

S14 Text: Practical sessions for Tandem repeated motifs search (pdf).

Bioinformatics and Genome Analyses

September 18 – December 15, 2017. Institut Pasteur Tunis
<https://webext.pasteur.fr/tekaia/BCGAIPT2017.html>

Aim of the session:

- 1- Connect to the server <https://tandem.bu.edu/trf/trf.html> and download the “Linux command line 64 bits” version into your `~/bin/` directory and rename it *trf*.
- 2- Make a directory TRF where to work during this session (same level as RBH).
- 3- Data to be considered: GSACE.dna, GCAGL.dna, and GZYRO.dna.
- 4- Search for repeated motifs in genes:
 - consider first, 2 simple examples: YDR204w.dna, YDR545.dna
 - search for tandem motifs using the *trf* server than the local *trf*.
- 5- Write a script to search for tandem motifs in all SACE gene sequences (included in `allsacedna.fasta` directory)
- 6- Write a script to show results into a table form.
- 7- Look for genes including megasatellites (Megasatellites are motifs with size ≥ 90 nt).

Reminder (G. Benson (1999) "Tandem repeats finder: a program to analyze DNA sequences". *NAR*:27 pp. 573-580.):

use: *trf file match mismatch delta pm pi minscore maxperiod*

where

file = dna sequence(s) input file (fasta)

match = matching weight

mismatch = mismatching penalty

delta = indel penalty

pm = match probability (whole number)

pi = indel probability (whole number)

minscore = minimum alignment score to report

(exp: 50 assuming perfect alignment. Need to align at least 5 characters to meet the minimum score (5 copies of 5 char motif)

maxPeriod = maximum period size to report

The recommended values for Match, Mismatch and Delta are 2, 7, and 7 respectively.

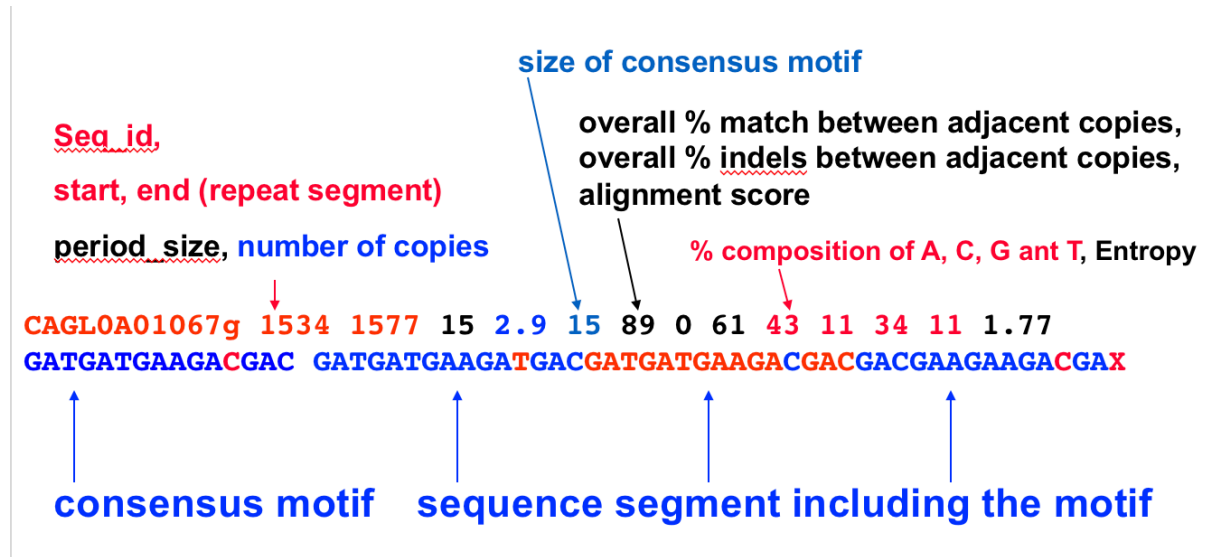
trf inputfile.fa 2 7 7 80 10 50 2000 -d -m -h

-d : output text file, -m: mask input seq and put “N” in repeated positions,

-h : no html output file

Example of command line:

trf CAGLOA01067g.fa 2 7 7 80 10 50 2000 -d -m -h
and corresponding output:



Other examples:

Seq_Id, start, end (repeat segment), period_size, number of copies, size of consensus motif, overall % match between adjacent copies, overall % indels between adjacent copies, alignment score, % composition of A, C, G ant T, Entropy, motif, segment

CAGLOA01067g 1534 1577 15 2.9 15 89 0 61 43 11 34 11 1.77 GATGATGAAGACGAC
GATGATGAAGATGACGATGATGAAGACGACGACGAAGAAGACGA

CAGLOA01193g 113 147 12 2.9 12 100 0 70 51 31 8 8 1.63 AGAACATCACAC AGAACATCACACAGAACATCACACAGAACATCACA

CAGLOA01001g 534 586 15 3.5 15 92 0 88 33 32 20 13 1.91 TCAGCAAGCTCAAGC
TCAGCAAGCTCAGGCTCAGCAAGCTCAAACCTCAGCAAGCTCAAGCTCAGCAAG

CAGLOA01001g 565 651 42 2.1 42 88 0 129 37 32 18 11 1.86 CAGCAAGCTCAAGCTCAGCAAGTACCACAGTACAGAACACAA
CAGCAAGCTCAAGCTCAGCAAGTACCACAGTACCGTACACAACAGCAAGCTCCAGCTCAGCAAGTACCACAGTATAGAACGCAACAG

CAGLOA01284g 1856 2048 51 3.8 50 73 14 162 30 6 34 28 1.83

TGGATGAGAACGGTGATTGTGACTATAGTTGTA AAAAAGACGGTGTTAAGT

TGGATGAGAACGGTGATTGTGACTATAGTTGTA AAAAATGGCGGTGTTAAGTTGGATGAAGACGGTAATTGTGATATGTCTGTAAAGA
GAACGCGGTCAGTTGGATGAGGATGGTATTGTGACTTGAGCTGTA AAGAAGATGGTGGTAAGTTAGATGGTAACGGTGATTGTGAT
TATAGTTGCAAAGAAGA

CAGLOA01284g 1856 2045 102 1.9 101 79 10 217 29 6 34 28 1.83

TGGATGAGAACGGTGATTGTGACTATAGCTGTA AAAAATGACGGTGGTAAGTTAGATGAAGACGGTAATTGTGATATGTCTGCAAAGAG
AACGCGGTCAGT

TGGATGAGAACGGTGATTGTGACTATAGTTGTA AAAAATGGCGGTGTTAAGTTGGATGAAGACGGTAATTGTGATATGTCTGTAAAGA
GAACGCGGTCAGTTGGATGAGGATGGTATTGTGACTTGAGCTGTA AAGAAGATGGTGGTAAGTTAGATGGTAACGGTGATTGTGAT
TATAGTTGCAAAGA

4- Search for repeated motifs in genes:

- consider first, 2 simple examples: YDR204w.dna, YDR545w.dna
- search for tandem motifs using the *trf* server than the local *trf*.

trf YDR204w.dna 2 7 7 80 10 50 2000 -d -m -h > YDR204w.trf

trf YDR545w.dna 2 7 7 80 10 50 2000 -d -m -h > YDR545w.trf

5- Write a script to search for tandem motifs in all SACE gene sequences (included in allsacedna.fasta directory). Output should be directed to gene_ident.trf.

```
#!/bin/sh
For file in `ls $path/allsacedna.fasta/*.dna`
do
FILE=`echo $file | sed -e "s/\./.*fasta\\//g" -e "s/\./dna//g"`
trf $file 2 7 7 80 10 50 2000 -d -m -h > $FILE.trf
done
```

6- concatenate results into a single file SACE.trf.

```
cat Y*.trf >> SACE.trf
```

7- Look for genes including megasatellites (Megasatellites are motifs with size ≥ 90 nt).

Following the detailed example shown with CAGLOA01067g.fa the size of the consensus tandemly repeated motif is at position 6.

```
#!/bin/perl
$IN=@ARGV[0]; #input file SACE.trf
$OUT="MEGASAT90.trf";
open (IN, "$IN");
open (OUT, ">$OUT");
while<IN>
{
@tab=split(/\s+/, $_);
if ($tab[5] >= 90 ) {print OUT $_; }
}
close(IN); close(OUT);
```

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