# **Bioinformatics one-liners**

Useful bash one-liners for useful for bioinformatics.

https://github.com/stephenturner/oneliners

#### Sources:

- http://sed.sourceforge.net/sed1line.txt
- <a href="https://github.com/lh3/seqtk">https://github.com/lh3/seqtk</a>
- http://lh3lh3.users.sourceforge.net/biounix.shtml
- <a href="http://genomespot.blogspot.com/2013/08/a-selection-of-useful-bash-one-liners.html">http://genomespot.blogspot.com/2013/08/a-selection-of-useful-bash-one-liners.html</a>
- <a href="http://biowize.wordpress.com/2012/06/15/command-line-magic-for-your-gene-annotations/">http://biowize.wordpress.com/2012/06/15/command-line-magic-for-your-gene-annotations/</a>
- http://genomics-array.blogspot.com/2010/11/some-unixperl-oneliners-for.html
- <a href="http://gettinggeneticsdone.blogspot.com/2013/10/useful-linux-oneliners-for-bioinformatics.html#comments">http://gettinggeneticsdone.blogspot.com/2013/10/useful-linux-oneliners-for-bioinformatics.html#comments</a>

#### awk, sed

Sum column 1 of file.txt:

```
awk '{sum+=$1} END {print sum}' file.txt
```

Number each line in file.txt:

```
sed = file.txt | sed 'N;s/\n/ /'
```

Get unique entries in file.txt based on column 1 (takes only the first instance):

```
awk '!arr[$2]++' file.txt
```

Print each line where the 5th field is equal to 'abc123':

```
awk '$5 == "abc123"' file.txt
```

Print each line where the 5th field is not equal to 'abc123':

```
awk '$5 == "abc123"' file.txt
```

Print each line whose 7th field matches the regular expression:

```
awk '$7 ~ /^[a-f]/' file.txt
```

Print each line whose 7th field *does not* match the regular expression:

```
awk '$7 ~ /^[a-f]/' file.txt
```

Replace all occurances of foo with bar in file.txt:

```
sed 's/foo/bar/g' file.txt
```

Convert a FASTQ file to FASTA:

```
sed -n '1~4s/^@/>/p;2~4p' file.fq > file.fa
```

Extract every 4th line starting at the second line (extract the sequence from FASTQ file):

```
sed -n '2~4p' file.fq
```

Basic sequence statistics. Print total number of reads, total number unique reads, percentage of unique reads, most abundant sequence, its frequency, and percentage of total in file.fq:

```
cat myfile.fq | awk '((NR-2)%4==0){read=$1;total++;count[read]++}END{for(read in
count){if(!max||count[read]>max)
{max=count[read];maxRead=read};if(count[read]==1){unique++}};print
total,unique,unique*100/total,maxRead,count[maxRead],count[maxRead]*100/total}'
```

Convert .bam back to .fastq:

```
samtools view file.bam | awk 'BEGIN {FS="\t"} {print "@" $1 "\n" $10 "\n+\n" $11}' > file.fq
```

Keep only top bit scores in blast hits (best bit score only):

```
awk '{ if(!x[$1]++) {print $0; bitscore=($14-1)} else { if($14>bitscore) print $0} }' blastout.txt
```

Keep only top bit scores in blast hits (5 less than the top):

```
awk '{ if(!x[$1]++) {print $0; bitscore=($14-6)} else { if($14>bitscore) print $0} }' blastout.txt
```

Trim leading whitespace in file.txt:

```
sed 's/^[ \t]*//' file.txt
```

Trim trailing whitespace in file.txt:

```
sed 's/[ \t]*$//'
```

Trim leading and trailing whitespace in file.txt:

```
sed 's/^[ \t]*//;s/[ \t]*$//' file.txt
```

Delete blank lines in file.txt:

```
sed '/^$/d'
```

# sort, uniq, cut, etc.

Count the number of unique lines in file.txt

```
cat file.txt | sort | uniq | wc -l
```

Find number of lines shared by 2 files:

```
sort file1 file2 | uniq -d
```

Find the most common strings in column 2:

```
cut -f2 file.txt | sort | uniq -c | sort -k1nr | head
```

Pick 10 random lines from a file:

```
shuf file.txt | head -n 10
```

Print rows where column 3 is larger than column 5 in file.txt:

```
awk '$3>$5' file.txt
```

Compute the mean of column 2:

```
awk '{x+=$2}END{print x/NR}' file.txt
```

Extract fields 2, 4, and 5 from file.txt:

```
awk '{print $2,$4,$5}' input.txt
```

Print all possible 3mer DNA sequence combinations:

```
echo {A,C,T,G}{A,C,T,G}{A,C,T,G}
```

Untangle an interleaved paired-end FASTQ file. If a FASTQ file has paired-end reads intermingled, and you want to separate them into separate /1 and /2 files, and assuming the /1 reads precede the /2 reads:

```
cat interleaved.fq |paste - - - - - - | tee >(cut -f 1-4 | tr "\t" "\n" > deinterleaved_1.fq) | cut -f 5-8 | tr "\t" "\n" > deinterleaved_2.fq
```

#### find, xargs, and GNU parallel

Download GNU parallel at <a href="https://www.gnu.org/software/parallel/">https://www.gnu.org/software/parallel/</a>.

Search for .bam files anywhere in the current directory recursively:

```
find . -name "*.bam"
```

Delete all .bam files:

```
find . -name "*.bam" | xargs rm
```

Rename all .txt files to .bak (backup \*.txt before doing something else to them, for example):

```
find . -name "*.txt" | sed "s/\.txt$//" | xargs -i echo mv \{\}.txt \{\}.bak | sh
```

Chastity filter raw Illumina data (grep reads containing :N:, append (-A) the three lines after the match containing the sequence and quality info, and write a new filtered fastq file):

```
find *fq | parallel "cat {} | grep -A 3 '^@.*[^:]*:N:[^:]*:' | grep -v '^\-\-$'
> {}.filt.fq"
```

Run FASTQC in parallel 12 jobs at a time:

```
find *.fq | parallel -j 12 "fastqc {} --outdir ."
```

Index your bam files in parallel, but only echo the commands (--dry-run) rather than actually running them:

```
find *.bam | parallel --dry-run 'samtools index {}'
```

### seqtk

Download seqtk at <a href="https://github.com/lh3/seqtk">https://github.com/lh3/seqtk</a>. Seqtk is a fast and lightweight tool for processing sequences in the FASTA or FASTQ format. It seamlessly parses both FASTA and FASTQ files which can also be optionally compressed by gzip.

#### Convert FASTQ to FASTA:

```
seqtk seq -a in.fq.gz > out.fa
```

Convert ILLUMINA 1.3+ FASTQ to FASTA and mask bases with quality lower than 20 to lowercases (the 1st command line) or to N (the 2nd):

```
seqtk seq -aQ64 -q20 in.fq > out.fa
seqtk seq -aQ64 -q20 -n N in.fq > out.fa
```

Fold long FASTA/Q lines and remove FASTA/Q comments:

```
seqtk seq -Cl60 in.fa > out.fa
```

Convert multi-line FASTQ to 4-line FASTQ:

```
seqtk seq -10 in.fq > out.fq
```

Reverse complement FASTA/Q:

```
seqtk seq -r in.fq > out.fq
```

Extract sequences with names in file name.lst, one sequence name per line:

```
seqtk subseq in.fq name.lst > out.fq
```

Extract sequences in regions contained in file reg.bed:

```
seqtk subseq in.fa reg.bed > out.fa
```

Mask regions in reg. bed to lowercases:

```
seqtk seq -M reg.bed in.fa > out.fa
```

Subsample 10000 read pairs from two large paired FASTQ files (remember to use the same random seed to keep pairing):

```
seqtk sample -s100 read1.fq 10000 > sub1.fq
seqtk sample -s100 read2.fq 10000 > sub2.fq
```

Trim low-quality bases from both ends using the Phred algorithm:

```
seqtk trimfq in.fq > out.fq
```

Trim 5bp from the left end of each read and 10bp from the right end:

```
seqtk trimfq -b 5 -e 10 in.fa > out.fa
```

Untangle an interleaved paired-end FASTQ file. If a FASTQ file has paired-end reads intermingled, and you want to separate them into separate /1 and /2 files, and assuming the /1 reads precede the /2 reads:

```
seqtk seq -l0 interleaved.fq | awk '{if ((NR-1) % 8 < 4) print >>
"deinterleaved_1.fq"; else print >> "deinterleaved_2.fq"}'
```

#### **GFF3 Annotations**

Print all sequences annotated in a GFF3 file.

```
cut -s -f 1,9 yourannots.gff3 | grep $'\t' | cut -f 1 | sort | uniq
```

Determine all feature types annotated in a GFF3 file.

```
grep -v '^#' yourannots.gff3 | cut -s -f 3 | sort | uniq
```

Determine the number of genes annotated in a GFF3 file.

```
grep -c $'\tgene\t' yourannots.gff3
```

Extract all gene IDs from a GFF3 file.

```
grep \' \ yourannots.gff3 | perl -ne '/ID=([^;]+)/ and printf("%s\n", $1)'
```

Print length of each gene in a GFF3 file.

```
grep \'\ yourannots.gff3 | cut -s -f 4,5 | perl -ne '@v = split(/\t/); printf("%d\n", \ v[1] - \ v[0] + 1)'
```

FASTA header lines to GFF format (assuming the length is in the header as an appended "\_length" as in <u>Velvet</u> assembled transcripts):

```
grep '>' file.fasta | awk -F "_" 'BEGIN{i=1; print "##gff-version 3"}{ print
$0"\t
BLAT\tEXON\t1\t"$10"\t95\t+\t.\tgene_id="$0";transcript_id=Transcript_"i;i++ }'
> file.gff
```

# Other generally useful aliases for your .bashrc

Get a prompt that looks like user@hostname:/full/path/cwd/:\$

```
export PS1="\u@\h:\w\\$ "
```

Never type cd .../... again:

```
alias ..='cd ..'
alias ...='cd .././'
alias ...='cd ../../'
alias ....='cd ../../../'
alias ....='cd ../../../'
```

Ask before removing or overwriting files:

```
alias mv="mv -i"
alias cp="cp -i"
alias rm="rm -i"
```

My favorite 1s aliases:

```
alias ls="ls -1p --color=auto"
alias l="ls -lhGgo"
alias ll="ls -lh"
alias la="ls -lhGgoA"
alias lt="ls -lhGgotr"
alias lS="ls -lhGgoSr"
alias l.="ls -lhGgod .*"
alias lhead="ls -lhGgo | head"
alias ltail="ls -lhGgo | tail"
alias lmore='ls -lhGgo | more'
```

Use cut on space- or comma- delimited files:

```
alias cuts="cut -d \" \""
alias cutc="cut -d \",\""
```

Pack and unpack tar.gz files:

```
alias tarup="tar -zcf"
alias tardown="tar -zxf"
```

Use mcd to create a directory and cd to it simultaneously:

```
function mcd { mkdir -p "$1" && cd "$1";}
```

Go up to the parent directory and list it's contents:

```
alias u="cd ..;ls"
```

Make grep pretty:

```
alias grep="grep --color=auto"
```

Refresh your .bashrc:

```
alias refresh="source ~/.bashrc"
```

#### Common typos:

```
alias mf="mv -i"
alias mroe="more"
```