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#!/usr/bin/perl
use strict;
use warnings;
use File::Basename;
# SCRIPT TO CALCULATE SOME GENE PARAMETERS
##### Spring 2016 - MLS - UNIL - Marie Zufferey
# USAGE :
# ./stat_CDS.pl my_cds.fasta outfile.txt
    -> my cds.fasta => file with the CDS
    -> outfile.txt
                   => file in which output written !!!! if already exist, will be overwritten !
# output as follow (tab-separated):
    -> the tag of the sequence, its length, the ratio of purine and the ratio of GC-content (for the
whole gene sequence):
               LENGTH
                           ratioGC
                                       ratioPu
   SEQ_TAG
my scds = sARGV[0];
my $outfile = $ARGV[1];
print("WARNING: outfile will be overwritten !\n");
system("rm -f $outfile");
system("touch $outfile");
# write the header in the output file
open(my $out, '>>', $outfile) or die("open: $$");
print($out "Seq_tag\tLength\tratioGC\tratioPu\n");
close($out);
my ($id, $seq, $nG, $nC, $nA, $size, $nPu, $nGC);
open(CDS, $cds) || die;
while(my $line = <CDS>){
    chomp($line);
    if(\frac{= - /^{}}{}){ # => it is the ID line
       if(length $seq){
                                 # we have a new ID (all IDs except the 1st)
           # calculate and write the sequence information in output file
           snG = (seq =~ tr/gG//);
           snC = (seq =  tr/cC//);
           nA = (seq =  tr/aA//);
           $size = length($seg);
           nPu = (nA+nG)/size;
           snGC = (snC+snG)/ssize;
           open(my $out, '>>', $outfile) or die("open: $$");
           print($out "$id\t$size\t$nGC\t$nPu\n" );
           close($out);
           # reinitialize the seq
           $seq = "";
        (sid = sline) =   s/^{/}; # store the id without the >  that starts the line
                   # => it is one seq line (can be separated with \n ...)
   $seq .= $line; # concatenate the sequence parts
   }
# do not forget the last sequence # TODO put in subroutine to avoid repetition of code...
# calculate and write the sequence information in output file
nG = (seq = tr/gG//);
snC = (seq =~ tr/cC//);
nA = (seq =  tr/aA//);
$size = length($seq);
nPu = (nA+nG)/size;
snGC = (snC+snG)/ssize;
open($out, '>>', $outfile) or die("open: $$");
print($out "$id\t$size\t$nGC\t$nPu\n" );
close($out);
close(CDS);
```

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#foreach my $pos(@query_id){
#    print("$pos\n");
#    open(my $queryF, '>>', $query_file) or die("open: $$");
#    print($queryF ">$pos\n$all_seq{$pos}\n" );
#    print $x;
#}
```