```
Modules
```

<u>numpy</u> <u>os</u> <u>random</u>

## **Functions**

#### main()

Generates random sequence data files.
datadir/datatype.fasta
...
>rseq\_{i}\_{energy}
sequence
structure
>rseq\_{i+1}\_{energy}

sequence

structure

111

# predict

/home/julian-zim/Desktop/proj/rnadeep/examples/predict.py

## Modules

<u>absl</u>

numpy

os

#### Functions

main()

# train ali (version v0.1)

index

 $/\!home/\!julian-zim/Desktop/proj/rnadeep/examples/train\_ali.py$ 

## Modules

<u>absl</u>

argparse

<u>os</u>

tensorflow

# Functions **Section** 10 miles

## main()

RNAdeep training interface.

## $\boldsymbol{parse\_rnadeep\_args}(p)$

Arguments that are used by RNAdeep.

 $\textbf{training} (\texttt{datatag, dbn\_dir, ali\_dir, spotmodel=None, basemodel=None, savedir='.', epochs=50, epoch0=0, batch\_size=4) \\$ 

## **train** (version v0.1)

index

/home/julian-zim/Desktop/proj/rnadeep/examples/train.py

## Modules

<u>absl</u>

argparse

<u>os</u>

# Functions

#### main()

RNAdeep training interface.

 $python\ train.py\ -d\ l30\ -t\ train\_data/fixlen30\_n100000.fa-train\ -v\ train\_data/fixlen30\_n100000.fa-valid\ -m\ 4\ -b\ 250\ -e\ 20\ --model-log-dir\ intermediate\_models\ -l\ intermediate\_models/sm4\_l30\_010/\ --epoch0\ 10\ 2>>\ sm4\_l30.out\ \&$ 

## $parse\_rnadeep\_args(p)$

Arguments that are used by RNAdeep.

training(datatag, ftrain, fvalid, spotmodel=None, basemodel=None, savedir='.', epochs=50, epoch0=0, batch size=4)

## Modules

absl

numpy

os

## **Functions**

main()

# rfam\_converter

index

/home/julian-zim/Desktop/proj/rnadeep/rnaconv/rfam converter.py

## **Modules**

**RNA** numpy shutil

SVS **textdistance** 

## Functions |

## convert\_rfam\_data(seed\_filepath, tree\_dirpath, outpath)

Calls the necessary functions to convert the whole rfam database into single files, preparing them to be used by

#### Parameters:

seed\_filepath (str): Path to the Rfam.seed file in STOCKHOLM format, containing the families tree\_dirpath (str): Path to the directory in which Rfam tree files in newick format (.seed\_tree) files are outpath (str): Path to the directory in which to save the converted data

# fix\_newick\_strings(tree dirpath, outpath)

Fixes newick strings by replacing every control character (e.g. '(', ')', ',', '.', ':') within a node name with an underscore.

Additionally, multifurcations are resolved and non-leaf node labels are removed. (These three steps are nessecary for SISSI to be able to parse the Rfam tree files.)

#### Parameters:

tree\_dirpath (str): path to the directory containing the tree files in newick string format outpath (str): path to the directory in which to save the trees in the fixed newick string format.

#### main()

#### obtain equilibrium frequencies(ali dirpath, neigh dirpath, outpath)

Extracts the equilibrium frequencies for unpaired single nucleotides and nucleotide pairs from an alignment, by counting the occurences of single nucleotides in unpaired site and saving them in a 4-vector, counting the occurences of nucleotide pairs in paired sites and saving them in a 16-vector, adding pseudocounts to both (+1 for each element) and normalizing in the end.

ali dirpath (str): Path to the directory containing the alignment files in CLUSTAL format neigh dirpath (str): Path to the directory containing the alignment consensus structure files in dot bracket notation format

outpath (str): Path to the directory in which to save the extracted unpaired single and paired nucleotide equilibrium frequencies

#### rescale\_newick\_strings(tree dirpath, ali dirpath, outpath)

Rescales the tree branch lengths for trees which corresponding sequence alignments sequences are over 95% similar with respect to their mean pairwise hamming distance, in order to increase the evolution rate when using the tree for evolutionary simulation.

## The rescale factor is 2.

#### Parameters:

 ${\tt tree\_dirpath~(str):~path~to~the~directory~containing~the~tree~files~in~newick~string~format}$ ali\_dirpath (str): path to the directory containing the alignment files in CLUSTAL format outpath (str): path to the directory in which to save the rescaled trees in the newick string format.

## stockholm\_to\_alignments(filepath, outpath)

Converts the alignments contained in the STOCKHOLM input file into CLUSTAL files.

filepath (str): Path to the Rfam.seed file in STOCKHOLM format, containing the families outpath (str): Path to the directory in which to save the extracted alignments in the CLUSTAL format

## **stockholm to neighbourhoods**(filepath, outpath)

Calls the necessary functions to convert the consensus structures contained in the STOCKHOLM input file into single files in the wuss and dbn formats, respectively.

#### Parameters

filepath (str): Path to the Rfam.seed file in STOCKHOLM format, containing the families outpath (str): Path to the directory in which to save the extracted consensus structures

## stockholm\_to\_wuss(filepath, outpath)

Converts the consensus structures contained in the STOCKHOLM input file into single files in the washington university secondary structure (wuss) format.

#### Parameters:

filepath (str): Path to the Rfam.seed file in STOCKHOLM format, containing the families outpath (str): Path to the directory in which to save the resulting wuss file

wuss to db(filepath, outpath)

Converts the consensus structures contained in the wuss input file into the dot bracket notation format

Parameters

filepath (str): Path to the file in wuss format, containing the secondary structure outpath (str): Path to the directory in which to save the resulting dot bracket notation file

data filter

/home/julian-zim/Desktop/proj/rnadeep/rnaconv/data filter.py

Modules

RNA numpy os sys

## Functions

filter\_data(ali\_dirpath, seq\_dirpath, neigh\_dbn\_dirpath, neigh\_ct\_dirpath, max\_dbrs\_deviation=20)

Filters the data generated by the data\_generator: In each alignment, every sequence is removed that deviates by over <max\_dbrs\_deviation> from the consensus structure that was used by SISSI to generate it. This is achieved by using RNAfold to predict the secondary structure and the base pair distance to compare it to the desired consensus structure. If this results in leaving less than 2 sequences in the alignment, essentially deleting it or turning it into a single sequence, the whole alignment file with the corresponding consensus structure and sequence

Note: If families were generated, the consensus structures used for the alignment generation were generated by RNAfold and saved into the neigh\_dirpath.

If only alignments were generated, the consensus structures used for the generation were provided by the user, most likely from a converted Rfam database, but then copied to the neigh\_dirpath anyway, for the sake of integrity.

Therefore, in both cases, neigh\_dirpath can be used to retrieve the desired consensus structures to compare the sequences with.

#### Parameters:

ali\_dirpath (str): Path to the directory of the generated alignments.
seq\_dirpath (str): Path to the directory of the generated or copied sequences.
neigh\_dbn\_dirpath (str): Path to the directory of the generated or copied dbn files.
neigh\_ct\_dirpath (str): Path to the directory of the generated ct files.
max\_dbrs\_deviation (int, None, optional): maximum allowed base pair distance deviation from the consensus structure in percent. Default is 20.

#### main()

 ${\bf obtain\_and\_compare\_equilibrium\_frequencies} (a {\it li\_dirpath, neigh\_dirpath, orig\_single\_freq\_dirpath, orig\_doublet\_freq\_dirpath, outpath})$ 

Extracts the equilibrium frequencies for unpaired single nucleotides and nucleotide pairs from the generated alignments and forms the differences to the already extracted equilibrium frequencies of the original alignments.

#### Parameters:

orig\_doublet\_freq\_dirpath (str): Path to the directory of the extracted doublet frequency files of the original alignments

outpath (str): Path to the directory in which to save the extracted unpaired single and paired nucleotide equilibrium frequencies and frequency differences

rfam filter

index /home/julian-zim/Desktop/proj/rnadeep/rnaconv/rfam filter.py

## Modules

<u>os</u> <u>sys</u>

## **Functions**

**filter\_rfam\_data**(ali\_dirpath, single\_freq\_dirpath, doublet\_freq\_dirpath, neigh\_wuss\_dirpath, neigh\_dbn\_dirpath, tree\_fixed\_dirpath, tree\_rescaled\_dirpath, max\_length=700)

Filters the alignments, consensus structures, frequencies and trees of a converted rfam database (not touching the original tree files and original Rfam.seed file):

Every data point (consisting of the four filetypes mentioned above) which alignment exceeds a certain length is removed.

#### Parameters

ali\_dirpath (str): Path to the directory of the converted alignments. single\_freq\_dirpath (str): Path to the directory of the extracted single frequency files. doublet\_freq\_dirpath (str): Path to the directory of the extracted doublet frequency files. neigh\_wuss\_dirpath (str): Path to the directory of the extracted wuss files. neigh\_dbn\_dirpath (str): Path to the directory of the extracted dbn files. tree\_fixed\_dirpath (str): Path to the directory of the fixed newick string tree files. tree\_rescaled\_dirpath (str): Path to the directory of the rescaled tree files. max\_length (int, None, optional): Maximum allowed length of an alignment. Default is 700.

main()

#### **Modules**

RNA os subprocess argparse random

#### **Functions**

#### **db to ct**(dbn, seq)

Converts the consensus structures contained in the dot bracket notation input file into the connect table format.

#### Parameters:

dbn (str): Secondary structure in dot bracket notation
seq (str): Sequence

generate\_alignment\_set(sissi\_filepath, number, tree\_dirpath, neigh\_dirpath, sfreq\_dirpath, dfreq\_dirpath, ali\_dirpath, outpath)
Generates <number> alternative alignments for each tree file in the given tree-directory, searching in the
respectively given consensus-structure-, single- & doublet-frequencies- and, optionally for readding
ndels, alignment-directories for files of the same name to use.

#### Parameters:

sissi\_filepath (str): Path to the compiled sissi099 file
number (int): The number of alignments to generate
tree\_dirpath (str): Path to a directory containing tree files in the newick string format ('.seed\_tree')
neigh\_dirpath (str): Path to a directory containing neighbourhood files in the dot-bracket notation format ('.dbn')
sfreq\_dirpath (str): Path to a directory containing files storing a single frequency vector ('.sfreq')
dfreq\_dirpath (str): Path to a directory containing files storing a doublet frequency vector ('.dfreq')
ali\_dirpath (str, None): Path to a directory containing alignment files in the clustal format ('.aln')
outpath (str): The Path to which to write the generated sequences, alignments & copied consensus structures

generate\_alignments(sissi\_filepath, number, tree\_filepath, neigh\_filepath, sfreq\_filepath, dfreq\_filepath, ali\_filepath, outpath)
Generates <number> alternative alignments for the given tree-, consensus-structure-, single- & doublet-frequenciesand, optionally for readding indels, alignment-file, using:

- RNAinverse to generate an ancestral sequence for the provided consensus structure
- SISSI simulate homologous sequence alignments (taking the generated ancestral sequence, provided tree, provided consensus structure and provided equilibrium frequencies as input).

#### Note:

The provided consensus structure will also be copied into an additional file per generated alignment, in order to create pairs of samples and tags to be used for training (during the process, the dbn files are converted to ct files, which are also saved).

The generated ancestral sequence will also be saved to maintain integrity.

#### Parameters:

sissi\_filepath (str): Path to the compiled sissi099 file
number (int): The number of alignments to generate
tree\_filepath (str): Path to a directory containing tree files in the newick string format ('.seed\_tree')
neigh\_filepath (str): Path to a directory containing neighbourhood files in the dot-bracket notation format ('.dbn')
sfreq\_filepath (str): Path to a directory containing files storing a single frequency vector ('.sfreq')
dfreq\_filepath (str): Path to a directory containing files storing a doublet frequency vector ('.dfreq')
ali\_filepath (str, None): Path to a directory containing alignment files in the clustal format ('.aln')
outpath (str): The Path to which to write the generated sequences, alignments & copied consensus structures

generate\_families(sissi\_filepath, number, min\_length, max\_length, tree\_filepath, sfreq\_filepath, dfreq\_filepath, outpath)

Generates <number> families for the given tree- and single- & doublet-frequencies-file, using:

- random ancestral sequences of uniformly distributed lengths up to <maxlength>
- RNAfold to predict secondary structures for these sequences to be used as consensus structures for the alignment generation
- SISSI simulate corresponding homologous sequence alignments (taking the random ancestral sequences, provided tree, predicted consensus structures, and provided equilibrium frequencies as input).

#### Note:

During the process, the generated dbn files are converted to ct files, which are also saved.

## Parameters:

sissi\_filepath (str): Path to the compiled sissi099 file
number (int): The number of families to generate
min\_length (int): Minimum allowed length of the ancestral sequences used to generate the families
max\_length (int): Maximum allowed length of the ancestral sequences used to generate the families
tree\_filepath (str): Path to a tree file in the newick string format ('.seed\_tree')
sfreq\_filepath (str): Path to a file containing a single frequency vector ('.sfreq')
dfreq\_filepath (str): Path to a file containing a doublet frequency vector ('.dfreq')
outpath (str): The path to which to write the generated families

generate\_family\_set(sissi\_filepath, number, min\_length, max\_length, tree\_dirpath, sfreq\_dirpath, dfreq\_dirpath, outpath)
Generates <number> RNA families of uniformaly distributed lengths up to <maxlength> for each tree file in the given
tree-directory, searching in the respectively given single- & doublet-frequencies-directories for files of the same
name to use.

For more information, refer to the  $\underline{\texttt{generate}\_\texttt{families}}(\texttt{)}$  function.

#### Parameters:

sissi\_filepath (str): Path to the compiled sissi099 file
number (int): The number of families to generate
min\_length (int): Minimum allowed length of the ancestral sequences used to generate the families
max\_length (int): Maximum allowed length of the ancestral sequences used to generate the families
tree\_dirpath (str): Path to a directory containing tree files in the newick string format ('.seed\_tree')
sfreq\_dirpath (str): Path to a directory containing files storing a single frequency vector ('.sfreq')
dfreq\_dirpath (str): Path to a directory containing files storing a doublet frequency vector ('.dfreq')
outpath (str): Path to which to write the generated families

## $\textbf{generate\_sequence\_structure\_pair} (length = 85, \\ min\_paired\_sites\_percent = 20)$

Repeatedly generates a random sequence and predicts its secondary structure using RNAfold, until the structure has at least min\_paired\_sites paired sites.

# sampling ali

index /home/julian-zim/Desktop/proj/rnadeep/rnadeep/sampling\_ali.py

### Modules

numpy

<u>os</u>

## Functions 5 3 2

draw ali sets(ali directory, dbn directory, splits=None)

parse\_families(ali\_dirpath, dbn\_dirpath)

Combines pairs of the same name of alignment CLUSTAL files and neighbourhood Dot Bracket String files found in the respective directories to be used for training.

Parameters:

ali\_dirpath (str): Path to the directory containing the alignment CLUSTAL files dbn\_dirpath (str): Path to the directory containing the neighbourhood Dot Bracket String files

parse\_family(ali\_filepath, dbn\_filepath)

Reads an alignment CLUSTAL file and neighbourhood Dot Bracket String file and combines them into a pair to be used for training.

Parameters:

ali\_filepath (str): Path to the alignment CLUSTAL file
dbn\_filepath (str): Path to the neighbourhood Dot Bracket String file

## models

index /home/julian-zim/Desktop/proj/rnadeep/rnadeep/models.py

## Modules

keras.api.\_v2.keras.layers

tensorflow

#### **Functions**

```
spotrna alignment models(model=1, use mask=True)
     Some modifications to Julia's SPOT-RNA implementations.
     Supposed to be a reimplementation of the models in the \,
     SPOT-RNA paper. If you find mistakes, please let us know!
     Overview:
             - Initial 3x3 convolution laver
             - ResNet blocks
             - Act./Norm.
             - 2D-BLSTM
             - Fully Connected blocks
             - Output masking layer (optional)
             - Output layer
     Aras:
             model: select the model (0-4)
             use_mask: for padded input/output (defaults to True!)
spotrna_models(model=1, use mask=True)
     Some modifications to Julia's SPOT-RNA implementations.
     Supposed to be a reimplementation of the models in the
     SPOT-RNA paper. If you find mistakes, please let us know!
     Overview:
             - Initial 3x3 convolution layer
             - ResNet blocks
             - Act./Norm.
             - 2D-BLSTM
             - Fully Connected blocks
             - Output masking layer (optional)
             - Output layer
             model: select the model (0-4)
             use_mask: for padded input/output (defaults to True!)
```

index

<u>numpy</u> <u>os</u> <u>random</u>

## Functions

```
draw_sets(fname, splits=None)
generate_random_structures(lengths)
rseq(l)
write_data_file(data, fname, mode='w')
        Save sequence/structure pairs for the given lengths.
write_fixed_len_data_file(seqlen, num, root=")
write_normal_len_data_file(central, std, num, root=")
write_uniform_len_data_file(minlen, maxlen, num, root=")
```

index

/home/julian-zim/Desktop/proj/rnadeep/rnadeep/encoding\_utils.py

# encoding\_utils

Mod<u>ules</u>

numpy

## **Functions**

```
base_pair_matrix(ss)
binary_encode(structure)
create_windows(sequences, window_size)
encode padded alignment matrix(alignments, max length=None)
encode padded sequence matrix(sequences, max length=None)
encode_padded_structure_matrix(structures, max length=None)
encode_sequence(sequences)
encode_sequence_matrix(sequences)
     Make a BP probability matrix with one-hot encoding of basepairs.
     NOTE: This only works if all sequences have the same length, otherwise
     you need to use: encode_padded_sequence_matrix
encode_sequence_windows(sequences, window size)
encode_structure(structures)
encode_structure_matrix(structures)
     Make a BP probability matrix with one-hot encoding of basepairs.
     NOTE: This only works if all sequences have the same length!
make pair table(ss, base=0, chars=['.'])
     Return a secondary struture in form of pair table.
       ss (str): secondary structure in dot-bracket format
       base (int, optional): choose between a pair-table with base 0 or 1
       chars (list, optional): a list of characters to be are ignored, default:
            11.11
     **Example:**
       base=0: ((..)). => [5,4,-1,-1,1,0,-1]
           i.e. start counting from 0, unpaired = -1
       base=1: ((...)). => [7,6,5,0,0,2,1,0]
            i.e. start counting from 1, unpaired = 0, pt[0]=len(ss)
     Returns:
       [list]: A pair-table
one_hot_encode(char)
one_hot_matrix(seq)
profile vec matrix(ali)
     Creates a profile matrix for the given alignment: For each cell a ij, the columns i and j or the alignment are
     combined by forming the outer product of two profile vectors for the two respective symbols at the current row
     index of the two columns and summing them all up. The two respective profile vectors created by the sheme defined in
     the base to ids dictionary variable.
```