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# **Chapter 1**

# ViennaRNA Package core - RNAlib

A Library for folding and comparing RNA secondary structures

#### Date

1994-2010

#### **Authors**

Ivo Hofacker, Peter Stadler, Ronny Lorenz and many more

#### **Table of Contents**

- Introduction
- Folding Routines Functions for Folding RNA Secondary Structures
- · Parsing and Comparing Functions to Manipulate Structures
- Utilities Odds and Ends
- Example A Small Example Program
- References

## 1.1 Introduction

The core of the Vienna RNA Package is formed by a collection of routines for the prediction and comparison of RNA secondary structures. These routines can be accessed through stand-alone programs, such as RNAfold, RNAdistance etc., which should be sufficient for most users. For those who wish to develop their own programs we provide a library which can be linked to your own code.

This document describes the library and will be primarily useful to programmers. However, it also contains details about the implementation that may be of interest to advanced users. The stand-alone programs are described in separate man pages. The

latest version of the package including source code and html versions of the documentation can be found at

http://www.tbi.univie.ac.at/~ivo/RNA/

# **Chapter 2**

# Folding Routines - Functions for Folding RNA Secondary Structures

#### **Table of Contents**

- Calculating Minimum Free Energy Structures
- Calculating Partition Functions and Pair Probabilities
- Searching for Predefined Structures
- Enumerating Suboptimal Structures
- Predicting hybridization structures of two molecules
- Predicting local structures of large sequences
- Predicting Consensus Structures from Alignment
- · Global Variables for the Folding Routines
- · Reading Energy Parameters from File

## 2.1 Calculating Minimum Free Energy Structures

The library provides a fast dynamic programming minimum free energy folding algorithm as described by Zuker & Stiegler (1981).

## Associated functions are:

```
float fold (char* sequence, char* structure);
```

Compute minimum free energy and an appropriate secondary structure of an RNA sequence.

```
float circfold (char* sequence, char* structure);
```

Compute minimum free energy and an appropriate secondary structure of an RNA sequence assuming it to be circular instead of linear.

Calculate the free energy of an already folded RNA.

Calculate the free energy of an already folded circular RNA.

```
void update_fold_params(void);
```

Recalculate energy parameters.

```
void free_arrays(void);
```

Free arrays for mfe folding.

#### See also

fold.h, cofold.h, 2Dfold.h, Lfold.h, alifold.h and subopt.h for a complete list of available functions.

## 2.2 Calculating Partition Functions and Pair Probabilities

Instead of the minimum free energy structure the partition function of all possible structures and from that the pairing probability for every possible pair can be calculated, using a dynamic programming algorithm as described by McCaskill (1990). The following functions are provided:

Compute the partition function Q of an RNA sequence.

```
void free_pf_arrays (void)
```

Free arrays from pf\_fold()

```
void update_pf_params (int length)
```

Recalculate energy parameters.

Get the centroid structure of the ensemble.

Get the centroid structure of the ensemble.

Get the mean base pair distance in the thermodynamic ensemble.

#### See also

part\_func.h, part\_func\_co.h, part\_func\_up.h, 2Dpfold.h, LPfold.h, alifold.h and MEA.h for a complete list of available functions.

## 2.3 Searching for Predefined Structures

We provide two functions that search for sequences with a given structure, thereby inverting the folding routines.

Find sequences with predefined structure.

Find sequence that maximizes probability of a predefined structure.

The following global variables define the behavior or show the results of the inverse folding routines:

```
char *symbolset
```

This global variable points to the allowed bases, initially "AUGC".

#### See also

inverse.h for more details and a complete list of available functions.

## 2.4 Enumerating Suboptimal Structures

Returns list of subopt structures or writes to fp.

Returns list of circular subopt structures or writes to fp.

```
SOLUTION *zukersubopt(const char *string);
```

Compute Zuker type suboptimal structures.

Sample secondary structure representatives from a set of distance classes according to their Boltzmann probability.

```
char *alipbacktrack (double *prob)
```

Sample a consensus secondary structure from the Boltzmann ensemble according its probability

```
char *pbacktrack(char *sequence);
```

Sample a secondary structure from the Boltzmann ensemble according its probability

```
char *pbacktrack_circ(char *sequence);
```

Sample a secondary structure of a circular RNA from the Boltzmann ensemble according its probability.

#### See also

subopt.h, part\_func.h, alifold.h and 2Dpfold.h for more detailed descriptions

## 2.5 Predicting hybridization structures of two molecules

The function of an RNA molecule often depends on its interaction with other RNAs. The following routines therefore allow to predict structures formed by two RNA molecules upon hybridization.

One approach to co-folding two RNAs consists of concatenating the two sequences and keeping track of the concatenation point in all energy evaluations. Correspondingly, many of the cofold() and co\_pf\_fold() routines below take one sequence string as argument and use the global variable cut\_point to mark the concatenation point. Note that while the *RNAcofold* program uses the '&' character to mark the chain break in its input, you should not use an '&' when using the library routines (set cut\_point instead).

In a second approach to co-folding two RNAs, cofolding is seen as a stepwise process. In the first step the probability of an unpaired region is calculated and in a second step this probability of an unpaired region is multiplied with the probability of an interaction between the two RNAs. This approach is implemented for the interaction between a long target sequence and a short ligand RNA. Function pf\_unstru() calculates the partition function over all unpaired regions in the input sequence. Function pf\_interact(), which calculates the partition function over all possible interactions between two sequences, needs both sequence as separate strings as input.

```
int cut_point
```

Marks the position (starting from 1) of the first nucleotide of the second molecule within the concatenated sequence.

Compute the minimum free energy of two interacting RNA molecules.

```
void free_co_arrays (void)
```

Free memory occupied by cofold()

## **Partition Function Cofolding**

To simplify the implementation the partition function computation is done internally in a null model that does not include the duplex initiation energy, i.e. the entropic penalty for producing a dimer from two monomers). The resulting free energies and pair probabilities are initially relative to that null model. In a second step the free energies can be corrected to include the dimerization penalty, and the pair probabilities can be divided into the conditional pair probabilities given that a re dimer is formed or not formed.

Calculate partition function and base pair probabilities.

```
void free_co_pf_arrays(void);
```

Free the memory occupied by co pf fold()

### Cofolding all Dimeres, Concentrations

After computing the partition functions of all possible dimeres one can compute the probabilities of base pairs, the concentrations out of start concentrations and sofar and soaway.

Compute Boltzmann probabilities of dimerization without homodimers.

```
ConcEnt *get_concentrations(double FEAB,
double FEAA,
double FEBB,
double FEA,
double FEB,
double * startconc)
```

Given two start monomer concentrations a and b, compute the concentrations in thermodynamic equilibrium of all dimers and the monomers.

## Partition Function Cofolding as a stepwise process

In this approach to cofolding the interaction between two RNA molecules is seen as a stepwise process. In a first step, the target molecule has to adopt a structure in which a binding site is accessible. In a second step, the ligand molecule will hybridize with a region accessible to an interaction. Consequently the algorithm is designed as a two step process: The first step is the calculation of the probability that a region within the target is unpaired, or equivalently, the calculation of the free energy needed to expose a region. In the second step we compute the free energy of an interaction for every possible binding site. Associated functions are:

Calculate the partition function over all unpaired regions of a maximal length.

```
void free_pu_contrib_struct (pu_contrib *pu)
```

Frees the output of function pf unstru().

```
int max_w,
char *cstruc,
int incr3,
int incr5)
```

Calculates the probability of a local interaction between two sequences.

```
void free_interact (interact *pin)
```

Frees the output of function pf\_interact().

#### See also

```
cofold.h, part_func_co.h and part_func_up.h for more details
```

## 2.6 Predicting local structures of large sequences

Local structures can be predicted by a modified version of the fold() algorithm that restricts the span of all base pairs.

The local analog to fold().

```
float aliLfold( const char **strings,
                char *structure,
                int maxdist)
float Lfoldz (const char *string,
              char *structure,
              int maxdist,
              int zsc,
              double min_z)
plist *pfl_fold (
            char *sequence,
            int winSize,
            int pairSize,
            float cutoffb,
            double **pU,
            struct plist **dpp2,
            FILE *pUfp,
            FILE *spup)
```

Compute partition functions for locally stable secondary structures (berni! update me)

#### See also

Lfold.h and LPfold.h for more details

## 2.7 Predicting Consensus Structures from Alignment

Consensus structures can be predicted by a modified version of the fold() algorithm that takes a set of aligned sequences instead of a single sequence. The energy function consists of the mean energy averaged over the sequences, plus a covariance term that favors pairs with consistent and compensatory mutations and penalizes pairs that cannot be formed by all structures. For details see Hofacker (2002).

Compute MFE and according consensus structure of an alignment of sequences.

Compute MFE and according structure of an alignment of sequences assuming the sequences are circular instead of linear.

```
void free_alifold_arrays (void)
```

Free the memory occupied by MFE alifold functions.

Calculate the free energy of a consensus structure given a set of aligned sequences.

```
struct pair_info
```

A base pair info structure.

```
double cv_fact
```

This variable controls the weight of the covariance term in the energy function of alignment folding algorithms.

```
double nc_fact
```

This variable controls the magnitude of the penalty for non-compatible sequences in the covariance term of alignment folding algorithms.

#### See also

alifold.h for more details

## 2.8 Global Variables for the Folding Routines

The following global variables change the behavior the folding algorithms or contain additional information after folding.

```
int noGU
```

Global switch to forbid/allow GU base pairs at all.

```
int no_closingGU
```

GU allowed only inside stacks if set to 1.

```
int noLonelyPairs
```

Global switch to avoid/allow helices of length 1.

```
int tetra_loop
```

Include special stabilizing energies for some tri-, tetra- and hexa-loops;.

```
int energy_set
```

0 = BP; 1=any mit GC; 2=any mit AU-parameter

```
float temperature
```

Rescale energy parameters to a temperature in degC.

```
int dangles
```

Switch the energy model for dangling end contributions (0, 1, 2, 3)

```
{\tt char} *nonstandards
```

contains allowed non standard base pairs

```
int cut_point
```

Marks the position (starting from 1) of the first nucleotide of the second molecule within the concatenated sequence.

```
float pf_scale
```

A scaling factor used by pf fold() to avoid overflows.

```
int fold_constrained
```

Global switch to activate/deactivate folding with structure constraints.

```
int do_backtrack
do backtracking, i.e.
```

```
char backtrack_type
```

A backtrack array marker for inverse fold()

include fold\_vars.h if you want to change any of these variables from their defaults.

#### See also

fold\_vars.h for a more complete and detailed description of all global variables and how to use them

## 2.9 Reading Energy Parameters from File

A default set of parameters, identical to the one described in Mathews et.al. (2004), is compiled into the library.

Alternately, parameters can be read from and written to a file.

```
void read_parameter_file (const char fname[])
```

Read energy parameters from a file.

```
void write_parameter_file (const char fname[])
```

Write energy parameters to a file.

To preserve some backward compatibility the RNAlib also provides functions to convert energy parameter files from the format used in version 1.4-1.8 into the new format used since version 2.0

Convert/dump a Vienna 1.8.4 formatted energy parameter file.

#### See also

read\_epars.h and convert\_epars.h for detailed description of the available functions

Next Page: Parsing and Comparing

## **Chapter 3**

# Parsing and Comparing - Functions to Manipulate Structures

## **Representations of Secondary Structures**

The standard representation of a secondary structure is the *bracket notation*, where matching brackets symbolize base pairs and unpaired bases are shown as dots. Alternatively, one may use two types of node labels, 'P' for paired and 'U' for unpaired; a dot is then replaced by '(U)', and each closed bracket is assigned an additional identifier 'P'. We call this the expanded notation. In Fontana et al. (1993) a condensed representation of the secondary structure is proposed, the so-called homeomorphically irreducible tree (HIT) representation. Here a stack is represented as a single pair of matching brackets labeled 'P' and weighted by the number of base pairs. Correspondingly, a contiguous strain of unpaired bases is shown as one pair of matching brackets labeled 'U' and weighted by its length. Generally any string consisting of matching brackets and identifiers is equivalent to a plane tree with as many different types of nodes as there are identifiers.

Bruce Shapiro (1988) proposed a coarse grained representation, which, does not retain the full information of the secondary structure. He represents the different structure elements by single matching brackets and labels them as 'H' (hairpin loop), 'I' (interior loop), 'B' (bulge), 'M' (multi-loop), and 'S' (stack). We extend his alphabet by an extra letter for external elements 'E'. Again these identifiers may be followed by a weight corresponding to the number of unpaired bases or base pairs in the structure element. All tree representations (except for the dot-bracket form) can be encapsulated into a virtual root (labeled 'R'), see the example below.

The following example illustrates the different linear tree representations used by the package. All lines show the same secondary structure.

```
a) .((((...(((...)))..((..)))).).
   (U) (((((U) (U) (U) (U) (U) P)P)P) (U) (U) (((U) (U)P)P)P) (U)P)P) (U)
b) (U) (((U2) ((U3)P3) (U2) ((U2)P2)P2) (U)P2) (U)
c) (((H) (H)M)B)
   ((((((H)S) ((H)S)M)S)B)S)
   (((((((H)S) ((H)S)M)S)B)S)E)
```

```
d) ((((((((H3)S3)((H2)S2)M4)S2)B1)S2)E2)R)
```

Above: Tree representations of secondary structures. a) Full structure: the first line shows the more convenient condensed notation which is used by our programs; the second line shows the rather clumsy expanded notation for completeness, b) HIT structure, c) different versions of coarse grained structures: the second line is exactly Shapiro's representation, the first line is obtained by neglecting the stems. Since each loop is closed by a unique stem, these two lines are equivalent. The third line is an extension taking into account also the external digits. d) weighted coarse structure, this time including the virtual root.

For the output of aligned structures from string editing, different representations are needed, where we put the label on both sides. The above examples for tree representations would then look like:

Aligned structures additionally contain the gap character '.'.

#### Parsing and Coarse Graining of Structures

Several functions are provided for parsing structures and converting to different representations.

```
char *expand_Full(const char *structure)
```

Convert the full structure from bracket notation to the expanded notation including root.

```
char *b2HIT (const char *structure)
```

Converts the full structure from bracket notation to the HIT notation including root.

```
char *b2C (const char *structure)
```

Converts the full structure from bracket notation to the a coarse grained notation using the 'H' 'B' 'I' 'M' and 'R' identifiers.

```
char *b2Shapiro (const char *structure)
```

Converts the full structure from bracket notation to the *weighted* coarse grained notation using the 'H' 'B' 'I' 'M' 'S' 'E' and 'R' identifiers.

```
char *expand_Shapiro (const char *coarse);
```

Inserts missing 'S' identifiers in unweighted coarse grained structures as obtained from b2C().

```
char *add_root (const char *structure)
```

Adds a root to an un-rooted tree in any except bracket notation.

```
char *unexpand_Full (const char *ffull)
```

Restores the bracket notation from an expanded full or HIT tree, that is any tree using only identifiers 'U' 'P' and 'R'.

```
char *unweight (const char *wcoarse)
```

Strip weights from any weighted tree.

```
void unexpand_aligned_F (char *align[2])
```

Converts two aligned structures in expanded notation.

```
void parse_structure (const char *structure)
```

Collects a statistic of structure elements of the full structure in bracket notation.

#### See also

RNAstruct.h for prototypes and more detailed description

#### **Distance Measures**

A simple measure of dissimilarity between secondary structures of equal length is the base pair distance, given by the number of pairs present in only one of the two structures being compared. I.e. the number of base pairs that have to be opened or closed to transform one structure into the other. It is therefore particularly useful for comparing structures on the same sequence. It is implemented by

Compute the "base pair" distance between two secondary structures s1 and s2.

For other cases a distance measure that allows for gaps is preferable. We can define distances between structures as edit distances between trees or their string representations. In the case of string distances this is the same as "sequence alignment". Given a set of edit operations and edit costs, the edit distance is given by the minimum sum of the costs along an edit path converting one object into the other. Edit distances like these always define a metric. The edit operations used by us are insertion, deletion and replacement of nodes. String editing does not pay attention to the matching of brackets,

while in tree editing matching brackets represent a single node of the tree. Tree editing is therefore usually preferable, although somewhat slower. String edit distances are always smaller or equal to tree edit distances.

The different level of detail in the structure representations defined above naturally leads to different measures of distance. For full structures we use a cost of 1 for deletion or insertion of an unpaired base and 2 for a base pair. Replacing an unpaired base for a pair incurs a cost of 1.

Two cost matrices are provided for coarse grained structures:

```
/* Niill.
                                         S,
                                               F.
               Η.
                     В,
                            I,
                                  Μ,
                                       1,
   { 0, 2, 2, 2, 2,
                                             1 } ,
                                                      /* Null replaced */
              0, 2, 2, 2, INF, INF},
2, 0, 1, 2, INF, INF},
                                                      /* H replaced */
        2,
                                                       /* B
        2,
                                                                  replaced */
                                                     /* I replaced */
/* I replaced */
/* M replaced */
/* S replaced */
/* E replaced */
       2, 2, 1, 0, 2, INF, INF},
        2, 2, 2, 2, 0, INF, INF},
1, INF, INF, INF, INF, 0, INF},
        2.
       1, INF, INF, INF, INF, INF,
                                              0 } ,
     ull, H, B, I, M, S, E */
0, 100, 5, 5, 75, 5, 5},
100, 0, 8, 8, 8, INF, INF},
5, 8, 0, 3, 8, INF, INF},
/* Null,
                                                       /* Null replaced */
                                                       /* H replaced */
/* B replaced */
                                                                 replaced */
       5, 8, 3, 0, 8, INF, INF},
                                                      /* I replaced */
        75, 8, 8, 8, 0, INF, INF}, 5, INF, INF, INF, INF, INF, O, INF},
                                                      /* M
/* S
       75,
                                                                  replaced */
                                                                  replaced */
        5, INF, INF, INF, INF, INF,
                                             0 } ,
                                                      /* E
                                                                  replaced */
```

The lower matrix uses the costs given in Shapiro (1990). All distance functions use the following global variables:

```
int cost_matrix;
```

Specify the cost matrix to be used for distance calculations.

```
int edit_backtrack;
```

Produce an alignment of the two structures being compared by tracing the editing path giving the minimum distance.

```
char *aligned_line[4];
```

Contains the two aligned structures after a call to one of the distance functions with edit backtrack set to 1.

## See also

utils.h, dist vars.h and stringdist.h for more details

**Functions for Tree Edit Distances** 

```
Tree *make_tree (char *struc)
```

Constructs a Tree ( essentially the postorder list ) of the structure 'struc', for use in tree\_edit\_distance().

```
float tree_edit_distance (Tree *T1, Tree *T2)
```

Calculates the edit distance of the two trees.

```
void free_tree(Tree *t)
```

Free the memory allocated for Tree t.

#### See also

dist vars.h and treedist.h for prototypes and more detailed descriptions

**Functions for String Alignment** 

```
swString *Make_swString (char *string)
```

Convert a structure into a format suitable for string\_edit\_distance().

```
\begin{tabular}{lll} float & string\_edit\_distance & (swString *T1, \\ & swString *T2) \end{tabular}
```

Calculate the string edit distance of T1 and T2.

### See also

dist\_vars.h and stringdist.h for prototypes and more detailed descriptions

**Functions for Comparison of Base Pair Probabilities** 

For comparison of base pair probability matrices, the matrices are first condensed into probability profiles which are the compared by alignment.

condense pair probability matrix into a vector containing probabilities for upstream paired, downstream paired and unpaired.

```
float profile_edit_distance ( const float *T1, const float *T2)
```

Align the 2 probability profiles T1, T2

### See also

ProfileDist.h for prototypes and more details of the above functions

Next Page: Utilities

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# **Chapter 4**

# **Utilities - Odds and Ends**

#### **Table of Contents**

- · Producing secondary structure graphs
- Producing (colored) dot plots for base pair probabilities
- Producing (colored) alignments
- RNA sequence related utilities
- · RNA secondary structure related utilities
- Miscellaneous Utilities

## 4.1 Producing secondary structure graphs

Produce a secondary structure graph in PostScript and write it to 'filename'.

Produce a secondary structure graph in PostScript including additional annotation macros and write it to 'filename'.

Produce a secondary structure graph in Graph Meta Language (gml) and write it to a file.

Produce a secondary structure graph in SStructView format.

Produce a secondary structure plot in SVG format and write it to a file.

Produce a secondary structure plot for further editing in XRNA.

```
int rna_plot_type
```

Switch for changing the secondary structure layout algorithm.

Two low-level functions provide direct access to the graph lauyouting algorithms:

Calculate nucleotide coordinates for secondary structure plot the Simple way

## See also

PS\_dot.h and naview.h for more detailed descriptions.

## 4.2 Producing (colored) dot plots for base pair probabilities

Produce a postscript dot-plot from two pair lists.

## See also

PS\_dot.h for more detailed descriptions.

## 4.3 Producing (colored) alignments

## 4.4 RNA sequence related utilities

Several functions provide useful applications to RNA sequences

Create a random string using characters from a specified symbol set.

Calculate hamming distance between two sequences.

```
void str_DNA2RNA(char *sequence);
```

Convert a DNA input sequence to RNA alphabet.

```
void str_uppercase(char *sequence);
```

Convert an input sequence to uppercase.

## 4.5 RNA secondary structure related utilities

```
char *pack_structure (const char *struc)
```

Pack secondary secondary structure, 5:1 compression using base 3 encoding.

```
char *unpack_structure (const char *packed)
```

Unpack secondary structure previously packed with pack structure()

```
short *make_pair_table (const char *structure)
```

Create a pair table of a secondary structure.

```
short *copy_pair_table (const short *pt)
```

Get an exact copy of a pair table.

## 4.6 Miscellaneous Utilities

```
void print_tty_input_seq (void)
```

Print a line to *stdout* that asks for an input sequence.

```
void print_tty_constraint_full (void)
```

Print structure constraint characters to stdout (full constraint support)

```
void print_tty_constraint (unsigned int option)
```

Print structure constraint characters to stdout.

```
int *get_iindx (unsigned int length)
```

Get an index mapper array (iindx) for accessing the energy matrices, e.g.

```
int *get_indx (unsigned int length)
```

Get an index mapper array (indx) for accessing the energy matrices, e.g.

Insert constraining pair types according to constraint structure string.

```
char *get_line(FILE *fp);
```

Read a line of arbitrary length from a stream.

Get a data record from stdin.

```
char *time_stamp (void)
```

Get a timestamp.

```
void warn_user (const char message[])
```

Print a warning message.

```
void nrerror (const char message[])
```

Die with an error message.

```
void init_rand (void)
```

Make random number seeds.

```
unsigned short xsubi[3];
```

Current 48 bit random number.

```
double urn (void)
```

get a random number from [0..1]

```
int int_urn (int from, int to)
```

Generates a pseudo random integer in a specified range.

```
void *space (unsigned size)
```

Allocate space safely.

Reallocate space safely.

#### See also

utils.h for a complete overview and detailed description of the utility functions

Next Page: Examples

## **Example - A Small Example Program**

The following program exercises most commonly used functions of the library.

The program folds two sequences using both the mfe and partition function algorithms and calculates the tree edit and profile distance of the resulting structures and base pairing probabilities.

```
#include <stdio.h>
#include <math.h>
#include "utils.h"
#include "fold_vars.h"
#include "fold.h"
#include "part_func.h"
#include "inverse.h"
#include "RNAstruct.h"
#include "treedist.h"
#include "stringdist.h"
#include "ProfileDist.h"
void main()
   *struct1, * struct2, * xstruc;
   float e1, e2, tree_dist, string_dist, profile_dist, kT;
   Tree *T1, *T2;
   swString *S1, *S2;
   float **pf1, **pf2;
   FLT_OR_DBL *bppm;
   /\star fold at 30C instead of the default 37C \star/
   temperature = 30.;
                            /* must be set *before* initializing */
   /\star allocate memory for structure and fold \star/
   struct1 = (char* ) space(sizeof(char)*(strlen(seq1)+1));
   e1 = fold(seq1, struct1);
   struct2 = (char*) space(sizeof(char)*(strlen(seq2)+1));
   e2 = fold(seq2, struct2);
                      /* free arrays used in fold() */
   /\star produce tree and string representations for comparison \star/
   xstruc = expand_Full(struct1);
```

```
T1 = make tree(xstruc);
  S1 = Make_swString(xstruc);
   free (xstruc):
   xstruc = expand_Full(struct2);
  T2 = make_tree(xstruc);
  S2 = Make_swString(xstruc);
  free (xstruc);
   /\star calculate tree edit distance and aligned structures with gaps \star/
   edit_backtrack = 1;
  tree_dist = tree_edit_distance(T1, T2);
   free_tree(T1); free_tree(T2);
  unexpand_aligned_F(aligned_line);
  /* same thing using string edit (alignment) distance */
   string_dist = string_edit_distance(S1, S2);
   free(S1); free(S2);
  printf("%s mfe=%5.2f\n%s mfe=%5.2f dist=%3.2f\n",
         aligned_line[0], e1, aligned_line[1], e2, string_dist);
  /\star for longer sequences one should also set a scaling factor for
     partition function folding, e.g: */
   kT = (temperature + 273.15) * 1.98717/1000.; /* kT in kcal/mol */
  pf_scale = exp(-e1/kT/strlen(seq1));
  /\star calculate partition function and base pair probabilities \star/
  e1 = pf_fold(seq1, struct1);
   /\star get the base pair probability matrix for the previous run of pf_fold() \star/
  bppm = export_bppm();
  pf1 = Make_bp_profile_bppm(bppm, strlen(seq1));
  e2 = pf_fold(seq2, struct2);
   /\star get the base pair probability matrix for the previous run of pf_fold() \star/
  bppm = export_bppm();
  pf2 = Make_bp_profile(strlen(seq2));
   free_pf_arrays(); /* free space allocated for pf_fold() */
  profile_dist = profile_edit_distance(pf1, pf2);
  printf("%s free energy=%5.2f\n%s free energy=%5.2f dist=%3.2f\n",
         aligned_line[0], e1, aligned_line[1], e2, profile_dist);
   free_profile(pf1); free_profile(pf2);
In a typical Unix environment you would compile this program using:
cc ${OPENMP_CFLAGS} -c example.c -I${hpath}
and link using
cc ${OPENMP_CFLAGS} -o example -L${lpath} -lRNA -lm
```

where *\${hpath}* and *\${lpath}* point to the location of the header files and library, respectively.

#### Note

As default, the RNAlib is compiled with build-in OpenMP multithreading support.

Thus, when linking your own object files to the library you have to pass the compiler specific *\${OPENMP\_CFLAGS}* (e.g. '-fopenmp' for **gcc**) even if your code does not use openmp specific code. However, in that case the *OpenMP* flags may be ommitted when compiling example.c

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## References

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## **Deprecated List**

Global base\_pair Do not use this variable anymore!

**Global centroid(int length, double \*dist)** This function is deprecated and should not be used anymore as it is not threadsafe!

**Global energy\_of\_circ\_struct(const char** \*string, const char \*structure) This function is deprecated and should not be used in future programs Use energy\_of\_circ\_structure() instead!

**Global energy\_of\_struct(const char \*string, const char \*structure)** This function is deprecated and should not be used in future programs! Use energy\_of\_structure() instead!

Global energy\_of\_struct\_pt(const char \*string, short \*ptable, short \*s, short \*s1)

This function is deprecated and should not be used in future programs! Use energy\_of\_structure\_pt() instead!

Global expHairpinEnergy(int u, int type, short si1, short sj1, const char \*string)

Use exp\_E\_Hairpin() from loop\_energies.h instead

Global expLoopEnergy(int u1, int u2, int type, int type2, short si1, short sj1, short sp1, short sq1)

Use exp\_E\_IntLoop() from loop\_energies.h instead

Global get\_plist(struct plist \*pl, int length, double cut\_off) { This function is deprecated and will be removed soon!} use assign\_plist\_from\_pr() instead!

**Global HairpinE(int size, int type, int si1, int sj1, const char \*string)** {This function is deprecated and will be removed soon.

**Global iindx** Do not use this variable anymore!

**Global init\_co\_pf\_fold(int length)** { This function is deprecated and will be removed soon!}

Global init\_pf\_fold(int length) This function is obsolete and will be removed soon!

**Global initialize\_cofold(int length)** {This function is obsolete and will be removed soon!}

**Global initialize\_fold(int length)** {This function is deprecated and will be removed soon!}

Global LoopEnergy(int n1, int n2, int type, int type\_2, int si1, int sj1, int sp1, int sq1) {This function is deprecated and will be removed soon.

**Global Make\_bp\_profile(int length)** This function is deprecated and will be removed soon! See Make\_bp\_profile\_bppm() for a replacement

**Global mean\_bp\_dist(int length)** This function is not threadsafe and should not be used anymore. Use mean\_bp\_distance() instead!

Global pr Do not use this variable anymore!

**Global PS\_dot\_plot(char** \***string, char** \***file)** This function is deprecated and will be removed soon! Use PS\_dot\_plot\_list() instead!

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## 8.1 Data Structures

Here are the data structures with brief descriptions:	
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cpair (This datastructure is used as input parameter in functions of PS_dot.c) 3 duplexT	8
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path_t	.3
weights of the energy parameters )	3,
and others )	
- 1 00t0 001_10t	. •

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enna RNA Package can be found here)	
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H/stringdist.h (Functions for String Alignment)	
H/subopt.h (RNAsubopt and density of states declarations)	
H/svm utils.h	
H/treedist.h (Functions for Tree Edit Distances )	
H/utils.h (Various utility- and helper-functions used throughout the Vienna RNA	
package)	. 140
lib/1.8.4_epars.h (Free energy parameters for parameter file conversion)	
lib/1.8.4_intloops.h (Free energy parameters for interior loop contributions need	
by the parameter file conversion functions )	
lib/intl11.h	
lib/intl11dH.h	. ??
lib/intl21.h	
lib/intl21dH.h	. ??
lib/intl22.h	. ??
lib/intl22dH.h	. ??
lib/list.h	. ??

## **Data Structure Documentation**

## 10.1 bondT Struct Reference

base pair

## 10.1.1 Detailed Description

base pair

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.2 bondTEn Struct Reference

base pair with associated energy

## 10.2.1 Detailed Description

base pair with associated energy

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.3 cofoldF Struct Reference

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.4 ConcEnt Struct Reference

The documentation for this struct was generated from the following file:

· H/data structures.h

#### 10.5 constrain Struct Reference

The documentation for this struct was generated from the following file:

· H/data structures.h

## 10.6 COORDINATE Struct Reference

this is a workarround for the SWIG Perl Wrapper RNA plot function that returns an array of type COORDINATE

## 10.6.1 Detailed Description

this is a workarround for the SWIG Perl Wrapper RNA plot function that returns an array of type COORDINATE

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.7 cpair Struct Reference

this datastructure is used as input parameter in functions of PS\_dot.c

## 10.7.1 Detailed Description

this datastructure is used as input parameter in functions of PS\_dot.c

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.8 duplexT Struct Reference

The documentation for this struct was generated from the following file:

· H/data\_structures.h

## 10.9 dupVar Struct Reference

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.10 folden Struct Reference

The documentation for this struct was generated from the following file:

• H/data\_structures.h

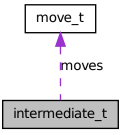
## 10.11 interact Struct Reference

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.12 intermediate\_t Struct Reference

Collaboration diagram for intermediate\_t:



The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.13 INTERVAL Struct Reference

sequence interval stack element used in subopt.c

## 10.13.1 Detailed Description

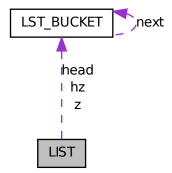
sequence interval stack element used in subopt.c

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.14 LIST Struct Reference

Collaboration diagram for LIST:

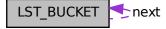


The documentation for this struct was generated from the following file:

· lib/list.h

## 10.15 LST\_BUCKET Struct Reference

Collaboration diagram for LST\_BUCKET:



The documentation for this struct was generated from the following file:

· lib/list.h

## 10.16 move\_t Struct Reference

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.17 PAIR Struct Reference

base pair data structure used in subopt.c

## 10.17.1 Detailed Description

base pair data structure used in subopt.c

The documentation for this struct was generated from the following file:

• H/data structures.h

## 10.18 pair\_info Struct Reference

A base pair info structure.

## 10.18.1 Detailed Description

A base pair info structure.

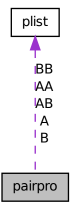
for each base pair (i,j) the structure lists: its probability 'p', an entropy-like measure for its well-definedness 'ent', and in 'bp[]' the frequency of each type of pair. 'bp[0]' contains the number of non-compatible sequences, 'bp[1]' the number of CG pairs, etc.

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.19 pairpro Struct Reference

Collaboration diagram for pairpro:



The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.20 paramT Struct Reference

The datastructure that contains temperature scaled energy parameters.

#### 10.20.1 Detailed Description

The datastructure that contains temperature scaled energy parameters.

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.21 path\_t Struct Reference

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.22 pf\_paramT Struct Reference

The datastructure that contains temperature scaled Boltzmann weights of the energy parameters.

#### 10.22.1 Detailed Description

The datastructure that contains temperature scaled Boltzmann weights of the energy parameters.

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.23 plist Struct Reference

this datastructure is used as input parameter in functions of PS\_dot.h and others

### 10.23.1 Detailed Description

this datastructure is used as input parameter in functions of PS\_dot.h and others The documentation for this struct was generated from the following file:

• H/data\_structures.h

#### 10.24 Postorder\_list Struct Reference

The documentation for this struct was generated from the following file:

• H/dist\_vars.h

## 10.25 pu\_contrib Struct Reference

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.26 pu\_out Struct Reference

The documentation for this struct was generated from the following file:

· H/data structures.h

## 10.27 sect Struct Reference

stack of partial structures for backtracking

#### 10.27.1 Detailed Description

stack of partial structures for backtracking

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.28 snoopT Struct Reference

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.29 SOLUTION Struct Reference

solution element from subopt.c

## 10.29.1 Detailed Description

solution element from subopt.c

The documentation for this struct was generated from the following file:

H/data\_structures.h

## 10.30 svm\_model Struct Reference

The documentation for this struct was generated from the following file:

· H/svm utils.h

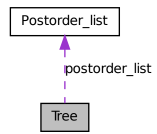
## 10.31 swString Struct Reference

The documentation for this struct was generated from the following file:

• H/dist\_vars.h

## 10.32 Tree Struct Reference

Collaboration diagram for Tree:



The documentation for this struct was generated from the following file:

• H/dist\_vars.h

## 10.33 TwoDfold\_solution Struct Reference

Solution element returned from TwoDfoldList.

#### 10.33.1 Detailed Description

Solution element returned from TwoDfoldList.

This element contains free energy and structure for the appropriate kappa (k), lambda (l) neighborhood The datastructure contains two integer attributes 'k' and 'l' as well as an attribute 'en' of type float representing the free energy in kcal/mol and an attribute 's' of type char\* containg the secondary structure representative,

A value of INF in k denotes the end of a list

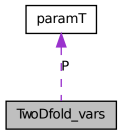
The documentation for this struct was generated from the following file:

• H/data\_structures.h

### 10.34 TwoDfold\_vars Struct Reference

Variables compound for 2Dfold MFE folding.

Collaboration diagram for TwoDfold\_vars:



## 10.34.1 Detailed Description

Variables compound for 2Dfold MFE folding.

The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.35 TwoDpfold\_solution Struct Reference

Solution element returned from TwoDpfoldList.

#### 10.35.1 Detailed Description

Solution element returned from TwoDpfoldList.

This element contains the partition function for the appropriate kappa (k), lambda (l) neighborhood The datastructure contains two integer attributes 'k' and 'l' as well as an attribute 'q' of type #FLT\_OR\_DBL

A value of INF in k denotes the end of a list

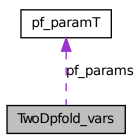
The documentation for this struct was generated from the following file:

• H/data\_structures.h

## 10.36 TwoDpfold\_vars Struct Reference

Variables compound for 2Dfold partition function folding.

Collaboration diagram for TwoDpfold\_vars:



## 10.36.1 Detailed Description

Variables compound for 2Dfold partition function folding.

The documentation for this struct was generated from the following file:

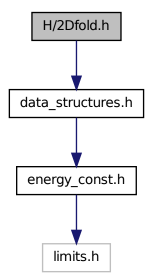
• H/data\_structures.h

## **File Documentation**

## 11.1 H/2Dfold.h File Reference

Compute the minimum free energy (MFE) and secondary structures for a partitioning of the secondary structure space according to the base pair distance to two fixed reference structures basepair distance to two fixed reference structures.

Include dependency graph for 2Dfold.h:



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#### **Functions**

TwoDfold\_vars \* get\_TwoDfold\_variables (const char \*seq, const char \*structure1, const char \*structure2, int circ)

Get a structure of type TwoDfold\_vars prefilled with current global settings.

- void destroy\_TwoDfold\_variables (TwoDfold\_vars \*our\_variables)
  - Destroy a TwoDfold\_vars datastructure without memory loss.
- TwoDfold\_solution \* TwoDfoldList (TwoDfold\_vars \*vars, int distance1, int distance2)

Compute MFE's and representative for distance partitioning.

• char \* TwoDfold\_backtrack\_f5 (unsigned int j, int k, int l, TwoDfold\_vars \*vars)

Backtrack a minimum free energy structure from a 5' section of specified length.

#### 11.1.1 Detailed Description

Compute the minimum free energy (MFE) and secondary structures for a partitioning of the secondary structure space according to the base pair distance to two fixed reference structures basepair distance to two fixed reference structures.

#### 11.1.2 Function Documentation

```
11.1.2.1 TwoDfold_vars* get_TwoDfold_variables ( const char * seq, const char * structure1, const char * structure2, int circ )
```

Get a structure of type TwoDfold\_vars prefilled with current global settings.

This function returns a datastructure of type TwoDfold\_vars. The data fields inside the TwoDfold\_vars are prefilled by global settings and all memory allocations necessary to start a computation are already done for the convenience of the user

#### Note

Make sure that the reference structures are compatible with the sequence according to Watson-Crick- and Wobble-base pairing

### See also

destroy\_TwoDfold\_variables(), TwoDfold(), TwoDfold\_circ

#### **Parameters**

seq	The RNA sequence
structure1	The first reference structure in dot-bracket notation
structure2	The second reference structure in dot-bracket notation
circ	A switch to indicate the assumption to fold a circular instead of linear RNA
	(0=OFF, 1=ON)

#### Returns

A datastructure prefilled with folding options and allocated memory

11.1.2.2 void destroy\_TwoDfold\_variables ( TwoDfold\_vars \* our\_variables )

Destroy a TwoDfold\_vars datastructure without memory loss.

This function free's all allocated memory that depends on the datastructure given.

#### See also

get TwoDfold variables()

#### **Parameters**

our	A pointer to the datastructure to be destroyed
variables	

11.1.2.3 TwoDfold\_solution\* TwoDfoldList ( TwoDfold\_vars \* vars, int distance1, int distance2 )

Compute MFE's and representative for distance partitioning.

This function computes the minimum free energies and a representative secondary structure for each distance class according to the two references specified in the datastructure 'vars'. The maximum basepair distance to each of both references may be set by the arguments 'distance1' and 'distance2', respectively. If both distance arguments are set to '-1', no restriction is assumed and the calculation is performed for each distance class possible.

The returned list contains an entry for each distance class. If a maximum basepair distance to either of the references was passed, an entry with k=l=-1 will be appended in the list, denoting the class where all structures exceeding the maximum will be thrown into. The end of the list is denoted by an attribute value of INF in the k-attribute of the list entry.

#### See also

get\_TwoDfold\_variables(), destroy\_TwoDfold\_variables(), TwoDfold\_solution

#### **Parameters**

vars	the datastructure containing all predefined folding attributes
distance1	maximum distance to reference1 (-1 means no restriction)
distance2	maximum distance to reference2 (-1 means no restriction)

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11.1.2.4 char\* TwoDfold\_backtrack\_f5 ( unsigned int j, int k, int l, TwoDfold\_vars \* vars )

Backtrack a minimum free energy structure from a 5' section of specified length.

This function allows to backtrack a secondary structure beginning at the 5' end, a specified length and residing in a specific distance class. If the argument 'k' gets a value of -1, the structure that is backtracked is assumed to reside in the distance class where all structures exceeding the maximum basepair distance specified in TwoDfoldList() belong to.

#### Note

The argument 'vars' must contain precalculated energy values in the energy matrices, i.e. a call to TwoDfoldList() preceding this function is mandatory!

#### See also

TwoDfoldList(), get\_TwoDfold\_variables(), destroy\_TwoDfold\_variables()

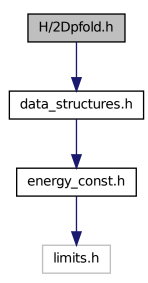
#### **Parameters**

j	The length in nucleotides beginning from the 5' end
k	distance to reference1 (may be -1)
1	distance to reference2
vars	the datastructure containing all predefined folding attributes

## 11.2 H/2Dpfold.h File Reference

Compute the partition function and stochastically sample secondary structures for a partitioning of the secondary structure space according to the base pair distance to two fixed reference structures.

Include dependency graph for 2Dpfold.h:



#### **Functions**

TwoDpfold\_vars \* get\_TwoDpfold\_variables (const char \*seq, const char \*structure1, char \*structure2, int circ)

Get a datastructure containing all necessary attributes and global folding switches.

TwoDpfold\_vars \* get\_TwoDpfold\_variables\_from\_MFE (TwoDfold\_vars \*mfe\_vars)

Get the datastructure containing all necessary attributes and global folding switches from a pre-filled mfe-datastructure.

void destroy\_TwoDpfold\_variables (TwoDpfold\_vars \*vars)

Free all memory occupied by a TwoDpfold\_vars datastructure.

TwoDpfold\_solution \* TwoDpfoldList (TwoDpfold\_vars \*vars, int maxDistance1, int maxDistance2)

Compute the partition function for all distance classes.

char \* TwoDpfold\_pbacktrack (TwoDpfold\_vars \*vars, int d1, int d2)

Sample secondary structure representatives from a set of distance classes according to their Boltzmann probability.

char \* TwoDpfold\_pbacktrack5 (TwoDpfold\_vars \*vars, int d1, int d2, unsigned int length)

Sample secondary structure representatives with a specified length from a set of distance classes according to their Boltzmann probability.

#### 11.2.1 Detailed Description

Compute the partition function and stochastically sample secondary structures for a partitioning of the secondary structure space according to the base pair distance to two fixed reference structures.

#### 11.2.2 Function Documentation

11.2.2.1 TwoDpfold\_vars\* get\_TwoDpfold\_variables ( const char \* seq, const char \* structure1, char \* structure2, int circ )

Get a datastructure containing all necessary attributes and global folding switches.

This function prepares all necessary attributes and matrices etc which are needed for a call of TwoDpfoldList. A snapshot of all current global model switches (dangles, temperature and so on) is done and stored in the returned datastructure. Additionally, all matrices that will hold the partition function values are prepared.

#### **Parameters**

seq	the RNA sequence in uppercase format with letters from the alphabet
	{AUCG}
structure1	the first reference structure in dot-bracket notation
structure2	the second reference structure in dot-bracket notation
circ	a switch indicating if the sequence is linear (0) or circular (1)

#### **Returns**

the datastructure containing all necessary partition function attributes

11.2.2.2 TwoDpfold\_vars\* get\_TwoDpfold\_variables\_from\_MFE ( TwoDfold\_vars \* mfe\_vars )

Get the datastructure containing all necessary attributes and global folding switches from a pre-filled mfe-datastructure.

This function actually does the same as get\_TwoDpfold\_variables but takes its switches and settings from a pre-filled MFE equivalent datastructure

#### See also

get\_TwoDfold\_variables(), get\_TwoDpfold\_variables()

#### **Parameters**

mfe_vars	the pre-filled mfe datastructure

#### Returns

the datastructure containing all necessary partition function attributes

11.2.2.3 void destroy\_TwoDpfold\_variables ( TwoDpfold vars \* vars )

Free all memory occupied by a TwoDpfold\_vars datastructure.

This function free's all memory occupied by a datastructure obtained from from get\_TwoDpfold\_variables() or get\_TwoDpfold\_variables\_from\_MFE()

#### See also

get\_TwoDpfold\_variables(), get\_TwoDpfold\_variables\_from\_MFE()

#### **Parameters**

vars	the datastructure to be free'd

11.2.2.4 TwoDpfold\_solution\* TwoDpfoldList ( TwoDpfold\_vars \* vars, int maxDistance1, int maxDistance2 )

Compute the partition function for all distance classes.

This function computes the partition functions for all distance classes according the two reference structures specified in the datastructure 'vars'. Similar to TwoDfoldList() the arguments maxDistance1 and maxDistance2 specify the maximum distance to both reference structures. A value of '-1' in either of them makes the appropriate distance restrictionless, i.e. all basepair distancies to the reference are taken into account during computation. In case there is a restriction, the returned solution contains an entry where the attribute k=l=-1 contains the partition function for all structures exceeding the restriction. A values of INF in the attribute 'k' of the returned list denotes the end of the list

#### See also

get\_TwoDpfold\_variables(), destroy\_TwoDpfold\_variables(), TwoDpfold\_solution

#### **Parameters**

	vars	the datastructure containing all necessary folding attributes and matrices
m	naxDis-	the maximum basepair distance to reference1 (may be -1)
	tance1	
m	naxDis-	the maximum basepair distance to reference2 (may be -1)
	tance2	

#### Returns

a list of partition funtions for the appropriate distance classes

11.2.2.5 char\* TwoDpfold\_pbacktrack ( TwoDpfold\_vars \* vars, int d1, int d2 )

Sample secondary structure representatives from a set of distance classes according to their Boltzmann probability.

If the argument 'd1' is set to '-1', the structure will be backtracked in the distance class where all structures exceeding the maximum basepair distance to either of the references reside.

#### Note

The argument 'vars' must contain precalculated partition function matrices, i.e. a call to TwoDpfoldList() preceding this function is mandatory!

#### See also

TwoDpfoldList()

#### **Parameters**

vars	the datastructure containing all necessary folding attributes and matrices
d1	the distance to reference1 (may be -1)
d2	the distance to reference2

#### **Returns**

a sampled secondary structure in dot-bracket notation

11.2.2.6 char\* TwoDpfold\_pbacktrack5 ( TwoDpfold\_vars \* vars, int d1, int d2, unsigned int length )

Sample secondary structure representatives with a specified length from a set of distance classes according to their Boltzmann probability.

This function does essentially the same as TwoDpfold\_pbacktrack with the only difference that partial structures, i.e. structures beginning from the 5' end with a specified length of the sequence, are backtracked

### Note

The argument 'vars' must contain precalculated partition function matrices, i.e. a call to TwoDpfoldList() preceding this function is mandatory!

This function does not work (since it makes no sense) for circular RNA sequences!

#### See also

TwoDpfold\_pbacktrack(), TwoDpfoldList()

#### **Parameters**

vars	the datastructure containing all necessary folding attributes and matrices
d1	the distance to reference1 (may be -1)
d2	the distance to reference2
length	the length of the structure beginning from the 5' end

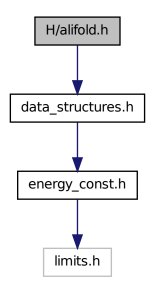
#### Returns

a sampled secondary structure in dot-bracket notation

#### 11.3 H/alifold.h File Reference

compute various properties (consensus MFE structures, partition function, Boltzmann distributed stochastic samples, ...) for RNA sequence alignments

Include dependency graph for alifold.h:



#### **Functions**

• void update\_alifold\_params (void)

Update the energy parameters for alifold function.

• float alifold (const char \*\*strings, char \*structure)

Compute MFE and according consensus structure of an alignment of sequences.

• float circalifold (const char \*\*strings, char \*structure)

Compute MFE and according structure of an alignment of sequences assuming the sequences are circular instead of linear.

• FLT\_OR\_DBL \* alipf\_export\_bppm (void)

Get a pointer to the base pair probability array.

void free\_alifold\_arrays (void)

Free the memory occupied by MFE alifold functions.

int get\_mpi (char \*Alseq[], int n\_seq, int length, int \*mini)

Get the mean pairwise identity in steps from ?to?(ident)

float \*\* readribosum (char \*name)

Read a ribosum or other user-defined scoring matrix.

float energy\_of\_alistruct (const char \*\*sequences, const char \*structure, int n\_-seq, float \*energy)

Calculate the free energy of a consensus structure given a set of aligned sequences.

void encode\_ali\_sequence (const char \*sequence, short \*S, short \*s5, short \*s3, char \*ss, unsigned short \*as, int circ)

Get arrays with encoded sequence of the alignment.

void alloc\_sequence\_arrays (const char \*\*sequences, short \*\*\*S, short \*\*\*S, short \*\*\*S, unsigned short \*\*\*a2s, char \*\*\*Ss, int circ)

Allocate memory for sequence array used to deal with aligned sequences.

void free\_sequence\_arrays (unsigned int n\_seq, short \*\*\*S, short \*\*\*S, short \*\*\*S, unsigned short \*\*\*a2s, char \*\*\*Ss)

Free the memory of the sequence arrays used to deal with aligned sequences.

float alipf\_fold (const char \*\*sequences, char \*structure, plist \*\*pl)

The partition function version of alifold() works in analogy to pf\_fold().

- float alipf circ fold (const char \*\*sequences, char \*structure, plist \*\*pl)
- FLT\_OR\_DBL \* export\_ali\_bppm (void)

Get a pointer to the base pair probability array.

char \* alipbacktrack (double \*prob)

Sample a consensus secondary structure from the Boltzmann ensemble according its probability

#### **Variables**

· double cv fact

This variable controls the weight of the covariance term in the energy function of alignment folding algorithms.

· double nc\_fact

This variable controls the magnitude of the penalty for non-compatible sequences in the covariance term of alignment folding algorithms.

#### 11.3.1 Detailed Description

compute various properties (consensus MFE structures, partition function, Boltzmann distributed stochastic samples, ...) for RNA sequence alignments

## 11.3.2 Function Documentation

11.3.2.1 void update\_alifold\_params (void)

Update the energy parameters for alifold function.

Call this to recalculate the pair matrix and energy parameters after a change in folding parameters like temperature

11.3.2.2 float alifold ( const char \*\* strings, char \* structure )

Compute MFE and according consensus structure of an alignment of sequences.

This function predicts the consensus structure for the aligned 'sequences' and returns the minimum free energy; the mfe structure in bracket notation is returned in 'structure'.

Sufficient space must be allocated for 'structure' before calling alifold().

#### **Parameters**

strings	A pointer to a NULL terminated array of character arrays		
structure	A pointer to a character array that may contain a constraining consensus		
	structure (will be overwritten by a consensus structure that exhibits the MFE)		

## Returns

The free energy score in kcal/mol

11.3.2.3 float circalifold ( const char \*\* strings, char \* structure )

Compute MFE and according structure of an alignment of sequences assuming the sequences are circular instead of linear.

# **Parameters**

strings	A pointer to a NULL terminated array of character arrays	
structure	A pointer to a character array that may contain a constraining consensus	
	structure (will be overwritten by a consensus structure that exhibits the MFE)	

#### Returns

The free energy score in kcal/mol

11.3.2.4 int get\_mpi ( char \* Alseq[], int n\_seq, int length, int \* mini )

Get the mean pairwise identity in steps from ?to?(ident)

Alseq	

n_seq	n_seq The number of sequences in the alignment	
length	The length of the alignment	
mini		

# Returns

The mean pairwise identity

11.3.2.5 float energy\_of\_alistruct ( const char \*\* sequences, const char \* structure, int n\_seq, float \* energy )

Calculate the free energy of a consensus structure given a set of aligned sequences.

## **Parameters**

sequences	The NULL terminated array of sequences	
structure	The consensus structure	
n_seq	The number of sequences in the alignment	
energy	A pointer to an array of at least two floats that will hold the free energies	
	(energy[0] will contain the free energy, energy[1] will be filled with the covari-	
	ance energy term)	

# Returns

free energy in kcal/mol

11.3.2.6 void encode\_ali\_sequence ( const char \* sequence, short \* S, short \* S, short \* S, char \* ss, unsigned short \* as, int circ )

Get arrays with encoded sequence of the alignment.

this function assumes that in S, S5, s3, ss and as enough space is already allocated (size must be at least sequence length+2)

se	quence	The gapped sequence from the alignment	
	S	pointer to an array that holds encoded sequence	
	s5	pointer to an array that holds the next base 5' of alignment position i	
	s3	pointer to an array that holds the next base 3' of alignment position i	
	SS		
	as		
	circ	assume the molecules to be circular instead of linear (circ=0)	

```
11.3.2.7 void alloc_sequence_arrays ( const char *** sequences, short *** S, short *** S, short *** S, short *** S, int circ )
```

Allocate memory for sequence array used to deal with aligned sequences.

Note that these arrays will also be initialized according to the sequence alignment given

## See also

free\_sequence\_arrays()

## **Parameters**

sequences	The aligned sequences	
S	A pointer to the array of encoded sequences	
S5	A pointer to the array that contains the next 5' nucleotide of a sequence position	
S3	A pointer to the array that contains the next 3' nucleotide of a sequence position	
a2s	A pointer to the array that contains the alignment to sequence position mapping	
Ss	A pointer to the array that contains the ungapped sequence	
circ	assume the molecules to be circular instead of linear (circ=0)	

11.3.2.8 void free\_sequence\_arrays ( unsigned int 
$$n\_seq$$
, short \*\*\*  $S$ , short

Free the memory of the sequence arrays used to deal with aligned sequences.

This function frees the memory previously allocated with alloc\_sequence\_arrays()

## See also

alloc\_sequence\_arrays()

n_seq	The number of aligned sequences	
S	A pointer to the array of encoded sequences	
S5	A pointer to the array that contains the next 5' nucleotide of a sequence position	
S3	A pointer to the array that contains the next 3' nucleotide of a sequence position	
a2s	A pointer to the array that contains the alignment to sequence position mapping	
Ss	A pointer to the array that contains the ungapped sequence	

```
11.3.2.9 float alipf_fold ( const char ** sequences, char * structure, plist ** pl )
```

The partition function version of alifold() works in analogy to pf\_fold().

Pair probabilities and information about sequence covariations are returned via the 'pi' variable as a list of pair\_info structs. The list is terminated by the first entry with pi.i = 0.

#### **Parameters**

sequences	
structure	
pl	

## **Returns**

```
11.3.2.10 float alipf_circ_fold ( const char ** sequences, char * structure, plist ** pl )
```

## **Parameters**

sequences	
structure	
pi	

## Returns

```
11.3.2.11 FLT_OR_DBL* export_ali_bppm ( void )
```

Get a pointer to the base pair probability array.

Accessing the base pair probabilities for a pair (i,j) is achieved by

```
FLT_OR_DBL *pr = export_bppm(); pr_ij = pr[iindx[i]-j];
```

# See also

```
get_iindx()
```

# Returns

A pointer to the base pair probability array

```
11.3.2.12 char* alipbacktrack ( double * prob )
```

Sample a consensus secondary structure from the Boltzmann ensemble according its probability

.

## **Parameters**

prob to be described (berni)

# Returns

A sampled consensus secondary structure in dot-bracket notation

# 11.3.3 Variable Documentation

11.3.3.1 double cv\_fact

This variable controls the weight of the covariance term in the energy function of alignment folding algorithms.

Default is 1.

11.3.3.2 double nc\_fact

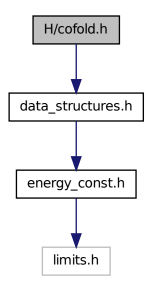
This variable controls the magnitude of the penalty for non-compatible sequences in the covariance term of alignment folding algorithms.

Default is 1.

# 11.4 H/cofold.h File Reference

MFE version of cofolding routines.

Include dependency graph for cofold.h:



# **Functions**

• float cofold (const char \*sequence, char \*structure)

Compute the minimum free energy of two interacting RNA molecules.

void free\_co\_arrays (void)

Free memory occupied by cofold()

• void update\_cofold\_params (void)

Recalculate parameters.

SOLUTION \* zukersubopt (const char \*string)

Compute Zuker type suboptimal structures.

• void get\_monomere\_mfes (float \*e1, float \*e2)

get\_monomer\_free\_energies

void export\_cofold\_arrays (int \*\*f5\_p, int \*\*c\_p, int \*\*fML\_p, int \*\*fM1\_p, int \*\*fc\_p, int \*\*indx\_p, char \*\*ptype\_p)

Export the arrays of partition function cofold.

• void initialize\_cofold (int length)

allocate arrays for folding

# 11.4.1 Detailed Description

MFE version of cofolding routines. This file includes (almost) all function declarations within the **RNAlib** that are related to MFE Cofolding... This also includes the Zuker suboptimals calculations, since they are implemented using the cofold routines.

## 11.4.2 Function Documentation

11.4.2.1 float cofold ( const char \* sequence, char \* structure )

Compute the minimum free energy of two interacting RNA molecules.

The code is analog to the fold() function. If cut\_point ==-1 results should be the same as with fold().

#### **Parameters**

sequence	The two sequences concatenated
structure	Will hold the barcket dot structure of the dimer molecule

## Returns

minimum free energy of the structure

# 11.4.2.2 SOLUTION\* zukersubopt ( const char \* string )

Compute Zuker type suboptimal structures.

Compute Suboptimal structures according to M. Zuker, i.e. for every possible base pair the minimum energy structure containing the resp. base pair. Returns a list of these structures and their energies.

#### **Parameters**

string RNA sequence	

# Returns

List of zuker suboptimal structures

11.4.2.3 void get\_monomere\_mfes ( float \* e1, float \* e2 )

get\_monomer\_free\_energies

Export monomer free energies out of cofold arrays

e1	A pointer to a variable where the energy of molecule A will be written to
e2	A pointer to a variable where the energy of molecule B will be written to

```
11.4.2.4 void export_cofold_arrays ( int ** f5_p, int ** c_p, int ** fML_p, int ** fML_p, int ** fmL_p, int ** fmL_p, char ** ptype_p)
```

Export the arrays of partition function cofold.

Export the cofold arrays for use e.g. in the concentration Computations or suboptimal secondary structure backtracking

#### **Parameters**

f5_p	A pointer to the 'f5' array, i.e. array conatining best free energy in interval
	[1,j]
c_p	A pointer to the 'c' array, i.e. array containing best free energy in interval [i,j]
	given that i pairs with j
fML_p	A pointer to the 'M' array, i.e. array containing best free energy in interval [i,j]
	for any multiloop segment with at least one stem
fM1_p	A pointer to the 'M1' array, i.e. array containing best free energy in interval
	[i,j] for multiloop segment with exactly one stem
fc_p	A pointer to the 'fc' array, i.e. array
indx_p	A pointer to the indexing array used for accessing the energy matrices
ptype_p	A pointer to the ptype array containing the base pair types for each possibility
	(i,j)

## 11.4.2.5 void initialize\_cofold ( int length )

allocate arrays for folding

# **Deprecated**

{This function is obsolete and will be removed soon!}

# 11.5 H/convert\_epars.h File Reference

Functions and definitions for energy parameter file format conversion.

# **Defines**

• #define VRNA\_CONVERT\_OUTPUT\_ALL 1U

Flag to indicate printing of a complete parameter set.

• #define VRNA\_CONVERT\_OUTPUT\_HP 2U

Flag to indicate printing of hairpin contributions.

• #define VRNA\_CONVERT\_OUTPUT\_STACK 4U

Flag to indicate printing of base pair stack contributions.

• #define VRNA\_CONVERT\_OUTPUT\_MM\_HP 8U

Flag to indicate printing of hairpin mismatch contribution.

#define VRNA CONVERT OUTPUT MM INT 16U

Flag to indicate printing of interior loop mismatch contribution.

#define VRNA CONVERT OUTPUT MM INT 1N 32U

Flag to indicate printing of 1:n interior loop mismatch contribution.

• #define VRNA\_CONVERT\_OUTPUT\_MM\_INT\_23 64U

Flag to indicate printing of 2:3 interior loop mismatch contribution.

• #define VRNA\_CONVERT\_OUTPUT\_MM\_MULTI 128U

Flag to indicate printing of multi loop mismatch contribution.

#define VRNA CONVERT OUTPUT MM EXT 256U

Flag to indicate printing of exterior loop mismatch contribution.

#define VRNA CONVERT OUTPUT DANGLE5 512U

Flag to indicate printing of 5' dangle conctribution.

#define VRNA CONVERT OUTPUT DANGLE3 1024U

Flag to indicate printing of 3' dangle contribution.

• #define VRNA CONVERT OUTPUT INT 11 2048U

Flag to indicate printing of 1:1 interior loop contribution.

#define VRNA CONVERT OUTPUT INT 21 4096U

Flag to indicate printing of 2:1 interior loop contribution.

• #define VRNA CONVERT OUTPUT INT 22 8192U

Flag to indicate printing of 2:2 interior loop contribution.

• #define VRNA CONVERT OUTPUT BULGE 16384U

Flag to indicate printing of bulge loop contribution.

• #define VRNA CONVERT OUTPUT INT 32768U

Flag to indicate printing of interior loop contribution.

• #define VRNA\_CONVERT\_OUTPUT\_ML 65536U

Flag to indicate printing of multi loop contribution.

#define VRNA\_CONVERT\_OUTPUT\_MISC 131072U

Flag to indicate printing of misc contributions (such as terminalAU)

#define VRNA\_CONVERT\_OUTPUT\_SPECIAL\_HP 262144U

Flag to indicate printing of special hairpin contributions (tri-, tetra-, hexa-loops)

#define VRNA CONVERT OUTPUT VANILLA 524288U

Flag to indicate printing of given parameters only

Note

This option overrides all other output options, except VRNA\_CONVERT\_OUTPUT\_-DUMP!

#define VRNA\_CONVERT\_OUTPUT\_NINIO 1048576U

Flag to indicate printing of interior loop asymmetry contribution.

#define VRNA\_CONVERT\_OUTPUT\_DUMP 2097152U

Flag to indicate dumping the energy contributions from the library instead of an input file.

#### **Functions**

void convert\_parameter\_file (const char \*iname, const char \*oname, unsigned int options)

Convert/dump a Vienna 1.8.4 formatted energy parameter file.

#### 11.5.1 Detailed Description

Functions and definitions for energy parameter file format conversion.

#### 11.5.2 Function Documentation

11.5.2.1 void convert\_parameter\_file ( const char \* *iname*, const char \* *oname*, unsigned int *options* )

Convert/dump a Vienna 1.8.4 formatted energy parameter file.

The options argument allows to control the different output modes.

Currently available options are:

VRNA\_CONVERT\_OUTPUT\_ALL, VRNA\_CONVERT\_OUTPUT\_HP, VRNA\_CONVERT\_-OUTPUT\_STACK

VRNA\_CONVERT\_OUTPUT\_MM\_HP, VRNA\_CONVERT\_OUTPUT\_MM\_INT, VRNA\_CONVERT\_OUTPUT\_MM\_INT\_1N

VRNA\_CONVERT\_OUTPUT\_MM\_INT\_23, VRNA\_CONVERT\_OUTPUT\_MM\_MULTI, VRNA\_CONVERT\_OUTPUT\_MM\_EXT

VRNA\_CONVERT\_OUTPUT\_DANGLE5, VRNA\_CONVERT\_OUTPUT\_DANGLE3, VRNA\_-CONVERT\_OUTPUT\_INT\_11

VRNA\_CONVERT\_OUTPUT\_INT\_21, VRNA\_CONVERT\_OUTPUT\_INT\_22, VRNA\_CONVERT\_OUTPUT\_BULGE

VRNA\_CONVERT\_OUTPUT\_INT, VRNA\_CONVERT\_OUTPUT\_ML, VRNA\_CONVERT\_-OUTPUT\_MISC

VRNA\_CONVERT\_OUTPUT\_SPECIAL\_HP, VRNA\_CONVERT\_OUTPUT\_VANILLA, VRNA\_-CONVERT\_OUTPUT\_NINIO

# VRNA\_CONVERT\_OUTPUT\_DUMP

The defined options are fine for bitwise compare- and assignment-operations, e. g.: pass a collection of options as a single value like this:

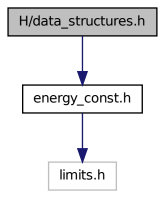
convert\_parameter\_file(ifile, ofile, option\_1 | option\_2 | option\_n)

iname	The input file name (If NULL input is read from stdin)
oname	The output file name (If NULL output is written to stdout)
options	The options (as described above)

# 11.6 H/data\_structures.h File Reference

All datastructures and typedefs shared among the Vienna RNA Package can be found here.

Include dependency graph for data\_structures.h:



This graph shows which files directly or indirectly include this file:



# **Data Structures**

struct plist

this datastructure is used as input parameter in functions of PS\_dot.h and others

• struct cpair

this datastructure is used as input parameter in functions of PS\_dot.c

struct COORDINATE

this is a workarround for the SWIG Perl Wrapper RNA plot function that returns an array of type COORDINATE

struct sect

stack of partial structures for backtracking

struct bondT

base pair

struct bondTEn

base pair with associated energy

struct paramT

The datastructure that contains temperature scaled energy parameters.

struct pf\_paramT

The datastructure that contains temperature scaled Boltzmann weights of the energy parameters.

struct PAIR

base pair data structure used in subopt.c

struct INTERVAL

sequence interval stack element used in subopt.c

struct SOLUTION

solution element from subopt.c

- struct cofoldF
- struct ConcEnt
- · struct pairpro
- struct pair\_info

A base pair info structure.

- struct move\_t
- struct intermediate\_t
- struct path\_t
- struct pu\_contrib
- struct interact
- struct pu\_out
- · struct constrain
- struct duplexT
- struct folden
- struct snoopT
- struct dupVar
- struct TwoDfold\_solution

Solution element returned from TwoDfoldList.

struct TwoDfold\_vars

Variables compound for 2Dfold MFE folding.

• struct TwoDpfold\_solution

Solution element returned from TwoDpfoldList.

struct TwoDpfold\_vars

Variables compound for 2Dfold partition function folding.

## **Defines**

• #define MAXALPHA 20

Maximal length of alphabet.

• #define MAXDOS 1000

Maximum density of states discretization for subopt.

#define FILENAME\_MAX\_LENGTH 80

Maximum length of filenames that are generated by our programs.

• #define FILENAME\_ID\_LENGTH 42

Maximum length of id taken from fasta header for filename generation.

## 11.6.1 Detailed Description

All datastructures and typedefs shared among the Vienna RNA Package can be found here

# 11.6.2 Define Documentation

# 11.6.2.1 #define FILENAME\_MAX\_LENGTH 80

Maximum length of filenames that are generated by our programs.

This definition should be used throughout the complete ViennaRNA package wherever a static array holding filenames of output files is declared.

# 11.6.2.2 #define FILENAME\_ID\_LENGTH 42

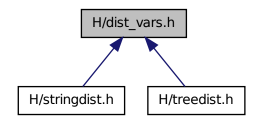
Maximum length of id taken from fasta header for filename generation.

this has to be smaller than FILENAME\_MAX\_LENGTH since in most cases, some suffix will be appended to the  $\ensuremath{\mathsf{ID}}$ 

# 11.7 H/dist vars.h File Reference

Global variables for Distance-Package.

This graph shows which files directly or indirectly include this file:



# **Data Structures**

- struct Postorder\_list
- struct Tree
- struct swString

# **Variables**

· int edit backtrack

Produce an alignment of the two structures being compared by tracing the editing path giving the minimum distance.

• char \* aligned\_line [4]

Contains the two aligned structures after a call to one of the distance functions with edit\_backtrack set to 1.

· int cost\_matrix

Specify the cost matrix to be used for distance calculations.

# 11.7.1 Detailed Description

Global variables for Distance-Package.

# 11.7.2 Variable Documentation

# 11.7.2.1 int edit\_backtrack

Produce an alignment of the two structures being compared by tracing the editing path giving the minimum distance.

set to 1 if you want backtracking

## 11.7.2.2 int cost\_matrix

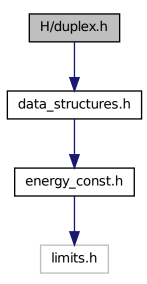
Specify the cost matrix to be used for distance calculations.

if 0, use the default cost matrix (upper matrix in example), otherwise use Shapiro's costs (lower matrix).

# 11.8 H/duplex.h File Reference

Duplex folding function declarations...

Include dependency graph for duplex.h:



# 11.8.1 Detailed Description

Duplex folding function declarations...

# 11.9 H/edit\_cost.h File Reference

global variables for Edit Costs included by treedist.c and stringdist.c

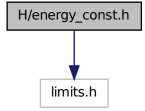
# 11.9.1 Detailed Description

global variables for Edit Costs included by treedist.c and stringdist.c

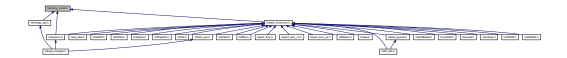
# 11.10 H/energy\_const.h File Reference

energy constants

Include dependency graph for energy\_const.h:



This graph shows which files directly or indirectly include this file:



# **Defines**

• #define GASCONST 1.98717

The gas constant.

• #define K0 273.15

0 deg Celsius in Kelvin

• #define INF (INT\_MAX/10)

Infinity as used in minimization routines.

• #define FORBIDDEN 9999

forbidden

• #define BONUS 10000

bonus contribution

• #define NBPAIRS 7

The number of distinguishable base pairs.

• #define TURN 3

The minimum loop length.

• #define MAXLOOP 30

The maximum loop length.

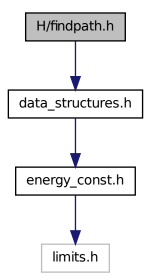
# 11.10.1 Detailed Description

energy constants

# 11.11 H/findpath.h File Reference

Compute direct refolding paths between two secondary structures.

Include dependency graph for findpath.h:



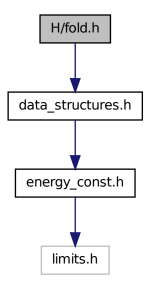
# 11.11.1 Detailed Description

Compute direct refolding paths between two secondary structures.

# 11.12 H/fold.h File Reference

MFE calculations and energy evaluations for single RNA sequences.

Include dependency graph for fold.h:



# **Functions**

- float fold (const char \*sequence, char \*structure)
  - Compute minimum free energy and an appropriate secondary structure of an RNA sequence.
- float energy\_of\_structure (const char \*string, const char \*structure, int verbosity\_level)

Calculate the free energy of an already folded RNA.

• int energy\_of\_structure\_pt (const char \*string, short \*ptable, short \*s, short \*s1, int verbosity level)

Calculate the free energy of an already folded RNA.

void free\_arrays (void)

Free arrays for mfe folding.

• void parenthesis\_structure (char \*structure, bondT \*bp, int length)

Create a dot-backet/parenthesis structure from backtracking stack.

void parenthesis\_zuker (char \*structure, bondT \*bp, int length)

Create a dot-backet/parenthesis structure from backtracking stack obtained by zuker suboptimal calculation in cofold.c.

void update\_fold\_params (void)

Recalculate energy parameters.

• float circfold (const char \*string, char \*structure)

Compute minimum free energy and an appropriate secondary structure of an RNA sequence assuming it to be circular instead of linear.

float energy\_of\_circ\_structure (const char \*string, const char \*structure, int verbosity\_-level)

Calculate the free energy of an already folded circular RNA.

void assign\_plist\_from\_db (plist \*\*pl, const char \*struc, float pr)

Create a plist from a dot-bracket string.

- int LoopEnergy (int n1, int n2, int type, int type\_2, int si1, int sj1, int sp1, int sq1)
- int HairpinE (int size, int type, int si1, int sj1, const char \*string)
- void initialize\_fold (int length)

Allocate arrays for folding

• float energy of struct (const char \*string, const char \*structure)

Calculate the free energy of an already folded RNA.

• int energy\_of\_struct\_pt (const char \*string, short \*ptable, short \*s, short \*s1)

Calculate the free energy of an already folded RNA.

• float energy\_of\_circ\_struct (const char \*string, const char \*structure)

Calculate the free energy of an already folded circular RNA.

## **Variables**

• int logML

if nonzero use logarithmic ML energy in energy\_of\_struct

• int uniq\_ML

do ML decomposition uniquely (for subopt)

int cut\_point

brief set to first pos of second seq for cofolding

• int eos\_debug

brief verbose info from energy\_of\_struct

# 11.12.1 Detailed Description

MFE calculations and energy evaluations for single RNA sequences. This file includes (almost) all function declarations within the RNAlib that are related to MFE folding...

#### 11.12.2 Function Documentation

11.12.2.1 float fold ( const char \* sequence, char \* structure )

Compute minimum free energy and an appropriate secondary structure of an RNA sequence.

The first parameter given, the RNA sequence, must be uppercase and should only contain an alphabet  $\Sigma$  that is understood by the RNAlib

(e.g. 
$$\Sigma = \{A, U, C, G\}$$
)

The second parameter, structure, must always point to an allocated block of memory with a size of at least strlen(sequence) + 1 Depending on the global variable fold\_constrained structure may contain a secondary structure constraint in an enhanced dot-bracket notation. The characters "  $\mid x < >$  " mark bases that are paired, unpaired, paired upstream, or downstream, respectively. Matching brackets " ( ) " denote base pairs, dots "." are used for unconstrained bases. Constrained folding works by assigning bonus energies to all structures that comply with the constraint.

After a successful call of fold(), a backtracked secondary structure (in dot-bracket notation) that exhibits the minimum of free energy will be written to the memory *structure* is pointing to. The function returns the minimum of free energy for any fold of the sequence given. Of course, the resulting free energy depends strongly on the energy model selected.

#### See also

circfold(), dangles, noLonelyPairs, noGU, no\_closingGU, tetra\_loop

#### **Parameters**

sequend	e RNA sequence
structur	A pointer to the character array where the secondary structure in dot-bracket
	notation will be written to

## **Returns**

the minimum free energy (MFE) in kcal/mol

11.12.2.2 float energy\_of\_structure ( const char \* string, const char \* structure, int verbosity\_level )

Calculate the free energy of an already folded RNA.

If verbosity level is set to a value >0, energies of structure elements are printed to stdout

# See also

energy of circ structure(), energy of structure pt()

	string	RNA sequence
str	ructure	secondary structure in dot-bracket notation
verb	osity level	a flag to turn verbose output on/off

## Returns

the free energy of the input structure given the input sequence in kcal/mol

11.12.2.3 int energy\_of\_structure\_pt ( const char \* string, short \* ptable, short \* s, short \* s, short \* s, int verbosity\_level )

Calculate the free energy of an already folded RNA.

If verbosity level is set to a value >0, energies of structure elements are printed to stdout

## See also

make\_pair\_table(), energy\_of\_struct()

## **Parameters**

string	RNA sequence
ptable	the pair table of the secondary structure
S	encoded RNA sequence
s1	encoded RNA sequence
verbosity level	a flag to turn verbose output on/off

# Returns

the free energy of the input structure given the input sequence in 10kcal/mol

11.12.2.4 void parenthesis\_structure ( char \* structure, bondT \* bp, int length )

Create a dot-backet/parenthesis structure from backtracking stack.

#### Note

This function is threadsafe

11.12.2.5 void parenthesis\_zuker ( char \* structure, bondT \* bp, int length )

Create a dot-backet/parenthesis structure from backtracking stack obtained by zuker suboptimal calculation in cofold.c.

# Note

This function is threadsafe

11.12.2.6 float circfold ( const char \* string, char \* structure )

Compute minimum free energy and an appropriate secondary structure of an RNA sequence assuming it to be circular instead of linear.

#### See also

fold(), dangles, noLonelyPairs, noGU, no\_closingGU, tetra\_loop

## **Parameters**

string	RNA sequence
structure	A pointer to the character array the secondary structure in dot-bracket nota-
	tion will be written to

## Returns

the minimum free energy (MFE) in kcal/mol

11.12.2.7 float energy\_of\_circ\_structure ( const char \* string, const char \* structure, int verbosity\_level )

Calculate the free energy of an already folded circular RNA.

If verbosity level is set to a value >0, energies of structure elements are printed to stdout

# See also

energy\_of\_struct(), energy\_of\_struct\_pt()

# **Parameters**

string	RNA sequence
structure	secondary structure in dot-bracket notation
verbosity	a flag to turn verbose output on/off
level	

# Returns

the free energy of the input structure given the input sequence in kcal/mol

11.12.2.8 void assign\_plist\_from\_db ( plist \*\* pl, const char \* struc, float pr )

Create a plist from a dot-bracket string.

The dot-bracket string is parsed and for each base pair an entry in the plist is created. The probability of each pair in the list is set by a function parameter.

The end of the plist is marked by sequence positions i as well as j equal to 0. This condition should be used to stop looping over its entries

This function is threadsafe

#### **Parameters**

pl	A pointer to the plist that is to be created
struc	The secondary structure in dot-bracket notation
pr	The probability for each base pair

11.12.2.9 int LoopEnergy ( int n1, int n2, int type, int type\_2, int si1, int sj1, int sp1, int sq1 )

# **Deprecated**

{This function is deprecated and will be removed soon.

```
Use E_IntLoop() instead!}
```

11.12.2.10 int HairpinE (int size, int type, int si1, int sj1, const char \* string)

# **Deprecated**

{This function is deprecated and will be removed soon.

```
Use E_Hairpin() instead!}
```

11.12.2.11 void initialize\_fold ( int length )

Allocate arrays for folding

.

# **Deprecated**

{This function is deprecated and will be removed soon!}

```
11.12.2.12 float energy_of_struct ( const char * string, const char * structure )
```

Calculate the free energy of an already folded RNA.

# Note

This function is not entirely threadsafe! Depending on the state of the global variable eos\_debug it prints energy information to stdout or not...

# **Deprecated**

This function is deprecated and should not be used in future programs! Use energy\_of structure() instead!

## See also

```
energy_of_structure, energy_of_circ_struct(), energy_of_struct_pt()
```

#### **Parameters**

string	RNA sequence
structure	secondary structure in dot-bracket notation

## **Returns**

the free energy of the input structure given the input sequence in kcal/mol

11.12.2.13 int energy\_of\_struct\_pt ( const char \* string, short \* ptable, short \* s, short \* s1 )

Calculate the free energy of an already folded RNA.

#### Note

This function is not entirely threadsafe! Depending on the state of the global variable eos\_debug it prints energy information to stdout or not...

# **Deprecated**

This function is deprecated and should not be used in future programs! Use energy\_of\_structure\_pt() instead!

#### See also

make\_pair\_table(), energy\_of\_structure()

# **Parameters**

strina	RNA sequence
	·
ptable	the pair table of the secondary structure
s	encoded RNA sequence
s1	encoded RNA sequence

# Returns

the free energy of the input structure given the input sequence in 10kcal/mol

11.12.2.14 float energy\_of\_circ\_struct ( const char \* string, const char \* structure )

Calculate the free energy of an already folded circular RNA.

# Note

This function is not entirely threadsafe! Depending on the state of the global variable eos\_debug it prints energy information to stdout or not...

# **Deprecated**

This function is deprecated and should not be used in future programs Use energy\_of\_circ\_structure() instead!

# See also

 $energy\_of\_circ\_structure(),\ energy\_of\_struct(),\ energy\_of\_struct\_pt()$ 

## **Parameters**

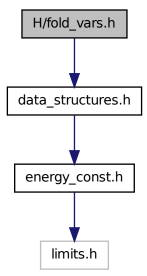
string	RNA sequence
structure	secondary structure in dot-bracket notation

## Returns

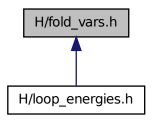
the free energy of the input structure given the input sequence in kcal/mol

# 11.13 H/fold\_vars.h File Reference

Here all all declarations of the global variables used throughout RNAlib. Include dependency graph for fold\_vars.h:



This graph shows which files directly or indirectly include this file:



## **Variables**

· int fold\_constrained

Global switch to activate/deactivate folding with structure constraints.

· int noLonelyPairs

Global switch to avoid/allow helices of length 1.

• int dangles

Switch the energy model for dangling end contributions (0, 1, 2, 3)

• int noGU

Global switch to forbid/allow GU base pairs at all.

• int no\_closingGU

GU allowed only inside stacks if set to 1.

int tetra\_loop

Include special stabilizing energies for some tri-, tetra- and hexa-loops;.

· int energy\_set

0 = BP; 1=any mit GC; 2=any mit AU-parameter

• int circ

backward compatibility variable.

int csv

generate comma seperated output

• int oldAliEn

use old alifold energies (with gaps)

• int ribo

use ribosum matrices

• char \* RibosumFile

warning this variable will vanish in the future ribosums will be compiled in instead

• char \* nonstandards

contains allowed non standard base pairs

· double temperature

Rescale energy parameters to a temperature in degC.

· int james rule

interior loops of size 2 get energy 0.8Kcal and no mismatches, default 1

int logML

use logarithmic multiloop energy function

· int cut\_point

Marks the position (starting from 1) of the first nucleotide of the second molecule within the concatenated sequence.

bondT \* base\_pair

Contains a list of base pairs after a call to fold().

FLT OR DBL \* pr

A pointer to the base pair probability matrix.

• int \* iindx

index array to move through pr.

double pf scale

A scaling factor used by pf\_fold() to avoid overflows.

· int do backtrack

do backtracking, i.e.

char backtrack\_type

A backtrack array marker for inverse\_fold()

# 11.13.1 Detailed Description

Here all all declarations of the global variables used throughout RNAlib.

# 11.13.2 Variable Documentation

# 11.13.2.1 int noLonelyPairs

Global switch to avoid/allow helices of length 1.

Disallow all pairs which can only occur as lonely pairs (i.e. as helix of length 1). This avoids lonely base pairs in the predicted structures in most cases.

# 11.13.2.2 int dangles

Switch the energy model for dangling end contributions (0, 1, 2, 3)

If set to 0 no stabilizing energies are assigned to bases adjacent to helices in free ends and multiloops (so called dangling ends). Normally (dangles = 1) dangling end energies are assigned only to unpaired bases and a base cannot participate simultaneously in two dangling ends. In the partition function algorithm pf\_fold() these checks are neglected. If dangles is set to 2, all folding routines will follow this convention. This

treatment of dangling ends gives more favorable energies to helices directly adjacent to one another, which can be beneficial since such helices often do engage in stabilizing interactions through co-axial stacking.

If dangles = 3 co-axial stacking is explicitly included for adjacent helices in mutli-loops. The option affects only mfe folding and energy evaluation (fold() and energy\_of\_structure()), as well as suboptimal folding (subopt()) via re-evaluation of energies. Co-axial stacking with one intervening mismatch is not considered so far.

Default is 2 in most algorithms, partition function algorithms can only handle 0 and 2

11.13.2.3 int tetra\_loop

Include special stabilizing energies for some tri-, tetra- and hexa-loops;.

default is 1.

11.13.2.4 int energy\_set

0 = BP; 1=any mit GC; 2=any mit AU-parameter

If set to 1 or 2: fold sequences from an artificial alphabet ABCD..., where A pairs B, C pairs D, etc. using either GC (1) or AU parameters (2); default is 0, you probably don't want to change it.

11.13.2.5 int circ

backward compatibility variable.

. this does not effect anything

11.13.2.6 char\* nonstandards

contains allowed non standard base pairs

Lists additional base pairs that will be allowed to form in addition to GC, CG, AU, UA, GU and UG. Nonstandard base pairs are given a stacking energy of 0.

11.13.2.7 double temperature

Rescale energy parameters to a temperature in degC.

Default is 37C. You have to call the update\_...\_params() functions after changing this parameter.

11.13.2.8 int cut\_point

Marks the position (starting from 1) of the first nucleotide of the second molecule within the concatenated sequence.

To evaluate the energy of a duplex structure (a structure formed by two strands), concatenate the to sequences and set it to the first base of the second strand in the concatenated sequence. The default value of -1 stands for single molecule folding. The cut\_point variable is also used by PS\_rna\_plot() and PS\_dot\_plot() to mark the chain break in postscript plots.

```
11.13.2.9 bondT* base_pair
```

Contains a list of base pairs after a call to fold().

base\_pair[0].i contains the total number of pairs.

## **Deprecated**

Do not use this variable anymore!

```
11.13.2.10 FLT_OR_DBL* pr
```

A pointer to the base pair probability matrix.

## **Deprecated**

Do not use this variable anymore!

```
11.13.2.11 int* iindx
```

index array to move through pr.

The probability for base i and j to form a pair is in pr[iindx[i]-j].

# Deprecated

Do not use this variable anymore!

```
11.13.2.12 double pf_scale
```

A scaling factor used by pf\_fold() to avoid overflows.

Should be set to approximately exp((-F/kT)/length), where F is an estimate for the ensemble free energy, for example the minimum free energy. You must call update\_pf\_params() after changing this parameter.

If pf\_scale is -1 (the default), an estimate will be provided automatically when computing partition functions, e.g. pf\_fold() The automatic estimate is usually insufficient for sequences more than a few hundred bases long.

#### 11.13.2.13 int do backtrack

do backtracking, i.e.

compute secondary structures or base pair probabilities

If 0, do not calculate pair probabilities in pf\_fold(); this is about twice as fast. Default is 1.

#### 11.13.2.14 char backtrack\_type

A backtrack array marker for inverse\_fold()

If set to 'C': force (1,N) to be paired, 'M' fold as if the sequence were inside a multi-loop. Otherwise ('F') the usual mfe structure is computed.

# 11.14 H/inverse.h File Reference

Inverse folding routines.

## **Functions**

• float inverse\_fold (char \*start, const char \*target)

Find sequences with predefined structure.

• float inverse\_pf\_fold (char \*start, const char \*target)

Find sequence that maximizes probability of a predefined structure.

# **Variables**

char \* symbolset

This global variable points to the allowed bases, initially "AUGC".

· float final\_cost

when to stop inverse\_pf\_fold()

• int give\_up

default 0: try to minimize structure distance even if no exact solution can be found

• int inv\_verbose

print out substructure on which inverse\_fold() fails

# 11.14.1 Detailed Description

Inverse folding routines.

#### 11.14.2 Function Documentation

11.14.2.1 float inverse\_fold ( char \* start, const char \* target )

Find sequences with predefined structure.

This function searches for a sequence with minimum free energy structure provided in the parameter 'target', starting with sequence 'start'. It returns 0 if the search was successful, otherwise a structure distance in terms of the energy difference between the search result and the actual target 'target' is returned. The found sequence is returned in 'start'. If give\_up is set to 1, the function will return as soon as it is clear that the search will be unsuccessful, this speeds up the algorithm if you are only interested in exact solutions.

#### **Parameters**

start	The start sequence
target	The target secondary structure in dot-bracket notation

#### Returns

The distance to the target in case a search was unsuccessful, 0 otherwise

11.14.2.2 float inverse\_pf\_fold ( char \* start, const char \* target )

Find sequence that maximizes probability of a predefined structure.

This function searches for a sequence with maximum probability to fold into the provided structure 'target' using the partition function algorithm. It returns  $-kT \cdot \log(p)$  where p is the frequency of 'target' in the ensemble of possible structures. This is usually much slower than inverse\_fold().

#### **Parameters**

start	The start sequence
target	The target secondary structure in dot-bracket notation

#### Returns

The distance to the target in case a search was unsuccessful, 0 otherwise

# 11.14.3 Variable Documentation

11.14.3.1 char\* symbolset

This global variable points to the allowed bases, initially "AUGC".

It can be used to design sequences from reduced alphabets.

# 11.15 H/Lfold.h File Reference

Predicting local MFE structures of large sequences.

#### **Functions**

- float Lfold (const char \*string, char \*structure, int maxdist)

  The local analog to fold().
- float aliLfold (const char \*\*strings, char \*structure, int maxdist)
- float Lfoldz (const char \*string, char \*structure, int maxdist, int zsc, double min\_z)

# 11.15.1 Detailed Description

Predicting local MFE structures of large sequences.

# 11.15.2 Function Documentation

11.15.2.1 float Lfold ( const char \* string, char \* structure, int maxdist )

The local analog to fold().

Computes the minimum free energy structure including only base pairs with a span smaller than 'maxdist'

## **Parameters**

string	
structure	
maxdist	

11.15.2.2 float aliLfold ( const char \*\* strings, char \* structure, int maxdist )

# **Parameters**

strings	
structure	
maxdist	

# Returns

11.15.2.3 float Lfoldz ( const char \* string, char \* structure, int maxdist, int zsc, double min\_z

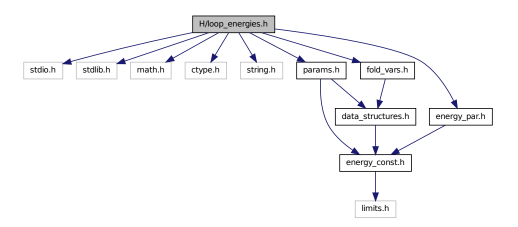
#### **Parameters**

string	
structure	
maxdist	
ZSC	
min_z	

# 11.16 H/loop\_energies.h File Reference

Energy evaluation for MFE and partition function calculations.

Include dependency graph for loop\_energies.h:



# **Functions**

- PRIVATE int E\_IntLoop (int n1, int n2, int type, int type\_2, int si1, int sj1, int sp1, int sq1, paramT \*P)
- PRIVATE int E\_Hairpin (int size, int type, int si1, int sj1, const char \*string, paramT \*P)
- PRIVATE int E\_Stem (int type, int si1, int sj1, int extLoop, paramT \*P)
- PRIVATE double exp\_E\_Stem (int type, int si1, int sj1, int extLoop, pf\_paramT \*P)
- PRIVATE double exp\_E\_Hairpin (int u, int type, short si1, short sj1, const char \*string, pf\_paramT \*P)

PRIVATE double exp\_E\_IntLoop (int u1, int u2, int type, int type2, short si1, short sj1, short sp1, short sq1, pf\_paramT \*P)

# 11.16.1 Detailed Description

Energy evaluation for MFE and partition function calculations. This file contains functions for the calculation of the free energy  $\Delta G$  of a hairpin- [E\_Hairpin()] or interior-loop [E\_IntLoop()].

The unit of the free energy returned is  $10^{-2}*kcal/mol$ 

In case of computing the partition function, this file also supplies functions which return the Boltzmann weights  $e^{-\Delta G/kT}$  for a hairpin- [ exp\_E\_Hairpin() ] or interior-loop [ exp\_E\_IntLoop() ].

# 11.16.2 Function Documentation

11.16.2.1 PRIVATE int E\_IntLoop ( int n1, int n2, int type, int  $type_2$ , int si1, int sj1, int sp1, int sg1, paramT \*P)

Compute the Energy of an interior-loop

This function computes the free energy  $\Delta G$  of an interior-loop with the following structure:

This general structure depicts an interior-loop that is closed by the base pair (X,Y). The enclosed base pair is (V,U) which leaves the unpaired bases a\_1-a\_n and b\_1-b\_n that constitute the loop. In this example, the length of the interior-loop is (n+m) where n or m may be 0 resulting in a bulge-loop or base pair stack. The mismatching nucleotides for the closing pair (X,Y) are:

5'-mismatch: a\_1 3'-mismatch: b m

and for the enclosed base pair (V,U):

5'-mismatch: b 1

3'-mismatch: a\_n

## Note

Base pairs are always denoted in 5'->3' direction. Thus the enclosed base pair must be 'turned arround' when evaluating the free energy of the interior-loop

#### See also

```
scale_parameters()
paramT
```

# Note

This function is threadsafe

#### **Parameters**

n1	The size of the 'left'-loop (number of unpaired nucleotides)
n2	The size of the 'right'-loop (number of unpaired nucleotides)
type	The pair type of the base pair closing the interior loop
type_2	The pair type of the enclosed base pair
si1	The 5'-mismatching nucleotide of the closing pair
sj1	The 3'-mismatching nucleotide of the closing pair
sp1	The 3'-mismatching nucleotide of the enclosed pair
sq1	The 5'-mismatching nucleotide of the enclosed pair
Р	The datastructure containing scaled energy parameters

# Returns

The Free energy of the Interior-loop in dcal/mol

```
11.16.2.2 PRIVATE int E_Hairpin ( int size, int type, int si1, int sj1, const char * string, paramT * P )
```

Compute the Energy of a hairpin-loop

To evaluate the free energy of a hairpin-loop, several parameters have to be known. A general hairpin-loop has this structure:

where X-Y marks the closing pair [e.g. a (G,C) pair]. The length of this loop is 6 as there are six unpaired nucleotides (a1-a6) enclosed by (X,Y). The 5' mismatching

nucleotide is a1 while the 3' mismatch is a6. The nucleotide sequence of this loop is "a1.a2.a3.a4.a5.a6"

#### Note

The parameter sequence should contain the sequence of the loop in capital letters of the nucleic acid alphabet if the loop size is below 7. This is useful for unusually stable tri-, tetra- and hexa-loops which are treated differently (based on experimental data) if they are tabulated.

#### See also

```
scale_parameters()
paramT
```

# Warning

Not (really) thread safe! A threadsafe implementation will replace this function in a future release!

Energy evaluation may change due to updates in global variable "tetra\_loop"

#### **Parameters**

size	The size of the loop (number of unpaired nucleotides)
type	The pair type of the base pair closing the hairpin
si1	The 5'-mismatching nucleotide
sj1	The 3'-mismatching nucleotide
string	The sequence of the loop
Р	The datastructure containing scaled energy parameters

#### Returns

The Free energy of the Hairpin-loop in dcal/mol

```
11.16.2.3 PRIVATE int E_Stem ( int type, int si1, int sj1, int extLoop, paramT * P )
```

Compute the energy contribution of a stem branching off a loop-region

This function computes the energy contribution of a stem that branches off a loop region. This can be the case in multiloops, when a stem branching off increases the degree of the loop but also *immediately interior base pairs* of an exterior loop contribute free energy. To switch the bahavior of the function according to the evaluation of a multiloop- or exterior-loop-stem, you pass the flag 'extLoop'. The returned energy contribution consists of a TerminalAU penalty if the pair type is greater than 2, dangling end contributions of mismatching nucleotides adjacent to the stem if only one of the si1, sj1 parameters is greater than 0 and mismatch energies if both mismatching nucleotides are positive values. Thus, to avoid incooperating dangling end or mismatch energies just pass a negative number, e.g. -1 to the mismatch argument.

This is an illustration of how the energy contribution is assembled:

Here, (X,Y) is the base pair that closes the stem that branches off a loop region. The nucleotides si1 and sj1 are the 5'- and 3'- mismatches, respectively. If the base pair type of (X,Y) is greater than 2 (i.e. an A-U or G-U pair, the TerminalAU penalty will be included in the energy contribution returned. If si1 and sj1 are both nonnegative numbers, mismatch energies will also be included. If one of sij or sj1 is a negtive value, only 5' or 3' dangling end contributions are taken into account. To prohibit any of these mismatch contributions to be incoorporated, just pass a negative number to both, si1 and sj1. In case the argument extLoop is 0, the returned energy contribution also includes the *internal-loop-penalty* of a multiloop stem with closing pair type.

#### See also

E\_MLstem()
E\_ExtLoop()

#### Note

This function is threadsafe

#### **Parameters**

type	The pair type of the first base pair un the stem
si1	The 5'-mismatching nucleotide
sj1	The 3'-mismatching nucleotide
extLoop	A flag that indicates whether the contribution reflects the one of an exterior
	loop or not
Р	The datastructure containing scaled energy parameters

#### Returns

The Free energy of the branch off the loop in dcal/mol

Compute the Boltzmann weighted energy contribution of a stem branching off a loop-region

This is the partition function variant of E\_Stem()

#### See also

E Stem()

#### Note

This function is threadsafe

### Returns

The Boltzmann weighted energy contribution of the branch off the loop

```
11.16.2.5 PRIVATE double exp_E_Hairpin ( int u, int type, short si1, short sj1, const char * string, pf_paramT * P )
```

Compute Boltzmann weight  $e^{-\Delta G/kT}$  of a hairpin loop

multiply by scale[u+2]

#### See also

```
get_scaled_pf_parameters()
pf_paramT
E_Hairpin()
```

### Warning

Not (really) thread safe! A threadsafe implementation will replace this function in a future release!

Energy evaluation may change due to updates in global variable "tetra\_loop"

#### **Parameters**

и	The size of the loop (number of unpaired nucleotides)
type	The pair type of the base pair closing the hairpin
si1	The 5'-mismatching nucleotide
sj1	The 3'-mismatching nucleotide
string	The sequence of the loop
Р	The datastructure containing scaled Boltzmann weights of the energy pa-
	rameters

### Returns

The Boltzmann weight of the Hairpin-loop

```
11.16.2.6 PRIVATE double exp_E_IntLoop ( int u1, int u2, int type, int type2, short si1, short sj1, short sg1, pf_paramT * P )
```

Compute Boltzmann weight  $e^{-\Delta G/kT}$  of interior loop

multiply by scale[u1+u2+2] for scaling

### See also

```
get_scaled_pf_parameters()
pf_paramT
E_IntLoop()
```

### Note

This function is threadsafe

### **Parameters**

The size of the 'left'-loop (number of unpaired nucleotides)
The size of the 'right'-loop (number of unpaired nucleotides)
The pair type of the base pair closing the interior loop
The pair type of the enclosed base pair
The 5'-mismatching nucleotide of the closing pair
The 3'-mismatching nucleotide of the closing pair
The 3'-mismatching nucleotide of the enclosed pair
The 5'-mismatching nucleotide of the enclosed pair
The datastructure containing scaled Boltzmann weights of the energy pa-
rameters

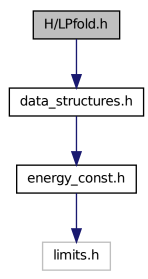
### Returns

The Boltzmann weight of the Interior-loop

# 11.17 H/LPfold.h File Reference

Function declarations of partition function variants of the Lfold algorithm.

Include dependency graph for LPfold.h:



#### **Functions**

- void update pf paramsLP (int length)
- plist \* pfl\_fold (char \*sequence, int winSize, int pairSize, float cutoffb, double \*\*pU, struct plist \*\*dpp2, FILE \*pUfp, FILE \*spup)

Compute partition functions for locally stable secondary structures (berni! update me)

- void putoutpU\_prob (double \*\*pU, int length, int ulength, FILE \*fp, int energies)

  Writes the unpaired probabilities (pU) or opening energies into a file.
- void putoutpU\_prob\_bin (double \*\*pU, int length, int ulength, FILE \*fp, int energies)

Writes the unpaired probabilities (pU) or opening energies into a binary file.

void init pf foldLP (int length)

Dunno if this function was ever used by external programs linking to RNAlib, but it was declared PUBLIC before.

### 11.17.1 Detailed Description

Function declarations of partition function variants of the Lfold algorithm.

#### 11.17.2 Function Documentation

11.17.2.1 void update\_pf\_paramsLP ( int length )

### **Parameters**

length

11.17.2.2 plist\* pfl\_fold ( char \* sequence, int winSize, int pairSize, float cutoffb, double \*\* pU, struct plist \*\* dpp2, FILE \* pUfp, FILE \* spup )

Compute partition functions for locally stable secondary structures (berni! update me)

pfl\_fold computes partition functions for every window of size 'winSize' possible in a RNA molecule, allowing only pairs with a span smaller than 'pairSize'. It returns the mean pair probabilities averaged over all windows containing the pair in 'pl'. 'winSize' should always be >= 'pairSize'. Note that in contrast to Lfold(), bases outside of the window do not influence the structure at all. Only probabilities higher than 'cutoffb' are kept.

If 'pU' is supplied (i.e is not the NULL pointer), pfl\_fold() will also compute the mean probability that regions of length 'u' and smaller are unpaired. The parameter 'u' is supplied in 'pup[0][0]'. On return the 'pup' array will contain these probabilities, with the entry on 'pup[x][y]' containing the mean probability that x and the y-1 preceding bases are unpaired. The 'pU' array needs to be large enough to hold n+1 float\* entries, where n is the sequence length.

If an array dpp2 is supplied, the probability of base pair (i,j) given that there already

exists a base pair (i+1,j-1) is also computed and saved in this array. If pUfp is given (i.e. not NULL), pU is not saved but put out imediately. If spup is given (i.e. is not NULL), the pair probabilities in pl are not saved but put out imediately.

#### **Parameters**

sequence	RNA sequence
winSize	size of the window
pairSize	maximum size of base pair
cutoffb	cutoffb for base pairs
рU	array holding all unpaired probabilities
dpp2	array of dependent pair probabilities
pUfp	file pointer for pU
spup	file pointer for pair probabilities

#### Returns

list of pair probabilities

11.17.2.3 void putoutpU\_prob ( double \*\* pU, int length, int ulength, FILE \* fp, int energies )

Writes the unpaired probabilities (pU) or opening energies into a file.

Can write either the unpaired probabilities (accessibilities) pU or the opening energies -log(pU)kT into a file

### **Parameters**

pU	pair probabilities
length	length of RNA sequence
ulength	maximum length of unpaired stretch
fp	file pointer of destination file
energies	switch to put out as opening energies

11.17.2.4 void putoutpU\_prob\_bin ( double \*\* pU, int length, int ulength, FILE \* fp, int energies )

Writes the unpaired probabilities (pU) or opening energies into a binary file.

Can write either the unpaired probabilities (accessibilities) pU or the opening energies -log(pU)kT into a file

### **Parameters**

рU	pair probabilities
length	length of RNA sequence
ulength	maximum length of unpaired stretch
fp	file pointer of destination file
energies	switch to put out as opening energies

11.17.2.5 void init\_pf\_foldLP ( int length )

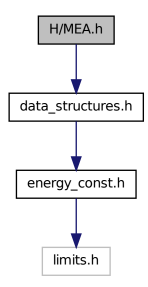
Dunno if this function was ever used by external programs linking to RNAlib, but it was declared PUBLIC before.

Anyway, never use this function as it will be removed soon and does nothing at all

### 11.18 H/MEA.h File Reference

Computes a MEA (maximum expected accuracy) structure.

Include dependency graph for MEA.h:



## **Functions**

float MEA (plist \*p, char \*structure, double gamma)
 Computes a MEA (maximum expected accuracy) structure.

### 11.18.1 Detailed Description

Computes a MEA (maximum expected accuracy) structure.

#### 11.18.2 Function Documentation

11.18.2.1 float MEA ( plist \*p, char \*structure, double gamma )

Computes a MEA (maximum expected accuracy) structure.

The algorithm maximizes the expected accuracy

$$A(S) = \sum_{(i,j)\in S} 2\gamma p_{ij} + \sum_{i\notin S} p_i^u$$

Higher values of  $\gamma$  result in more base pairs of lower probability and thus higher sensitivity. Low values of  $\gamma$  result in structures containing only highly likely pairs (high specificity). The code of the MEA function also demonstrates the use of sparse dynamic programming scheme to reduce the time and memory complexity of folding.

## 11.19 H/mm.h File Reference

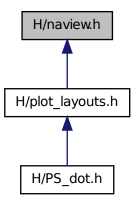
Several Maximum Matching implementations.

### 11.19.1 Detailed Description

Several Maximum Matching implementations. This file contains the declarations for several maximum matching implementations

# 11.20 H/naview.h File Reference

This graph shows which files directly or indirectly include this file:

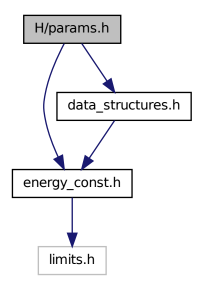


# 11.20.1 Detailed Description

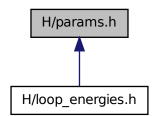
# 11.21 H/params.h File Reference

Several functions to obtain (pre)scaled energy parameter data containers.

Include dependency graph for params.h:



This graph shows which files directly or indirectly include this file:



### **Functions**

pf\_paramT \* get\_scaled\_pf\_parameters (void)

get a datastructure of type pf\_paramT which contains the Boltzmann weights of several energy parameters scaled according to the current temperature

### 11.21.1 Detailed Description

Several functions to obtain (pre)scaled energy parameter data containers.

### 11.21.2 Function Documentation

11.21.2.1 pf\_paramT\* get\_scaled\_pf\_parameters ( void )

get a datastructure of type pf\_paramT which contains the Boltzmann weights of several energy parameters scaled according to the current temperature

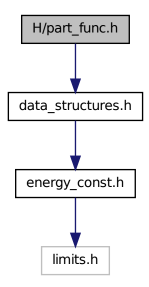
### **Returns**

The datastructure containing Boltzmann weights for use in partition function calculations

# 11.22 H/part\_func.h File Reference

Partition function of single RNA sequences.

Include dependency graph for part\_func.h:



### **Functions**

• float pf\_fold (const char \*sequence, char \*structure)

Compute the partition function Q of an RNA sequence.

• float pf\_circ\_fold (const char \*sequence, char \*structure)

Compute the partition function of a circular RNA sequence.

• char \* pbacktrack (char \*sequence)

Sample a secondary structure from the Boltzmann ensemble according its probability

• char \* pbacktrack\_circ (char \*sequence)

Sample a secondary structure of a circular RNA from the Boltzmann ensemble according its probability.

void free\_pf\_arrays (void)

Free arrays from pf\_fold()

• void update\_pf\_params (int length)

Recalculate energy parameters.

• FLT\_OR\_DBL \* export\_bppm (void)

Get a pointer to the base pair probability array.

void assign\_plist\_from\_pr (plist \*\*pl, FLT\_OR\_DBL \*probs, int length, double cutoff)

Create a plist from a probability matrix.

int get\_pf\_arrays (short \*\*S\_p, short \*\*S1\_p, char \*\*ptype\_p, FLT\_OR\_DBL \*\*qb\_p, FLT\_OR\_DBL \*\*qm\_p, FLT\_OR\_DBL \*\*q1k\_p, FLT\_OR\_DBL \*\*q1n\_p)

Get the pointers to (almost) all relavant computation arrays used in partition function computation.

char \* get centroid struct pl (int length, double \*dist, plist \*pl)

Get the centroid structure of the ensemble.

char \* get centroid struct pr (int length, double \*dist, FLT OR DBL \*pr)

Get the centroid structure of the ensemble.

double mean bp distance (int length)

Get the mean base pair distance of the last partition function computation.

double mean\_bp\_distance\_pr (int length, FLT\_OR\_DBL \*pr)

Get the mean base pair distance in the thermodynamic ensemble.

• void bppm to structure (char \*structure, FLT OR DBL \*pr, unsigned int length)

Create a dot-bracket like structure string from base pair probability matrix.

char bppm\_symbol (const float \*x)

Get a pseudo dot bracket notation for a given probability information.

void init\_pf\_fold (int length)

Allocate space for pf\_fold()

- char \* centroid (int length, double \*dist)
- double mean\_bp\_dist (int length)

get the mean pair distance of ensemble

- double expLoopEnergy (int u1, int u2, int type, int type2, short si1, short sj1, short sp1, short sq1)
- double expHairpinEnergy (int u, int type, short si1, short sj1, const char \*string)

### Variables

· int st back

a flag indicating that auxilary arrays are needed throughout the computations which are necessary for stochastic backtracking

### 11.22.1 Detailed Description

Partition function of single RNA sequences. This file includes (almost) all function declarations within the **RNAlib** that are related to Partion function folding...

### 11.22.2 Function Documentation

11.22.2.1 float pf\_fold ( const char \* sequence, char \* structure )

Compute the partition function Q of an RNA sequence.

If structure is not a NULL pointer on input, it contains on return a string consisting of the letters " . , | { } ( ) " denoting bases that are essentially unpaired, weakly paired, strongly paired without preference, weakly upstream (downstream) paired, or strongly up- (down-)stream paired bases, respectively. If  $fold_constrained$  is not 0, the structure string is interpreted on input as a list of constraints for the folding. The character "x" marks bases that must be unpaired, matching brackets " ( ) " denote base pairs, all other characters are ignored. Any pairs conflicting with the constraint will be forbidden. This is usually sufficient to ensure the constraints are honored. If  $fold_constraint_constrain$ 

#### Note

The global array pr is deprecated and the user who wants the computed base pair probabilities for further computations is advised to use the function export bppm()

#### See also

pf\_circ\_fold(), bppm\_to\_structure(), export\_bppm()

### **Parameters**

sequence	The RNA sequence to be computed
structure	A pointer to a char array where a base pair probability information might be
	stored in a pseudo-dot-bracket notation (might be NULL, too)

### Returns

The Gibbs free energy of the ensemble ( $G = -RT \cdot \log(Q)$ ) in kcal/mol

11.22.2.2 float pf\_circ\_fold ( const char \* sequence, char \* structure )

Compute the partition function of a circular RNA sequence.

#### See also

pf fold()

#### **Parameters**

sequence	The RNA sequence to be computed
structure	A pointer to a char array where a base pair probability information might be
	stored in a pseudo-dot-bracket notation (might be NULL, too)

#### **Returns**

The Gibbs free energy of the ensemble (  $G = -RT \cdot \log(Q)$ ) in kcal/mol

```
11.22.2.3 char* pbacktrack ( char * sequence )
```

Sample a secondary structure from the Boltzmann ensemble according its probability

#### **Parameters**

sequence	The RNA sequence

#### Returns

A sampled secondary structure in dot-bracket notation

```
11.22.2.4 char* pbacktrack_circ ( char * sequence )
```

Sample a secondary structure of a circular RNA from the Boltzmann ensemble according its probability.

This function does the same as pbacktrack() but assumes the RNA molecule to be circular

### **Parameters**

```
sequence The RNA sequence
```

### Returns

A sampled secondary structure in dot-bracket notation

```
11.22.2.5 void update_pf_params ( int length )
```

Recalculate energy parameters.

Call this function to recalculate the pair matrix and energy parameters after a change in folding parameters like temperature

```
11.22.2.6 FLT_OR_DBL* export_bppm ( void )
```

Get a pointer to the base pair probability array.

Accessing the base pair probabilities for a pair (i,j) is achieved by

```
FLT_OR_DBL *pr = export_bppm(); pr_ij = pr[iindx[i]-j];
```

#### See also

get iindx()

#### Returns

A pointer to the base pair probability array

11.22.2.7 void assign\_plist\_from\_pr ( plist \*\* pl, FLT\_OR\_DBL \* probs, int length, double cutoff )

Create a plist from a probability matrix.

The probability matrix given is parsed and all pair probabilities above the given threshold are used to create an entry in the plist

The end of the plist is marked by sequence positions i as well as j equal to 0. This condition should be used to stop looping over its entries

#### Note

This function is threadsafe

#### **Parameters**

pl	A pointer to the plist that is to be created
probs	The probability matrix used for creting the plist
length	The length of the RNA sequence
cutoff	The cutoff value

Get the pointers to (almost) all relavant computation arrays used in partition function computation.

### **Parameters**

S_p	A pointer to the 'S' array (integer representation of nucleotides)
S1_p	A pointer to the 'S1' array (2nd integer representation of nucleotides)
ptype_p	A pointer to the pair type matrix
· —	A pointer to the Q <sup>B</sup> matrix
qm_p	A pointer to the Q <sup>M</sup> matrix
q1k_p	A pointer to the 5' slice of the Q matrix ( $q1k(k) = Q(1,k)$ )
qln_p	A pointer to the 3' slice of the Q matrix ( $qln(l) = Q(l,n)$ )

#### Returns

Non Zero if everything went fine, 0 otherwise

11.22.2.9 char\* get\_centroid\_struct\_pl ( int *length*, double \* *dist*, plist \* *pl* )

Get the centroid structure of the ensemble.

This function is a threadsafe replacement for centroid() with a 'plist' input

The centroid is the structure with the minimal average distance to all other structures

$$< d(S) > = \sum_{(i,j) \in S} (1 - p_{ij}) + \sum_{(i,j) \notin S} p_{ij}$$

Thus, the centroid is simply the structure containing all pairs with  $p_i j > 0.5$  The distance of the centroid to the ensemble is written to the memory addressed by dist.

#### **Parameters**

length	The length of the sequence
dist	A pointer to the distance variable where the centroid distance will be written
	to
pl	A pair list containing base pair probability information about the ensemble

#### Returns

The centroid structure of the ensemble in dot-bracket notation

11.22.2.10 char\* get\_centroid\_struct\_pr ( int length, double \* dist, FLT\_OR\_DBL \* pr )

Get the centroid structure of the ensemble.

This function is a threadsafe replacement for centroid() with a probability array input

The centroid is the structure with the minimal average distance to all other structures

$$< d(S) > = \sum_{(i,j) \in S} (1 - p_{ij}) + \sum_{(i,j) \notin S} p_{ij}$$

Thus, the centroid is simply the structure containing all pairs with  $p_i j > 0.5$  The distance of the centroid to the ensemble is written to the memory addressed by *dist*.

#### **Parameters**

lei	ngth	The length of the sequence
	dist	A pointer to the distance variable where the centroid distance will be written
		to
	pr	A upper triangular matrix containing base pair probabilities (access via iindx
		get_iindx() )

#### Returns

The centroid structure of the ensemble in dot-bracket notation

11.22.2.11 double mean\_bp\_distance ( int length )

Get the mean base pair distance of the last partition function computation.

### See also

mean\_bp\_distance\_pr()

#### **Parameters**

length

### Returns

mean base pair distance in thermodynamic ensemble

11.22.2.12 double mean\_bp\_distance\_pr ( int length, FLT\_OR\_DBL \* pr )

Get the mean base pair distance in the thermodynamic ensemble.

This is a threadsafe implementation of mean\_bp\_dist()!

$$\langle d \rangle = \sum_{a,b} p_a p_b d(S_a, S_b)$$

this can be computed from the pair probs  $p_i j$  as

$$\langle d \rangle = \sum_{ij} p_{ij} (1 - p_{ij})$$

#### Note

This function is threadsafe

#### **Parameters**

length	The length of the sequence
pr	The matrix containing the base pair probabilities

### Returns

The mean pair distance of the structure ensemble

11.22.2.13 void init\_pf\_fold ( int length )

Allocate space for pf\_fold()

### **Deprecated**

This function is obsolete and will be removed soon!

11.22.2.14 char\* centroid ( int length, double \* dist )

#### **Deprecated**

This function is deprecated and should not be used anymore as it is not threadsafe!

#### See also

get\_centroid\_struct\_pl(), get\_centroid\_struct\_pr()

```
11.22.2.15 double mean_bp_dist ( int length )
```

get the mean pair distance of ensemble

### **Deprecated**

This function is not threadsafe and should not be used anymore. Use mean\_bp\_distance() instead!

11.22.2.16 double expLoopEnergy ( int u1, int u2, int type, int type2, short si1, short sj1, short sp1, short sq1)

### **Deprecated**

Use exp\_E\_IntLoop() from loop\_energies.h instead

11.22.2.17 double expHairpinEnergy ( int u, int type, short si1, short sj1, const char \*string )

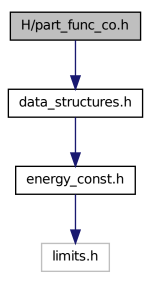
### **Deprecated**

Use exp\_E\_Hairpin() from loop\_energies.h instead

# 11.23 H/part\_func\_co.h File Reference

Partition function for two RNA sequences.

Include dependency graph for part\_func\_co.h:



### **Functions**

- cofoldF co\_pf\_fold (char \*sequence, char \*structure)
  - Calculate partition function and base pair probabilities.
- FLT\_OR\_DBL \* export\_co\_bppm (void)

Get a pointer to the base pair probability array.

void free\_co\_pf\_arrays (void)

Free the memory occupied by co\_pf\_fold()

• void update\_co\_pf\_params (int length)

Recalculate energy parameters.

 void compute\_probabilities (double FAB, double FEA, double FEB, struct plist \*prAB, struct plist \*prA, struct plist \*prB, int Alength)

Compute Boltzmann probabilities of dimerization without homodimers.

ConcEnt \* get\_concentrations (double FEAB, double FEAA, double FEBB, double FEA, double FEB, double \*startconc)

Given two start monomer concentrations a and b, compute the concentrations in thermodynamic equilibrium of all dimers and the monomers.

• plist \* get\_plist (struct plist \*pl, int length, double cut\_off)

DO NOT USE THIS FUNCTION ANYMORE.

void init\_co\_pf\_fold (int length)

DO NOT USE THIS FUNCTION ANYMORE.

#### **Variables**

· int mirnatog

Toggles no intrabp in 2nd mol.

• double F\_monomer [2]

Free energies of the two monomers.

### 11.23.1 Detailed Description

Partition function for two RNA sequences. As for folding one RNA molecule, this computes the partition function of all possible structures and the base pair probabilities. Uses the same global pf\_scale variable to avoid overflows.

To simplify the implementation the partition function computation is done internally in a null model that does not include the duplex initiation energy, i.e. the entropic penalty for producing a dimer from two monomers). The resulting free energies and pair probabilities are initially relative to that null model. In a second step the free energies can be corrected to include the dimerization penalty, and the pair probabilities can be divided into the conditional pair probabilities given that a re dimer is formed or not formed.

After computing the partition functions of all possible dimeres one can compute the probabilities of base pairs, the concentrations out of start concentrations and sofar and soaway.

Dimer formation is inherently concentration dependent. Given the free energies of the monomers A and B and dimers AB, AA, and BB one can compute the equilibrium concentrations, given input concentrations of A and B, see e.g. Dimitrov & Zuker (2004)

### 11.23.2 Function Documentation

11.23.2.1 cofoldF co\_pf\_fold ( char \* sequence, char \* structure )

Calculate partition function and base pair probabilities.

This is the cofold partition function folding. The second molecule starts at the cut\_point nucleotide.

### **Parameters**

sequence	Concatenated RNA sequences
structure	Will hold the structure or constraints

### Returns

cofoldF structure containing a set of energies needed for concentration computations.

```
11.23.2.2 FLT_OR_DBL* export_co_bppm ( void )
```

Get a pointer to the base pair probability array.

Accessing the base pair probabilities for a pair (i,j) is achieved by

```
FLT_OR_DBL *pr = export_bppm(); pr_ij = pr[iindx[i]-j];
```

#### See also

get iindx()

#### Returns

A pointer to the base pair probability array

11.23.2.3 void compute\_probabilities ( double *FAB*, double *FEA*, double *FEB*, struct plist \* prAB, struct plist \* prB, int Alength )

Compute Boltzmann probabilities of dimerization without homodimers.

Given the pair probabilities and free energies (in the null model) for a dimer AB and the two constituent monomers A and B, compute the conditional pair probabilities given that a dimer AB actually forms. Null model pair probabilities are given as a list as produced by <a href="mailto:assign\_plist\_from\_pr(">assign\_plist\_from\_pr()</a>, the dimer probabilities 'prAB' are modified in place.

#### **Parameters**

FAB	free energy of dimer AB
FEA	free energy of monomer A
FEB	free energy of monomer B
prAB	pair probabilities for dimer
prA	pair probabilities monomer
prB	pair probabilities monomer
Alength	Length of molecule A

11.23.2.4 ConcEnt\* get\_concentrations ( double FEAB, double FEAA, double FEBB, double FEA, double \*\* startconc\*)

Given two start monomer concentrations a and b, compute the concentrations in thermodynamic equilibrium of all dimers and the monomers.

This function takes an array 'startconc' of input concentrations with alternating entries for the initial concentrations of molecules A and B (terminated by two zeroes), then computes the resulting equilibrium concentrations from the free energies for the dimers. Dimer free energies should be the dimer-only free energies, i.e. the FcAB entries from the cofoldF struct.

### **Parameters**

FEAB	Free energy of AB dimer (FcAB entry)
FEAA	Free energy of AA dimer (FcAB entry)
FEBB	Free energy of BB dimer (FcAB entry)
FEA	Free energy of monomer A
FEB	Free energy of monomer B
startconc	List of start concentrations [a0],[b0],[a1],[b1],,[an][bn],[0],[0]

#### **Returns**

ConcEnt array containing the equilibrium energies and start concentrations

```
11.23.2.5 plist* get_plist ( struct plist * pl, int length, double cut_off )
```

DO NOT USE THIS FUNCTION ANYMORE.

### **Deprecated**

{ This function is deprecated and will be removed soon!} use assign\_plist\_from\_pr() instead!

11.23.2.6 void init\_co\_pf\_fold ( int length )

DO NOT USE THIS FUNCTION ANYMORE.

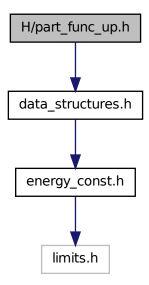
### **Deprecated**

{ This function is deprecated and will be removed soon!}

# 11.24 H/part\_func\_up.h File Reference

Partition Function Cofolding as stepwise process.

Include dependency graph for part\_func\_up.h:



### **Functions**

- pu\_contrib \* pf\_unstru (char \*sequence, int max\_w)
  - Calculate the partition function over all unpaired regions of a maximal length.
- interact \* pf\_interact (const char \*s1, const char \*s2, pu\_contrib \*p\_c, pu\_contrib \*p\_c2, int max\_w, char \*cstruc, int incr3, int incr5)

Calculates the probability of a local interaction between two sequences.

void free\_interact (interact \*pin)

Frees the output of function pf\_interact().

• void free\_pu\_contrib\_struct (pu\_contrib \*pu)

Frees the output of function pf\_unstru().

### 11.24.1 Detailed Description

Partition Function Cofolding as stepwise process. In this approach to cofolding the interaction between two RNA molecules is seen as a stepwise process. In a first step, the target molecule has to adopt a structure in which a binding site is accessible. In a second step, the ligand molecule will hybridize with a region accessible to an interaction. Consequently the algorithm is designed as a two step process: The first step is the calculation of the probability that a region within the target is unpaired, or equivalently,

the calculation of the free energy needed to expose a region. In the second step we compute the free energy of an interaction for every possible binding site.

#### 11.24.2 Function Documentation

```
11.24.2.1 pu_contrib* pf_unstru ( char * sequence, int max_w )
```

Calculate the partition function over all unpaired regions of a maximal length.

You have to call function pf\_fold() providing the same sequence before calling pf\_unstru(). If you want to calculate unpaired regions for a constrained structure, set variable 'structure' in function 'pf\_fold()' to the constrain string. It returns a pu\_contrib struct containing four arrays of dimension [i = 1 to length(sequence)][j = 0 to u-1] containing all possible contributions to the probabilities of unpaired regions of maximum length u. Each array in pu\_contrib contains one of the contributions to the total probability of being unpaired: The probability of being unpaired within an exterior loop is in array pu\_contrib->E, the probability of being unpaired within a hairpin loop is in array pu\_contrib->H, the probability of being unpaired within an interior loop is in array pu\_contrib->I and probability of being unpaired within a multi-loop is in array pu\_contrib->M. The total probability of being unpaired is the sum of the four arrays of pu\_contrib.

This function frees everything allocated automatically. To free the output structure call free pu contrib().

#### **Parameters**

sequence	
max_w	

#### Returns

```
11.24.2.2 interact* pf_interact( const char * s1, const char * s2, pu_contrib * p_c, pu_contrib * p_c2, int max_w, char * cstruc, int incr3, int incr5)
```

Calculates the probability of a local interaction between two sequences.

The function considers the probability that the region of interaction is unpaired within 's1' and 's2'. The longer sequence has to be given as 's1'. The shorter sequence has to be given as 's2'. Function pf\_unstru() has to be called for 's1' and 's2', where the probabilities of being unpaired have to be given in 'p\_c' and 'p\_c2', respectively. If you do not want to include the probabilities of being unpaired for 's2' set 'p\_c2' to NULL. If variable 'cstruc' is not NULL, constrained folding is done: The available constrains for intermolecular interaction are: '.' (no constrain), 'x' (the base has no intermolecular interaction) and '|' (the corresponding base has to be paired intermolecularily).

The parameter 'w' determines the maximal length of the interaction. The parameters 'incr5' and 'incr3' allows inclusion of unpaired residues left ('incr5') and right ('incr3') of the region of interaction in 's1'. If the 'incr' options are used, function pf\_unstru() has to be called with w=w+incr5+incr3 for the longer sequence 's1'.

It returns a structure of type interact which contains the probability of the best local interaction including residue i in Pi and the minimum free energy in Gi, where i is the position in sequence 's1'. The member Gikjl of structure interact is the best interaction between region [k,i] k < i in longer sequence 's1' and region [j,l] j < l in 's2'. Gikjl\_wo is Gikjl without the probability of beeing unpaired.

Use free\_interact() to free the returned structure, all other stuff is freed inside pf\_interact().

#### **Parameters**

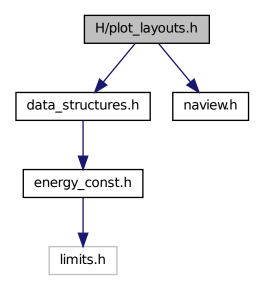
s1	
s2	
<i>p_c</i>	
p_c2	
max_w	
cstruc	
incr3	
incr5	

#### Returns

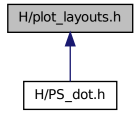
# 11.25 H/plot\_layouts.h File Reference

Secondary structure plot layout algorithms.

Include dependency graph for plot\_layouts.h:



This graph shows which files directly or indirectly include this file:



### **Defines**

#define VRNA\_PLOT\_TYPE\_SIMPLE 0
 Definition of Plot type simple

• #define VRNA PLOT TYPE NAVIEW 1

Definition of Plot type Naview

• #define VRNA\_PLOT\_TYPE\_CIRCULAR 2

Definition of Plot type Circular

#### **Functions**

- int simple\_xy\_coordinates (short \*pair\_table, float \*X, float \*Y)
   Calculate nucleotide coordinates for secondary structure plot the Simple way
- int simple\_circplot\_coordinates (short \*pair\_table, float \*x, float \*y)
   Calculate nucleotide coordinates for Circular Plot

#### **Variables**

• int rna\_plot\_type

Switch for changing the secondary structure layout algorithm.

### 11.25.1 Detailed Description

Secondary structure plot layout algorithms. c Ronny Lorenz The ViennaRNA Package

### 11.25.2 Define Documentation

11.25.2.1 #define VRNA\_PLOT\_TYPE\_SIMPLE 0

Definition of Plot type simple

This is the plot type definition for several RNA structure plotting functions telling them to use **Simple** plotting algorithm

#### See also

```
rna_plot_type, PS_rna_plot_a(), PS_rna_plot(), svg_rna_plot(), gmlRNA(), ssv_-
rna_plot(), xrna_plot()
```

#### 11.25.2.2 #define VRNA\_PLOT\_TYPE\_NAVIEW 1

Definition of Plot type Naview

This is the plot type definition for several RNA structure plotting functions telling them to use **Naview** plotting algorithm

#### See also

```
rna_plot_type, PS_rna_plot_a(), PS_rna_plot(), svg_rna_plot(), gmlRNA(), ssv_-
rna_plot(), xrna_plot()
```

#### 11.25.2.3 #define VRNA\_PLOT\_TYPE\_CIRCULAR 2

Definition of Plot type Circular

This is the plot type definition for several RNA structure plotting functions telling them to produce a **Circular plot** 

#### See also

```
rna_plot_type, PS_rna_plot_a(), PS_rna_plot(), svg_rna_plot(), gmlRNA(), ssv_-
rna_plot(), xrna_plot()
```

#### 11.25.3 Function Documentation

```
11.25.3.1 int simple_xy_coordinates ( short * pair_table, float * X, float * Y )
```

Calculate nucleotide coordinates for secondary structure plot the Simple way

#### See also

```
make_pair_table(), rna_plot_type, simple_circplot_coordinates(), naview_xy_coordinates(), PS_rna_plot_a(), PS_rna_plot, svg_rna_plot()
```

#### **Parameters**

pair_table	The pair table of the secondary structure
X	a pointer to an array with enough allocated space to hold the x coordinates
Y	a pointer to an array with enough allocated space to hold the y coordinates

### Returns

length of sequence on success, 0 otherwise

```
11.25.3.2 int simple_circplot_coordinates ( short * pair_table, float * x, float * y )
```

Calculate nucleotide coordinates for Circular Plot

This function calculates the coordinates of nucleotides mapped in equal distancies onto a unit circle.

### Note

In order to draw nice arcs using quadratic bezier curves that connect base pairs one may calculate a second tangential point  $P^t$  in addition to the actual  $\mathbb{R}^2$  coordinates. the simplest way to do so may be to compute a radius scaling factor rs in the interval [0,1] that weights the proportion of base pair span to the actual length of the sequence. This scaling factor can then be used to calculate the coordinates for  $P^t$ , i.e.  $P^t_x[i] = X[i] * rs$  and  $P^t_y[i] = Y[i] * rs$ .

#### See also

make pair table(), rna plot type, simple xy coordinates(), naview xy coordinates(),

PS\_rna\_plot\_a(), PS\_rna\_plot, svg\_rna\_plot()

#### **Parameters**

pair_table	The pair table of the secondary structure
X	a pointer to an array with enough allocated space to hold the x coordinates
У	a pointer to an array with enough allocated space to hold the y coordinates

#### Returns

length of sequence on success, 0 otherwise

#### 11.25.4 Variable Documentation

11.25.4.1 int rna\_plot\_type

Switch for changing the secondary structure layout algorithm.

Current possibility are 0 for a simple radial drawing or 1 for the modified radial drawing taken from the *naview* program of Bruccoleri & Heinrich (1988).

### Note

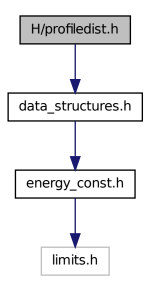
To provide thread safety please do not rely on this global variable in future implementations but pass a plot type flag directly to the function that decides which layout algorithm it may use!

### See also

VRNA\_PLOT\_TYPE\_SIMPLE, VRNA\_PLOT\_TYPE\_NAVIEW, VRNA\_PLOT\_TYPE\_-CIRCULAR

# 11.26 H/profiledist.h File Reference

Include dependency graph for profiledist.h:



#### **Functions**

- float profile\_edit\_distance (const float \*T1, const float \*T2)

  Align the 2 probability profiles T1, T2
- float \* Make\_bp\_profile\_bppm (FLT\_OR\_DBL \*bppm, int length)

  condense pair probability matrix into a vector containing probabilities for upstream paired, downstream paired and unpaired.
- void print\_bppm (const float \*T)
   print string representation of probability profile
- void free\_profile (float \*T)
   free space allocated in Make\_bp\_profile
- float \* Make\_bp\_profile (int length)

### 11.26.1 Detailed Description

### 11.26.2 Function Documentation

11.26.2.1 float profile\_edit\_distance ( const float \* T1, const float \* T2 )

Align the 2 probability profiles T1, T2

This is like a Needleman-Wunsch alignment, we should really use affine gap-costs ala Gotoh

11.26.2.2 float\* Make\_bp\_profile\_bppm ( FLT\_OR\_DBL \* bppm, int length )

condense pair probability matrix into a vector containing probabilities for upstream paired, downstream paired and unpaired.

This resulting probability profile is used as input for profile\_edit\_distance

#### **Parameters**

bppm	A pointer to the base pair probability matrix
length	The length of the sequence

#### Returns

The bp profile

11.26.2.3 void free\_profile (float \* T)

free space allocated in Make\_bp\_profile

Backward compatibility only. You can just use plain free()

11.26.2.4 float\* Make\_bp\_profile ( int length )

#### Note

This function is NOT threadsafe

### See also

Make\_bp\_profile\_bppm()

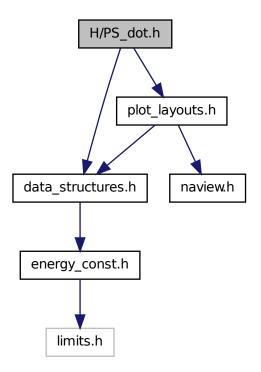
#### **Deprecated**

This function is deprecated and will be removed soon! See Make\_bp\_profile\_bppm() for a replacement

### 11.27 H/PS\_dot.h File Reference

Various functions for plotting RNA secondary structures, dot-plots and other visualizations.

Include dependency graph for PS\_dot.h:



### **Functions**

- int PS\_rna\_plot (char \*string, char \*structure, char \*file)

  Produce a secondary structure graph in PostScript and write it to 'filename'.
- int PS\_rna\_plot\_a (char \*string, char \*structure, char \*file, char \*pre, char \*post)

Produce a secondary structure graph in PostScript including additional annotation macros and write it to 'filename'.

- int gmlRNA (char \*string, char \*structure, char \*ssfile, char option)

  Produce a secondary structure graph in Graph Meta Language (gml) and write it to a file
- int ssv\_rna\_plot (char \*string, char \*structure, char \*ssfile)

  Produce a secondary structure graph in SStructView format.
- int svg\_rna\_plot (char \*string, char \*structure, char \*ssfile)
  - Produce a secondary structure plot in SVG format and write it to a file.
- int xrna plot (char \*string, char \*structure, char \*ssfile)

Produce a secondary structure plot for further editing in XRNA.

• int PS\_dot\_plot\_list (char \*seq, char \*filename, plist \*pl, plist \*mf, char \*comment)

Produce a postscript dot-plot from two pair lists.

 int aliPS\_color\_aln (const char \*structure, const char \*filename, const char \*seqs[], const char \*names[])

PS\_color\_aln for duplexes.

• int PS\_dot\_plot (char \*string, char \*file)

Wrapper to PS\_dot\_plot\_list.

### 11.27.1 Detailed Description

Various functions for plotting RNA secondary structures, dot-plots and other visualizations.

#### 11.27.2 Function Documentation

```
11.27.2.1 int PS_rna_plot ( char * string, char * structure, char * file )
```

Produce a secondary structure graph in PostScript and write it to 'filename'.

Note that this function has changed from previous versions and now expects the structure to be plotted in dot-bracket notation as an argument. It does not make use of the global base\_pair array anymore.

#### **Parameters**

string	The RNA sequence
structure	The secondary structure in dot-bracket notation
file	The filename of the postscript output

#### Returns

1 on success, 0 otherwise

```
11.27.2.2 int PS_rna_plot_a ( char * string, char * structure, char * file, char * pre, char * post )
```

Produce a secondary structure graph in PostScript including additional annotation macros and write it to 'filename'.

Same as PS\_rna\_plot() but adds extra PostScript macros for various annotations (see generated PS code). The 'pre' and 'post' variables contain PostScript code that is verbatim copied in the resulting PS file just before and after the structure plot. If both arguments ('pre' and 'post') are NULL, no additional macros will be printed into the PostScript.

### **Parameters**

string	The RNA sequence
structure	The secondary structure in dot-bracket notation
file	The filename of the postscript output
pre	PostScript code to appear before the secondary structure plot
post	PostScript code to appear after the secondary structure plot

#### **Returns**

1 on success, 0 otherwise

11.27.2.3 int gmlRNA ( char \* string, char \* structure, char \* ssfile, char option )

Produce a secondary structure graph in Graph Meta Language (gml) and write it to a file.

If 'option' is an uppercase letter the RNA sequence is used to label nodes, if 'option' equals X' or X' the resulting file will coordinates for an initial layout of the graph.

#### **Parameters**

string	The RNA sequence
structure	The secondary structure in dot-bracket notation
ssfile	The filename of the gml output
option	The option flag

#### **Returns**

1 on success, 0 otherwise

11.27.2.4 int ssv\_rna\_plot ( char \* string, char \* structure, char \* ssfile )

Produce a secondary structure graph in SStructView format.

Write coord file for SStructView

### **Parameters**

string	The RNA sequence
structure	The secondary structure in dot-bracket notation
ssfile	The filename of the ssv output

#### Returns

1 on success, 0 otherwise

11.27.2.5 int svg\_rna\_plot ( char \* string, char \* structure, char \* ssfile )

Produce a secondary structure plot in SVG format and write it to a file.

#### **Parameters**

string	The RNA sequence
structure	The secondary structure in dot-bracket notation
ssfile	The filename of the svg output

#### Returns

1 on success, 0 otherwise

11.27.2.6 int xrna\_plot ( char \* string, char \* structure, char \* ssfile )

Produce a secondary structure plot for further editing in XRNA.

#### **Parameters**

string	The RNA sequence
structure	The secondary structure in dot-bracket notation
ssfile	The filename of the xrna output

#### Returns

1 on success, 0 otherwise

11.27.2.7 int PS\_dot\_plot\_list ( char \* seq, char \* filename, plist \* pl, plist \* mf, char \* comment )

Produce a postscript dot-plot from two pair lists.

This function reads two plist structures (e.g. base pair probabilities and a secondary structure) as produced by <a href="mailto:assign\_plist\_from\_pr">assign\_plist\_from\_pr</a>() and <a href="mailto:assign\_plist\_from\_db">assign\_plist\_from\_db</a>() and <a href="prop-produces">produces</a> a postscript "dot plot" that is written to 'filename'.

Using base pair probabilities in the first and mfe structure in the second plist, the resulting "dot plot" represents each base pairing probability by a square of corresponding area in a upper triangle matrix. The lower part of the matrix contains the minimum free energy structure.

#### See also

assign\_plist\_from\_pr(), assign\_plist\_from\_db()

#### **Parameters**

seq	The RNA sequence
filename	A filename for the postscript output
pl	The base pair probability pairlist
mf	The mfe secondary structure pairlist
comment	A comment

#### **Returns**

1 if postscript was successfully written, 0 otherwise

```
11.27.2.8 int PS_dot_plot ( char * string, char * file )
```

Wrapper to PS dot plot list.

Produce postscript dot-plot

Reads base pair probabilities produced by pf\_fold() from the global array pr and the pair list base\_pair produced by fold() and produces a postscript "dot plot" that is written to 'filename'. The "dot plot" represents each base pairing probability by a square of corresponding area in a upper triangle matrix. The lower part of the matrix contains the minimum free energy

#### Note

DO NOT USE THIS FUNCTION ANYMORE SINCE IT IS NOT THREADSAFE

#### **Deprecated**

This function is deprecated and will be removed soon! Use PS\_dot\_plot\_list() instead!

## 11.28 H/read\_epars.h File Reference

Functions to read and write energy parameter sets from/to files.

### **Functions**

void read\_parameter\_file (const char fname[])

Read energy parameters from a file.

void write\_parameter\_file (const char fname[])

Write energy parameters to a file.

## 11.28.1 Detailed Description

Functions to read and write energy parameter sets from/to files.

### 11.28.2 Function Documentation

11.28.2.1 void read\_parameter\_file ( const char fname[] )

Read energy parameters from a file.

#### **Parameters**

mane i me pam to me me comamino me energy parameters	fname	The path to the file containing the energy parameters	
--	-------	---	--

11.28.2.2 void write\_parameter\_file ( const char fname[] )

Write energy parameters to a file.

#### **Parameters**

fname	A filename	(path) f	or the	file	where	the	current	energy	parameters	will	be
	written to										

# 11.29 H/RNAstruct.h File Reference

Parsing and Coarse Graining of Structures.

#### **Functions**

char \* b2HIT (const char \*structure)

Converts the full structure from bracket notation to the HIT notation including root.

char \* b2C (const char \*structure)

Converts the full structure from bracket notation to the a coarse grained notation using the 'H' 'B' 'I' 'M' and 'R' identifiers.

char \* b2Shapiro (const char \*structure)

Converts the full structure from bracket notation to the weighted coarse grained notation using the 'H' 'B' 'I' 'M' 'S' 'E' and 'R' identifiers.

char \* add\_root (const char \*structure)

Adds a root to an un-rooted tree in any except bracket notation.

char \* expand\_Shapiro (const char \*coarse)

Inserts missing 'S' identifiers in unweighted coarse grained structures as obtained from b2C().

char \* expand\_Full (const char \*structure)

Convert the full structure from bracket notation to the expanded notation including root.

char \* unexpand\_Full (const char \*ffull)

Restores the bracket notation from an expanded full or HIT tree, that is any tree using only identifiers 'U' 'P' and 'R'.

• char \* unweight (const char \*wcoarse)

Strip weights from any weighted tree.

void unexpand\_aligned\_F (char \*align[2])

Converts two aligned structures in expanded notation.

void parse\_structure (const char \*structure)

Collects a statistic of structure elements of the full structure in bracket notation.

#### **Variables**

```
    int loop_size [STRUC]
        contains a list of all loop sizes.
```

• int helix size [STRUC]

contains a list of all stack sizes.

• int loop\_degree [STRUC]

contains the corresponding list of loop degrees.

· int loops

contains the number of loops ( and therefore of stacks ).

· int unpaired

contains the number of unpaired bases.

· int pairs

contains the number of base pairs in the last parsed structure.

# 11.29.1 Detailed Description

Parsing and Coarse Graining of Structures. Example:

#### 11.29.2 Function Documentation

```
11.29.2.1 char* b2HIT ( const char * structure )
```

Converts the full structure from bracket notation to the HIT notation including root.

#### **Parameters**

```
structure
```

#### **Returns**

11.29.2.2 char\* b2C ( const char \* structure )

Converts the full structure from bracket notation to the a coarse grained notation using the 'H' 'B' 'I' 'M' and 'R' identifiers.

#### **Parameters**

structure

#### Returns

11.29.2.3 char\* b2Shapiro ( const char \* structure )

Converts the full structure from bracket notation to the *weighted* coarse grained notation using the 'H' 'B' 'I' 'M' 'S' 'E' and 'R' identifiers.

## **Parameters**

structure

#### Returns

11.29.2.4 char\* add\_root ( const char \* structure )

Adds a root to an un-rooted tree in any except bracket notation.

# **Parameters**

structure

#### Returns

11.29.2.5 char\* expand\_Shapiro ( const char \* coarse )

Inserts missing 'S' identifiers in unweighted coarse grained structures as obtained from b2C().

#### **Parameters**

coarse

#### **Returns**

```
11.29.2.6 char* expand_Full ( const char * structure )
```

Convert the full structure from bracket notation to the expanded notation including root.

#### **Parameters**

```
structure
```

## Returns

```
11.29.2.7 char* unexpand_Full ( const char * ffull )
```

Restores the bracket notation from an expanded full or HIT tree, that is any tree using only identifiers 'U' 'P' and 'R'.

#### **Parameters**

```
ffull
```

## Returns

11.29.2.8 char\* unweight ( const char \* wcoarse )

Strip weights from any weighted tree.

#### **Parameters**

```
wcoarse
```

## Returns

11.29.2.9 void unexpand\_aligned\_F ( char \* align[2] )

Converts two aligned structures in expanded notation.

Takes two aligned structures as produced by tree\_edit\_distance() function back to bracket notation with '\_' as the gap character. The result overwrites the input.

#### **Parameters**

align

11.29.2.10 void parse\_structure ( const char \* structure )

Collects a statistic of structure elements of the full structure in bracket notation.

The function writes to the following global variables: loop\_size, loop\_degree, helix\_size, loops, pairs, unpaired

## **Parameters**

structure

## Returns

# 11.29.3 Variable Documentation

11.29.3.1 int loop\_size[STRUC]

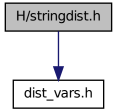
contains a list of all loop sizes.

loop\_size[0] contains the number of external bases.

# 11.30 H/stringdist.h File Reference

Functions for String Alignment.

Include dependency graph for stringdist.h:



## **Functions**

• swString \* Make\_swString (char \*string)

Convert a structure into a format suitable for string\_edit\_distance().

float string\_edit\_distance (swString \*T1, swString \*T2)

Calculate the string edit distance of T1 and T2.

# 11.30.1 Detailed Description

Functions for String Alignment.

## 11.30.2 Function Documentation

11.30.2.1 swString \* Make\_swString ( char \* string )

Convert a structure into a format suitable for string\_edit\_distance().

## **Parameters**

string	

#### Returns

11.30.2.2 float string\_edit\_distance ( swString \* 71, swString \* 72 )

Calculate the string edit distance of T1 and T2.

#### **Parameters**

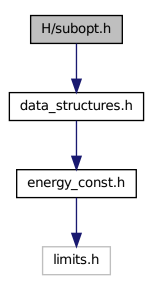
T1	
T2	

# Returns

# 11.31 H/subopt.h File Reference

RNAsubopt and density of states declarations.

Include dependency graph for subopt.h:



# **Functions**

- SOLUTION \* subopt (char \*seq, char \*sequence, int delta, FILE \*fp)

  Returns list of subopt structures or writes to fp.
- SOLUTION \* subopt\_circ (char \*seq, char \*sequence, int delta, FILE \*fp)

  Returns list of circular subopt structures or writes to fp.

## **Variables**

- int subopt\_sorted
  - Sort output by energy.
- double print\_energy

printing threshold for use with logML

# 11.31.1 Detailed Description

RNAsubopt and density of states declarations.

#### 11.31.2 Function Documentation

```
11.31.2.1 SOLUTION* subopt ( char * seq, char * sequence, int delta, FILE * fp )
```

Returns list of subopt structures or writes to fp.

This function produces **all** suboptimal secondary structures within 'delta' \* 0.01 kcal/mol of the optimum. The results are either directly written to a 'fp' (if 'fp' is not NULL), or (fp==NULL) returned in a SOLUTION \* list terminated by an entry were the 'structure' pointer is NULL.

#### **Parameters**

seq	
sequence	
delta	
fp	

#### Returns

```
11.31.2.2 SOLUTION* subopt_circ ( char * seq, char * sequence, int delta, FILE * fp )
```

Returns list of circular subopt structures or writes to fp.

This function is similar to <a href="subopt">subopt</a>() but calculates secondary structures assuming the RNA sequence to be circular instead of linear

#### **Parameters**

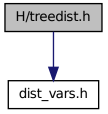
seq	
sequence	
delta	
fp	

## Returns

# 11.32 H/treedist.h File Reference

Functions for Tree Edit Distances.

Include dependency graph for treedist.h:



## **Functions**

• Tree \* make\_tree (char \*struc)

Constructs a Tree ( essentially the postorder list ) of the structure 'struc', for use in tree\_edit\_distance().

• float tree\_edit\_distance (Tree \*T1, Tree \*T2)

Calculates the edit distance of the two trees.

void print\_tree (Tree \*t)

Print a tree (mainly for debugging)

void free\_tree (Tree \*t)

Free the memory allocated for Tree t.

# 11.32.1 Detailed Description

Functions for Tree Edit Distances.

## 11.32.2 Function Documentation

11.32.2.1 Tree\* make\_tree ( char \* struc )

Constructs a Tree ( essentially the postorder list ) of the structure 'struc', for use in  $tree\_edit\_distance()$ .

#### **Parameters**

struc may be any rooted structure representation.

#### Returns

```
11.32.2.2 float tree_edit_distance ( Tree * 71, Tree * 72 )
```

Calculates the edit distance of the two trees.

#### **Parameters**

T1	
T2	

#### Returns

```
11.32.2.3 void free_tree ( Tree * t )
```

Free the memory allocated for Tree t.

#### **Parameters**

```
t
```

# 11.33 H/utils.h File Reference

Various utility- and helper-functions used throughout the Vienna RNA package.

## **Defines**

- #define VRNA\_INPUT\_ERROR 1U
  - Output flag of <a href="mailto:get\_input\_line">get\_input\_line</a>(): "An ERROR has occured, maybe EOF".
- #define VRNA\_INPUT\_QUIT 2U
  - Output flag of get\_input\_line(): "the user requested quitting the program".
- #define VRNA\_INPUT\_MISC 4U
  - Output flag of get\_input\_line(): "something was read".
- #define VRNA\_INPUT\_FASTA\_HEADER 8U

Input/Output flag of get\_input\_line():

if used as input option this tells get\_input\_line() that the data to be read should comply
with the FASTA format.

• #define VRNA INPUT SEQUENCE 16U

Input flag for get\_input\_line():

Tell get\_input\_line() that we assume to read a nucleotide sequence.

• #define VRNA INPUT CONSTRAINT 32U

Input flag for get\_input\_line():

Tell get\_input\_line() that we assume to read a structure constraint.

#define VRNA\_INPUT\_NO\_TRUNCATION 256U

Input switch for get\_input\_line(): "do not trunkate the line by eliminating white spaces at end of line"

• #define VRNA\_INPUT\_NO\_REST 512U

Input switch for read\_record(): "do fill rest array".

• #define VRNA INPUT NO SPAN 1024U

Input switch for read\_record(): "never allow data to span more than one line".

• #define VRNA\_INPUT\_NOSKIP\_BLANK\_LINES 2048U

Input switch for read\_record(): "do not skip empty lines".

• #define VRNA INPUT BLANK LINE 4096U

Output flag for read\_record(): "read an empty line".

• #define VRNA\_INPUT\_NOSKIP\_COMMENTS 128U

Input switch for get\_input\_line(): "do not skip comment lines".

• #define VRNA INPUT COMMENT 8192U

Output flag for read\_record(): "read a comment".

• #define VRNA\_CONSTRAINT\_PIPE 1U

pipe sign '\' switch for structure constraints (paired with another base)

#define VRNA CONSTRAINT DOT 2U

dot '

#define VRNA\_CONSTRAINT\_X 4U

'x' switch for structure constraint (base must not pair)

#define VRNA\_CONSTRAINT\_ANG\_BRACK 8U

angle brackets '<', '>' switch for structure constraint (paired downstream/upstream)

#define VRNA\_CONSTRAINT\_RND\_BRACK 16U

round brackets '(',')' switch for structure constraint (base i pairs base j)

#define VRNA\_CONSTRAINT\_MULTILINE 32U

constraint may span over several lines

#define VRNA\_CONSTRAINT\_NO\_HEADER 64U

do not print the header information line

#define VRNA\_CONSTRAINT\_ALL 128U

placeholder for all constraining characters

• #define MIN2(A, B) ((A) < (B) ? (A) : (B))

Get the minimum of two comparable values.

• #define MAX2(A, B) ((A) > (B) ? (A) : (B))

Get the maximum of two comparable values.

• #define MIN3(A, B, C) (MIN2( (MIN2((A),(B))) ,(C)))

Get the minimum of three comparable values.

#define MAX3(A, B, C) (MAX2( (MAX2((A),(B))) ,(C)))

Get the maximum of three comparable values.

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#### **Functions**

void \* space (unsigned size)

Allocate space safely.

void \* xrealloc (void \*p, unsigned size)

Reallocate space safely.

void nrerror (const char message[])

Die with an error message.

void warn\_user (const char message[])

Print a warning message.

void init rand (void)

Make random number seeds.

· double urn (void)

get a random number from [0..1]

• int int urn (int from, int to)

Generates a pseudo random integer in a specified range.

• char \* time\_stamp (void)

Get a timestamp.

char \* random\_string (int I, const char symbols[])

Create a random string using characters from a specified symbol set.

• int hamming (const char \*s1, const char \*s2)

Calculate hamming distance between two sequences.

• int hamming\_bound (const char \*s1, const char \*s2, int n)

Calculate hamming distance between two sequences up to a specified length.

char \* get line (FILE \*fp)

Read a line of arbitrary length from a stream.

unsigned int get\_input\_line (char \*\*string, unsigned int options)

Retrieve a line from 'stdin' savely while skipping comment characters and other features This function returns the type of input it has read if recognized.

unsigned int read\_record (char \*\*header, char \*\*sequence, char \*\*\*rest, unsigned int options)

Get a data record from stdin.

• char \* pack structure (const char \*struc)

Pack secondary secondary structure, 5:1 compression using base 3 encoding.

char \* unpack\_structure (const char \*packed)

Unpack secondary structure previously packed with pack\_structure()

• short \* make\_pair\_table (const char \*structure)

Create a pair table of a secondary structure.

• short \* copy\_pair\_table (const short \*pt)

Get an exact copy of a pair table.

short \* alimake\_pair\_table (const char \*structure)

Pair table for snoop align.

short \* make pair table snoop (const char \*structure)

returns a newly allocated table, such that: table[i]=j if (i.j) pair or 0 if i is unpaired, table[0] contains the length of the structure.

int bp\_distance (const char \*str1, const char \*str2)

Compute the "base pair" distance between two secondary structures s1 and s2.

void print\_tty\_input\_seq (void)

Print a line to stdout that asks for an input sequence.

void print\_tty\_input\_seq\_str (const char \*s)

Print a line with a user defined string and a ruler to stdout.

void print\_tty\_constraint\_full (void)

Print structure constraint characters to stdout (full constraint support)

void print\_tty\_constraint (unsigned int option)

Print structure constraint characters to stdout.

• void str\_DNA2RNA (char \*sequence)

Convert a DNA input sequence to RNA alphabet.

• void str\_uppercase (char \*sequence)

Convert an input sequence to uppercase.

int \* get\_iindx (unsigned int length)

Get an index mapper array (iindx) for accessing the energy matrices, e.g.

• int \* get indx (unsigned int length)

Get an index mapper array (indx) for accessing the energy matrices, e.g.

 void constrain\_ptypes (const char \*constraint, unsigned int length, char \*ptype, int \*BP, int min\_loop\_size, unsigned int idx\_type)

Insert constraining pair types according to constraint structure string.

## **Variables**

• unsigned short xsubi [3]

Current 48 bit random number.

#### 11.33.1 Detailed Description

Various utility- and helper-functions used throughout the Vienna RNA package.

# 11.33.2 Define Documentation

#### 11.33.2.1 #define VRNA\_INPUT\_FASTA\_HEADER 8U

Input/Output flag of <a href="mailto:get\_input\_line">get\_input\_line</a>():

if used as input option this tells get\_input\_line() that the data to be read should comply with the FASTA format.

the function will return this flag if a fasta header was read

11.33.2.2 #define VRNA\_INPUT\_SEQUENCE 16U

Input flag for get\_input\_line():

Tell get\_input\_line() that we assume to read a nucleotide sequence.

11.33.2.3 #define VRNA\_INPUT\_CONSTRAINT 32U

Input flag for get\_input\_line():

Tell get\_input\_line() that we assume to read a structure constraint.

11.33.2.4 #define VRNA\_CONSTRAINT\_DOT 2U

dot '.

'switch for structure constraints (no constraint at all)

11.33.3 Function Documentation

11.33.3.1 void\* space ( unsigned size )

Allocate space safely.

## **Parameters**

oled The size of the memory to be allocated in bytes		size	The size of the memory to be allocated in bytes
--	--	------	---

#### **Returns**

A pointer to the allocated memory

11.33.3.2 void\* xrealloc (void \* p, unsigned size )

Reallocate space safely.

#### **Parameters**

р	A pointer to the memory region to be reallocated
size	The size of the memory to be allocated in bytes

#### Returns

A pointer to the newly allocated memory

11.33.3.3 void nrerror ( const char message[])

Die with an error message.

#### See also

warn\_user()

#### **Parameters**

message The error message to be printed before exiting with 'FAILURE'

11.33.3.4 void warn\_user ( const char message[] )

Print a warning message.

Print a warning message to stderr

## **Parameters**

message	The warning message	
---------	---------------------	--

11.33.3.5 double urn ( void )

get a random number from [0..1]

#### Note

Usually implemented by calling erand48().

## Returns

A random number in range [0..1]

11.33.3.6 int int\_urn ( int from, int to )

Generates a pseudo random integer in a specified range.

## **Parameters**

from	The first number in range
to	The last number in range

#### Returns

A pseudo random number in range [from, to]

11.33.3.7 char\* time\_stamp ( void )

Get a timestamp.

Returns a string containing the current date in the format

Generated on Sat Jan 14 2012 15:51:55 for RNAlib-2.0.0 by Doxygen

Fri Mar 19 21:10:57 1993

#### Returns

A string containing the timestamp

11.33.3.8 char\* random\_string ( int I, const char symbols[] )

Create a random string using characters from a specified symbol set.

#### **Parameters**

1	The length of the sequence
symbols	The symbol set

#### **Returns**

A random string of length 'I' containing characters from the symbolset

11.33.3.9 int hamming (const char \*s1, const char \*s2)

Calculate hamming distance between two sequences.

Calculate the number of positions in which

#### **Parameters**

s1	The first sequence
s2	The second sequence

## Returns

The hamming distance between s1 and s2

11.33.3.10 int hamming\_bound (const char \*s1, const char \*s2, int n)

Calculate hamming distance between two sequences up to a specified length.

This function is similar to <a href="hamming">hamming</a>() but instead of comparing both sequences up to their actual length only the first 'n' characters are taken into account

#### **Parameters**

s1	The first sequence
s2	The second sequence

#### **Returns**

The hamming distance between s1 and s2

```
11.33.3.11 char* get_line ( FILE * fp )
```

Read a line of arbitrary length from a stream.

Returns a pointer to the resulting string. The necessary memory is allocated and should be released using *free()* when the string is no longer needed.

#### **Parameters**

fp A file pointer to the stream where the function should read from

#### Returns

A pointer to the resulting string

```
11.33.3.12 unsigned int get_input_line ( char ** string, unsigned int options )
```

Retrieve a line from 'stdin' savely while skipping comment characters and other features. This function returns the type of input it has read if recognized.

An option argument allows to switch between different reading modes.

Currently available options are:

 $\verb| #VRNA_INPUT_NOPRINT_COMMENTS|, VRNA_INPUT_NOSKIP_COMMENTS|, \verb| #VRNA_INPUT_NOELIM_WS_SUFFIX| \\$ 

pass a collection of options as one value like this:

```
get_input_line(string, option_1 | option_2 | option_n)
```

If the function recognizes the type of input, it will report it in the return value. It also reports if a user defined 'quit' command (@-sign on 'stdin') was given. Possible return values are:

VRNA\_INPUT\_FASTA\_HEADER, VRNA\_INPUT\_ERROR, VRNA\_INPUT\_MISC, VRNA\_INPUT\_QUIT

#### **Parameters**

string	A pointer to the character array that contains the line read
options	A collection of options for switching the functions behavior

#### Returns

A flag with information about what has been read

```
11.33.3.13 unsigned int read_record ( char ** header, char ** sequence, char *** rest, unsigned int options )
```

Get a data record from stdin.

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This function may be used to obtain complete datasets from stdin. A dataset is always defined to contain at least a sequence. If data on stdin starts with a fasta header, i.e. a line like

```
>some header info
```

then read\_record() will assume that the sequence that follows the header may span over several lines. To disable this behavior and to assign a single line to the argument 'sequence' one can pass VRNA\_INPUT\_NO\_SPAN in the 'options' argument. If no fasta header is read in the beginning of a data block, a sequence must not span over multiple lines!

Unless the options VRNA\_INPUT\_NOSKIP\_COMMENTS or VRNA\_INPUT\_NOSKIP\_BLANK\_LINES are passed, a sequence may be interrupted by lines starting with a comment character or empty lines.

A sequence is regarded as completely read if it was either assumed to not span over multiple lines, a secondary structure or structure constraint follows the sequence on the next line or a new header marks the beginning of a new sequence...

All lines following the sequence (this includes comments) and not initiating a new dataset are available through the line-array 'rest'. Here one can usually find the structure constraint or other information belonging to the current dataset. Filling of 'rest' may be prevented by passing VRNA INPUT NO REST to the options argument.

#### Note

This function will exit any program with an error message if no sequence could be read!

The main purpose of this function is to be able to easily parse blocks of data from stdin in the header of a loop where all calculations for the appropriate data is done inside the loop. The loop may be then left on certain return values, e.g.:

```
char *id, *seq, **rest;
int i;
while(!(read_record(&id, &seq, &rest, 0) & (VRNA_INPUT_ERROR | VRNA_INPUT_QUIT))){
   if(id) printf("%s\n", id);
   printf("%s\n", seq);
   if(rest)
     for(i=0;rest[i];i++)
        printf("%s\n", rest[i]);
}
```

In the example above, the while loop will be terminated when read\_record() returns either an error or a user initiated guit request.

As long as data is read from stdin, the id is printed if it is available for the current block of data. The sequence will be printed in any case and if some more lines belong to the current block of data each line will be printed as well.

## Note

Do not forget to free the memory occupied by header, sequence and rest!

#### **Parameters**

header	A pointer which will be set such that it points to the header of the record
sequence	A pointer which will be set such that it points to the sequence of the record
rest	A pointer which will be set such that it points to an array of lines which also
	belong to the record
options	Some options which may be passed to alter the behavior of the function, use
	0 for no options

#### Returns

A flag with information about what the function actually did read

11.33.3.14 char\* pack\_structure ( const char \* struc )

Pack secondary secondary structure, 5:1 compression using base 3 encoding.

Returns a binary string encoding of the secondary structure using a 5:1 compression scheme. The string is NULL terminated and can therefore be used with standard string functions such as strcmp(). Useful for programs that need to keep many structures in memory.

#### **Parameters**

struc	The secondary structure in dot-bracket notation

#### Returns

The binary encoded structure

11.33.3.15 char\* unpack\_structure ( const char \* packed )

Unpack secondary structure previously packed with pack\_structure()

Translate a compressed binary string produced by pack\_structure() back into the familiar dot-bracket notation.

#### **Parameters**

packed The binary encoded packed secondary structure	
--	--

# Returns

The unpacked secondary structure in dot-bracket notation

11.33.3.16 short\* make\_pair\_table ( const char \* structure )

Create a pair table of a secondary structure.

Returns a newly allocated table, such that table[i]=j if (i.j) pair or 0 if i is unpaired, table[0] contains the length of the structure.

#### **Parameters**

structure	The secondary structure in dot-bracket notation

## Returns

A pointer to the created pair\_table

11.33.3.17 short\* copy\_pair\_table ( const short \* pt )

Get an exact copy of a pair table.

## **Parameters**

pt	The pair table to be copied

#### Returns

A pointer to the copy of 'pt'

11.33.3.18 short\* alimake\_pair\_table ( const char \* structure )

Pair table for snoop align.

11.33.3.19 short\* make\_pair\_table\_snoop ( const char \* structure )

returns a newly allocated table, such that: table[i]=j if (i.j) pair or 0 if i is unpaired, table[0] contains the length of the structure.

The special pseudoknotted H/ACA-mRNA structure is taken into account.

11.33.3.20 int bp\_distance ( const char \* str1, const char \* str2 )

Compute the "base pair" distance between two secondary structures s1 and s2.

The sequences should have the same length. dist = number of base pairs in one structure but not in the other same as edit distance with open-pair close-pair as move-set

#### **Parameters**

str1	First structure in dot-bracket notation
str2	Second structure in dot-bracket notation

## Returns

The base pair distance between str1 and str2

```
11.33.3.21 void print_tty_input_seq ( void )
```

Print a line to stdout that asks for an input sequence.

There will also be a ruler (scale line) printed that helps orientation of the sequence positions

```
11.33.3.22 void print_tty_input_seq_str ( const char * s )
```

Print a line with a user defined string and a ruler to stdout.

(usually this is used to ask for user input) There will also be a ruler (scale line) printed that helps orientation of the sequence positions

#### **Parameters**

s A user defined string that will be printed to stdout

```
11.33.3.23 void print_tty_constraint_full ( void )
```

Print structure constraint characters to stdout (full constraint support)

```
11.33.3.24 void print_tty_constraint ( unsigned int option )
```

Print structure constraint characters to stdout.

(constraint support is specified by option parameter)

Currently available options are:

```
VRNA CONSTRAINT PIPE (paired with another base)
```

VRNA CONSTRAINT DOT (no constraint at all)

VRNA\_CONSTRAINT\_X (base must not pair)

VRNA\_CONSTRAINT\_ANG\_BRACK (paired downstream/upstream)

VRNA CONSTRAINT RND BRACK (base i pairs base j)

pass a collection of options as one value like this:

```
print_tty_constraint(option_1 | option_2 | option_n)
```

## **Parameters**

option Option switch that tells which constraint help will be printed

```
11.33.3.25 void str_DNA2RNA ( char * sequence )
```

Convert a DNA input sequence to RNA alphabet.

This function substitudes T and t with U and u, respectively

#### **Parameters**

sequence The sequence to be converted	
---------------------------------------	--

11.33.3.26 void str\_uppercase ( char \* sequence )

Convert an input sequence to uppercase.

#### **Parameters**

sequence	The sequence to be converted

11.33.3.27 int\* get\_iindx ( unsigned int length )

Get an index mapper array (iindx) for accessing the energy matrices, e.g.

in partition function related functions.

Access of a position "(i,j)" is then accomplished by using

```
(i,j) \sim iindx[i]-j
```

This function is necessary as most of the two-dimensional energy matrices are actually one-dimensional arrays throughout the ViennaRNAPackage

Consult the implemented code to find out about the mapping formula;)

#### See also

```
get_indx()
```

#### **Parameters**

<i>length</i>   Th	e length of t	he RNA sequ	uence
--------------------	---------------	-------------	-------

#### **Returns**

The mapper array

11.33.3.28 int\* get\_indx ( unsigned int length )

Get an index mapper array (indx) for accessing the energy matrices, e.g.

in MFE related functions.

Access of a position "(i,j)" is then accomplished by using

```
(i,j) \sim indx[j]+i
```

This function is necessary as most of the two-dimensional energy matrices are actually one-dimensional arrays throughout the ViennaRNAPackage

Consult the implemented code to find out about the mapping formula;)

#### See also

get\_iindx()

## **Parameters**

length	The length of the RNA sequence

#### Returns

The mapper array

11.33.3.29 void constrain\_ptypes ( const char \* constraint, unsigned int length, char \* ptype, int \* BP, int min\_loop\_size, unsigned int idx\_type )

Insert constraining pair types according to constraint structure string.

#### See also

get\_indx(), get\_iindx()

#### **Parameters**

constraint	The structure constraint string
length	The actual length of the sequence (constraint may be shorter)
ptype	A pointer to the basepair type array
min_loop	The minimal loop size (usually TURN )
size	
idx_type	Define the access type for base pair type array (0 = indx, 1 = iindx)

## 11.33.4 Variable Documentation

11.33.4.1 unsigned short xsubi[3]

Current 48 bit random number.

This variable is used by urn(). These should be set to some random number seeds before the first call to urn().

# See also

urn()

# 11.34 lib/1.8.4\_epars.h File Reference

Free energy parameters for parameter file conversion.

#### 11.34.1 Detailed Description

Free energy parameters for parameter file conversion. This file contains the free energy parameters used in ViennaRNAPackage 1.8.4. They are summarized in:

D.H.Mathews, J. Sabina, M. ZUker, D.H. Turner "Expanded sequence dependence of thermodynamic parameters improves prediction of RNA secondary structure" JMB, 288, pp 911-940, 1999

Enthalpies taken from:

A. Walter, D Turner, J Kim, M Lyttle, P M"uller, D Mathews, M Zuker "Coaxial stckaing of helices enhances binding of oligoribonucleotides.." PNAS, 91, pp 9218-9222, 1994

D.H. Turner, N. Sugimoto, and S.M. Freier. "RNA Structure Prediction", Ann. Rev. Biophys. Biophys. Chem. 17, 167-192, 1988.

John A.Jaeger, Douglas H.Turner, and Michael Zuker. "Improved predictions of secondary structures for RNA", PNAS, 86, 7706-7710, October 1989.

L. He, R. Kierzek, J. SantaLucia, A.E. Walter, D.H. Turner "Nearest-Neughbor Parameters for GU Mismatches...." Biochemistry 1991, 30 11124-11132

A.E. Peritz, R. Kierzek, N, Sugimoto, D.H. Turner "Thermodynamic Study of Internal Loops in Oligoribonucleotides..." Biochemistry 1991, 30, 6428--6435

# 11.35 lib/1.8.4\_intloops.h File Reference

Free energy parameters for interior loop contributions needed by the parameter file conversion functions.

#### 11.35.1 Detailed Description

Free energy parameters for interior loop contributions needed by the parameter file conversion functions.

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