# Lost API documentation

Lost Members

June 1, 2010

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# Chapter 1

# An introduction to the lost framework

#### 1.1 The lost framework

The lost framework is a collection of utilities and models for working with biological sequences written in PRISM/B-Prolog. The framework tries to unify all these bits and pieces and to provide a way to integrate them.

### 1.2 Obtaining a copy of the lost framework

The lost framework can be obtained using *git* if you have an account on the *mox* server. Assuming that you have *git* installed on your local machine, to get a copy you need to clone the central repository:

#### \$ git clone ssh://your-username@mox.ruc.dk/var/git/lost.git

It is also possible to clone the repository directly from mox if you want to work there, rather than on your local machine. To do this, just type

#### \$ git clone /var/git/lost.git

In the following we will use \$LOST to refer to top-most directory of your copy of the framework.

### 1.3 Configuring your copy of the lost framework

After obtaining the lost framework, a little configuration is needed to get started. You will need to edit the file \$LOST/lost.pl. In the beginning of the file there are two important facts you may need to change,

```
lost_config(prism_command,'prism').
lost_config(lost_base_directory, '/change/to/local/lost/dir/').
lost_config(platform, windows_or_unix).
```

The option prism\_command should point to a the main PRISM executable binary. If it is in your \$PATH\$ then you can usually leave it unchanged. lost\_base\_directory should be the full path of the directory (including trailing /) containing the lost.pl file. Note, that even on windows platforms you should use forward slash rather than backslash in the path specification. The value of platform should be either windows or unix.

To get started you can examine and run example.pl which is located in the in the \$LOST/scripts/ directory.

#### 1.3.1 File structure layout of the lost framework

The lost framework is divided into several parts:

- \$LOST/scripts/: Contains scripts that run models. These scripts are just small Prolog programs. The scripts does not consult models directly, but instead use predicates that the framework provides.
- \$LOST/models/: Contains all the biological sequence models. Each model is located in its own subdirectory of \$TLOST/models/ the name of which is also used as name of the model.
- \$LOST/lib/: Contains shared libraries used multiple models.
- \$LOST/data/: Contains sequences and data to be used by the models. Annotations etc. generated by model invoked by the framework stored in this directory and given the extension .gen
- \$LOST/tmp/: Is used for temporary storage of files bt models.

# Chapter 2

# Creating lost models

#### 2.1 Lost model conventions

Each model is located in its own subdirectory of the \$LOST/models/ directory, henceforth called \$MODELS. So for instance, the sample model called sample\_model1 is located in \$MODELS/sample\_model1/. In the following, we will refer to the directory where a particular model resides as \$MODEL.

Models are allowed to consult files with paths relative to the \$MODEL directory, but should under normal circumstances only directly consult file which are located in the \$MODEL directory or a subdirectory of it. The exception to this is the file \$LOST/lost.pl. Consulting this file gives access to all the shared APIs (See chapter 3).

To integrate into the framework each model must provide a file called interface.pl, which must be located in the same directory as the model. In the file interface.pl, one or more interface predicates are defined. These predicates serve as entry points of using the model.

#### 2.1.1 Model interface predicates

This section describes predicates, that when implemented in the interface.pl file provided by a model, allows the model provide functionalities that integrate into the general framework.

For common tasks these predicate are named according to the following convention:

- annotate/3: Do inference with the model and produce a "annotation" as output. The annotation take different forms depending on context.
- train/3: Training the model on the input data and save a switch file or similar as output.
- $\bullet$  generate/3: Generate some data using the model.

It is possible to use the other predicate names than these, but it is *strongly* recommended to use these names whenever it makes sense.

The general form of these predicates is that they take a list of input files, a list of options and the name of an output file. They generally read the input files, do some processing and then write a result to the output file. The type of processing depends on the predicate. The predicate annotate/3 is meant to be used when running the model in prediction mode, so for instance, it might produce a file containing the viterbi path for a given input sequence. The predicate train/3 is meant to be used when training the model and the output file would usually be the learned parameters of the trained model. With the generate/3 predicate, the model is supposed to be used in a generative fashion.

Models should declare the *type* associated to the arguments of these predicates. These declarations include

- The formats of input files, see section ??.
- The format of the output file, see section ??.
- Options and default values for options, see section 2.1.2.
- Optionally, a set of possible values any declared option.

Interface predicates have the following form,

predicate\_name(+InputFileNames, +Options, +OutputFilename)

The predicate takes the following arguments:

- InputFileNames: Is a list of filenames (with absolute paths), each containing an input to the model. There is no restriction on the format of the files, but the model should declare what file formats it expects. Predicates to parse a wide range of fileformats are supplied in the *io* API (see section 3.2).
- Options: Is a list of facts on the form, where the functor works as key and the first argument serves as the value of the option, egg. key(Value). This list is used to parameterize the model in various ways. To check the value of a given option, the predicate get\_option (see section ??) should be used. The options must be declared as described in section .
- OutputFilename: Full filename which the result should be saved to. The model is expected to save the resulting annotation to this file before the completion of annotate. The *io* API contains some common predicates for saving annotations (see section 3.2).

Depending the the predicate\_name the predicates will do conceptually different things, but there is a common pattern: The model reads a processes the files given InputFileNames. It is then the responsibility of the predicate to save the annotation to OutputFilename. If the model for some reason is unable to write the annotation to OutputFilename, then it should throw an exception.

#### 2.1.2 Declaration of options for interface predicates

Options to interface predicates should be declared in the interface.pl file. An option is declared by adding a fact

lost\_option(+InterfaceGoal,+OptionName,+DefaultValue,+Description).

- InterfaceGoal: InterfaceGoal is the functor name of defined interface predicate.
- OptionName: An atom for the name of key. This is the functor identifying this particular option.
- DefaultValue: If the option is left unspecified then it will take this value.
- Description: A textual description of the purpose of the option. Should be in single quotes.

As example, consider an option declaration:

```
lost_option(annotate,debug,no,'Enable/disable debug prints').
```

In this declaration, we state that the goal annotate take the option debug. If this option is not a member of the Options list, then it will be added with the default value no before annotate is called by the framework.

For instance, assuming the model is called  $my\_model$ , it might be called from a script with

```
get_annotation_file(my_model,['/some/path/infile'],[],FileOut),
```

which will start a new PRISM process and invoke

```
annotate(['/some/path/infile'],[debug(no)],FileOut),
```

in \$MODELS/my\_model/interface.pl. Note that the option debug has been inserted with the default value. A default value is only inserted if the option is left unspecified.

#### Declaring option values

If the values of an option can be enumerated, then the model should provide a declaration of by defining a predicate,

```
lost_option_values(+Predicate, +OptionName, -ValuesList).
```

In our example above, the option debug might have two possible values, no and yes, so a declaration might look like,

```
lost_option_values(annotate, debug, [yes,no]).
```

#### Option checking

When a model is invoked by the framework, options are checked before invoking the model. If a model is called with an option which has not been declared by the model, then a warning will be issued. Similarly, if a declared option for a predicate also has a lost\_option\_values declaration and the a predicate is called with the option set to value which is not part of the declaration, the a warning is issued.

#### 2.1.3 Declaration of file formats for interface predicates

#### Input file formats

For each model interface predicate the format of input files and output files should be declared as facts in the interface.pl file for the model. To declare the format of the input files, a fact,

```
lost_input_format(PredicateName, [ Format1, .., FormatN ]).
```

where PredicateName is the name of the predicate to which the declaration belong. The second argument is a list of format identifiers as described in chapter ??. For instance, we may have a declaration,

```
lost_input(annotate,
    [ text(prolog(sequence(dna))), text(prolog(sequence(amino_acids)))]).
```

The last entry of the list of formats may have the special star quantifier, which works similar to a kleene star: It states that the last entry is a placeholder for between zero or more entries of the specified type. For instance,

```
lost_input(annotate,
    [ text(prolog(sequence(dna))), star(text(prolog(sequence(dna))))]).
```

specifies that annotate takes one or more input files of the type text(prolog(sequence(dna))).

#### Output file formats

Contrary to the input file formats, the output file format can depend on the options given to the model. That means, the model may produce files in different output formats depending on the values of the options it is called with.

The output format is thus declared with a predicate,

```
lost_output_format(PredicateName,Options, Format)
```

This predicate should specify the relation between a set of Options and a particular Format for the interface predicate PredicateName. Format should unify to one of the formats described in chapter ??.

#### Checking of file formats

The framework checks that file formats are declared according to the scheme described above. If a model neglects to declare file formats or declares them wrongly, then a warning will be issued.

# Chapter 3

# Lost shared APIs

To use the lost APIs, the file \$LOST/lost.pl must be consulted. To make the models independent of the absolute path of the \$LOST directory, they should consult it with a path relative to the model path (e.g. :- ['../../lost.pl'].). Then, APIs, which are located in the \$LOST/lib directory can be consulted using the goal,

lost\_include\_api(+APIName)

where APIName is the name of a Prolog file located in the \$LOST/lib/ directory except the .pl extension.

#### 3.1 The interface API: interface

The API provides the interface to lost models following the conventions described in section 2.1.

Running a model is done using the goal run\_model

run\_model(Model,annotate(InputFiles,Options,OutputFile)

In this case we run the models annotate goal. This is used to retrieve the annotation generated by a specified Model invoked with the specified InputFiles and Options. InputFiles is a list of file names containing input for the model. Options is a list of facts on the form key(Value), where key is the name of an option and Value is a concrete value. As a consequence of calling this predicate the output from the model is written to the file OutputFile. If the model has been previously invoked with exactly the same InputFiles and Options, then OutputFile will unify with the outcome of the previous invokation. In this way, the results of running a model are cached.

The generated annotation files are named according to a convention, where all annotation files will be placed in the \$LOST/data/ directory and the Filename is construed according to the following convention:

{Model}\_{GoalFunctor}\_{Id}.gen

The first time a model OutputFile is generated, the file annotation.idx will be created in \$LOST/data/. This file serves as a database to map the filename of a generated annotation file to the model, input files and options that generated the particular annotation. This database file contains Prolog facts of the form

fileid(Id,Filename,Model,Goal,Options,InputFiles).

The annotation index is automatically maintained by get\_annotation\_file/4 and should not be edited by hand.

If an annotation for a particular run of a model is not present then get\_annotation\_file/4 will start a new PRISM process that invokes the annotate predicate provided be the model interface.pl file. By the contract of model conventions, the model will generate the annotation and save it to the file indicated by the provided filename.

#### 3.1.1 Moving generated files around

Once a file has been generated, you may wish the move or rename the file. Doing this using normal system commands will fail to the update the annotation index and as result the file may be regenerated, even if it was already generated once. Instead, you should use

```
move_data_file(+OldFilename, +NewFilename)
```

This predicate makes sure that the annotation index is updated accordingly.

# 3.2 Input-Output API: io

In this module (io.pl), several predicates are defined to manipulate \*.seq files:

- loading information from files that extracts from a file data information used as input of models (sequence annotation for example);
- saving information into a file;
- and maybe more.

#### 3.2.1 Loading information from files

- load\_annotation\_from\_file(++Type\_Info,++Options,++File,--Annotation): Generate from File a sequence of Annotation. It is assumed that File is composed of terms. Type\_Info is used to specify what format of information into file
  - sequence means that information is stored into a list. For example,
    data(Key\_Index,1,10,[a,t,c,c,c..]).

db means that information is represented by a set of range that specified specific zone (coding region for example)

```
gb(Key_Index,1,10).
```

For each Type\_Info, several options are available represented by the list Options. Options available for sequence:

- [] (default): data list is the  $2^{th}$  argument of the terms and these lists of data are appended;
- data\_position(Num) specified that data list is  $Num^{th}$  argument of term;
- range(Min, Max) extracts from the list of the complete annotation the sublist from position Min to position Max;
- all\_lists: generate a list of each data list by term. Warning: range(Min, Max) is not support by this option.

File Example toto.seq:

```
data(Key_Index,1,5,[1,2,3,4,5]).
data(Key_Index,6,10,[6,7,8,9,10]).
data(Key_Index,11,15,[11,12,13,14,15]).
```

Results of request are:

```
| ?- load_annotation_from_file(sequence,[data_position(4)],'toto.seq',R).
R = [1,2,3,4,5,6,7,8,9,10,11,12,13,14,15] ?
| ?- load_annotation_from_file(sequence,[data_position(4),range(4,10)],'toto.seq',R).
R = [4,5,6,7,8,9,10] ?
load_annotation_from_file(sequence,[data_position(4),all_lists],'toto.seq',R).
R = [[1,2,3,4,5],[6,7,8,9,10],[11,12,13,14,15]] ?
```

Options available for db:

- [] (default): first and the second element of the term defined a range.
   A list of 0-1 values is generated, 0 when you are outside ranges and 1 you are inside at least one;
- in\_db(Letter) replaces the default value 1 by Letter;
- out\_db(Letter) replaces the default value 0 by Letter;
- range\_position(Min,Max) allows to specify the position argument number of a term of the minimal and maximal value of term;
- range(Min, Max) extracts from the list of the complete annotation the sublist from position Min to position Max;

File Example toto.seq:

```
gb(3,5).
gb(7,9).
gb(8,11). % Overlap;)

Results of request are:

| ?- load_annotation_from_file(db,[],'toto.seq',R).
R = [0,0,1,1,1,0,1,1,1,1,1] ?
| ?- load_annotation_from_file(db,[in_db(c),out_db(nc)],'toto.seq',R).
R = [nc,nc,c,c,c,nc,c,c,c,c,c]?
| ?- load_annotation_from_file(db,[range(3,7)],R).
R = [1,1,1,0,1] ?
| ?- load_annotation_from_file(db,[in_db(c),out_db(nc),range(8,16)],'toto.seq',R).
R = [c,c,c,nc,nc,nc,nc,nc]?
```

• get\_sequence\_from\_file(++File,++Options,--Sequence): Generate from File of data a sequence (read: a list) of data Sequence. This predicate is a more efficient implementation of load\_annotation\_from\_file\4 with the type option sequence when several ranges of data have to be computed (option ranges)

We assumed that data facts of File are composed of facts with the format:

```
Term =.. [_Data_Functor,_Key,Range_Min,Range_Max,Data]
File Example toto.seq:
data(toto,1,10,[1,2,3,4,5,6,7,8,9,10]).
data(toto,11,20,[11,12,13,14,15,16,17,18,19,20]).
data(toto,21,30,[21,22,23,24,25,26,27,28,29,30]).

Result of the request is:
| ?- get_data_from_file('toto.seq',[],R).
R = [1,2,3,4,5,6,7,8,9,10,11,12,13,14,15,16,17,18,19,20,
```

Different options are available to specify the format of the data term:

- data\_position(Pos) to specify an other position for the data;
- left\_position(Left) to specify an other position for the left bound
  of the range that specifies the position of the data into the data file.
  When this left bound is not available, the keyword none for Pos can
  be used:

21,22,23,24,25,26,27,28,29,30] ?

 right\_position(Right) to specify an other position for the right bound of the range that specifies the position of the data into the data file. When this right bound is not available, the keyword none for Pos can be used.

```
File Example toto2.seq:
```

```
data(1,10,[1,2,3,4,5,6,7,8,9,10]).
data(11,20,[11,12,13,14,15,16,17,18,19,20]).
data(21,30,[21,22,23,24,25,26,27,28,29,30]).
```

Result of the request is:

To ask for specific ranges of data, two options are available:

- range(Min, Max) computes the sequence of data between Min and Max;
- ranges(List\_Ranges) computes sequences of data for a list of ranges with a format [[Min1,Max1],[Min2,Max2],...]. Note that the ranges should be ordered, i.e Min1 ≤ Min2 must be satisfied.

Result of the request is:

```
| ?- get_data_from_file('toto.seq',[range(5,10)],R).
R = [5,6,7,8,9,10] ?
| ?- get_data_from_file('toto.seq',[ranges([5,10],[5,8],[9,15])],R).
R = [[5,6,7,8,9,10],[5,6,7,8],[9,10,11,12,13,14,15]] ?
```

#### 3.2.2 Saving information to files

The io API also contains some predicates for saving sequences to a file.

• save\_annotation\_to\_sequence\_file(+KeyIndex,+ChunkSize,+Annotation,+File): Saves the data in the list given by Annotation to the file File in *sequence* format which is number of Prolog facts, on the form:

```
data(KeyIndex,1,5, [a,t,c,c,g]).
```

The first argument KeyIndex is used as an identifier. ChunkSize is the number of elements from Annotation to store with each fact. The second and third argument of a stored fact, corresponds to the start and end position in Annotation. The fourth argument of the fact is a list containing the relevant elements of the Annotation list. Note that if the the length of Annotation is not a multiple of ChunkSize, then the last fact stored will have a shorter range.

#### 3.2.3 Memory mapping of sequence files

Some times it can be convenient load a sequence file into memory and then be able to extract regions of the sequence on demand. The io API contains a method for doing just this.

The predicate to map a file in the format text(prolog(sequence(\_))) into memory is

```
load_sequence(+SequenceID, +Filename)
```

This will load the sequence from the file given by Filename and associate with the identifier SequenceID.

```
get_sequence_range(+SequenceID, +Min, +Max, -Data)
```

The predicate <code>get\_sequence\_range/4</code> can then be used to extract a part of the sequence. This is fairly efficient, but depends on the size of the data terms, |t|, in the file. The running time of the predicate is bounded by  $O(2|t| \times n)$ , where n = Max - Min.

#### 3.2.4 Splitting a Prolog file into multiple files

When working with large data files, it can sometimes be practical to split a large file with Prolog facts into multiple files, each having a distinct subset of the facts in the original file. The io API contains the predicate split\_file/2 for doing this:

The n facts in the InputFile will be divided into  $m=n/{\tt FactsPerFile}$  output files.

#### 3.3 The Stats API: stats

In this module (stats.pl), several predicates are defined to automatically compute frequencies, probabilities of occurrences of nucleotides, codons ... This computation is based on a simple counting method given a simple input data.

Variable Data\_Type is used to specify the type of the input data: Data type available are:

- nucleotide where input data are composed of letter from  $\{a, c, g, t\}$ ;
- codon where a codon is represented by a list of three letters from  $\{a, c, g, t\}$ ;
- amino\_acid where input data are composed of letter from  $\{a, c, d, e, f, g, h, i, k, l, m, n, p, q, r, s, t, v, w, y\}$ ;

• length. This type is used to compute frequencies and statistics about length of specific region of a genome.

Here is the description of main predicates of stats.pl

- stats(++Data\_Type,++Options,++Data,++Input\_Counting,-Result)
  Data\_Type setting define on which counting procedure is initialized. User
  must take care then to give the right Data. For {nucleotide,codon,amino\_acid}
  data type, Data must be a Prolog list with the right element, For length
  data type, Data must be a list of ranges represented by a list of two integer
  values ([Min,Max]). Two options are available:
  - order (Num) defines the size of the past stored to perform the counting
  - past(List) allows to give as input of the computation a previous past. This option is useful to make connection between two counting computations.

Input\_Counting is used as well to perform a counting computation on an input data divided into a set of lists. By default, counting computation is initialized when is a variable Input\_Counting. However, the counting can restart from Input\_Counting if this variable is unified to the result of a previous result. Finally, Result has the following format:

```
[(Past1,[(Elt1,Count11),(Elt2,Count12),...]),
(Past2,[(Elt1,Count21),(Elt2,Count22),...]),
...]
```

For example let consider this following data:

```
data('U00096',1,5,[a,g,c,t,t]).
data('U00096',6,10,[c,c,c,c,c]).
```

Here is the result of this following request

```
([c,a],[(a,0),(c,0),(g,0),(t,0)]),
([c,c],[(a,0),(c,3),(g,0),(t,1)]),
([c,g],[(a,0),(c,1),(g,0),(t,0)]),
([c,t],[(a,0),(c,0),(g,0),(t,1)]),
([g,a],[(a,0),(c,0),(g,0),(t,0)]),
([g,c],[(a,0),(c,0),(g,0),(t,1)]),
([g,g],[(a,0),(c,1),(g,0),(t,0)]),
([g,t],[(a,0),(c,0),(g,0),(t,1)]),
([t,a],[(a,0),(c,0),(g,0),(t,1)]),
([t,c],[(a,0),(c,0),(g,0),(t,1)]),
([t,g],[(a,0),(c,0),(g,0),(t,0)]),
([t,t],[(a,0),(c,0),(g,0),(t,0)]),
([t,t],[(a,0),(c,1),(g,0),(t,0)]),
([t,t],[(a,0),(c,1),(g,0),(t,0)]),
]?
```

• stats(++Data\_Type,++Options,++Data,++Input\_Counting,-Past,-Result) stats/6 computes exactly the same counting. This predicate allows only to record into Past the last past of the computing. This predicate is usefull to compute statistics on a series of data input. For example, this following request

can be used to obtain counting information on the previous data.

• normalize(++Data\_Type,++List\_Counting,-Probabilities) From the result of a counting procedure, normalize/3 modified List\_Counting to compute domains and probabilities distributions.

For example, here is the result of several requests:

To get a representation of the statistics where each count is represented as a single fact, the predicate

```
build_stat_facts(+Stats,-StatsFacts)
```

can be useful. For instance,

```
| ?- stats(nucleotide,[],[a,g,c,t,t],_,R), build_stat_facts(R,F).
R = [([],[(a,1),(c,1),(g,1),(t,2)])]
F = [stat([a],1),stat([c],1),stat([g],1),stat([t],2)] ?
```

### 3.4 The regex API

The regex API provides POSIX-like regular expressions for matching Prolog strings and atoms. A regular expression is created using

```
re_compile(+Regex, -CompiledRegex)
```

The input argument Regex is an atom or a list of symbols representing a particular regular expression. The output argument CompiledRegex is used in subsequent matching with this regular expression.

Matching with a regular expression is then done using

```
re_match(+CompiledRegex,+String,-Matches)
```

The goal re\_match/3 is true whenever the CompiledRegex matches the String. The String may be an atom or a list of symbols. If the goal is true, then Matches is a list containing the matching parts of the String corresponding to particular match groups (paranthesized sub-expressions) of the regular expression.

Currently the regular expressions can be expressed using a significant subset of the well-known POSIX regular expression operators. The following operators are supported:

- Concatenation: is expressed by the concatenation of two sub expressions. For instance, the regular expression ab matches an a followed by ab.
- Alternation: is expressed with the vertical bar (1). For instance the expression a|b matches either a or b.
- Repetition: Repetition operators indicate the number of times a preceding regular expression may be matched. The following repetition operators are supported:
  - representing the Kleene star, meaning that the preceding expression is matched zero or more times.
  - + the preceding expression one or more times.
  - ? The preceding expression zero times or one time.

Note that repetition binds stronger than alternation and alternation binds stronger than concatenation. To enforce a different binding order, enclosing sub-expressions in parentheses (which binds strongest) are used.

Parenthesised sub-expressions can also be used to create *match groups*. An outer-most paranthesized sub-expression, where the paranthesis are superflouos is interpreted as a match group. When an expression is matched, then the part of the string matched by the paranthesized expression is extracted.

Additionally, ranges are supported by the regular expression syntax. A range is expressed using square brackets. For instance, <code>[a-z0-9]</code> is range expression which matches a lower case alphanumeric character or a digit.

### 3.5 The genecode API

This API contains a set of genetic code tables. Each table defines a relation between codons and amino acids. The main predicate of the API is

```
genecode(+Code, ?Codon, ?AminoAcid)
```

The argument code is an integer specifying the genetic code to use. Currently, only genetic code 11 (bacterial) is supported. Codon is a list of of three nucleotides symbols from the alphabet [a,g,c,t]. The argument AminoAcid is a symbol from the alphabet

```
[a,c,d,e,f,g,h,i,k,l,m,n,p,q,r,s,t,v,w,y,*]
```

Where all symbols except \* represents an amino acid (the name of which starts which that letter). The special symbol \* signifies a stop codon in the specified genetic code.

Other predicates in the genecode API include

```
genecode_start_codons(+GeneCode,-StartCodons)
```

The genecode\_start\_codons/2 predicate unifies StartCodons with a list of all valid start codons for the given GeneCode.

```
genecode_stop_codons(+GeneCode,-StopCodons)
```

The genecode\_stop\_codons/2 predicate unifies StopCodons with a list of all valid stop codons for the given GeneCode.

# 3.6 The sequence API: sequence

The sequence API is library containing predicates for doing common tasks with biological sequences.

#### 3.6.1 Complementing DNA sequences

A complementary dna sequence can be achieved using the predicate

```
dna_seq_complement(+NucleotideSeq,-ComplNucleotideSeq)
```

When given the input sequence NucleotideSeq is a list of symbols from the alphabet [a,g,c,t], then ComplNucleotideSeq is unified to a list of complementary symbols, e.g.

```
| ?- dna_seq_complement([a,g,a,c,t,a],X).
X = [t,c,t,g,a,t] ?
yes
```

Typically, when complementing a DNA sequence, we are actually interested in the *reverse* complement. To achieve this, use the built-in predicate <code>reverse/2</code> before <code>dna\_seq\_complement/2</code>.

#### 3.6.2 Translating a DNA sequence to amino acids

The predicate,

```
dna_amino_acid(+Genecode, ?DNA_Sequence,?AminoAcids)
```

Translates a DNA sequence to a sequence of amino acids or vice versa; produces all possible DNA sequences for a given amino acids sequence. Genecode is an integer for the genetic code to be used in the translation. DNA\_Sequence is a list of symbols from the alphabet [a,g,c,t] and length of the list is expected to be divisible by three. AminoAcids is a list of symbols from the alphabet,

```
[a,c,d,e,f,g,h,i,k,l,m,n,p,q,r,s,t,v,w,y,*].
```

# Chapter 4

# Models

Pre-defined models are introduced. These models are used to build more complex models.

### 4.1 Parsers of Biological Data

To extract information from different Biological database, several parsers have been designed to parse report of analyses (Blast, Easygene and Genemark) and database (Genbank). These parsers generated a series of Prolog terms that can be used after that input of different probabilistic models. script\_parser.pl collects different scripts to generate different data files.

#### 4.1.1 Parser\_fna

\*.fna file of Genbank is composed of a complete genome in the FASTA format. Parser\_fna permits to parse this \*fna.file from Genbank and generate list of terms. These terms store the genome into a Prolog list composed of  $\{a,c,g,t\}$ . Two scripts are implemented:

```
parser_fna(++Name_FNA_File,++Name_GBK_File,++Options) and
parser_fna(++Name_FNA_File,++Name_GBK_File,++Options,--OutputFile).
Note that *.gbk file is necessary as well. This file is used to automatically extract genome information (Genbank key and size of genome).
```

Output format of the generated terms are: data(Genebank\_Key,Start,End,List\_of\_Data). Option list(Number) can be used to divide the complete genome into several lists with a length defined by the parameter Number. Example with the E.Coli K12 genome:

```
>gi|48994873|gb|U00096.2| Escherichia coli .....
AGCTTTTCATTCTGACTGCAACGGGCAATATGTCTCTGTGTGG.....
.....
Result by default:
```

```
%>gi|48994873|gb|U00096.2| Escherichia coli .... data('U00096',1,4639675,[a,g,c,t,t,t, ...]).

Result when list(280) option is used

%>gi|48994873|gb|U00096.2| Escherichia ..... data('U00096',1,280,[a,g,c,t,...]).
data('U00096',281,560,[c,c,c,...]).
```

#### 4.1.2 Parser\_ptt

A PTT file is the NCBI format for representation of a protein table. Typically, ptt files are composed of information about known or predicted protein coding genes in a genome. Each such gene is represented by a single line where tabulation separated fields represent various properties of the gene such as the position, length, name and protein product. The parser\_ptt model parses PTT files and generates a Prolog term in the output file for each gene in the input (ptt) file. We use the format identifier, text(ptt), for PTT files, which is also the input format of parser\_ptt. The output format is text(prolog(ranges(gene))). The model does not declare any options.

Consider the following extract of a PTT file:

```
Location Strand Length PID Gene Synonym Code COG Product 190..255 + 21 1786182 thrL b0001 - - thr operon leader peptide
```

The first line is just a header describing the names of the individual fields, but the second is an entry for a particular gene. The corresponding Prolog term generated by the parser for this gene is then

```
gb(190,255,'+',1,[
    gene_name(thrL),length(21),pid(1786182),
    synonym(b0001),code('-'),cog('-'),
    product('thr operon leader peptide')]
).
```

Additionaly, the file \$SCRIPTS/script\_parser.pl implements two helper predicates to make it easy to use this parser:

```
parser_ptt(++Name_PTT_File) and
parser_ptt(++Name_PTT_File,,--OutputFile).
```

#### 4.1.3 Parser\_Easygene

This parser allows to generate from the GFF format of Easygene prediction several Prolog terms with the following format: eg(Left,Right,Dir,Frame,[]).

```
Two scripts are implemented: parser_easygene(++Report_Name) and
```

```
parser_easygene(++Report_Name) and
parser_easygene(++Report_Name,--OutputFile).
```

#### Parser\_Genemark

This parser allows to generate from a report of Genemark.HMM of Easygene prediction several Prolog terms with the following format: gm(Left,Right,Dir,Frame,[]). Genemark report of prediction coordinates is parsed.

```
Two scripts are implemented:
parser_genemark(++Report_Name) and
parser_genemark(++Report_Name,--OutputFile).
```

#### 4.1.5Parser\_Blast

This parser translated into Prolog terms the XML report of Tblastn. The parser generates fact for each hits detected of the XML report.

#### 4.2 Models for measurement and statistical reports

#### 4.2.1 accuracy\_report

The model accuracy\_report can be used the produce a report of various measures of the accuracy of particular gene predictions compared with a golden standard such a genebank.

To use the model, you need to call get\_annotation\_file/4 with the following arguments,

```
get_annotation_file(accuracy_report,
    [ReferenceFile, PredictionFile],
     option(reference_functor, RefFunctor),
     option(prediction_functor, PredFunctor),
     option(start, StartPos),
     option(end, EndPos)
    ],
    OutputFile),
```

ReferenceFile must be the full path to a file with facts representing the "correct predictions". PredictionFile must be the full path to a file with facts representing the predictions. Both files must be a a db type format, with facts on the following form,

```
functor(To, From, Strand, ReadingFrame, Name).
```

Each such fact represent a prediction in the PredictionFile or correct gene in the in ReferenceFile. The functor is a any given functor, but the ReferenceFile and the PredictionFile should use different functors. The To argument represents the position in the genome where the prediction begins (inclusive) and the From argument represents the end position of the prediction

ReadingFrame is a integer in the range  $\{1,2,3\}$  Strand is either + for the forward strand or - for the reverse strand.

get\_annotation\_file for the accuracy\_report model must be called with
four mandatory options:

- reference\_functor: The functor used in the ReferenceFile
- prediction\_functor: The functor used in the PredictionFile
- start: An integer corresponding to the beginning of the range on which accuracy should be measured.
- end: An integer corresponding to the end of the range (inclusive) on which accuracy should be measured.

#### 4.2.2 The range\_stats model

The range\_stats model calculates statistics for each gene or similar from a file in the format text(prolog(ranges(gene))). The output file is also in the text(prolog(ranges(gene))) format, with an entry for each range in the original file, but with some statistics facts appended to the ExtraList.

The model takes two input files,

- A ranges file in text(prolog(ranges(gene))) format.
- A data file in text(prolog(sequence(gene))) format.

The model can generate statistics basic nucleotide frequencies, amino acids frequencies and length. The nucleotide sequence for the range is automatically extracted from the *data file* and translated or reverse complemented as necessary. It is possible to adjust which kinds of statistics that should be generated by supplying the model with relevant options:

- amino\_acid\_stats: Whether to add statistics based on amino acid frequency. Values are yes (default) or no.
- nucleotide\_stats: Whether to add statistics based on nucleotide frequency. Values are yes (default) or no.
- length\_stats: Whether to add statistics based on the length of the range. Values are yes (default) or no.
- max\_nucleotide\_order(N): Is an integer that determines the maximal order of nucleotide n-grams to calculate statistics for. Given this option the model will calculate statistics for nucleotide n-grams of size N+1 down to size 1. The default value is 1, which that single nucleotide frequency and di-nucleotide frequency will be calculated.

- max\_amino\_acid\_order(N): Is an integer that determines the maximal order of n-grams to calculate statistics for. Given this option the model will calculate statistics for amino acid n-grams of size N+1 down to size 1. The default value is 0 (e.g. only single amino acid frequencies).
- genecode: An integer which identifies the gene code table to use when translating to amino acid sequence. Default is 11 (bacterial).

Depending the on the options for the model a subset of the following facts are appended to the ExtraList of each gene in the output file:

- nucleotide\_stats(L) where L is list of elements of the type stat(Seq,Freq) and Seq is list of nucleotides signifying a particular n-gram (n being the length of list) and Freq is the frequency of the n-gram in the gene.
- amino\_acid\_stats(L) where L is a list of elements of the type stat(Seq,Freq) and Seq is list of amino acid symbols signifying a particular n-gram (n being the length of list) and Freq is the frequency of the n-gram in the gene.
- normalized\_gene\_length(F) where F is a number in the range  $0\cdots 1$  representing the proportion of the length of the range relative to the longest gene in the file. For instance, if the longest gene in the file is 2000 nucleotides long and the gene in question is 1000 nucleotides long, then the normalized gene length will be 1000/2000 = 0.5.

# 4.3 Prediction/Gene filter models

Common for these models is that they all take files in the text(prolog(ranges(\_))) format and produce output in the same format. Usually, the output will be a subset of the entries from the input file.

#### 4.3.1 longest\_predication\_per\_stop\_codon

Given an input file with gene predictions, this model selects for each stop codon the longest matching prediction. Only the longest prediction for each stop coden will be written to the output file.

For the goal annotate/3, the following files and options are expected:

- Input files: text(prolog(ranges(gene)))
- Output file: text(prolog(ranges(gene)))

The model only takes one option, file\_functor, which is used to set the functor to identify the gene predictions in the input file. If the input file contains only facts with the same functor the default value auto can be used and the model will infer the functor automatically.

#### 4.3.2 best\_prediction\_per\_stop\_codon

Given an input file with gene predictions this model selects, for each stop codon, the prediction with the highest score and write such predictions to the output file. The score is for each prediction is assumed to be present the Extra list argument of the prediction facts in the input file.

For the goal annotate/3, the following files and options are expected:

- Input files: text(prolog(ranges(gene)))
- Output file: text(prolog(ranges(gene)))

The model only takes two option:

- file\_functor: Is used to set the functor to identify the gene predictions in the input file. If the input file contains only facts with the same functor the default value auto can be used and the model will infer the functor automatically.
- score\_functor: Is used to specify how the score is represented in the Extra list. For instance, if the Extra list looks like [...,score(0.75),..], then the score\_functor option should be set to score. The argument of the score functor (0.75 in the example) should be a number or at least something comparable with the standard less-than-or-equal-to Prolog operators.

#### 4.3.3 gene\_filter

The gene\_filter model is a model that filters a set of genes based on various matching criteria. The input to the model is a file in text(prolog(ranges(gene))) format containing the list of genes to be filtered and a file in text(prolog(sequence(dna))) that includes the nucleotide sequence for the entire genome to which the genes belong.

The model supports filtering based on a wide range of criteria, specified using the options given to the model. The model is invoked using get\_annotation\_file/4. The options supported by the model are:

- match\_frames: A list of valid frames. Genes in other frames will be filtered. The list must be a subset of the *default value*: [1,2,3].
- match\_strands: A list of valid strands. Genes occurring an other strand will be filtered. The list must be a subset of the default value: ['+', '-'].
- exact\_match\_extra\_fields: A list of Prolog terms terms each of which must unify with an element of the Extra list of the gene terms. Genes which do not are filtered.
- exact\_no\_match\_extra\_fields: A list of Prolog terms terms any of which may not unify with any element of the Extra list of the gene terms. Genes which do are filtered.

- regex\_match\_extra\_fields: A list of regular field names in the Extra list and regular expressions to match those fields. For example, a regular expression matcher for the name field would be specified as name('^y.\*\$'), here matching genes with a name starting with the letter y. The syntax of the regular expressions are as specified in section 3.4. Gene tems that do not match each of the specified regular expressions are filtered.
- regex\_no\_match\_extra\_fields: A list of field names in the extra and regular expressions to match those fields. If a gene is matched by any of these regular expression, it will be filtered.
- match\_protein\_coding: Must be either yes or no. Default is no. If set to yes, this will filter gene terms which do not have a valid start and stop codon. The codon table to be used is specified using the the genecode option.
- genecode: An integer specifying what genecode to use. Default is 11, which is the bacterial gene code. See 3.5 for a list of valid gene code tables.
- invert\_results: Must be either yes or no. Default is no. If set to yes, then only the otherwise non-filtered results are filtered.

### 4.4 Model that integrate other programs

#### 4.4.1 Genemark

To be documented...

#### 4.4.2 Glimmer3

To be documented...

#### 4.5 Various models

#### 4.5.1 hard\_to\_find\_genes

The model hard\_to\_find\_genes ranks a given set of genes according to how many genefinders that find the gene.

The input to the model is a list of files, of which the first element is taken to the *golden standard* - a file containing a set of true genes. The rest of the n files in the input file list are files that contain gene predictions of various genefinders (one genefinder per input file). All the files are expected to be in the text(prolog(ranges(gene))) format.

For each gene in the *golden standard* file, the model checks which of the gene finders predicts this gene. It writes each of these genes to the output file, but with facts added to the Extra list. The added facts are

- found\_by\_genefinders(F): Where F is a list of n entries containing
  - 1 if the *n*th gene finder predicts the gene.
  - 0 if the nth gene finder did not predict the gene.
- gene\_finding\_difficulty\_score(S): Where S is a decimal number between zero and one representing the inverse of the proportion of genefinders that predicts this gene. E.g. if it is zero, no gene finders predicts the gene, and if it is one, then all n gene finders predicts the gene.

The model take a number of options:

- start: Is a positive integer indicating the start of the range in which to consider genes. Only genes starting/ending after this position are included in the output file. The default value min is taken to mean the start of the file, e.g. position 1.
- end: Is a positive integer indicating the end of the range in which to consider genes. Only genes with a right border below this value included in the output file. The default value max is taken to mean the maximal position of any gene in the *golden standard* file.
- gene\_match\_criteria: Is used to determine what is required for a gene prediction to *match* a gene. If the value is set to start\_and\_stop (default), then both start codon and stop codon needs to match. If the value is set to stop then only the stop codon needs to match.

# Chapter 5

# File formats

The chapter documents the different types of file formats used in the lost framework. The naming of file formats follow a simple scheme using nested Prolog functors where the outermost functor is the most general and the innermost is the most specific description of the file format. For instance, consider the naming for a DNA sequence expressed as a Prolog file,

```
text(prolog(sequence(dna)))
```

The outermost functor text specifies that the file is a text file, and the next functor prolog says that the text file contains Prolog code. The next one, sequence, specifies the type of data (sequence data) we expect to be expressed in the prolog facts finally, the innermost functor dna specifies the type of sequence that we are dealing with.

# 5.1 text(prolog(\_))

text(prolog(\_)) formats contain Prolog facts. The functor and arity of those facts cannot be determined by knowing that it is text(prolog(\_)), but needs specification using further embbed functors.

#### 5.1.1 text(prolog(sequence(\_)))

This format should be specified for sequence data expressed as prolog facts. For instance, a file of this format may contain facts like the ones below, which specifies the alphabet,

```
data(alphabet,1,10, [a,b,c,d,e,f,g,h,i,j]).
data(alphabet,11,20],[k,l,m,n,o,p,q,r,s,t]).
data(alphabet,21,27,[u,v,w,x,y,z]).
```

The form of these facts are,

functor(Identifier,To,From,SequenceElementList).

The format does not dictate the functor of the facts (e.g. data), nor the size of the data list in the fourth argument. However, the range expressed by To and From should correspond to the numer of elements in SequenceElementList.

#### text(prolog(sequence(dna)))

Like the above format but restricts the alphabet of the data elements in the SequenceElementList to the set  $\{a,g,c,t\}$ .

```
text(prolog(sequence(rna)))
```

Like the above format but restricts the alphabet of the data elements in the SequenceElementList to the set  $\{a,g,c,u\}$ .

```
text(prolog(sequence(amino_acids)))
```

Like the above format but restricts the alphabet of the data elements in the SequenceElementList to the set {a,c,d,e,f,g,h,i,k,l,m,n,p,q,r,s,t,v,w,y}.

#### 5.1.2 text(prolog(ranges(\_)))

The format consists of Prolog facts, each of which contain an annotation for a particular range of some sequence

```
F =.. [ Functor, SequenceId, LeftEnd, RightEnd | _ ]
```

The functor may vary depending on the type of annotation but is expected to be same within a file. Similarly, SequenceId is an atom which serves as sequence identifier. The next two arguments, LeftEnd and RightEnd are positive integers which specifies the range relative to the sequence. Both are inclusive. It must be the case that LeftEnd  $\leq$  RightEnd.

```
text(prolog(ranges(gene)))
```

This is a compatible subtype text(prolog(ranges(\_))). The format consists of Prolog facts, each of which contain an gene annotation for a particular range of some DNA sequence (all facts refer to the same sequence).

The facts are on the form,

 $\verb|functor(SequenceId, LeftEnd, RightEnd, Strand, Frame, ExtraList)|.\\$ 

The functor may vary depending on the type of annotation but is expected to be same within a file. The first argument SequenceId is a sequence identifer. Is is recommended that the Genbank accession id is used where possible. The next two arguments, LeftEnd and RightEnd are positive integers which specifies the range of the annotation relative to the DNA sequence. Both are inclusive. It must be the case that LeftEnd  $\leq$  RightEnd. The Strand argument is '+' for the forward strand or'-' for the reverse strand. The Frame argument is one

of  $\{1,2,3\}$ . The final argument, ExtraList is a list possibly containing extra information. Each element in the ExtraList is on the form F = ... [ Key, Value ] where Key must be unique in the list. There are no restrictions on Value.

#### 5.1.3 text(prolog(prism\_switches))

This is the format used by PRISM to save and load parameter files.

#### 5.1.4 text(ptt)

TODO ptt files

#### 5.1.5 text(easygene\_report)

TODO ptt files

#### 5.1.6 text(fasta(SequenceType)

A sequence in FASTA format begins with a single-line description, followed by lines of sequence data. The description line is distinguished from the sequence data by a greater-than (" $\dot{\iota}$ ") symbol in the first column. The word following the " $\dot{\iota}$ " symbol is the identifier of the sequence, and the rest of the line is the description (both are optional). There should be no space between the " $\dot{\iota}$ " and the first letter of the identifier. It is recommended that all lines of text be shorter than 80 characters. The sequence ends if another line starting with a " $\dot{\iota}$ " appears; this indicates the start of another sequence.

The format identifier text(fasta(\_)) is used to refer to any type of file in FASTA format, but we distinguish between different subtypes with SequenceType. A ground value for SequenceType may be one of the following:

- fasta: Specifies generic fasta format. This is the same as specifying text(fasta(\_)).
- fna: fasta nucleic acid.
- ffn: FASTA nucleotide coding regions. Contains coding regions for a genome.
- ffa: fasta amino acid.

# Chapter 6

# Predicates listing

### 6.1 /lost.pl

- $lost\_config(++Type,++Value)$
- lost\_include\_api(++ApiName)
- lost\_include\_script(++ScriptName)

# 6.2 /lib/accuracy.pl

- accuracy\_stats(++RefFunctor,++PredFunctor,++Start,++End,-OutputFile)
- sensitivity(++TruePos,++FalseNeg,-Sens)
- specificity(++TruePos,++FalsePos,-Spec)
- specificity\_traditional(++TrueNeg,++FalsePos,-ST)
- $\bullet$  correlation\_coefficient (++TruePos,++FalsePos,++TrueNeg,++FalseNeg,-CC)
- simple\_matching\_coefficient(++TruePos,++FalsePos,++TrueNeg,++FalseNeg,-SMC)
- $\bullet \ \, average\_conditional\_probability(++TruePos,++FalsePos,++TrueNeg,++FalseNeg,-ACP) \\$
- $\bullet$  approximate\_correlation (++TruePos,++FalsePos,++TrueNeg,++FalseNeg,-AC)
- count\_genes(++Functor,++Start,++End,-Count)
- correct\_genes(++RefFunctor,++PredFunctor,++Start,++End,-Correct\_Genes)

- missing\_genes(++RefFunctor,++PredFunctor,++Start,++End,-Missing\_Genes)
- wrong\_genes(++RefFunctor,++PredFunctor,++Start,++End,-Wrong\_Genes)
- number\_of\_correct\_genes(++RefFunctor,++PredFunctor,++Start,++End,-Num\_Correct)
- number\_of\_missing\_genes(++RefFunctor,++PredFunctor,++Start,++End,-Num\_Missing)
- number\_of\_wrong\_genes(++RefFunctor,++PredFunctor,++Start,++End,-Num\_Wrong)
- gene\_finding\_difficulty\_report(++RefFunctor,++PreducFunctors,++RangeMin,++RangeMax
- gene\_difficulty\_score(++NumGeneFinders,++NumFoundGene,-DifficultyScore)
- report\_nstats(++RefFunctor,++PredicFunctor,++Start,++End)
- strand(??Strand)
- reading\_frame(??Frame)

GenePredictionDifficultyList)

### 6.3 /lib/autoAnnotations.pl

- prismAnnot(++File)
- prismAnnot(++File,++CompilationOptions)
- viterbiAnnot(++Goal,-Proba)

# 6.4 /lib/chmm.pl

- init\_store
- init\_constraint\_store(++Constraint,-Store)
- forward\_store(++Store)
- $get\_store(++S)$
- init\_constraint\_stores(++Constraints,-Stores)
- $\bullet \ \, init\_constraint\_store(++Constraint,-Store)\\$
- $\bullet \ \ check\_constraints(++StateUpdate)$
- check\_each\_constraints(++StateUpdate,++Constraints,++StoreBefores,-StoreAfters)
- $\bullet$  constraint\_check(++Constraint,++StateUpdate,++StoreBefore,-StoreAfter)

### 6.5 /lib/fasta.pl

- Maybe to merge with the fna parser?
- fasta\_load\_sequence(++InputFile,++SequenceId,-FastaHeader,-Sequence)
- fasta\_save\_sequence(++OutputFile,++SequenceData,++Header)
- atom\_code\_list(++Codes,,-Atoms)

### 6.6 /lib/genecode.pl

- genecode(++TableNum,??Codon,??Amino\_Acid)
- genecode\_start\_codon(++TableNum,??Codon,??Amino\_Acid)
- genecode\_stop\_codon(++TableNum,??Codon)
- genecode\_start\_codons(++GeneCode,-StartCodons)
- $\bullet \ genecode\_stop\_codons(++GeneCode, -StopCodons) \\$

# 6.7 /lib/genedb.pl

• To remove??

# 6.8 /lib/interface.pl

- >(++Call)
- get\_annotation\_file(++Model,++Inputs,++Options,-FileName)
- run\_model(++Model,++Inputs,++Options,-FileName)
- train\_model(++Model,++TrainingDataFiles,++Options,-SavedParamsFile)
- get\_option(++Options,-KeyValue)
- get\_option(++Options,-Key, -Value)
- launch\_prism\_process(++PrismFile,-Goal)
- lost\_tmp\_directory(-Dir)
- lost\_models\_directory(-Dir)
- lost\_data\_directory(-Dir)
- $\bullet$  lost\_model\_directory(++Model,-Dir)

- lost\_model\_parameters\_directory(++Model,-Dir)
- lost\_model\_annotations\_directory(++Model,-Dir)
- lost\_model\_interface\_file(++Model,-ModelFile)
- lost\_model\_parameter\_index\_file(++Model,-IndexFile)
- lost\_annotation\_index\_file(++Index)
- lost\_data\_file(++SequenceID,-SequenceFile)
- is\_generated\_file(++File)
- lost\_file\_index\_get\_filename(++Index,++Model,++InputFiles,++Options,-Filename)
- $\bullet \ \, check\_valid\_model\_call(++Model,++InterfacePredicate,++Arity,++Options)\\$
- declared\_model\_options(++Model,++Goal,-DeclaredOptions)
- $\bullet \ lost\_interface\_input\_formats(++Model,++InterfacePredicate,-Formats)\\$
- $\bullet$  lost\_interface\_defines\_output\_format(++Model,-InterfacePredicate)
- $\bullet \ lost\_model\_option\_values(++Model,++InterfacePredicate,-OptionName,-Value) \\$
- rm\_gen
- $\bullet$  rm\_tmp
- list\_lost\_models(-Models)

# 6.9 /lib/io.pl

- $\bullet$  get\_data\_from\_file(++File,++Options,-Data)
- get\_data\_from\_terms(++Terms,++Options,-Data)
- empty\_lists(??List)
- load\_annotation\_from\_file(++Type,++Options,++File,-Annotation)
- $\bullet \ \ sequence\_terms\_to\_annotations(++Options,++ListTerms,-Annotation)$
- $\bullet \ \ db\_terms\_to\_annotations(++Options,++ListTerms,-Annotation)$
- $\bullet$  get\_next\_range(++Range\_Position,++DB,-Range)
- $\bullet$  save\_annotation\_to\_sequence\_file (++KeyIndex,++ChunkSize,++Annotation,-File)

- load\_sequence(++SeqId,-FileName)
- unload\_sequence(++SeqId)
- $\bullet \ \ get\_sequence\_range(++SeqId,++Min,++Max,-Data)\\$
- terms\_from\_file(++File,-Terms)
- terms\_to\_file(++File,++Terms)
- file\_functor(++FileName,-Functor)
- $\bullet \ \, \text{spilt\_file}(++\text{Filename},++\text{ChunkSize},++\text{OutputFilePrefix},++\text{OutputFileSuffix})\\$

### 6.10 /lib/misc\_utils.pl

- $\bullet \ \, atom\_concat\_list(++List,\!-Atom) \\$
- atom\_list\_code\_list(++Atoms,-Codes)
- $inlists\_nth0(++List,++Nth,-ListElt)$
- flexible\_append(++List1,++List2,-List)
- flatten\_once(++List,-ListFlatten)
- map(++Functor,++InList,-OutList) :-
- list\_head(++List,-Head)
- list\_tail(++List,-Tail)
- rotate\_list\_vector(++ListOfLists,-RotatedList)
- $\bullet$  replace(++Symbol,++Replacement,++Inlist,-Outlist)
- match\_tail(++InputList,-HeadOfInputList,++TailOfInputList)
- $not\_member(++Elt,++List)$
- terms\_has\_rule\_with\_head(++Terms,++Functor,Arity)
- abs(++Elt,-Abs) RK: Not ISO predicate?
- min(++Elt1,++El2,-Min) RK: Not ISO predicate?
- max(++Elt1,++Elt2,-Max) RK: Not ISO predicate?
- $list_max(++List,-Max)$
- list\_min(++List,-Min)
- atom2integer(++Atom,-Integer)

- check\_or\_fail(++Goal,++Error)
- check\_or\_warn(++Check,++Error)
- dirname(++Filename,-DirPartAtom)
- $\bullet \ \operatorname{overlaps}(++\operatorname{Start1},++\operatorname{End1},++\operatorname{Start2},++\operatorname{End2})$
- sum\_range\_list(++Ranges,-Sum)
- upper\_lower(??UpperLower,??UpperLower)
- is\_upper\_case\_alphanumeric(??Code)
- is\_lower\_case\_alphanumeric(??Code)

### 6.11 /lib/regex.pl

- re\_compile(++RegexAtom,-Regex): compiles an atom representing a regular expression to a Prolog list representation
- re\_match(++Regex,++Atom,-Matches): tries to match Atom with the regular expression Regex

# 6.12 /lib/sequence.pl

- Remark: maybe to merge in one api for this kind of stuff ??(merge with genecode and some predicate of stats)
- dna\_seq\_complement(++List,-ListComplement)
- $\bullet$  dna\_complement(++Nuc,-Complement)
- dna\_translate(++GeneCode,??NucList,??AAList)

# 6.13 /lib/stats.pl

- stats(++Type,++Options,++Data,++InputCounting,-Result)
- get\_init(++Type,-InitCounting)
- update\_counting(++Datum,++Past,++InputCounting,-CountingUpdate)
- get(++Type,-Value) RK: to merge with genecode
- $\bullet \ \ normalize(++Type,++Counting,-Probabilities)\\$
- sum\_list(++List,-Sum) RK: already define
- build\_stat\_facts(++Counting,-ListFacts)

- $\bullet$  normalize\_stat\_facts(++Facts,-NormalizedFacts)
- count\_all(++Facts,-Count)

### 6.14 /lib/utils\_parser\_report.pl

- parser\_line(++EntryCodes,++ListTokens)
- parser\_line(++EntryCodes,++IgnoredChars,++ListTokens)
- is\_number(++ListCodes)
- readline(++Stream,-CodeList)
- read\_tab(++Stream,-CodeList)
- read\_token(++Stream,-NSpaces,-CodeList)
- read\_n\_spaces(++Stream,++Number)

### 6.15 /lib/xml.pl

- xml2pl(++Input,-Output)
- pl2xml(++Document,-Codes)
- xml-parse(+Controls,+?Chars,?+Document): parses Chars to/from a data

# 6.16 /models

- accuracy\_report
- best\_prediction\_per\_stop\_codon
- chunk\_aa\_conservation
- chunk\_translator
- cluster\_gene\_finder
- consorf\_genefinder
- format\_converter
- gene\_filter
- genebank\_annotator
- genemark
- genemark\_genefinder

- $\bullet$  glimmer3
- $\bullet~{\rm hard\_to\_find\_genes}$
- $\bullet \ \, \mbox{hein\_genefinder}$
- $\bullet \ longest\_prediction\_per\_stop\_codon$
- $\bullet$  lost\_genefinder
- mork\_hmm
- $\bullet \ \ orf\_annotator$
- $\bullet \ \, {\rm orf\_chopper}$
- $\bullet \ parser\_blastxml$
- parser\_easygene
- $\bullet \ parser\_fna$
- $\bullet \ \, parser\_genemark$
- $\bullet \ parser\_ptt$
- $\bullet$  range\_stats
- $\bullet$  sample\_model1
- $\bullet \ sample\_model2$

# Chapter 7

# Notes and stuff

#### 7.1 Feature wish list

• Division of models into models (probabilistic models) and nodes (which are just data processing).

•

- Document gene database file format and make sure that all models adhere to this format.
- Option, input and output formats for all models. (Matthieu, Christian)
- Data outside git (Christian)
- Models should lost\_tmp\_directory for intermediary files.
- Document all of undocumented models. (Matthieu, Christian)

Some things that we have discussed, but decided not to do

• Do something to avoid multiple declaration of lost\_include\_api everywhere... (Matthieu).

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