# CIAlign - Clean and Interpret Alignments

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CIAlign is a command line tool which performs various functions to clean and analyse a multiple sequence alignment (MSA).

The tool is designed to be highly customisable, allowing users to specify exactly which functions to run and which settings to use. It is also transparent, generating a clear log file and alignment markup showing exactly how the alignment has changed and what has been removed by which function.

This allows the user to:

- Remove sources of noise from their MSA
  - Remove insertions which are not present in the majority of sequences
  - Remove sequences below a threshold number of bases or amino acids
  - Crop poorly aligned sequence ends
  - Remove columns containing only gaps
  - Remove sequences above a threshold level percentage of divergence from the majority
- Generate consensus sequences
- Visualise alignments
  - Generate image files showing the alignment before and after using CIAlign cleaning functions and showing which columns and rows have been removed
  - Draw sequence logos
  - Visualise coverage at each position in the alignment
- Analyse alignment statistics
  - Generate a similarity matrix showing the percentage identity between each sequence pair
- Unalign the alignment
- Replace U's by T's

### Requirements

- python >= 3.6
- matplotlib >= 2.1.1
- numpy >= 1.16.3
- scipy >= 1.3.0

### Installation

The easiest way to install CIAlign is using pip3:

The current release of CIAlign can also be downloaded directly using this link,

If you download the package directly, you will also need to add the CIAlign directory to your PATH environment variable as described here

### Usage

### Basic Usage

CIAlign --infile INFILE --outfile\_stem STEM --inifile my\_config.ini

### Parameters

Parameters can be specified in the command line or in a config file using the naming system below.

A template config file is provided in CIAlign/templates/ini\_template.txt - edit this file and provide the path to the –inifile argument. If this argument is not provided command line arguments and defaults will be used.

Parameters passed in the command line will take precedence over config file parameters, which take precedence over defaults.

Command help can be accessed by typing CIAlign --help

Parameter	Description	Default
infile	Path to input alignment file in FASTA format	None
inifile	Path to config file	None
outfile_stem	Prefix for output files, including the path to the output directory	CIAlign
silent	Do not print progress to the screen	False
all	Use all available functions with default parameters	False
help	Show all available parameters with an explanation	None
version	Show the version	None

Beside these main parameters, the use of every function and corresponding thresholds can be specified by the user by adding parameters to the command line or by setting them in the configuration file. Available functions and their parameters will be specified in the following section.

CIAlign always produces a log file, specifying which functions have been run with witch parameters and what has been removed. It also outputs a file that only specifies what has been removed with the original column positions and the sequence names.

#### Output files:

- OUTFILE\_STEM\_log.txt general log file
- $\bullet$  <code>OUTFILE\_STEM\_removed.txt</code> removed columns positions and sequence names text file

# Cleaning an MSA

Each of these steps will be performed sequentially in the order specified in the table below.

The "cleaned" alignment after all steps have been performed will be saved as OUTFILE\_STEM\_cleaned.fasta

Parameter	Description	Default Value
remove_divergent	Remove sequences with $<=$ N proportion of positions at which the most common base $/$ amino acid in the alignment is present	False
remove_divergent_minperc	Minimum proportion of positions which should be identical to the most common base $/$ amino acid in order to be preserved	0.75
remove_insertions	Remove insertions found in $<=50\%$ of sequences from the alignment	False
insertion_min_size	Only remove insertions >= this number of residues	3

Parameter	Description	Default Value
insertion_max_size	Only remove insertions <= this number of residues	100
$insertion\_min\_flank$	Minimum number of bases on either side of an insertion to classify it as an insertion	5
crop_ends	Crop the ends of sequences if they are poorly aligned	False
crop_ends_mingap_perc	Minimum proportion of the sequence length (excluding gaps) that is the threshold for change in gap numbers.	0.05
crop_ends_redefine_perc	Proportion of the sequence length (excluding gaps) that is being checked for change in gap numbers to redefine start/end.	0.1
remove_short	Remove sequences $<=$ N bases $/$ amino acids from the alignment	False
remove_min_length	Sequences are removed if they are shorter than this minimum length, excluding gaps.	50
keep_gaponly	Keep gap only columns in the alignment	True

# Generating a Consensus Sequence

This step generates a consensus sequence based on the cleaned alignment. If no cleaning functions are performed, the consensus will be based on the input alignment. For the "majority" based consensus sequences, where the two most frequent characters are equally common a random character is selected.

### Output files:

- OUTFILE\_STEM\_consensus.fasta the consensus sequence only
- OUTFILE\_STEM\_with\_consensus.fasta the cleaned alignment plus the consensus

Parameter	Description	Default
make_consensus	Make a consensus sequence based on the cleaned alignment	False
consensus_type	Type of consensus sequence to make - can be majority, to use the most common character at each position in the consensus, even if this is a gap, or majority_nongap, to use the most common non-gap character at each position	majority
consensus_keep_gaps	If there are gaps in the consensus (if majority_nongap is used as consensus_type), should these be included in the consensus (True) or should this position in the consensus be deleted (False)	False
consensus_name	Name to use for the consensus sequence in the output fasta file	consensus

# Unaligning the Alignment

This function simply removes the gaps from the input or output alignment and creates and unaligned file of the sequences. Output files:

- OUTFILE\_STEM\_unaligned\_input.fasta unaligned sequences of input alignment
- OUTFILE\_STEM\_unaligned\_output.fasta unaligned sequences of output alignment

Parameter	Description	Default
unalign_input	Generates a copy of the input alignment with no gaps	False

Parameter	Description	Default
unalign_output	Generates a copy of the output alignment with no gaps	False

# Replacing U's by T's

This function replaces the U nucleotides by T nucleotides without disturbing the sequence names.

### Output files:

- OUTFILE\_STEM\_T\_input.fasta input alignment with T's instead of U's
- OUTFILE\_STEM\_T\_output.fasta output alignment with T's instead of U's

Parameter	Description	Default
replace_input	Generates a copy of the input alignment with T's instead of U's	False
replace_output	Generates a copy of the output alignment with T's instead of U's	False

# Visualising Alignments

Each of these functions produces some kind of visualisation of your alignment.

# Mini Alignments

These functions produce "mini alignments" - images showing a small representation of your whole alignment, so that gaps and poorly aligned regions are clearly visible.

### Output files:

- OUTFILE\_STEM\_input.png (or svg, tiff, jpg) the input alignment
- OUTFILE\_STEM\_output.png (or svg, tiff, jpg) the cleaned output alignment
- OUTFILE\_STEM\_markup.png (or svg, tiff, jpg) the input alignment with deleted rows and columns marked

Parameter	Description	Default
plot_input	Plot a mini alignment - an image representing the input alignment	False
plot_output	Plot a mini alignment - an image representing the output alignment	False
plot_markup	Draws the input alignment but with the columns and rows which have been removed by each function marked up in corresponding colours	False
$-plot_dpi$	DPI for mini alignments	300
plot_format	Image format for mini alignments - can be png, svg, tiff or jpg	png
$-plot\_width$	Mini alignment width in inches	5
plot_height	Mini alignment height in inches	3

### Sequence logos

These functions draw sequence logos representing your output (cleaned) alignment. If no cleaning functions are specified, the logo will be based on your input alignment.

 $Output\_files:$ 

- OUTFILE\_STEM\_logo\_bar.png (or svg, tiff, jpg) the alignment represented as a bar chart
- OUTFILE\_STEM\_logo\_text.png (or svg, tiff, jpg) the alignment represented as a standard sequence logo using text

Parameter	Description	Default
make_sequence_logo	Draw a sequence logo	False
sequence_logo_type	Type of sequence logo - $bar/text/both$	bar
sequence_logo_dpi	DPI for sequence logo	300
sequence_logo_font	Font (see NB below) for bases $/$ amino acids in a text based sequence logo	monospace
sequence_logo_nt_per_row	Number of bases / amino acids to show per row in the sequence logo, where the logo is too large to show on a single line	50
sequence_logo_filetype	Image file type to use for the sequence logo - can be png, svg, tiff or jpg	png

NB: to see available fonts on your system, run CIAlign —list\_fonts\_only and view CIAlign\_fonts.png

### Coverage Plots

This function plots the number of non-gap residues at each position in the alignment.

Output file:

- OUTFILE\_STEM\_input\_coverage.png (or svg, tiff, jpg) image showing the input alignment coverage
- OUTFILE\_STEM\_output\_coverage.png (or svg, tiff, jpg) image showing the output alignment coverage

Parameter	Description	Default
plot_coverage_input	Plot the coverage of the input MSA	False
plot_coverage_output	Plot the coverage of the output MSA	False
$plot\_coverage\_dpi$	DPI for coverage plot	300
$-plot\_coverage\_height$	Height for coverage plot (inches)	3
$-plot\_coverage\_width$	Width for coverage plot (inches)	5
plot_coverage_colour	Colour for coverage plot (hex code or name)	$\#007\mathrm{bf5}$
plot_coverage_filetype	File type for coverage plot (png, svg, tiff, jpg)	png

# **Analysing Alignment Statistics**

These functions provide additional analyses you may wish to perform on your alignment.

### Similarity Matrices

Generates a matrix showing the proportion of identical bases / amino acids between each pair of sequences in the MSA. Output file:

- OUTFILE\_STEM\_input\_similarity.tsv similarity matrix for the input file
- OUTFILE\_STEM\_output\_similarity.tsv similarity matrix for the output file

Parameter	Description	Default
make_similarity_matrix_input	Make a similarity matrix for the input alignment	False
make_similarity_matrix_output	Make a similarity matrix for the output alignment	False
make_simmatrix_keepgaps	Include positions with gaps in either or both sequences in the similarity calculation	False
make_simmatrix_dp	Number of decimal places to display in the similarity matrix output file	4
make_simmatrix_minoverlap	Minimum overlap between two sequences to have non-zero similarity in the similarity matrix	1