# ConCat v-10 Manual

# Alignment concatenation and analysis utility

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# **Contents**

1.Introduction	3
1.1. Input/Output	4
1.2. ConCat Block	
2.Usage/Options	5
2.1. ConCat-import	
2.1.1. ConCat-import Usage	
2.1.2. Import sequence from NCBI	
2.1.2. Import sequence from NGBI	
2.1.4. Import set of CDS/mRNA sequence for specific taxa to create gene alignment	
2.1.4. Import set of CD3/IndvA sequence for specific taxa to create gene anginnenc	
2.2.1. –ftype and –otype	
2.2.2. –convert	
2.2.3. –spell	
2.2.4block	
2.2.5. –RNA	
2.2.6. –pipe	
2.2.7. –shannon	
2.2.8rcv	
2.2.9. –OV	
2.2.10GC	
2.2.11RY	
2.2.12addT and -remT	12
2.2.13inc and -exc	
2.2.14. –pbin	13
2.2.15rbin, -ebin, -gcbin	
2.2.16. –ugcbin	13
2.3. ConCat-analyze Options	13
2.3.1fevol	
2.3.2rembin	14
233 _ugchin	12

3.Liscence/Help Desk/Citations	15
4.Copyright	15

## 1. Introduction

ConCat is a biopython based alignment concatenation utility designed to obtain alignment super matrix from list of alignment files. It is divided into three modules (ConCat-import, ConCat-build, ConCat-analyze). ConCat-import module provides functions for CDS and mRNA sequence import from NCBI and dynamic sequence alignment options with multiple input/output format options. ConCat-build module is for concatenation, annotation handling and storage. ConCat-analyze module conducts post analysis of super matrix obtained from ConCat-build module. Workflow of ConCat is shown below:

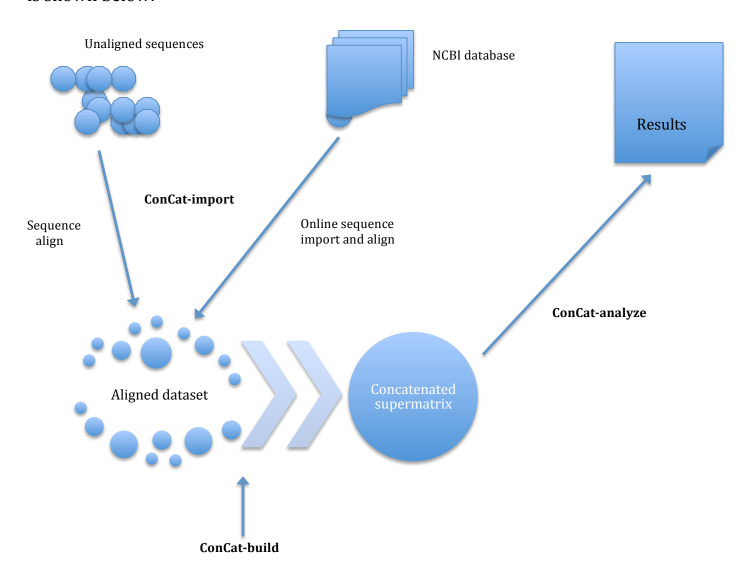


Fig 1. ConCat Workflow

ConCat produces rich character annotation obtained by user-selected functions applied during concatenation process. The program can handle input file in fasta, phylip-interleaved, phylip-relaxed and nexus format. ConCat has several alignment processing functions that helps user to create and store publication ready files. It extracts genome IDs (if available) from the taxon IDs and stores it in an excel sheet in a publication ready format. Furthermore, it scans for the spelling mistakes in the taxon name within the alignment files, identifies missing taxa in each alignment and stores these information in nexus output as Taxset data. Moreover, it allows user to select or reject specific taxa while performing concatenation, edit taxa name by adding or removing user supplied nomenclature group and creates useful bins by classifying different input alignment file datasets according to user defined criterion.

# 1.1. Input/Output

Creating, handling and storing annotation is one of the most important features of ConCat, which is why the nexus is a preferred input format in ConCat. Although, ConCat allows multiple input/output formats including fasta (.fas), nexus (.nex), phylip-interleaved (.phy) and phylip-relaxed (.phy). ConCat has "-CA" and "-convert" arguments to handle non-nexus input files. These arguments allows user to first convert non-nexus input files to nexus format and then run the analysis or select to convert and run the analysis simultaneously. ConCat also allows user to select fasta, nexus, phylip-interleaved and phylip-relaxed output formats.

### 1.2. ConCat block

ConCat allows users to define the alignment file type, define files to run RNA structure mapping and supply user defined RNA structure through ConCat block option. Structure of ConCat block is shown below:

```
begin ConCat;
    Ali_type = DNA;
    RNA_type = False;
    RNA_Struc = Species_Name, (((..(...))....).),6;
end;
```

ConCat block can be defined in nexus alignment input file after the first line that contains #NEXUS. This block has three variables and each carries special set of information.

**Ali\_type** variable takes alignment type information from user. User can allow ConCat to create RaxML partition file by defining the alignment type in ConCat block. ConCat takes the user defined alignment type from each file and extract their corresponding nucleotide positions from the concatenated super matrix to create a final RaxML partition file. This variable also allows ConCat to distinguish between DNA and amino acid alignment data while calculating alignment entropy and RCV values.

**RNA\_type** variable allows user to label the alignment files for which ConCat is supposed to create RNA structure data and map it over the concatenated alignment matrix. ConCat uses RNAfold package to obtain the RNA structure data from the consensus sequence obtained from the input alignment file. Program will skip this step if RNAfold program is not installed on the users system.

**RNA\_Struc** variable allows user to supply predefined RNA structure to map over the concatenated super matrix data. It contains three sub-variables Species\_name, structure and structure starting position. So if user has following alignment file:

Homo\_sapiens ACTAGATACAGATACGATCAGATCA Gorilla\_gorilla ACTAGATAGAGAAACGATCAGATCA Macaca\_mulata ACTAGATACAGAAACGAACCGCTCA

And Homo\_sapiens RNA structure "(((..(...))...)" starting at position 5<sup>th</sup> then the ideal way of RNA\_Struc variable representation is

RNA\_Struc = Homo\_sapiens, (((..(...))....).), 5;

# 2. Usage/Options

# 2.1. ConCat-import

ConCat-import module allows user to import multi-taxa sequences from NCBI database and provides dynamic options to perform sequence alignment.

# 2.1.1. ConCat-import usage

Place all the unaligned sequence files in Align directory and run

python ConCat-import.py

Muscle is set as default alignment package. User can choose to run Mafft by using –pkg argument and supply alignment argument through –args argument.

```
python ConCat-import.py -pkg mafft -args "--localpair --maxiterate 1000"
```

ConCat-import module also allows user to supply distinct mafft arguments for each sequence file through –argf argument while conducting batch alignment. Arguments can be supplied by via authority text file. File format is shown below.

```
File1.fas = --localpair --maxiterate 1000
File2.fas = --globalpair --maxiterate 1000
File3.fas = --retree 2 --maxiterate 2
```

Now run:

Python ConCat-import.py -pkg mafft -argf authority.txt

ConCat-import usage descriptions are given below:

```
-cds CDS
                     Takes gene name via text file for CDS import
-mrna MRNA
                     Takes gene name via text file for mRNA import
-orgn ORGN
                     Takes organism group name to extract sequence data
-pkg {muscle,mafft}
                     Select alignment program
                     Arguments to run MAFFT. EXAMPLE: "--retree 2
-args ARGS
                     --maxiterate 10"
                     Include if you want to run different models for
-sep
                     different alignment files
-argf ARGF
                     Takes argument file as input
```

# 2.1.2. Import sequences from NCBI

CDS and mRNA sequences can be directly fetched from NCBI through –cds and –mrna arguments. To obtain CDS sequence of Eutherian BRCA1, CENPJ and BRCA2 gene use

python ConCat-import.py –cds authority.txt –orgn Eutheria

authority.txt file contains list of genes in following format

BRCA1 CENPJ BRCA2

Each gene name should be in a new line.

This function imports longest available CDS sequence for all the Eutherian mammals from NCBI database, performs coding sequence alignment using muscle alignment program and saves it in ConCat Input directory. ConCat scans for out-frame indels, automatically identifies the correct coding frame for the given sequence and adjusts it accordingly before performing alignment. Use the following argument to import mRNA sequences:

python ConCat-import.py -mrna authority.txt -orgn Eutheria

Sometimes we face annotation issues while extracting sequences from NCBI database. For example if we intent to fetch CEP63 CDS/mRNA sequence for Eutherian mammals by using above step, we might end up skipping Loxodonta africana and Ailuropoda melanoleuca since their CEP63 gene sequence is stored with the id name LOC100669045 and LOC100464770 respectively. To handle this situation ConCat allows user to import vertebrate sequences using orthologue method.

python ConCat-import.py -cds authority.txt -ortho Homo\_sapiens

This argument will fetch CDS sequence for all the vertebrates that has the respective gene sequence annotated in NCBI by performing orthologue search using Human gene id. Same method can be used to extract mRNA sequence.

## 2.1.3. Import all CDS/mRNA sequence for a set of taxa

This functionality can be used in a situation where all the coding sequences encoded in a group of organism are required.

python ConCat-import.py -pull authority.txt

"authority.txt" file contains list of all the taxa to be for CDS import. Each taxon should be placed in a newline:

Homo sapiens Gorilla gorilla Felis catus

This will produce three files with the name "Homo\_sapiens.fas", "Gorilla\_gorilla.fas" and "Felis\_catus.fas" as an output having all their corresponding NCBI annotated CDS sequences.

# 2.1.4. Import set of CDS/mRNA sequence for specific taxa to create gene alignment

python ConCat-import.py –gcds authority.txt –orgn Homo\_sapiens & python ConCat-import.py –gmrna authority.txt –orgn Homo\_sapiens

These arguments will fetch Homo sapiens CDS/mRNA sequences for a set of genes supplied via authority.txt file and creates a gene CDS/mRNA alignment as an output using muscle package.

For example, if your authority.txt file contains following gene ids:

Gene1

Gene2

Gene3

and run

python ConCat-import.py -gcds authority.txt -orgn Homo\_sapiens

Then the program will output an alignment:

Gene1 ATGGTGACT......

Gene2 ATG---ACT......

Gene3 ATGGTG---.....

#### 2.2. ConCat-build

Steps to run ConCat via command line:

- a) Open terminal and move to the ConCat directory.
- b) Store your input files in the "Input" directory
- c) If input files are in nexus format the run

python ConCat.py

or else run

python ConCat.py -CA -ftype phylip-relaxed -otype phylip-relaxed

for phylip-relaxed input and output files. Select –ftype and –otype argument inputs accordingly.

### 2.3. ConCat-build options

#### 2.3.1. –ftype, –otype and –CA

-ftype and -otype arguments are used for passing input and output file type respectively. Bothe arguments allow user to select from 5 options (fasta, nexus, phylip, phylip-interleaved, phylip-relaxed). If a user has input files in fasta format and requires output super matrix file in phylip-relaxed format, then the typical command line operation will be

python ConCat-build.py -CA -ftype fasta -otype phylip-relaxed

-CA argument first converts all the –ftype file format files to nexus file format, which is then imported by ConCat-build module. Output is produced in nexus as well as in – otype file format.

ConCat uses nexus as default input format when the -ftype argument is not supplied.

#### 2.3.2. –convert

-convert argument allows user to convert between different file formats.

python ConCat-build.py -convert -ftype fasta -otype nexus

# 2.3.3. –spell

Spelling errors are common in taxon naming. ConCat-build –spell argument checks for possible spelling mistakes by comparing the taxon name in the input alignment files.

#### 2.3.4. -block

By passing –block argument user tells ConCat-build module to check ConCat block in all the alignment files. ConCat-build creates RaxML partition file if Ali\_type variable is intiated in the ConCat block in the input alignment files.

-block argument is important when user wants to create partition file, supply RNA structure or create and map RNA structure on the final super matrix data. Therefore, it

is important to initiate –block argument when defining ConCat block in the input alignment file or using –RNA argument.

Example:

python ConCat-build.py -CA -ftype fasta -otype phylip-relaxed -block -RNA

#### 2.3.5. -RNA

-RNA argument can be used to create and map RNA structures from the input alignment file. ConCat uses RNAfold program to generate RNA structure for the input files that has ConCat block RNA\_type variable set as True. Program checks for the RNAfold program on the users system and skips this step if the RNAfold program is not installed.

#### 2.3.6. –pipe

-pipe argument is very useful when user has database IDs attached with the taxa name separated by pipes.

#### Example:

Cricetulus\_griseus|NM\_001246726.1 Nannospalax\_galili|XM\_008839441.1 Microtus\_ochrogaster|XM\_005368436.1 Rattus\_norvegicus|NM\_207592.1

-pipe argument allows ConCat-build module to extract these IDs and store it as a Taxset in nexus output file. It also creates a publication ready database ID excel sheet as output.

#### 2.3.7. -shannon

-shannon argument allows user to obtain an estimate of variability within each alignment input files. ConCat has separate modules to calculate Shannon entropy for DNA and protein alignments.

For protein alignment, ConCat first divides each amino acid into 12 unique groups on the basis of their chemical similarity.

```
'u': ['D', 'E']
'b': ['R', 'K']
'i': ['I', 'V']
'l': ['L', 'M']
'f': ['F', 'W', 'Y']
'n': ['N', 'Q']
```

For DNA sequence A, C, G and T forms their own group. Gaps (-) are considered as a separate group in entropy calculation. Finally the entropy values for a particular alignment position is calculated by using following equation:

$$\xi = -\sum_{i=1}^{n} p_i \log p_i$$

Where  $p_i$  is the fraction of residue of amino acid group type and n is the number of amino acid group types. These values are averaged over the alignment length to obtain overall entropy.

#### 2.3.8. -rcv

Relative Composition Variability (RCV) value is a good indication of alignment quality. Similar to Entropy calculation, ConCat has separate modules for amino acid and DNA alignment RCV calculations. Similar to entropy calculation function, RCV calculation for amino acid alignment is performed by grouping amino acids into 12 groups on the basis of their chemical composition. A, C, G and T are the only groups taken into account while performing RCV calculation for DNA alignment.

$$RCV = \sum_{i=1}^{n} (|A_i - A^*| + |T_i - T^*| + |C_i - C^*| + |G_i - G^*|) / n \times t$$

where  $A_i$ ,  $T_i$ ,  $C_i$ , and  $G_i$  are the numbers of each nucleotide for the  $i^{th}$  taxon.  $A^*$ ,  $T^*$ ,  $C^*$ , and  $G^*$  are averages across the n taxa, and t is the number of sites. Constant sites were excluded for all  $\chi^2$  and RCV calculations.

#### 2.3.9. -OV

To calculate observed variability (OV), all sequences for a given position are compared in a pair-wise fashion. Mismatches are scored as 1 and matches as 0; the mean value amongst all the comparisons for a given position is used as the measure of character variability in the subsequent data sorting:

$$OV = \sum_{p=1}^{k} d_{ij} / p$$

Here k is the number of pair-wise comparisons made for a given position and  $d_{ij}$  is the score of character variability in each pair-wise comparison made (can be either 0 or 1). If n is the number of aligned sequences which do not have a gap at the given alignment position, then  $k = (n^2 - n)/2$ .

#### 2.3.10. -GC

This argument calculates GC content of each input alignment file.

#### 2.3.11. -RY

-RY argument asks user for filenames to conduct RY coding either on all the coding positions or for 3<sup>rd</sup> coding position. File names can be supplied though a text file having one file name in each line and RY coding position separated by commas.

ASPM.nex, 3 CENPJ.nex, all WDR62.nex, all

ConCat-build extracts these filenames and selects the RY coding position by matching it with the concatenated supermatrix nexus Charsets. The program performes RY coding for 3<sup>rd</sup> position if the position supplied by user is "3", whereas it performes RY coding for all position if the position supplied by user is "all".

#### 2.3.12. -addT and -remT

-addT and -remT arguments are used for editing taxa name while performing concatenation. -addT takes taxon genus, class, order, phylum and kingdom names as input from Taxanomy.csv file. Use -addT Phyum to add phylum name to an existing taxa name.

#### python ConCat-build.py -addT Phylum

This will extract Phylum names of each taxon from Taxanomy.scv file and add it to the corresponding taxon names. Use –remT to remove the last segment of taxa name from the final concatenated super matrix object.

python ConCat-build.py -remT

#### 2.3.13. -inc and -exc

-inc and -exc arguments allows user to select specific set of taxa from input alignment files to perform concatenation. These two arguments take input from and authority file. -inc argument limits the concatenation process for set of taxa supplied by the user via authority file whereas -exc argument limits the concatenation process for set of taxa which are absent in the authority file.

#### 2.3.14. -pbin

-pbin argument invokes a function that creates multiple 0-25, 25-75 and 75-100 percentile bins (defined as ConCat\_Bin) by characterizing the input alignment data on the basis of their corresponding RCV, GC and Entropy values.

This will generate 0-25, 25-75 and 75-100 percentile bins each for the data characterized by their corresponding RCV, GC and entropy values.

#### 2.3.15. -rbin, -ebin, -gcbin

These binning functions takes bin range from user as input and creates one bin of each argument type for the range supplied by user.

python ConCat-build.py -rcv -GC -shannon -rbin 0.1-0.3 -ebin 0.2-0.4 -gcbin 35-50

# 2.3.16. -ugcbin

-ugcbin argument allows user to set values for bin partitioning. If user enters –ugcbin 20 then the program creates and populates 5 bins with range 0-20, 20-40, 40-60, 60-80 and 80-100, whereas if 15 is supplied then the program forms 7 bins with range 0-15, 15-30, 30-45, 45-60, 60-75, 75-90, 90-100. –pbin argument is required to run – ugcbin.

python ConCat-build.py -GC -pbin -ugcbin 15

# 2.3 ConCat-analyze

ConCat-analyze takes nexus as input file format.

2.3.1. -fevol

-fevol argument allows user to eliminate fast evolving sites from the alignment either by manual selection of sites or by conduction fast evolving site search available in ConCat program.

Using ConCat-analyze to detect fast evolving sites

python ConCat-analyze.py -i Combined.nex -fevol -OV 0.9 -o out.phy -fout phylip-relaxed

Here –OV argument is used to define OV cutoff threshold to detect fast evolving sites. – i argument is for giving input alignment file and –o is to define output file name. Output file formats can be defined through -fout argument (Nexus is set as default).

#### 2.3.2. -- rembin

--rembin argument tells ConCat to read ConCat bins from input alignment file and initiate bin selection based alignment editing module. User can select to remove alignment regions in 0-25<sup>th</sup>, 25<sup>th</sup> -75<sup>th</sup> or 75<sup>th</sup> -100 percentile RCV, GC or/and Entropy bin range. To remove alignment regions those are in 25<sup>th</sup> -75<sup>th</sup> percentile RCV and GC bin range:

python ConCat-analyze.py -i Combined.nex --rembin -RCVrem 25-75 -GCrem 25-75 -o out.nex

--rembin argument is import to initiate -RCVrem, -GCrem and -ENTrem (Selection based on entropy bins) module.

### 2.3.3. -ugcbin

-ugcbin is a --rembin independent argument that allows user to input any range of GC values in order to remove the corresponding alignment regions from the input alignment file.

# 3. Liscence/Help Desk/Citations

ConCat v1.0 was developed by Ambuj Kumar in 2014. It is implemented in python and available at <a href="https://github.com/Ambuj-UF/ConCat-1.0">https://github.com/Ambuj-UF/ConCat-1.0</a>. It can be distributed and/or modified under the terms of the GNU General Public License as published by the Free Software Foundation; either 2 of the license, or (at your option) any later version. This program is distributed with the hope that it will be useful, but WITHOUT ANY WARRANTY; without even the implied warranty of MERCHANTABILITY or FITNESS FOR A PARTICULAR PURPOSE. See the GNU General Public License for more details. You should have received a copy of the GNU General Public License along with this program; if not, write to the Free Software Foundation, Inc., 675 Mass Ave, Cambridge, MA 02139, USA.

If you have any problems, error-reports or other questions about ConCat, feel free and write an email to <a href="mailto:ambuj@ufl.edu">ambuj@ufl.edu</a>.

Reference: Manuscript under preparation...

# 4. Copyright

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