Acquiring Retrotransposon Data From UTSC Table Browser

1) From UTSC acquire retrotransposon data with the following inputs. Select filter and enter the retrotransposon class enclosed in asterisks; (e.g. *I1*):



2) Click get output and ensure settings are as follows on the following page and click get sequence:

Sequence Retrieval Region Options:
Add 0 extra bases upstream (5') and 0 extra downstream (3')
Note: if a feature is close to the beginning or end of a chromosome and upstream/ochromosome.
Sequence Formatting Options:
 All upper case. All lower case. Mask repeats: to lower case to N
get sequence cancel

3) Navigate to the location in the cloned repository containing the file,

```
retrobase_commands-0.1-py3.whl.
```

Pip install using:

```
pip3 install retrobase commands-0.1-py3.whl
```

4) Call the command

```
retrobase_commands translate --input_file=<input_file> --
output_file=<output_file> superfamily=<superfamily>
```

Where:

- <input file> is the path to the file you downloaded from UCSC table browser
- -<output file> is the path to create the output file
- -<superfamily> is the retrotransposon superfamily/class

5) Install local blast from

https://ftp.ncbi.nlm.nih.gov/blast/executables/blast+/LATEST/

6) Create a local blast database using:

```
makeblastdb -in <input file> -parse_seqids -blastdb_version 5 -title
"<database title>" -dbtype prot
```

- 7) Get protein sequences in FASTA format from uniprot representing known retrotransposon protein sequences.
- 8) Use sequences acquired in step 7 as query sequences in psi-blast queries against the database created in step 6 using the following command:

```
psiblast -query <RT -protein-FASTA> -db <database-title> -out
<output-file-name> -outfmt 5 -max target seqs=1000000
```

Where:

- -<RT-protein-FASTA> is a protein sequence obtained in step 7
- -<database-title> is the title of the database created in step 6
- -<output-file-name> is the name of the psi-blast output that must be formatted: rotein
 name>_psiblast.txt
- -max_target_seqs set to a very large number ensures all sequences with e-values under the threshold will be included
- 9) Create a JSON file relating DNA records, translated protein sequences, predicted protein labels and associated data using the retrobase_commands command:

```
retrobase_commands assign_proteins --dna_input_file=<dna_input_file>
--protein_input_file=rotein_input_file> --
psiblast_directory=<psiblast_file_path> --outputfile=<outputfile> --
superfamily=<superfamily> --protein_names=protein_names> --
genome=<genome>
```

Where:

- -<dna_input_file> is the path to the path and filename to the fasta file output from UTSC table browser
- ------FASTA created by the tanslate command
- -<psiblast_directory> is the path to the files output by psiblast query. These file names must be formatted
- -<outputfile> is the name of the JSON file to be created
- -<superfamily> is the name of the retrotransposon class
- ------cprotein_names> is the names of the proteins queried using psiblast. The names must be written
 the same as those in the psiblast output filenames. Multiple protein names should be input by
 repeating the option as so: --protein_names=Gag --protein_names=Pol --protein_names=Pro
- -<genome> is the name of the genome from which the original retrotransposon sequences came (e.g. hg_38)

10) Add the path to the Retrobase base directory to the python path

11) Upload records to database using the retrobase_commands command:

```
retrobase_commands upload_records --input_file=<input_file> --
path_to_django_settings>
```

Where:

-<input_file> is the path and filename to the JSON file created in step 9

12) Acquire protein function data from Uniprot using the command:

```
retrobase_commands enter_uniprot_data --accession_ids=<accession_id>
--protein_names=retrobase_commands enter_uniprot_data --accession_ids=<accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id></accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><accession_id><ac
```

Where:

- -<accession_id> is the accession ID from of the Uniprot entry from which the standard sequence of a given protein came. For multiple accession IDs, simply use the option multiple times in the command. Multiple accession IDs must be listed in **the same order** as the protein names
- -------collected (must be written as in Retrobase database). For multiple protein_names, simply use the option multiple times in the command. Multiple protein names must be listed in the same order as the accession IDs.