# PLNCPRO User Manual

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### **Essential requirements**

- Operating System
  - Linux
- Software
  - Python 2.7
  - NCBI BLAST
  - framefinder(part of Estate package; provided with plncpro)
  - GNU C Library (glibc 2.12 or higher)
- Additional python modules
  - NumPy
  - SciPy
  - Biopython
  - Scikit-learn

#### **Setup**

- Install Python 2.7 and the required modules
- Download and extract plncpro.1.0.tar.gz from here
- Make *framefinder* executable
  - Go to directory plncpro/lib/estate
  - \$ sudo make
  - Copy framefinder executable to plncpro/lib/framefinder
  - \$ cp bin/framefinder ../framefinder
- Put the blast binaries in folder plncpro/lib/blast/bin
- Create a protein database to be used with blastx (swissprot recommended)
- Run the required program from command line using \$ python "script.py"

#### **Usage and examples**

 prediction.py: To label lncRNAs and mRNAs. This file reads an input file containing sequences and then classifies the sequences as coding or non-coding. It uses a model generated by build.py to make classifications. It outputs a file containing class label and class probabilities for each sequence.

**Usage:** python prediction.py -i input\_fasta\_file -o output\_directory -p output\_file\_name -t number\_of\_threads -d path\_to\_blastdb -m model\_file

#### **Parameters:**

-p,--prediction\_out output file name-i,--infile input sequence file

-m,--model model file

-o,--outdir output directory name -d,--db path to blast database

#### **OPTIONAL**

-t,--threads number of threads [default: 4]

-l,--labels path to the files containing labels(it outputs

classification accuracy)

-r,--remove\_temp clean up intermediate files

-v,--verbose show more messages on screen

--min\_len specifiy min\_length to filter input files

--noblast Don't use blast features

-no\_ff Don't use framefinder features

--qcov\_hsp specify query coverage parameter for blast

[default:30]

--blastres path to blast result for input file

#### **Examples:**

a.) \$ python prediction.py -i sample\_data/test/neg.fa -p pred\_res -o sample\_preds -m sample\_out/sample\_model -d lib/blastdb/sprotdb/sprotdb -t 10

Above command will label the sequences in the 'neg.fa' file using 10 threads. The output files will be written to the 'sample\_preds' directory and 'pred\_res' will contain the predicted class with probabilistic score. Each sequence predicted as mRNA will be labelled as 1 and lncRNAs will be labelled as 0.

b.) \$ python prediction.py -i sample\_data/test/neg.fa -p pred\_res -o sample\_preds -m sample\_out/sample\_model -d lib/blastdb/sprotdb/sprotdb -t 10 --min\_len 500

This command is same as above but it will use sequences with length greater than or equal to 500 bp for prediction.

2. **build.py**: used to build model using the given training data (mRNA/lncRNA transcripts). This file reads two labelled datasets containing coding and non-coding transcripts. Then it makes a random forest based classification model and saves the model, which can be used later to predict unknown sequences.

**Usage:** python build.py -p mRNAs\_fasta -n lncRNAs\_fasta -m output\_model\_name -t number\_of\_threads -o output\_dir -d path\_to\_blast\_database

#### **Parameters:**

-p,pos	mRNA sequence file
-n,neg	lncRNA sequence file
-m,model	output model name
-o,outdir	output directory name
-d	path to blast database

**OPTIONAL** 

-t,--threads number of threads [default: 4]
-k,--num\_trees number of trees [default: 1000]
-r,--remove\_temp clean up intermediate files

-v,--verbose show more messages

--min\_len specifiy min\_length to use for prediction

--noblast Don't use blast features

--no\_ff Don't use framefinder features

--qcov\_hsp specify query coverage parameter for blast

[default:30]

--pos\_blastres path to blast result for mRNA input file --neg\_blastres path to blast result for lncRNA input file

#### **Examples:**

a.) \$ python build.py -p sample\_data/train/pos.fa -n sample\_data/train/neg.fa -o sample\_out -m sample\_model -d lib/blastdb/sprotdb/sprotdb -t 10

NOTE: This constructs a model using the mRNA sequences in the 'pos.fa' file and lncRNA in 'neg.fa'. The program outputs the model in the file 'sample\_model' in 'sample\_out' directory. To use this model for predictions simply give the path to this model file as the -m,--model argument in *prediction.py*, as below:

\$ python prediction.py -i test.fa -out prediction\_out -p prediction\_file -m sample\_out/sample\_model -d path\_to\_blast\_db

b.) \$ python build.py -p sample\_data/train/pos.fa -n sample\_data/train/neg.fa -o sample\_out -m sample\_model -d lib/blastdb/sprotdb/sprotdb -t 10 --min\_len 300

This command will use all sequences from neg.fa and pos.fa having length greater than or equal to 300 bp for constructing the model.

3. **predtoseq.py**: used to extract mRNA or lncRNA sequences from PLNCPRO output file. This file reads a prediction output file and extracts sequences from a given class. User can specify class and probability cut-off and extract desired transcript sequences.

**usage**: python predtoseq.py -f fasta\_file -o outputfile -p PLNCPRO\_prediction\_file -l required\_label

## **PARAMETERS**

-f	input fasta file	
-0	output fasta file name	
-p	path to file containg predictions by PLNCPRO	
OPTIO	NAL	
-1	label of the required sequences (0 for lncRNA;	
1 for mRNA) [default:0]		
-S	class probability cutoff (extract sequences with	
probability greater than or equal to s)		
min	specifiy min_length of sequences [default:0]	
max	specifiy min_length of sequences [default:Inf]	

#### **Description of files**

- a. build.py: this file reads two labelled datasets containing coding and non-coding transcripts. Then it makes a random forest based classification model and saves the model, which can be used later to predict unknown sequences.
- b. prediction.py: this file reads an input file containing sequences and then classifies the sequences as coding or non-coding. It uses a model generated by build.py to make classifications. It outputs a file containing class label and class probabilities for each sequence.
- c. predtoseq.py: this file reads a prediction output file and extracts sequences from a given class. User can specify class and probability cut-off and extract desired transcript sequences.
- d. blastparse.py: this file reads output of blastx program, run with "-outfmt '6 qseqid sseqid pident evalue qcovs qcovhsp score bitscore qframe sframe", and extracts features from it.
- e. extractfeatures.py: this file extracts trimer frequency and lengths from input fasta sequence.
- f. ffparse.py: this file reads output from framefinder and extract features.
- g. mergefeatures.py: this file merges all the features generated from blastpare.py, extractfeatures.py and ffparse.py in to single feature file.
- h. buildmodel.py: this file reads an input file containing features and labels and outputs a random forest classification model
- i. predict.py this file reads an input feature file and predicts its label using a model.

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