# **EM2Tools**

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**CHAPTER** 

**ONE** 

## **GETTING STARTED**

## 1.1 Requirements

The EM2Tools package requires at least Python 3.6 with Biopython, pandas and gffpandas packages.

## 1.2 Installation

## 1.2.1 Linux

1. Download and unzip the master branch zip file:

```
\label{lem:master.zip} wget \ https://github.com/fplewniak/EM2Tools/archive/master.zip \\ unzip \ master.zip
```

2. Install the EM2Tools Python package and the required Python packages with pip:

```
pip install EM2Tools-master/.
```

## THE EM2TOOLS API REFERENCE

## 2.1 The "seq\_record" module

Extension module to the Biopython Bio.SeqRecord module

```
class em2lib.seq_record.SeqRecordEM2 (seq, **kwargs)
```

Extension to Biopython SeqRecord class

```
add_feature (**kwargs)
```

Adds a feature to the current record according to arguments passed as \*\*kwargs.

Parameters kwargs – keyword arguments to pass to SeqFeatureEM2 class

overlap (start, end=None, strand=0)

Retrieves features that overlap a given position range.

#### **Parameters**

- **strand** strand specification of features to be returned. If strand is 0, then features on both strands are returned. If feature.strand is 0, then all strands will match.
- start start of range
- end end of range, if None, then end=start

**Returns** a list of overlaping features

## feature\_after (position, strand=0, nearest=False)

Retrieves the features immediately after (but not overlaping) the specified position, on one strand or both. If nearest is True, then only the nearest ones are returned.

#### **Parameters**

- nearest if True, only the nearest features are returned. This only makes sense when strand is 0
- **strand** strand specification of features to be returned. If strand is 0, then features on both strands are returned
- position the position

**Returns** a list of features after the specified position

### feature before (position, strand=0, nearest=False)

Retrieves the features immediately before (but not overlaping) the specified position, on one strand or both. If nearest is True, then only the nearest ones are returned.

- nearest if True, only the nearest features are returned. This only makes sense when strand is 0
- **strand** strand specification of features to be returned. If strand is 0, then features on both strands are returned
- position the position

**Returns** a list of features before the specified position

## surrounding\_features (position, strand=0, nearest=False)

Retrieves all the features around a given position but not overlaping it. If nearest is True, then only the nearest features are returned.

#### **Parameters**

- position the position
- nearest if True, only the nearest features are returned.
- **strand** strand specification of features to be returned. If strand is 0, then features on both strands are returned

**Returns** a list of features around the specified position

join (other=None, offset=0, keepself=True)

Joins two SeqRecordEM2 objects into a new one representing the resulting merged sequence

#### **Parameters**

- **keepself** if True and overlapping subsequences are different, then keep sequence from self record, otherwise keep the sequence of other record.
- other the other SeqRecordEM2 object
- **offset** the offset of the two sequences. If the value is negative, then the two sequences overlap.

Returns the result of merging records as a new SeqRecordEM2 object

**stitch** (other, fpos\_in\_self, fpos\_in\_other, feature\_length, orientation=1, \*\*kwargs)

Stitches two records, that is, joins them according to an overlapping feature. The sequences may or may not overlap. If not, Ns or Xs are added to fill the gap. If they overlap, a warning is issued if sequences do not correspond exactly. The new record keeps track of the two original records as Features. By convention, the self record should contain the start position of the feature on the forward strand, the other contains the end position on the forward strand if orientation=1 or on the reverse strand if orientation=-1. If orientation is -1, then the other record is reversed/complemented before stitching and the position of the overlapping feature is modified accordingly. It is the user's responsibility to provide the records in the right order.

- other the other SeqRecordEM2 object to stitch
- **fpos\_in\_self** feature position in self record (start position of feature)
- **fpos\_in\_other** feature position in other record (end position of feature), according to FeatureLocation conventions for end position requiring that length = end start, it is not included in the feature.
- feature\_length feature length
- **orientation** the orientation of the other record relative to the self, either 1 if it is in the same orientation, -1 if other needs to be reversed before stitching, 0 if stranded but unknown, None for proteins

 kwargs – any additional parameters that may be passed to the constructor of the stitching feature in the new record

Returns the stitched record as a new SeqRecordEM2 object

reverse\_complement (id=False, name=False, description=False, features=True, annotations=False, letter annotations=True, dbxrefs=False)

Reverse-complement the record adjusting features and their positions accordingly. The record id is conserved but if name is not specified 'reversed' is appended. All other arguments are passed and handled by the parent method. Note that the main goal for this method is to replace SeqRecord and Seq objects by their SeqRecordEM2 and SeqEM2 equivalents when reverse/complementing.

#### **Parameters**

- id the id for the reversed record
- name the name for the reversed record
- **description** the description for the reversed record
- features keep and adjust location of features if True
- annotations keep annotations if True
- letter\_annotations keep letter\_annotations if True
- dbxrefs keep dbxrefs if True

**Returns** a reversed copy of the record

```
orfs_to_features (start=['ATG'], stop=None, filter=None, add=False)
```

Determines all open reading frames in a sequence record. All the returned ORFs have a length that is a multiple of 3. Thus, for sequences without any stop codon, 3 ORFs are returned, one for each frame. Both strands are examined but it is possible to filter the ORFs by length, frame, etc. with the FeatureFilter defined by the filter argument.

## **Parameters**

- start a list of accepted start codons
- stop a list of accepted stop codons
- filter a FeatureFilter object defining a filter to select ORFs according to some criteria
- add if True, the selected ORFs are added to the record's features

Returns a list of ORFs as SeqFeatureEM2 objects

## 2.2 The "seq" module

Extension of Bio.Seq.Seq class from Biopython to add or improve functionalities

```
class em2lib.seq.SeqEM2 (data, seqtype)
```

SeqEM2 class providing extension to Bio.Seq.Seq class of BioPython package.

#### classmethod dna(data)

Creates a DNA sequence

**Parameters** data – the sequence string

Returns a SeqEM2 DNA instance

## classmethod protein(data)

Creates a protein sequence

**Parameters** data – the sequence string

Returns a SeqEM2 protein instance

## is\_protein()

Tests whether sequence was created as a protein

**Returns** boolean, True if sequence was created as a protein, False otherwise.

## length\_in\_range (minlength=None, maxlength=None)

Checks whether the sequence length is with the specified range.

#### **Parameters**

- minlength lower length bound
- maxlength upper length bound

Returns True if sequence length is within specified range, False otherwise

### re\_search(regex)

Searches a sequence using a regular expression

**Parameters** regex – the regular expression

**Returns** a list of re.Match instances

#### search (pattern)

Searches sequence for a pattern specified with a fuzznuc or fuzzpro like syntax

Parameters pattern – the pattern to be searched for that is converted into a regular expression

**Returns** a list of re.Match objects

```
get_orfs (start=['ATG'], stop=None)
```

Determines all open reading frames in a sequence. It only examines the forward strand. All the returned ORFs have a length that is a multiple of 3. Thus, for sequences without any stop codon, 3 ORFs are returned, one for each frame.

#### **Parameters**

- **start** list of accepted start codon or None if ORFs do not need to start at a start codon.
- stop list of accepted stop codons

**Returns** a set of tuples (start, end) of orfs where start is the starting position of the orf and end its ending position, not including the stop codon

## 2.3 The "seq\_feature" module

Extension of Bio.SeqFeature module from Biopython to add or improve functionalities

```
class em2lib.seq_feature.SeqFeatureEM2 (parent=None, **kwargs)
```

SeqFeatureEM2 class providing extension to Bio.SeqFeature.SeqFeature class of BioPython package.

#### length\_in\_range (minlength=None, maxlength=None)

Checks whether the feature length is with the specified range.

#### **Parameters**

- minlength lower length bound
- maxlength upper length bound

**Returns** True if feature length is within specified range, False otherwise

#### lies within (start, end)

Determines whether feature lies entirely with the specified range. Fuzzy positions are turned into integers.

#### **Parameters**

- **start** start of range either int or ExactPosition
- end end of range either int or ExactPosition

**Returns** True if feature boundaries lie with the specified range.

## covers (start, end)

Determines whether feature covers the whole range specified by start and end

#### **Parameters**

- start start of range either int or ExactPosition
- end end of range either int or ExactPosition, if None then end=start

**Returns** True if feature covers the specified range

```
overlaps (start, end=None)
```

Determines whether feature overlaps a position range.

#### **Parameters**

- **start** start of range either int or ExactPosition
- end end of range either int or ExactPosition

**Returns** True if feature overlaps range

```
intersect (other, **kwargs)
```

Creates a new feature which is the intersection of feature and another one

Parameters other – the other feature

move (offset)

Moves a feature by a certain offset

Parameters offset – offset by which the feature must be moved

#### class em2lib.seq\_feature.FeatureFilter

A class for the definition and application of a filter to a list of features. May be used to retrieve features from a record or any list of features according to length, location, type, strand and frame. It is possible to combine several criteria into one single filter.

```
keep(keep=True)
```

Set the keep attribute of the filter. If True, the features corresponding to the specified criteria will be kept, otherwise, they will be discarded.

**Parameters** keep – boolean, True if features consistent with the criteria should be kept.

**Returns** the current filter

type (feat\_type=None)

Set the type of feature.

**Parameters feat\_type** – the type of feature to select

Returns the current filter

length (minlength=None, maxlength=None)

Set the length range of the feature.

- minlength the minimum length of the feature or 0 if None
- maxlength the maximum length of the feature or no limit if set to None

Returns the current filter

## strand(strand=0)

Set the strand of the feature.

**Parameters** strand – the strand of the feature

**Returns** the current filter

frame (frame=0, strand=1)

Set the frame of the filter. By default, this is on the forward strand unless the strand is set to -1.

#### **Parameters**

- frame the frame 0, 1 or 2
- strand the strand 1 or 1

**Returns** the current filter

covers (start=None, end=None)

Set a region that must be covered by the feature.

#### **Parameters**

- **start** start position of the region
- end end position of the region

**Returns** the current filter

overlaps (start=None, end=None)

Set a region that must overlap the feature.

#### **Parameters**

- start start position of the region
- end end position of the region

Returns the current filter

## lies\_within (start=None, end=None)

Set a region within which the feature must lie.

#### **Parameters**

- **start** start position of the region
- end end position of the region

**Returns** the current filter

## type\_applies (feature)

Test if type criterion applies to the feature and return a boolean stating whether the feature should be kept or rejected. If the criterion has not been set, then True is returned.

**Parameters** feature – the feature to test

Returns True the feature must be kept

## length\_applies (feature)

Test if length criterion applies to the feature and return a boolean stating whether the feature should be kept or rejected. If the criterion has not been set, then True is returned.

**Parameters** feature – the feature to test

**Returns** True the feature must be kept

## covers\_applies (feature)

Test if covers criterion applies to the feature and return a boolean stating whether the feature should be kept or rejected. If the criterion has not been set, then True is returned.

**Parameters** feature – the feature to test

**Returns** True the feature must be kept

### overlaps\_applies (feature)

Test if overlaps criterion applies to the feature and return a boolean stating whether the feature should be kept or rejected. If the criterion has not been set, then True is returned.

**Parameters** feature – the feature to test

Returns True the feature must be kept

## lies\_within\_applies (feature)

Test if lies\_within criterion applies to the feature and return a boolean stating the feature should be kept or rejected. If the criterion has not been set, then True is returned.

**Parameters** feature – the feature to test

**Returns** True the feature must be kept

### location\_applies (feature)

Test if location criterion applies to the feature and return a boolean stating whether the feature should be kept or rejected. If the criterion has not been set, then True is returned.

Parameters feature - the feature to test

Returns True the feature must be kept

## strand\_applies (feature)

Test if strand criterion applies to the feature and return a boolean stating whether the feature should be kept or rejected. If the criterion has not been set, then True is returned.

Parameters feature – the feature to test

**Returns** True the feature must be kept

## frame\_applies (feature)

Test if frame criterion applies to the feature and return a boolean stating whether the feature should be kept or rejected. If the criterion has not been set, then True is returned.

Parameters feature - the feature to test

**Returns** True the feature must be kept

#### apply (features)

Test if all defined criteria apply to the features in the list and return the list of features corresponding to the specified criteria.

Parameters features – the list of features to filter

**Returns** the filtered list of features

## 2.4 The "seq utils" module

Extension module to the Biopython Bio.SeqUtils module

em2lib.seq\_utils.ambiguous2string(code, protein=False)

Converts an ambiguous residue into a string with all compatible unambiguous residues. If the input code is not ambiguous, it is returned without any conversion.

#### **Parameters**

- **code** the input code to be converted into a list of residues.
- protein True if residue is amino-acid

**Returns** a string corresponding to the unambiguous residues compatible with the input code

em2lib.seq\_utils.isambiguous(code, protein=False)

Checks code is an ambiguous residue specification or not.

#### **Parameters**

- code the input code that must be checked for ambiguity
- protein True if code is amino-acid code

Returns boolean, True if code is ambiguous, False otherwise

em2lib.seq\_utils.pattern2regex (pattern, protein=False)

Converts a fuzznuc or fuzzpro-like pattern into a regular expression that can be used to search a sequence string.

- [ABC] => any of ABC residues,
- {ABC} => any residue except ABC,
- <ABC... => start of sequence,
- ... ABC> => end of sequence,
- A(n)(ABC)(n) => repeat residue or subsequence n times,
- A(n,m)(ABC)(n,m) => repeat residue or subsequence from n up to m times.

#### **Parameters**

- pattern the pattern definition (string)
- protein True if pattern applies to a protein sequence, False otherwise.

**Returns** the regular expression pattern as a string

## class em2lib.seq utils.SeqFilter

A class for the creation of a sequence filter to specify filtering criteria and applying the filter to a list of sequence records.

length (minlength=None, maxlength=None)

Minimal and maximal length specification

## **Parameters**

- minlength minimal accepted length
- maxlength maximal accepted length

**Returns** SeqFilter instance

#### pattern(pattern=None)

pattern specification

Parameters pattern – pattern that must be in the sequence

**Returns** SeqFilter instance

name (name=None)

sequence record name specification

Parameters name - name regular expression

**Returns** SeqFilter instance

keep (keep=True)

Boolean defining whether the matching sequences must be kept (True) or removed (False)

**Parameters** keep – True to keep positive sequences, False to remove them

**Returns** SeqFilter instance

### length\_applies (rec)

test whether length criterion applies to the sequence record

**Parameters** rec – the sequence record to test

**Returns** boolean True if criterion applies or False otherwise

### pattern\_applies(rec)

test whether parameter criterion applies to the sequence record

**Parameters** rec – the sequence record to test

**Returns** boolean True if criterion applies or False otherwise

## name\_applies (rec)

test whether name criterion applies to the sequence record

**Parameters** rec – the sequence record to test

**Returns** boolean True if criterion applies or False otherwise

## apply (records)

Filters a list of SeqRecords instances, keeping only records satisfying the specified criteria of length, match of a pattern, name specification. It is possible to invert the filtering process by setting the keep boolean to False and thus only keep records which do not satisfy the criteria.

Parameters records – list of SeqRecord instances to apply

**Returns** the filtered list of records

**class** em2lib.seq utils.**GFF** (*feature list=None*, *input df=None*)

Manipulation of features based upon gffpandas package

```
add_feature_list (feature_list=None)
```

Adds a list of feature to the list of an existing GFF object

Parameters feature\_list - list of features to add to DataFrame

**Returns** the GFF object with feature list appended

## static df\_from\_feature(feature)

Create a pandas DataFrame from a feature (SeqFeatureEM2 or SeqFeature)

Parameters feature – the feature to convert into a dataframe

**Returns** the resulting dataframe

#### to feature list (parents=None)

Converts features in a GFF object into a list of SeqFeatureEM2 objects

**Parameters** parents – list of references to parent SeqRecord objects or a single parent reference if all features are defined in the same parent. If it is a list, it should be of the same length as the dataframe, repeating references as needed to get the right number.

Returns a list of SegFeatureEM2 objects

## 2.5 The "argparse\_em2" module

Extension module to the standard argparse module. Adding custom actions and argument verification methods.

```
class em2lib.argparse_em2.GetList(option_strings, dest, nargs='+', **kwargs)
```

An argparse custom action to return a list from an argument containing a list of elements and/or file names. Files are supposed to contain one element of the list per line. There can be more than one file and the argument may take a combination of elements and files. In all cases, the returned list will contain all the specified elements without any checking for redundancy. If you need a non redundant set instead of a list, then use GetSet action instead.

## static arg2list(values)

This method converts the argument values containing elements and/or files containing elements into a list of elements.

**Parameters values** – argument values, this is supposed to be a list of arguments or None (returns an empty list)

**Returns** the list of elements or an empty list if the argument was None

```
class em2lib.argparse_em2.GetSet (option_strings, dest, nargs='+', **kwargs)
```

An argparse custom action to return a set from an argument containing a list of elements and/or file names. Files are supposed to contain one element of the list per line. There can be more than one file and the argument may take a combination of elements and files. In all cases, the returned set will contain all the specified elements keeping only one copy of each element. If you do not want to remove redundancy, then use GetList action instead.

```
static arg2set(values)
```

This method converts the argument values containing elements and/or files containing elements into a set of elements.

**Parameters values** – argument values, this is supposed to be a list of arguments or None (returns an empty set)

**Returns** the set of elements or an empty set if the argument was None

## 2.6 The "table" module

Some utilities for Table manipulation, comparison, etc. using pandas.DataFrame module.

```
class em2lib.table.Table
```

A class adding some utilities for easier DataFrame manipulation, comparison, etc.

```
static get_common_keys (first, other, keys_first=None, keys_other=None)
```

Return the index keys in common between first DataFrame and other DataFrame

## **Parameters**

• **first** – the first DataFrame object to compare

- other the other DataFrame object to compare
- **keys\_first** a list of references to the columns defining indices in first DataFrame. None refers to the original index
- **keys\_other** a list of references to the columns defining indices in other DataFrame. None refers to the original index

**Returns** DataFrame, the keys that are common between the two DataFrames.

Return rows with the same key values. Key columns from the other DataFrame object are droppped by default but can be kept if drop is set to False.

### **Parameters**

- first the first DataFrame object to compare
- other the other DataFrame object to compare
- **keys\_first** a list of references to the columns defining keys in first DataFrame. None refers to the original index
- **keys\_other** a list of references to the columns defining keys in other DataFrame. None refers to the original index
- **lsuffix** str, Suffix to add to first column names in case of redundancy.
- rsuffix str, Suffix to add to right column names in case of redundancy.
- **drop** True by default. Drop the other key columns in the resulting DataFrame.

**Returns** DataFrame with the same keys as first DataFrame, containing the rows that have the same key values between the two DataFrames.

static get\_keys\_not\_in (first, other, keys\_first=None, keys\_other=None)
Return the keys in first DataFrame which are not in other DataFrame

#### **Parameters**

- **first** the first DataFrame object to compare
- other the other DataFrame object to compare
- **keys\_first** a list of references to the columns defining keys in first DataFrame. None refers to the original index
- **keys\_other** a list of references to the columns defining keys in other DataFrame. None refers to the original index

**Returns** DataFrame, the keys that are in first DataFrame but not in the other one.

static get\_rows\_not\_in (first, other, keys\_first=None, keys\_other=None)
Return a DataFrame with rows with keys that are in first but not in other.

- first the first DataFrame object to compare
- other the other DataFrame object to compare
- **keys\_first** a list of references to the columns defining keys in first DataFrame. None refers to the original index

• **keys\_other** – a list of references to the columns defining keys in other DataFrame. None refers to the original index

**Returns** DataFrame, the rows whose keys are in first DataFrame but not in the other one.

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