactly as in the case of the previous section, with a command such as

```
KPopTwistDB -i T Classes -i t Test -d Classes -o d Test-vs-Classes -v
```

that allows us to compute all the distances of each test sequence from each of the "training" classes. However, given that in this case there is a very large number of distances to be computed, another and perhaps more appropriate strategy would have been not to merge the twisted test files into a single Test.KPopTwisted file, but rather to compute the distances from the twisted classes for each of the chunks with commands such as

```
KPopTwistDB -i T Classes -i t Test.aa -d Classes -o d Test-vs-Classes.aa
```

and then merge together all the distance files, as in

```
KPopTwistDB -a d Test-vs-Classes.aa -a d Test-vs-Classes.ab ... -o d Test-vs-Classes -v
```

One way or another, once a file Test-vs-Classes. KPopDMatrix containing all the distances has been generated, we can run

```
KPopTwistDB -i d Test-vs-Classes -s Test.aa Test-vs-Classes -v
```

in order to produce the usual textual summary of the distances. The resulting file will be ~118 MB in size.

And quite likely, one would also wish to run something like the following commands:

```
cat Test-vs-Classes.KPopSummary.txt | awk 'function remove_spaces(name, s){split(name,s,"/"); return gensub("[
_]","","g",s[1])"/"s[2]"/"s[3]} BEGIN{nr=0; while (getline < "lineages.csv") {++nr; if (nr>1) {split($0,s,",");
t[remove_spaces(gensub("^BHR/","Bahrain/",1,s[1]))]=s[2]}}} {printf $1"\t\""t[substr($1,2,length($1)-2)]"\""; for
(i=2;i<=NF;++i) printf "\t"$i; printf "\n"}' > Test-vs-Classes.KPopSummary.Truth.txt
cat Test-vs-Classes.KPopSummary.Truth.txt | awk -F '\t' 'function strip_quotes(s){return substr(s,2,length(s)-2)}
{one=strip_quotes($2); two=strip_quotes($7); print one"\t"two"\t"(substr(two,1,length(one)+1)) > "/dev/null"; if
($2!=$7&&substr(two,1,length(one)+1)!=one".") ++ko; else ++ok}
END{printf("%d\t%d\t%.3g%\t%d\t%.3g%\n",ok+ko,ok,ok/(ok+ko)*100,ko,ko/(ok+ko)*100)}'
```

to, first, annotate the summary with the original "true" classification of the sequences, and, second, generate a one-line summary of the results. The last command will produce something like:

```
641847 611770 95.3% 30077 4.69%
```

meaning that, of the 641,847 sequences present in the Test set, 611,770 (95.3%) were correctly classified, while 30,077 (4.69%) were not.

Note that as not all the classes describing lineages are disjoint, here we consider a classification correct if it is the same as, or a sublineage of, the original one (for instance, B.1.1 would be accepted if the initial category is B.1).

Finally, as discussed in more detail in our bioRxiv preprint, for this example you might wish to tune the definition of the distance by modifying the metric used. In order to do so, you would replace the command to compute distances we used previously with something like the following:

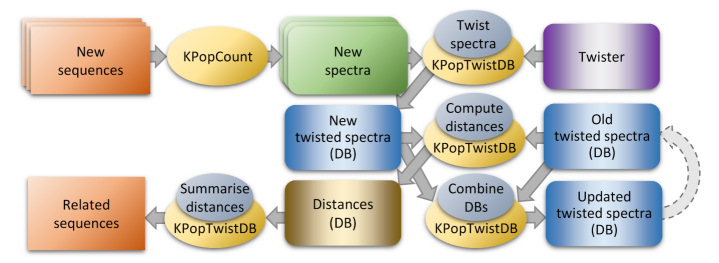
```
KPopTwistDB -m "sigmoid(1,1,5,5)" -i T Classes -i t Test -d Classes -o d Test-vs-Classes.sigmoid_1_1_5_5 -v
```

That takes into account in a different way the estimated importance of the different directions in twisted space, resulting in slightly better overall predictions.

5.2. Relatedness engine

The previous examples have all showcased KPOP 's capability to encode sequences as vectors in twisted space and find the closest neighbours there. This correctly suggests that by using KPOP one can also easily generate a "relatedness engine" out of a classification — i.e., a system finding the most similar sequences in a large database.

The overall strategy is illustrated in the following figure, which assumes that we have already generated a classifying twister (purple box) and a database of twisted sequences (second box from the top on the right) to start with:



Briefly, we can generate and twist new spectra (top left), find out their nearest neighbours in the database of twisted sequences (central line and bottom left), and add the new twisted spectra to the database (bottom right).

Here we illustrate the approach taking the previous exercise on COVID-19 as a starting point. For the sake of simplicity, we'll take as existing database the twisted test sequences (half of the GISAID database) in the file Test.KPopTwisted. The classifier will be the one we generated in the previous section and contained in the file Classes.KPopTwister. The command

```
cat Train/C.36.3.fasta | fasta-tabular | shuf | head -1 | awk -F '\t' '{print $1 > "/dev/stderr"; print ">"$1"\n"$2}' | KPopCount -k 10 -f /dev/stdin -l "C.36.3" | KPopTwistDB -i T Classes -i t Classes -k /dev/stdin -d Test --keep-at-most 300 -s Test Related
```

will then randomly select one sequence from the c.36.3 lineage and find the 300 sequences closest to it. A summary in the usual format will be output to file Related.KPopSummary.txt. Note that loading this specific database in memory is going to take long as the file is ~8.4 G, but one only needs to do it once when sequences are processed in batches.

5.3. Pseudo-phylogenetic trees

