## Filtering with SAMTools

#### $\odot$ Reservations

Use our summer school reservation (CoreNGS-Fri) when submitting batch jobs to get higher priority on the Is6 normal queue today:

```
sbatch --reservation=CoreNGS-Fri <batch file>.slurm
idev -m 180 -N 1 -A OTH21164 -r CoreNGS-Fri
```

- Overview
  - Setup
- References
  - SAM header fields
  - SAM flags field
  - SAM CIGAR string
- Exercises
  - ° Analyzing the CIGAR string for indels
  - Filtering by location range
  - About mapping quality
  - Filtering for high-quality reads

## Overview

As we have seen, the SAMTools suite allows you to manipulate the SAM/BAM files produced by most aligners. There are many sub-commands in this suite, but the most common and useful are:

- 1. Convert text-format SAM files into binary BAM files (samtools view) and vice versa
- 2. Sort BAM files by reference coordinates (samtools sort)
- 3. Index BAM files that have been sorted (samtools index)
- 4. Filter alignment records based on BAM flags, mapping quality or location (samtools view)

Since BAM files are binary, they can't be viewed directly using standard Unix file viewers such as more, less and head. We have seen how samtools view can be used to binary-format BAM files into text format for viewing. But samtools view also has options that let you do powerful filtering of the output. We focus on this filtering capability in this set of exercises.

The most common samtools view filtering options are:

- -q N only report alignment records with mapping quality of at least N (>= N).
- -f 0xXX only report alignment records where the specified flags are all set (are all 1) ° you can provide the flags in decimal, or as here as hexadecimal
- -F 0xXX only report alignment records where the specified flags are all cleared (are all 0)

## Setup

Login to Is6.tacc.utexas.edu and start an idev session.

#### Start an idev session

```
idev -m 180 -N 1 -A OTH21164 -r CoreNGS-Fri
# or
idev -m 90 -N 1 -A OTH21164 -p development
```

Next set up a directory for these exercises, and copy an indexed BAM file there. This is the yeast\_pe.sort.bam file from our The Basic Alignment Workflow exercises.

#### Setup for samtools exercises

```
mkdir -p $SCRATCH/core_ngs/samtools
cd $SCRATCH/core ngs/samtools
cp $CORENGS/catchup/for_samtools/* .
```

## References

Handy links

- The SAM format specification
  - especially section 1.4 alignment section fields
- Manual for SAMTools
  - especially the 1st section on samtools view.

## SAM header fields

The 11 SAM alignment record required fields (Tab-delimited).

Col	Field	Туре	Regexp/Range	Brief description
1	QNAME	String	[!-?A-~]{1,255}	Query template NAME read name from fastq
2	FLAG	Int	$[0, 2^{16} - 1]$	bitwise FLAGs
3	RNAME	String	\* [!-()+-<>-~][!-~]*	Reference sequence NAME contig + start
4	POS	Int	$[0, 2^{29}-1]$	1-based leftmost mapping POSition = locus
5	MAPQ	Int	$[0, 2^8 - 1]$	MAPping Quality
6	CIGAR	String	\* ([0-9]+[MIDNSHPX=])+	CIGAR string use this to find end coordinate
7	RNEXT	String	\* = [!-()+-<>-~][!-~]*	Ref. name of the mate/next segment
8	PNEXT	Int	$[0, 2^{29}-1]$	Position of the mate/next segment
9	TLEN	Int	$[-2^{29}+1, 2^{29}-1]$	observed Template LENgth insert size, if paired
10	SEQ	String	\* [A-Za-z=.]+	segment SEQuence
11	QUAL	String	[!-~]+	ASCII of Phred-scaled base QUALity+33

## SAM flags field

Meaning of each bit (flag) in the SAM alignment records flags field (column 2). The most commonly flags are denoted in red.

Е	Bit		
Decima	l Hex	Description	-
1	0x1	template having multiple segments in sequencing	1 = part of a read pair
<b>2</b>	0x2	each segment properly aligned according to the aligner	1 = "properly" paired
4	0x4	segment unmapped read <u>did</u> map = 0	1 = read did <u>not</u> map
8	0x8	next segment in the template unmapped	1 = mate did <u>not</u> map
16	0x10	SEQ being reverse complemented <u>plus</u> strand read = 0	1 = minus strand read
32	0x20	SEQ of the next segment in the template being reverse complemented	1 = mate on minus strand
64	0x40	the first segment in the template	1 = R1 read
128	0x80	the last segment in the template	1 = R2 read
256	0x100	secondary alignment	1 = secondary alignment
512	0x200	not passing filters, such as platform/vendor quality controls	
1024	0x400	PCR or optical duplicate	1 = marked as duplicate
2048	0x800	supplementary alignment	1 = maps to ALT contig

## SAM CIGAR string

Format of the CIGAR string in column 6 of SAM alignment records.

# Ref CTGGCCATTATCTC--GGTGGTAGGACATGGCATGCCC Read aaATGTCGCGGTG.TAGGAggatcc 2S5M2I4M1D5M6S

	Op	$\mathbf{BAM}$	Description		
	М	0	alignment match (can be a sequence match or mismatch)		
	I	1	insertion to the reference		
	D	2	deletion from the reference <i>"N" indicates splicing event in</i>		
	Ν	3	skipped region from the reference <b>RNAseq BAMs</b>		
	S	<b>4</b>	soft clipping (clipped sequences present in SEQ)		
*	Н	5	hard clipping (clipped sequences NOT present in $SEQ$ )		
*	Ρ	6	padding (silent deletion from padded reference)		
*	=	7	sequence match *Rarer / newer		
*	X	8	sequence mismatch		

CIGAR = "Concise Idiosyncratic Gapped Alignment Report"

## Exercises

## Analyzing the CIGAR string for indels

Suppose we want to know how many alignments included insertions or deletions (*indels*) versus the reference. Looking at the CIGAR field definition table above, we see that insertions are denoted with the character I and deletions with the character D. We'll use this information, along with samtools view, cut and grep, to count the number of indels.

```
mkdir -p $SCRATCH/core_ngs/samtools
cd $SCRATCH/core_ngs/samtools
cp $CORENGS/catchup/for_samtools/* .
```

We'll do this first exercise one step at a time to get comfortable with piping commands. Start by just looking at the first few alignment records of the **BAM** file:

```
module load biocontainers # takes a while
module load samtools
cd $SCRATCH/core_ngs/samtools
samtools view yeast_pe.sort.bam | head
```

With all the line wrapping, it looks pretty ugly. Still, you can pick out the CIGAR string in column 6. Let's select just that column with cut:

```
samtools view yeast_pe.sort.bam | cut -f 6 | head -20
```

Next, make sure we're only looking at alignment records that represent mapped reads. The -F 0x4 option says to filter records where the 0x4 flag (read un mapped) is 0, resulting it only mapped reads being output.

```
samtools view -F 0x4 yeast_pe.sort.bam \mid cut -f 6 \mid head -20
```

Now we use grep with the pattern '[ID]' to select lines (which are now just CIGAR strings) that have Insert or Deletion operators. The brackets ([]) denote a character class pattern, which matches any of the characters inside the brackets. Be sure to ask for Perl regular expressions (-P) so that this character class syntax is recognized.

```
samtools view -F 0x4 yeast_pe.sort.bam | cut -f 6 | grep -P '[ID]' | head
```

Ok, this looks good. We're ready to run this against all the alignment records, and count the results:

#### Count reads that mappedi with indels

samtools view -F 0x4 yeast\_pe.sort.bam | cut -f 6 | grep -P '[ID]' | wc -l

There are 6697 such records.

What is that in terms of the rate of indels? For that we need to count the total number of mapped reads. Here we can just use the -c (count only) option to s amtools view.

```
Count all mapped reads
samtools view -c -F 0x4 yeast_pe.sort.bam
```

There should be 547664 mapped alignments.

Knowing these two numbers we can just divide them, using **awk** (remember, **bash** only does integer arithmetic). Because we're not piping anything in to **aw**, any body we specify won't be executed. So we do the math in the **BEGIN** section:

awk 'BEGIN{ print 100\*6697/547664,"%" }'

The result should be 1.22283 %.

In addition to the rate, converted to a percentage, we also output the literal percent sign (%). The double quotes (") denote a literal string in **awk**, and the comma between the number and the string says to separate the two fields with whatever the default Output Field Separator (OFS) is. By default, both **a** wk's Input Field Separator (FS) is whitespace (any number of space and Tab characters) and its Output Field Separator is a single space.

So what if we want to get fancy and do all this in a one-liner command? We can use **echo** and **backtick evaluation** to put both numbers in a string, then pipe that string to **awk**. This time we use **awk** body code, and refer to the two **whitespace**-separated fields being passed in: total count (**\$1**) and indel count (**\$2**).

#### One-liner for calculating BAM indel rate

```
echo "`samtools view -F 0x4 -c yeast_pe.sort.bam` \
    $( samtools view -F 0x4 yeast_pe.sort.bam | cut -f 6 | grep -P '[ID]' | wc -l )" \
    | awk '{ print 100 * $2/$1,"%" }'
```

awk also has a printf function, which can take the standard formatting commands (see https://en.wikipedia.org/wiki /Printf\_format\_string#Type\_field).

awk 'BEGIN{ printf("%.2f %%\n", 100\*6697/547664) }'

#### Notes:

- The **printf** arguments are enclosed in parentheses since it is a true function.
  - The 1st argument is the *format string*, enclosed in double quotes.
    - The %.2f format specifier says to output a floating point number with 2 digits after the decimal place.
    - The %% format specifier is used to output a single, literal percent sign.
    - Unlike the standard print statement, the printf function does not automatically append a newline to the output, so \n is added to the format string here.
- Remaining arguments to printf are the values to be substituted for the format specifiers (here our percentage computation).

### Filtering by location range

Sometimes you just want to examine a subset of reads in detail. Once you have a *sorted and indexed* BAM, you can use the coordinate filtering options of *samtools view* to do this. Here are some examples:

 $\oslash$ 

When you're interested in *mapped reads* (which is true most of the time) be sure to specify the **-F 0x4** option, which says to filter records where the **0x4** flag (read *unmapped*) is **0**, resulting it only *mapped* reads being output.

```
mkdir -p $SCRATCH/core_ngs/samtools
cd $SCRATCH/core_ngs/samtools
cp $CORENGS/catchup/for_samtools/* .
module load biocontainers
module load samtools
```

cd \$SCRATCH/core\_ngs/samtools

 $\odot$ 

```
# count the number of reads mapped to chromosome 2 (chrII)
samtools view -c -F 0x4 yeast_pe.sort.bam chrII
```

# count the number of reads mapped to chromosomes 1 or M (chrI, chrM)
samtools view -c -F 0x4 yeast\_pe.sort.bam chrI chrM

# count the number of reads mapped to chromosomes 1 that overlap coordinates 1000-2000 samtools view -c -F 0x4 yeast\_pe.sort.bam chrI:1000-2000

# since there are only 20 reads in the chrI:1000-2000 region, examine them individually samtools view -F 0x4 yeast\_pe.sort.bam chrI:1000-2000

```
# look at a subset of field for chrI:1000-2000 reads
# 2=flags, 3=contig, 4=start, 5=mapping quality, 6=CIGAR, 9=insert size
samtools view -F 0x4 yeast_pe.sort.bam chrI:1000-2000 | cut -f 2-6,9
```

#### samtools view -L <bed\_file>

You can also provide a **BED**-format file with one record for each desired overlap region: **samtools view -L <bed\_file>**. This is a quick way to perform one of the functions of **bedtools intersect**.

Since you will find yourself wanting to interpret the flags field, and it's easier to do that when it is represented as hexadecimal, let's use awk to help do that.

```
# look at a subset of field for chrI:1000-2000 reads
# 2=flags, 3=contig, 4=start, 5=mapping quality, 6=CIGAR, 9=insert size
samtools view -F 0x4 yeast_pe.sort.bam chrI:1000-2000 | cut -f 2-6,9 | \
    awk 'BEGIN{FS=OFS="\t"}
    {$1 = sprintf("0x%x", $1); print}'
```

Notes:

- If a command line is continued on more than one line, the line continuation character (1) is used.
- There is a sprintf (string print formatted) function in awk, just as there is in many modern programming languages.
- We use sprintf to re-format the 1st input field (here the cut flags) in hex
  - using the *format directive* (%x)
  - also adding a literal "0x" prefix to denote the numeric base is hexadecimal
- The results of **sprintf** are stored back into the 1st field (\$1), replacing the original value.
- awk's print statement is then used with no other arguments to print all its input fields, including the changed \$1.

#### Exercise: How many of the chrl:1000-2000 alignments are from properly paired mapped reads?

Properly paired reads have the **0x2** flag set (1). *All* our reads also have the **0x1** flag set because they are paired-end reads. Mapped reads will have the **0x4** flag cleared (0), and properly paired mapped reads will have the **0x4** flag cleared (0) as well (*mate not unmapped* = mate mapped). So *all mapped proper pairs* will end in **0x3** = **0b0011** (binary).

Also, using '\$' in a grep pattern anchors the pattern to the *end of the line* (only patterns matching before a linefeed are printed). Here's one way to do it, building on our last command line:

```
samtools view -F 0x4 yeast_pe.sort.bam chrI:1000-2000 | awk '
BEGIN{FS=OFS="\t"}
{$2 = sprintf("0x%x", $2); print $2}' | grep -c '3$'
```

Note that here we don't need the line continuation character because the newline after the first single quote is part of the script command string.

samtools view -c -F 0x4 -f 0x2 yeast\_pe.sort.bam chrI:1000-2000

## About mapping quality

Mapping qualities are a measure of how likely a given sequence alignment to its reported location is correct. If a read's mapping quality is low (especially if it is zero, or *mapQ 0* for short) the *read maps to multiple locations on the genome* (they are *multi-hit* or *multi-mapping* reads), and we can't be sure whether the reported location is the correct one.

Aligners also differ in whether they report alternate alignments for multi-hit reads. Some things to keep in mind:

- Alternate locations for a mapped read will be flagged as secondary (flag 0x100)
- While they often provide valuable information, secondary reads must be filtered for some downstream applications
   e.g., ChIP-seq peak calling and variant analysis with GATK.
- When secondary reads are reported, the total number of alignment records in the BAM file is greater than the number of reads in the input FAST( files!
  - ° this affects how the true mapping rate must be calculated
  - true mapping rate = (pirmary mapped reads) / (total BAM file sequences secondary mapped reads)

Here are some examples of how different aligners handle reporting of multi-hit reads and their mapping qualities:

- bwa aln (global alignment) and bowtie2 with default parameters (both --local default end-to-end mode) report at most one location for a read that maps
  - this will be the location with the *best* mapping quality and alignment
  - if a given read maps equally well to multiple locations, these aligners pick one location at random
    - bwa aln will always report a 0 mapping quality for these multi-hit reads
    - bowtie2 will report a low mapping quality (< 10), based on the complexity (information content) of the sequence</p>
- bwa mem (local alignment) can always report more than one location for a mapped read
  - its definition of a *secondary* alignment is different (and a bit non-standard)
    - if one part of a read maps to one location and another part maps somewhere else (e.g. because of RNA splicing), the longer alignment is marked as primary and the shorter as secondary.
    - there is no way to disable reporting of secondary alignments with bwa mem.
      - but they can be filtered from the sorted BAM with -F 0x100 (secondary alignment flag = 0).
- bowtie2 can be configured to report more than one location for a mapped read
  - the -k N option says to report up to N alignments for a read
- most transcriptome-aware RNA-seq aligners by default report more than one location for a mapped read
  - e.g. hisat2, star, tophat2.
  - · when reads are quantified (counted with respect to genes), multiply-mapped reads can be counted fractionally
    - e.g. if a read maps to 5 genes, it can be counted as 1/5 for each of the genes

## Filtering for high-quality reads

Using our yeast\_pe.sort.bam file, let's do some some quality filtering.

```
mkdir -p $SCRATCH/core_ngs/samtools
cd $SCRATCH/core_ngs/samtools
cp $CORENGS/catchup/for_samtools/* .
module load biocontainers
module load samtools
```

Exercise: Use samtools view with -F, -f and -q options to create a BAM containing only mapped, properly paired, high-quality (mapQ 20+) reads. Call the output file yeast\_pe.sort.filt.bam.

Note that we use the -b flag to tell samtools view to output BAM format (not the SAM format text we've been looking at).

solution
<pre>samtools view -b -F 0x4 -f 0x2 -q 20 yeast_pe.sort.bam &gt; yeast_pe.sort.filt.bam # or</pre>
samtools view -b -F 0x4 -f 0x2 -q 20 -o yeast_pe.sort.filt.bam yeast_pe.sort.bam

Exercise: How many records are in the filtered BAM compared to the original? How many read pairs does this represent?

samtools view -c

```
solution
```

```
samtools view -c yeast_pe.sort.bam
samtools view -c yeast_pe.sort.filt.bam
# or to be really fancy...
echo "`samtools view -c yeast_pe.sort.bam` \
    `samtools view -c yeast_pe.sort.filt.bam`" | \
    awk '{printf "%d original reads\n", $1;
    printf "%d Q20 reads (%d read pairs)\n", $2, $2/2;
    printf "%0.2f%% high-quality reads\n", 100*$2/$1}'
```

There were 1,184,360 alignment records in the original BAM, and only 456,890 in the quality-filtered BAM, around 38% of our starting reads.

Since we have only properly paired reads, the filtered BAM will contain equal numbers of both R1s and R2s. So the number of read pairs is 456890/2 or 228451.

#### Exercise: How many primary aligned reads (0x100 = 0) are in the bwa\_local.sort.dup.bam file?

## Combining SAM flags

samtools view only pays attention to one -F or -f option, so to specify more than one flag value you need to combine them into one number. For example, combining 0x100 and 0x4 yields 0x104.

You want both the secondary (0x100) and unmapped (0x4) flags to be 0.

#### solution

```
samtools view -F 0x104 -c bwa_local.sort.dup.bam
```

#### There are 874791 primary alignments.

#### solution

```
samtools view -F 0x4 -f 0x100 -c bwa_local.sort.dup.bam
```

And 133209 secondary alignments.