RSATWSPortType

Service Web services for the Regulatory Sequence Analysis Tools (RSAT). Tools developed by

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interface developed by Olivier Sand

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Server Address http://localhost/rsat/web_services/RSATWS.cgi

Method retrieve_seq

Description Returns upstream, downstream or coding

DNA sequences for list of query genes.

Parameters Input Parameters

output Return type. Accepted values:

'server' (result is stored on a file on the server), 'client' (result is directly transferred to the client), or 'both'. Default is

'both'

Type = string

organism Organism. Words need to be

(required) underscore separated (example:

Escherichia_coli_K12).

Type = string

query A list of query genes

(required) Type = string

noorf Prevent overlap with upstream

open reading frames (ORF).

Type = string

from Inferior limit of the region to

retrieve. Default is organism dependant (example:

Saccharomyces cerevisiae =

-800). Type = int

to Superior limit of the region to

retrieve. Default is '-1'.

Type = int

feattype Type of genome features to

load. Supported: CDS, mRNA,

tRNA, rRNA. Type = string

type Sequence type. Supported:

upstream, downstream, ORF (unspliced open reading frame).

format Sequence format. Supported:

IG (Intelligenetics), WC (wconsensus), raw, FastA

Type = string

all Return sequences for all the

genes of the organism.

Type = string

lw Line width (0 for whole

sequence on one line).

Type = int

label Field(s) to be used in the

sequence label. Multiple fields can be specified, separated by commas. Supported: id, name,

organism_name,

sequence_type, current_from, current_to, ctg, orf_strand, reg_left, reg_right. Default:

name.
Type = string

label_sep Separator between the label

fields. Default: | (pipe

character). Type = string

nocom No comments. Only the

identifier and the sequence are returned. By default, the comment indicates the ORF and upstream sequence

coordinates. Type = string

repeat Use the repeat masked version

of the genome. Attention: repeated regions are annotated

for some genomes only.

Type = string

imp_pos Admit imprecise positions.

Type = string

Output Parameters

server Location of the result file on the

server. This can be used as input for a further request.

Type = string

command The stand alone command

executed on the server.

Type = string

client The result sequence(s).

Description

Mask repeated fragments of an input sequence.

Parameters

Input Parameters

output Return type. Accepted

values: 'server' (result is stored on a file on the server), 'client' (result is directly transferred to the client), or 'both'. Default is

'both'.

Type = string

sequence Sequence to purge. You

need to supply either this parameter or the next one

(tmp_infile). Type = string

tmp_infile Name of the file with input

sequence on the server. You need to supply either this parameter or the previous

one (sequence). Type = string

format Sequence format.

Supported: IG (Intelligenetics), WC (wconsensus), raw, FastA.

Type = string

match_length Minimal match length.

Default is 40. Type = int

mismatch Number of mismatches

allowed. Default is 3.

Type = int

str Discard duplications on the

direct strand only (1str) or on the reverse complement as well (2str). Default is 2str.

Type = string

delete Delete repeats instead of

masking them. Recognized

value is 'del'. Type = string

mask_short Mask (replace by N

characters) sequences shorter than the specified

length. Type = int

Output Parameters

server

Location of the result file on the server. This can be used as input for a further request.

command The stand alone command

executed on the server.

Type = string

client The result sequence(s).

Type = string

Method oligo_analysis

Description Analysis of the statistical significance of all

the oligomers of a given size in a sequence. Commonly used to detect over-represented oligonucleotides in a set of promoter

sequences.

Parameters Input Parameters

output Return type. Accepted

values: 'server' (result is stored on a file on the server), 'client' (result is directly transferred to the client), or 'both'. Default is 'both'.

Type = string

sequence Input sequence. You need to

supply either this parameter or the next one (tmp_infile).

Type = string

tmp_infile Name of the file with input

sequence on the server. You need to supply either this parameter or the previous

one (sequence). Type = string

format Input sequence format.

Supported: IG

(Intelligenetics), WC (wconsensus), raw, fasta.

Default is fasta Type = string

length Oligomer length.

(required) Type = int

organism Organism. Words need to be

underscore separated

(required) underscor (example:

Escherichia_coli_K12).

background model: Type of

sequences used as background model for estimating expected oligonucleotide frequencies. Supported: upstream,

upstreamL, upstream-noorf,

intergenic, input Type = string

stats List of statistics to return.

Supported:occ, mseq, freq, proba, ratio, zscore, like, pos,

rank

Type = string

noov No overlapping. Disable the

detection of overlapping matches for self-overlapping patterns (ex TATATA,

GATAGA). Type = string

str Oligonucleotide occurrences

found on both stands are summed (2str) or not (1str).

Default is 2str. Type = string

sort Sort oligomers according to

overrepresentation.

Type = string

lth Lower threshold on some

parameter. Format='param

value'

Type = string

Output Parameters

server Location of the result file on the

server. This can be used as input for a further request.

Type = string

command The stand alone command

executed on the server.

Type = string

client The result discovered

pattern(s). Type = string

Method gene_info

Description Get information about genes.

Parameters Input Parameters

organism Organism. Words need to be (required) underscore separated (example:

Escherichia_coli_K12).

Type = string

List of gene(s) for which you query

(required) want info on.

Type = string

full Full match only (no substring

matching). Type = string

Do not print the query at the noquery

begining of each line.

Type = string

descr Match query against the

description, too, not just against gene ID and name.

Type = string

Feature type (CDS, mRNA, tRNA, rRNA, scRNA). feattype

Type = string

Output Parameters

command The stand alone command

executed on the server.

Type = string

client The result gene(s) info(s).