



**Manual**

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**Citations:**

**Fedosov A.E.**, Achaz G., Puillandre N. 2019. Revisiting use of DNA characters in taxonomy with Mold - a tree independent algorithm to retrieve diagnostic nucleotide characters from monolocus datasets. *BioRxiv*. DOI: 10.1101/838151.

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## Operating systems

Mac OSX, Windows, and Linux are supported

## System requirements

Python 2.7 installed

## Input

### DATAFILE

The input file is in the text (or any compatible) format, in which each line corresponds to one sequenced specimen and contains three space-separated records:

1. species name
2. focus taxon name
3. nucleotide sequence (with all the sequences aligned across the data set).

The names of the taxa to be diagnosed correspond to the **second** column.

### EXAMPLE

Genera of the family Conidae (Gastropoda) – Puillandre et al. 2014

```
Conasprella_alisi Conasprella TATAAGATTTTGGCTTTTACCTCCTGCCCTTCTTTTACTCCTTTCTTCAGCT
Conasprella_alisi Conasprella TATAAGATTTTGGCTTTTACCTCCTGCCCTTCTTTTACTCCTTTCTTCAGCT
Conasprella_alisi Conasprella TATAAGATTTTGGCTTTTACCTCCTGCTCTTCTTTTACTCCTTTCTTCAGCT
Conasprella_alisi Conasprella TATAAGATTTTGGCTTTTACCTCCTGCCCTTCTTTTACTCCTTTCTTCAGCT
Conasprella_alisi Conasprella TATAAGATTTTGGCTTTTACCTCCTGCCCTTCTTTTACTCCTTTCTTCAGCT
Conasprella_baileyi Conasprella TATAAGATTTTGACTTTTGCCCTCCGGCCCTTCTTTTACTTCTTTCTTCAGCC
Conasprella_baileyi Conasprella TATAAGATTTTGACTTTTGCCCTCCGGCCCTTCTTTTACTTCTTTCTTCAGCC
Conasprella_boholensis Conasprella TATAAGATTTTGACTTTTACCTCCTGCGCTTCTTTTACTTCTTTCTTCAGCT
Conasprella_boucheti Conasprella TATAAGATTTTGACTTTTACCTCCCGCACTTCTTTTACTTCTTTCTTCAGCT
Conasprella_comatosa Conasprella TATAAGATTTTGACTTTTACCTCCTGCGTTGCTTCTACTCTTATCTTCAGCT
Conasprella_coriolisi Conasprella TATAAGATTTTGACTTTTACCCCTGCGTTGCTTCTACTCCTATCTTCAGCT
Conus_adamsonii Conus TATGAGTTTTTGGCTTCTTCTCCTGCGCTTTTACTCCTTCTGTCTTCGGCT
Conus_variegatus Conus TATAAGTTTCTGGCTTCTTCTCCTGCACTTTTACTTCTTTTATCATCAGCT
Conus_anemone Conus TATAAGTTTCTGGCTTCTTCTCCTGCTTTGTTGCTTCTTCTATCGTCTGCT
Lilliconus_sagei Lilliconus TATAAGCTTCTGACTTTTACCTCCTGCTTTATTACTTTTATTGCTTCTGCT
Lilliconus_sagei Lilliconus TATAAGCTTCTGACTTTTACCTCCTGCTTTATTACTTTTATTGCTTCTGCT
Profundiconus_barazeri Profundiconus CATGAGTTTGTGATTATTACCTCCTGCTTTATTACTTTTGTATCATCAGCT
Profundiconus_maribelae Profundiconus TATAAGCTTTTGATTATTACCTCCTGCTTTATTACTTTTATTATCATCAGCT
Profundiconus_loyalti Profundiconus TATAAGTTTCTGGTTATTACCTCCTGCTTTATTGCTTTTATTATCCTCAGCT
Profundiconus_neocaledonicus Profundiconus CATAAGCTTTTGACTATTACCTCCTGCTTTATTACTTTTATTATCATCAGCT
Profundiconus_neocaledonicus Profundiconus CATAAGCTTTTGACTATTACCTCCTGCTTTATTACTTTTATTATCATCAGCT
Profundiconus_neocaledonicus Profundiconus CATAAGCTTTTGACTATTACCTCCTGCTTTATTACTTTTATTATCATCAGCT
Pygmaeonus_traillii Pygmaeonus TATAAGTTTCTGGCTTTTACCTCCTTCTCTTTTATTGCTTTTAGCATCTGCT
Californiconus_californicus Californiconus TATAAGCTTTTGACTTTTACCCCTGCTTTGTTATTACTTCTATCATCAGCT
```

### PARAMETERS FILE

Contains all the parameters that should be provided after '=' without a space.

### 1. INPUT / OUTPUT

-INPUT\_FILE – input alignment file with complete path.  
 -OUTPUT\_FILE – name of the output file with complete path

### 2. TAXON PARAMETERS (NO DEFAULTS - no parameters entered will lead to an error).

#### qTAXA (**Focus taxa**)

Arguments:

[ Taxon1 , Taxon2 , Taxon3 ]    A comma separated list of taxa to be diagnosed in square brackets and without spaces.  
 ALL                                if all taxa in the dataset are to be diagnosed.  
 >N                                if all taxa with more than N sequences available (where N is a natural number) to be diagnosed.

#### Taxon\_rank (**Taxon rank**):

Arguments:

1                                for species  
 2                                for supraspecific taxa

### 3. ADVANCED PARAMETERS FOR pDNC RECOVERY

For explanation see 'Review of MolD' below or Fedosov *et al.* 2019. If you don't want to set them, don't enter anything after '=', and the defaults will be used.

Cutoff                                Number of the informative positions to be considered, integer, (default **100**).  
 NumberN                                Number of ambiguously called nucleotides allowed, integer (default **5**).  
 Number\_of\_iterations                Number recursions of MolD, integer (default **10000**).  
 MaxLen1                                Maximum length of the raw pDNCs, integer (default **12**).  
 MaxLen2                                Maximum length of the refined pDNCs, integer (default **7**).

### 4. PARAMETERS OF ARTIFICIAL DATASETS (only sDNSs).

Pdiff                                Percent difference between original and modified sequence, integer (default **1** for species-level taxa, **3** for for supraspecific taxa).  
 PrSeq                                Proportion of sequences in the dataset to be modified, integer (default **0.5** or **0.25** for species-level, depending on the number of input records, and **0.1** for supraspecific taxa).  
 NmaxSeq                                Max number of sequences per taxon to modify. Integer (default 20)  
  
 Scoring                                Sets threshold of sDNC robustness scoring (default stringent). 100 artificial datasets are created to score the sDNC. If the sDNC remains diagnostic in requested (defined by value of threshold), or higher number of artificial datasets in **TWO** consecutive runs, the sDNC is output.

Arguments:

lousy                                66  
 moderate                                75  
 stringent                                90  
 very\_stringent                        95

## Run

Run in terminal:

```
python /PATH_TO/MolD_sDNC.py -i /PATH_TO/MolD_parameters
```

## Review of the MoID

(For term definition and theoretical background see:

**Fedosov A.E.**, Achaz G., Puillandre N. 2019. Revisiting use of DNA characters in taxonomy with MoID - a tree independent algorithm to retrieve diagnostic nucleotide characters from monolocus datasets. *BioRxiv*. DOI: 10.1101/838151 )

The MoID algorithm is divided into five consecutive steps. At **first** step sequences are sorted by taxon (as defined by the column 2 of the input) and the positions conserved within each taxon are identified.

At the **second** step, each of the positions shared by all focus taxon sequences, is assigned a **cut-off** value, which corresponds to the number of sequences in the alignment with different (from the focus taxon) nucleotide at this position. The positions that are conserved across the entire data set have a minimum cut-off value of 0 (i.e. non-informative). The positions that correspond to type 1 characters (see Fedosov et al. 2019, Fig. 1) and allow to immediately diagnose the focus taxon, have a maximum cut-off value (equal to the total number of non-focus-taxa sequences in the data set). The output of the second step, **CPP** is a subset of taxon shared positions with highest cut-off values. The length of this subset (parameter **cutoff**) is by default set to 100.

The third step contains main functionality of the MoID algorithm implemented in two recursive functions. The **step\_reduction\_complist** function initiates a raw DNC, and picks up positions from CPP one-by-one in random order. For each picked position the non-focus taxa sequences that differ at this position from the focus taxon sequences are identified and excluded from further comparisons, and the picked position is appended to the raw DNC. Thus, the set of non-focus taxa sequences is reduced step-by-step until its length equals zero, a condition at which the function terminates, and the raw DNC is output. The output raw DNC allows unambiguous differentiation of the focus taxon members in an analyzed data set, but it is usually redundant (i.e. it includes more positions than minimum necessary). The function

**step\_reduction\_complist** is run repeatedly, where the parameter **Number\_of\_iterations** (default 10,000) defines the number of iterations that generate a pool of raw DNCs. The 500 shortest non-identical raw DNCs, each comprising no more than a predefined number of nucleotide positions (parameter **Maxlen1**, default 12), make up input of the **RemoveRedundantPositions** function. The latter function removes redundant positions from the raw DNCs by picking and discarding positions in each DNC one-by-one, and each time checking, whether the thus shortened combination remains diagnostic for the focus taxon. The non-identical refined DNCs are retained, if their length is equal to or less than a pre-defined threshold value (parameter **Maxlen2**, default 7). Each of the refined DNCs therefore defines a minimal and sufficient condition that a nucleotide sequence (and a corresponding specimen) belongs to the focus taxon (pDNCs). The steps 3 is executed 5 times to overcome random effect, and to ensure thorough sampling of the informative positions.

Two pDNCs may overlap by one or several nucleotide positions, or share no positions; in the latter case the two pDNCs are termed independent pDNCs (see Fedosov et al. 2019). In the case that all identified pDNCs share one or more nucleotide positions (i.e. no independent combinations are identified), such position(s) present in all pDNCs are termed key positions. The key position(s) are crucial for diagnosing a taxon, because a substitution at this position even in one sequence attributed to a focus-taxon would immediately make the focus-taxon impossible-to-diagnose with the selected genetic marker. On the contrary, when  $n$  independent

DNCs are recovered,  $n$  substitutions would be needed to make a focus taxon undiagnosable; the likelihood of the latter scenario is obviously much lower. At the **fourth** step the set of refined pDNCs is analyzed to output 25 shortest pDNCs, a set of independent pDNCs, or (if present), key position(s). In the case that no pDNCs were recovered for a pre-defined set of DNA sequences, an exception is raised.

At **fifth** step the set of pDNCs is converted into the sDNC that fulfills pre-defined requirements of robustness (for detailed explanation of the nature of both see fedosov et al. 2019). The robustness of a DNC is tested by generating 100 artificial DNA sequence datasets (derived from the original dataset, but different from it), and checking in how many of them the tested DNC will remain 1) shared by all focus taxon sequences, and 2) diagnostic for the focus taxon. When the DNC is tested 1 mismatch is allowed in the positions involved in the sDNC in a focus taxon sequences: in this case the sDNC will still be considered valid, if remaining positions constitute a valid DNC. To generate artificial datasets a pre-defined number of random nucleotide substitutions are introduced at random sites of nucleotide sequences without phylogenetic pattern and not at the sites recognized as key positions. The defaults are like:

- -for species **every other** sequence in the alignment is modified (***PrSeq* =0.5**) if the total number of records does not exceed **500**, alternatively, every fourth sequence in the alignment is modified (***PrSeq* =0.25**). The number of introduced substitutions is such to make a derived sequence 1% divergent from the original one (***Pdiff*=1**).
- -for genera and higher taxa only 10% of the sequences are modified (***PrSeq*=0.1**). The number of introduced substitutions is such to make a derived sequence 3% divergent from the original one (***Pdiff*=3**).

Thus each sDNC is scored by assigning a number from 0 (when a DNC failed in all 100 artificial datasets) to 100 (when it worked for all 100). This algorithm is used repeatedly: one informative position is appended to the DNC and then it is scored, if the desired criteria of robustness (see below) are met the generated sDNC is output, if not, another position is appended and thus extended DNC is scored again, and so on. If the length of DNC exceeds the arbitrarily selected threshold of 25 positions, and the criteria of robustness still not fulfilled, the best scoring sDNC is output with a warning.