# **AAF User Manual**

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# Introduction

AAF (alignment and assembly-free) is a free software package that reconstructs a phylogeny from next-generation sequencing data without assembly and alignment. This software package directly analyzes raw sequencing reads from all of the samples simutaneously and generates a distance matrix based on the proportion of shared k-mers between each sample and reconstruct a phylogeny based on the distance matrix, using comprehensively tested statistical methods. See Fan et al BMC genomics (in revision) for detailed description of the method.

AAF is mainly designed for large eukaryotes genomes (>10<sup>6</sup> base pairs). The whole reconstruction process is separated into two major steps: 1) k-mer counting and 2) distance calculation and phylogeny reconstruction. Two separate python scripts implement the two steps respectively: 1) **aaf\_phylokmer.py 2) aaf\_distance.py.** There are 3 more optional scripts in the AAF package: **aaf\_tip.py** trims excessive tips of the phylogeny generated due to sequencing error and incomplete coverage; **nonparametric\_bootstrap.py** and **parametric\_bootstrap.R** provide estimates of bootstrap support for the resulting phylogenies constructed. The use of these scripts is described below.

Two tutorials are provided to illustrate the usage of this software package. The quick start tutorial primarily explains data organization and script usage, using a dummy dataset (provided in the package) with 10 species and short genomes ( $\sim$ 800kb). The second tutorial (in a separate file: Tutorial\_21tropicaltrees.pdf, included in the package) demonstrates possible issues encountered when dealing with large datasets of real next-gen DNA sequence data, with special focus on parameter selection including k and filtering. This tutorial uses whole genomic data collected from 21 tropical tree genomes as described in Fan et al BMC Genomics (in revision). There is a separate section detailing the reasoning for optimal parameter selection at then end of the manual.

For the most recent version of AAF, please visit https://sourceforge.net/projects/aaf-phylogeny/

# Requirements

AAF can be used on a UNIX system (Linux, OsX...) with Python 2.6 and higher 2.X versions, but NOT Python 3.0+), and g++/gcc compilers. Biopython (<a href="http://biopython.org/wiki/Main\_Page">http://biopython.org/wiki/Main\_Page</a>) is required for the non-parametric bootstrap, and R (<a href="http://cran.r-project.org/">http://cran.r-project.org/</a>) and the R package 'ape' are required for the parametric bootstrap.

# Installation

- 0. Decompress the zip file downloaded from <a href="http://sourceforge.net/projects/aaf-phylogeny">http://sourceforge.net/projects/aaf-phylogeny</a> with the most recent version.
- 1. Compile kmer\_count(x) and kmer\_merge as follows. "path\_to\_AAF" stands for your path to the AAF folder generated by decompressing AAF.tar.gz.
  - a. path to AAF/AAF\$ cd phylokmer
  - b. path to AAF/AAF/phylokmer\$ make
  - c. Add kmer count(x) and kmer merge to your PATH or working directory
- 2. Compile fitch\_kmerX, consense and treedist
  - a. path to AAF/AAF\$ cd phylip src

```
b. path_to_AAF/AAF/phylip_src$ make allc. Add fitch kmerX and consense to your PATH or working directory
```

# **Quick start tutorial**

This tutorial demonstrates the installation of AAF and its application on a dummy dataset with 10 species and short genomes (~800kb).

## Demonstration with dummy dataset

1) Decompress the AAF package.

```
2) Move to the phylokmer directory and compile kmer_count, kmer_countx, and kmer_merge path_to_AAF/AAF$ cd phylokmer path_to_AAF/AAF/phylokmer$ make path to AAF/AAF/phylokmer$ cp kmer count kmer countx kmer merge ../
```

#### 3) Compile fitch kmerX

```
path_to_AAF/AAF/phylokmer$ cd ../phylip_src
path_to_AAF/AAF/phylip_src$ make all
path_to_AAF/AAF/phylip_src$ cp fitch_kmerX consense ../
path to AAF/AAF/phylip src$ cd ..
```

#### 4) k-mer counting

path to AAF/AAF/\$ python aaf phylokmer.py -k 21 -d data -G 2

### 5) Constructing the phylogenetic tree

```
path_to_AAF/AAF/$ python aaf_distance.py -i data/phylokmer.dat.gz -o test -t 2 -G 2
-f data/kmer diversity.wc
```

#### 6) Tip correction (optional)

```
path_to_AAF/AAF/$ python aaf_tip.py -i test.tre -k 21 --tip data/tip_info_test.txt
-f data/kmer diversity.wc
```

#### 7) Non-parametric bootstrap (optional)

```
path_to_AAF/AAF/$ python nonparametric_bootstrap.py -k 21 -t 2 -d data --S1 2 --S2 2
```

#### 8) Parametric bootstrap (optional)

Set your working directory to the AAF folder and change the parameters in the "set parameters" section, including nboot, k, filter, info.file and n.table.file.

#### Within R console or terminal:

```
> source("parametric bootstrap.R")
```

#### Description of output files

- 1) phylokmer.dat.gz (aaf\_phylokmer.py): This output file will be inside the data folder. It starts with header information including *k*-mer size, filter frequency and sample list. After the header is a table with frequencies of a given *k*-mer from each sample on one line in alphabetical order of the sample names.
- 2) kmer\_diversity.wc (aaf\_phylokmer.py): Inside the data folder. It contains the total *k*-mer diversity for each sample per line in alphabetical order of the sample names.
- 3) test nshare.csv (aaf distance.py): In the current working directory. It contains the number of shared k-

- mers for each pair of samples. This file is needed for the parametric bootstrap.
- 4) test.tre (aaf distance.py): In the current working directory. It is the phylogeny you want!
- 5) test.dist (aaf\_distance.py): In the current working directory. It is the distance matrix upon which fitch kmerX infers the phylogeny (test.tre in this case).
- 6) tip test.tre (aaf tip.py): In the current working directory. It is the tree after tip correction.
- 7) consensus\_trees\_read\_nonparametric (nonparametric\_bootstrap.py): In the current working directory. This file contains all the trees that were generated after each resampling of the reads.
- 8) consensus\_trees\_total\_nonparametric (nonparametric\_bootstrap.py): In the current working directory. This file contains all the trees that were generated after both resampling of the reads and the *k*-mer table counted from those reads.
- 9) consensus\_trees\_table\_nonparametric (nonparametric\_bootstrap.py): In the current working directory. This file contains all the trees that were generated after each resampling of the real *k*-mer table calculated from the original reads when the resampling of reads is skipped.
- 10) consensus\_read\_nonparametric.tre (nonparametric\_bootstrap.py): In the current working directory. This is the consensus tree generated from consensus\_trees\_read\_nonparametric by consense in PHYLIP.
- 11) consensus\_total\_nonparametric.tre (nonparametric\_bootstrap.py): In the current working directory. This is the consensus tree generated from consensus\_trees\_total\_nonparametric by consense in PHYLIP.
- 12) consensus\_table\_nonparametric.tre (nonparametric\_bootstrap.py): In the current working directory. This is the consensus tree generated from consensus\_trees\_table\_nonparametric by consense in PHYLIP.
- 13) consensus\_outfile\_read\_nonparametric (nonparametric\_bootstrap.py): In the current working directory. This file contains the bootstrap ratio for the branches in consensus read nonparametric.tre
- 14) consensus\_outfile\_total\_nonparametric (nonparametric\_bootstrap.py): In the current working directory. This file contains the bootstrap ratio for the branches in consensus total nonparametric.tre
- 15) consensus\_outfile\_table\_nonparametric (nonparametric\_bootstrap.py): In the current working directory. This file contains the bootstrap ratio for the branches in consensus table nonparametric.tre
- 16) consensus\_\*\_\*\_parametric (parametric\_boostrap.R): In the current working directory. See descriptions of their nonparametric counterparts.

# **Usage and options**

#### 1) aaf phylokmer.py

This script takes raw reads, count the k-mers from each sample and generate a shared k-mer table with k-mers that are present in at least two samples and their frequencies in each sample.

Usage: aaf\_phylokmer.py [options]

### Options:

--version show program's version number and exit

-h, --help show this help message and exit

-k KLEN k-mer length, default = 25

-t NTHREADS number of threads to use, default = 1

-n FILTER k-mer filtering threshold, default = 1

-f SEQFORMAT format of input files, FA|FQ, default = FA

-o OUTFILE output file, default = phylokmer.dat.gz

-d DATADIR directory containing the data, default = data/

-G MEMSIZE total memory limit (in GB), default = 4

-W withKmer include k-mers in the shared k-mer table

-s only print commands, do not run them

#### Detailed description of options:

- -k KLEN: *k*-mer size. Larger *k* will decrease the probability of two identical *k*-mers from different parts of the genome (*k*-mer homoplasy) while increase the probability of *k*-mers containing sequencing errors and multiple evolutionary events such as substitutions or indels. See more details in the parameter selection section. Set at 25 by default.
- -t NTHREADS: number of threads to use. Depends on how many cores are available on your machine. Set at 1 by default.
- -n FILTER: how many times a *k*-mer needs to be in the sample to be counted as present. This serves as the filter for singletons, which could be the result of sequencing error. See more details about the parameter selection section. Set at 1 by default.
- -f SEQFORMAT: format of the sequence files, FA or FQ. The default is set as FA.
- -o OUTFILE: output filename. If you would like your output file to be compressed, provide a name that ends with .gz. Otherwise it will not be compressed. The default output file is phylokmer.dat.gz, which is compressed.
- -d DATADIR: directory containing the data. Users should strictly follow the data structure required by AAF. Sequence files for each sample need to be in one directory named after that sample. Therefore, there will be N directories for N samples and the name of the directories will be the names displayed in the final phylogenetic tree. All the sample folders should be placed into the same directory, which will be your data directory requested by aaf\_phylokmer.py. Accepted extensions for sequence files include: .fa(sta)(.gz), .fq(.gz), and .fastq(.gz). See the "data" directory in the package as an example.
- -G MEMSIZE: the total memory allowance. Each kmer\_count thread has G/t memory allowance. Set at 4G by default.
- -W WITHKMER: to include k-mers in the shared k-mer table. When the final goal is to construct a phylogeny, we do not need to know the specific patterns of each k-mer. Therefore by default in the shared k-mer table only the frequencies of k-mers are kept. However if there's downstream analysis of k-mers with a certain pattern, k-mers need to be kept. Use -W to keep the k-mers.
- -s SIM: This will print out the commands that are going to run without executing them.

# 2) aaf distance.py

This script takes the shared k-mer table generated from aaf\_phylokmer.py, calculates the distance matrix and reconstructs the phylogeny based on it.

Usage: aaf-distance.py [options] -i <input filename>

### Options:

--version show program's version number and exit

-h, --help show this help message and exit

-i IPTF input file, default = phylokmer.dat(.gz)

-t NTHREADS number of threads to use, default = 1

-G MEMSIZE max memory to use (in GB), default = 1 -o OTPF prefix of the output files, default = aaf

-f COUNTF k-mer diversity file, default = kmer diversity.wc

## Detailed description of options:

-i IPTF: input file. The shared *k*-mer table generated from aaf\_phylokmer.py. The default file is phylokmer.dat(.gz)

- -t NTHREADS: number of threads to use. Depends on how many cores are available on your machine. Set at 1 by default.
- -G MEMSIZE: the total memory allowance in GB. Set at 4G by default.
- -o OTPF: prefix of the output files, including the distance matrix(.dist) and the phylogenetic tree(.tre). Default is set as "aaf".
- -f COUNTF: we file generated from kmer\_count. This file contains the *k*-mer diversity of each sample. The default is kmer\_diversity.wc

# 3) aaf\_tip.py

This script takes the phylogenetic tree generated from aaf\_distance.py and trims the excessive tips resulting from sequencing errors and incomplete coverage. Note that you need to prepare the tip setting file, which includes the coverage, read length and sequencing error of each sample before running the script.

Usage: aaf\_tip.py [options] -i <input tree file> -k <kmer size> --tip <information for tip correction>

### Options:

--version show program's version number and exit

-h, --help show this help message and exit

-i IPTF tree file to be trimmed

-k KLEN k-mer size used for constructing the input tree

--tip=TIP FILE tip setting file, default = tip file test.txt

-n k-mer filtering was on for tree construction

-f COUNTF k-mer diversity file, default = kmer diversity.wc

## Detailed description of options:

-i IPTF: input tree file. The tree file whose tips you would like to trim.

-k KLEN: the *k* that was used to construct the input tree.

--tip TIP\_FILE: To trim the excess tips caused by incomplete coverage and sequencing errors requires additional info on the coverage, read length and sequencing error of each sample. Put this information into a tab delimited text file in the format of tip\_info\_test.txt. See suggestions on estimation of coverage and sequencing error in Parameter Selection section.

-n: add it to the command if filter was used during the tree construction.

-f COUNTF: we file generated from kmer\_count. This file contains the *k*-mer diversity of each sample. The default is kmer\_diversity.wc

### 4) nonparametric\_bootstrap.py

This script takes the sequence data and reconstructs the phylogeny with bootstrap value for each node.

Usage: nonparametric bootstrap.py [options]

### Options:

-h, --help show this help message and exit

-k KLEN k-mer length, default = 25

-t NTHREADS number of threads to use, default = 1

-n FILTER k-mer filtering threshold, default = 1

-f SEQFORMAT format of input files, FA|FQ, default = FA

-o OUTFILE k-mer table name, default = phylokmer.dat.gz

-d DATADIR directory containing the data, default = data/

-G MEMSIZE total memory limit (in GB), default = 4

--S1=STAGE1 number of resampling of the reads, default = 0

--S2=STAGE2 number of resampling of each total kmer table, default = 0

Detailed description of options:

- -k KLEN: k-mer size. Set at 25 by default.
- -t NTHREADS: number of threads to use. Depends on how many cores are available on your machine. Set at 1 by default.
- -n FILTER: how many times a *k*-mer needs to be in the sample to be counted as present. Set at 1 by default.
- -f SEQFORMAT: Format of the sequence files, FA or FQ. The default is set as FA.
- -o OUTFILE: file name of the merged *k*-mer table. If you would like your *k*-mer table to be compressed, provide a name that ends with .gz. Otherwise it will not be compressed. The default output file is phylokmer.dat.gz, which is compressed.
- -d DATADIR: directory containing the data.
- -G MEMSIZE: the total memory allowance. Each kmer\_count thread has G/t memory allowance. Set at 4G by default.
- --S1: number of times to resample the reads for each sequence file. This is the first stage of our two-stage bootstrap. This bootstrap result shows the variance in sequencing error and incomplete coverage. Set at 0 by default, which means skip the first stage of bootstrap and only resample the *k*-mer table.
- --S2: number of times to resample the total *k*-mer table generated from one instance of resampling of the reads. If --S1 is set to be 0, the resampling is on the real *k*-mer table generated from the original data. Set at 0 by default, which means skipping this step.

## 5) parametric\_bootstrap.R

When it takes too long to bootstrap over large datasets, switch to the parametric bootstrap. This R script provides estimation of the variances in the two steps. It requires:

Info file: containing read length, sequencing error and coverage and used in aaf\_tip.py, default = tip info test.txt

nshare file: containing the number of shared kmers generated by aaf\_distance.py (ends with nshare.csv), default = test\_nshare.csv

nreadboot: number of replicates, default = 10

k: k-mer length used in previous steps, default = 21

i.filter: filter threshold used, default =1.

# **Parameter Selection**

Here we provide some guidelines in parameter selection using the dataset with 21 tropical trees as an example.

#### a. Optimal k

As we described in the manuscript, the selection of *k* is a trade-off between avoiding multiple

mutations on one k-mer (which favors shorter k) and decreasing the chances of k-mer homoplasy (which favors longer k). For the primate dataset in the manuscript, we plot the theoretical predictions of the proportion of shared k-mers, ph, calculated from the observed frequency distribution of k-mers and the ph calculated without homoplasy (Fig. 2D) to help view the effect of different choices of k. This procedure led to the selection of k that corresponded to an accurate phylogeny. Therefore this figure serves as a good indicator for optimal k, and this choice can be further proved by constructing phylogeny with k-mer lengths larger than optimal, in order to check the phylogenetic consistency.

To plot the *ph* vs. *k* figure for your dataset, here is a checklist for the genome information that is needed:

- i. Sample names
- ii. Coverage
- iii. Genome size
- iv. GC content
- v. d (genetic distance)
- vi.  $O_k$

We are aware that this information might not be all available, and we provide coarse calculation methods for some of the categories.

### ii. Coverage

There are multiple ways of estimating the sequencing coverage of your next-gen sequencing data. (1) If the genome size is known, coverage = total bp / genome size. (2) If the genome size is unknown, we can estimate the coverage by plotting the k-mer frequency distribution: "if a large fraction of k-mers occur c times, we can estimate the sequencing coverage to be approximately c and derive an estimate of the genome size from c and the total length of the reads." (Marcais and Kingsford 2011)). c will be the k-mer coverage. To get the base pair coverage, you need correct c using base\_coverage = c \* read\_length / (read\_length - k-mer\_size + 1) (see https://groups.google.com/forum/#!topic/bgi-soap/xKS39Nz4SCE). (3) When the coverage is low or sequencing error rate is high, there will be no clear peak in the k-mer frequency distribution at c. This is actually the case for all the tropical tree species in our dataset except  $Ficus\ vasculosa\ (FV)$ . A coarse estimation of the k-mer coverage will be the total number of k-mers (including multiple copies of the same k-mer) divided by k-mer diversity (number of k-mer that shows up at least once). Some assemblers (such as velvet, SOAPdenovo) report estimation of k-mer coverage as well.

Coverage information is also needed for tip correction.

#### iii. Genome size

You can try to check the genome size in databases such as Plant DNA C-values Database (<a href="http://data.kew.org/cvalues/">http://data.kew.org/cvalues/</a>) or Animal Genome Size Database (<a href="http://www.genomesize.com/">http://www.genomesize.com/</a>). If it is not available for your species, you can do a rough estimate using total base pair divided by coverage.

#### iv. GC content

There are many tools to calculate the GC content of your samples. In the AAF package we have provided our own, gc.py in the utils folder. Biopython needs to be preinstalled.

#### v. d (Genetic distance)

The genetic distance of the group (average number of mutations per base pair) is used to set the scale of the vertical axis in the ph vs. k figure. Because the figure is used to find k on the horizontal axis, the

conclusions from this figure about selecting k are mostly independent of the selection of d, so this selection does not need to be very accurate. A reasonable strategy is to guess d, or use the default 0.1, to select k. The subsequent phylogeny construction will give a good estimate of d from the distance matrix, which then could be used to plot the figure.

## vi. $Q_k$

There is more than one way of calculating the frequency distribution of k-mers. One of the easiest ways is to turn on the --stats option while counting the k-mers using jellyfish(Marcais and Kingsford 2011). However the maximum k that jellyfish can handle is 31. For k>31, use kmer\_countx to count the k-mers, then calculate the frequency distribution of k-mers from the pkdat files (the output file of kmer\_count( $k\le25$ ) and kmer\_countx(k>25)) using the pkdat2hist.py in the utils folder that is provided in the tutorial folder.

After gathering all the information, we generated the ph vs. k figure for the 21 tropical trees dataset (Fig. S5) using the R code phVSk.R in the utils folder. The trend for all the red lines (estimated ph based on the  $Q_k$  for each species) stabilized for  $k \ge 25$ , and the difference between the red lines and the black dashed line continued to decrease with larger k. Therefore, we constructed phylogenies for k from 25 to 31, and because the phylogenies were identical for  $k \ge 27$  (Fig.7), we selected 27 as the optimal k for the tropical trees dataset. The same phylogenetic topology was also obtained when k-mers were filtered to remove singletons. For k greater than 31, the topology within the *Ficus* group showed some small changes. We suspect that this is due to the loss of sensitivity to evolutionary changes when selecting k-mer lengths too long, especially for relatively small genomes (as the Ficus group has genome sizes less than half those of the other species).

To plot the ph vs. k figure for your dataset, simply replace the genome information for the tropical trees with the information for your own dataset in the beginning of the R script phVSk.R.

#### b. Filter or not

In deciding whether or not to filter *k*-mers (i.e., only including *k*-mers if they occur at least twice in a taxon), it is necessary to know the balance between loss of information through false *k*-mers that caused by sequencing errors if there is no filtering, and loss of information through removing true singletons if there is filtering (Fig. 5 in the manuscript). If there is a large range of coverages among taxa within a dataset, it is best to decide whether or not to filter based upon the taxon with the lowest coverage, because there is a large negative consequence of filtering low-coverage taxa (Fig. 5). For the tropical trees, we chose not to filter because more than half of the species have coverage less than 5.

# c. Tip trimming (optional)

Information needed for tip trimming/correction includes: coverage, read length, and sequencing error. You should be able to get the performance of your sequencing platform from your sequencing company. For example Illumina claims that the error rate for Genome Analyzer is about 1% (http://res.illumina.com/documents/products/datasheets/datasheet\_genomic\_sequence.pdf).

For formatting the required file, see tip\_info\_tt.txt in the tropical\_trees folder containing the tip information for the tropical trees.

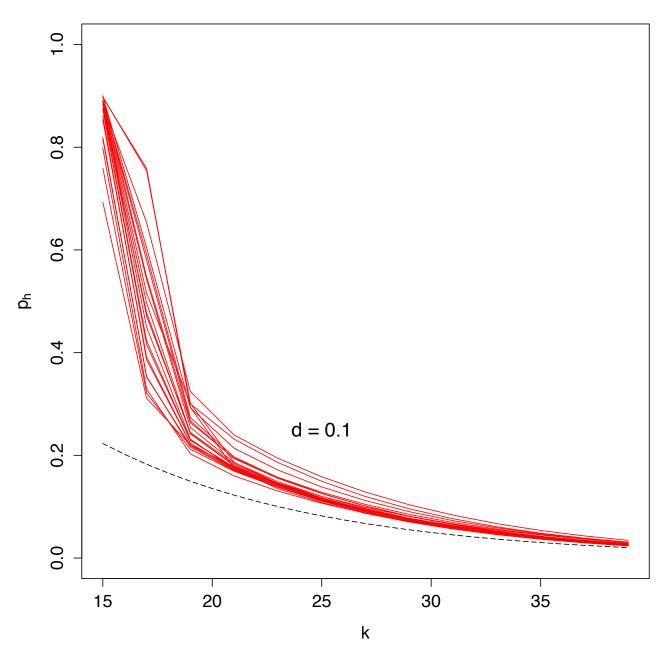
### d. Bootstrap

i. Nonparametric vs. parametric bootstraps

Nonparametric bootstrapping can be computationally intensive when the dataset is large (>100G in total). If you think it takes too long, switch to the parametric one. Also with large genomes, the bootstrap value tends to stay as 100%.

#### ii. Correction factor

Set to be 2. The correction factor in the R script is from Equation 11 that estimates the variation caused by sequencing error and incomplete coverage. Detailed simulations showed that the formula sometimes under-estimates and sometimes over-estimates the true standard deviation in the distance between taxa by as much as 50%. This occurs because of the complexities of accounting mathematically for the correlations among *k*-mers that occur on the same reads. The correction factor set to 2 provides a conservative bootstrap (i.e., one that is not going to improperly inflate the support for nodes), be insuring that the true variation caused by sequencing error and incomplete coverage is never underestimated.



**Figure S5.** Theoretical predictions of the proportion of shared k-mers,  $p_h$ , calculated from the observed

frequency distribution of k-mers,  $Q_k$ , for the tropical trees dataset ranging in genome size from 250M to 2Gbp assuming the true distance between taxa is d = 0.1 (divergence time 94Mya).

# **Reference:**

Marcais G, Kingsford C. 2011. A fast, lock-free approach for efficient parallel counting of occurrences of k-mers. *Bioinformatics* **27**: 764–770.