# Bedtools: Analyzing your aligned experiment

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## Use bedtools coverage to create a signal track

A *signal track* is a **bedGraph** (**BED3+**) file with an entry for every base in a defined set of regions (see https://genome.ucsc.edu/goldenpath/help /bedgraph.html). **bedGraph** files can be visualized in the Broad's **IGV** (Integrative Genomics Viewer) application (https://software.broadinstitute.org /software/igv/download) or in the **UCSC Genome Browser** (https://genome.ucsc.edu/).

The **bedtools coverage** function (https://bedtools.readthedocs.io/en/latest/content/tools/coverage.html), with the **-d** (per-base **depth**) option produces output that can be made into a **bedGraph**. Here we'll analyze the per-base coverage of yeast RNAseq reads in our merged yeast gene regions.

Make sure you're in an idev session, then prepare a directory for this exercise.

#### Prepare for bedtools coverage

```
idev -m 120 -N 1 -A OTH21164 -r CoreNGSday5
module load biocontainers
module load bedtools

mkdir -p $SCRATCH/core_ngs/bedtools_coverage
cp $CORENGS/catchup/bedtools_merge/merged*bed $SCRATCH/core_ngs/bedtools_coverage/
cp $CORENGS/yeast_rnaseq/yeast_mrna.sort.filt.bam* $SCRATCH/core_ngs/bedtools_coverage/
```

Then calling **bedtools coverage** is easy. The "A" file will be our gene regions, and the "B" file will be the yeast RNAseq reads. We also use the **-d** (perbase **depth**) and **-s** (force "strandedness") options.

```
cds; cd core_ngs/bedtools_coverage
bedtools coverage -s -d -a merged.good.sc_genes.bed -b yeast_mrna.sort.filt.bam > yeast_mrna.gene_coverage.txt
wc -l yeast_mrna.gene_coverage.txt # 8,829,317 lines!
```

It will complain a bit because our genes file includes the yeast plasmid "2-micron" but the RNAseq BAM doesn't include that contig. We'll ignore that warning.

Let's look at coverage for gene YAL067C:

cat yeast\_mrna.gene\_coverage.txt | grep -P 'YAL067C' | head -50

Will look like this:

chrI	7234	9016	YAL067C 1	-	1	0
chrI	7234	9016	YAL067C 1	-	2	0
chrI	7234	9016	YAL067C 1	-	3	0
chrI	7234	9016	YAL067C 1	-	4	0
chrI	7234	9016	YAL067C 1	-	5	0
chrI	7234	9016	YAL067C 1	-	6	0
chrI	7234	9016	YAL067C 1	-	7	0
chrI	7234	9016	YAL067C 1	-	8	0
chrI	7234	9016	YAL067C 1	-	9	0
chrT	7234	9016	YAL067C 1	_	10	ñ
chrT	7234	9016	VAL.067C 1	_	11	0
chrT	7221	9016	VAL.067C 1	_	10	0
chr.T	7001	9010 0016	VALOGIC 1	-	12	0
CHT1	1234	ANTP	YALUO/C I	-	13	U
chrI	7234	9016	YAL067C 1	-	14	0
chrI	7234	9016	YAL067C 1	-	15	0
chrI	7234	9016	YAL067C 1	-	16	0
chrI	7234	9016	YAL067C 1	-	17	1
chrI	7234	9016	YAL067C 1	-	18	1
chrI	7234	9016	YAL067C 1	-	19	1
chrI	7234	9016	YAL067C 1	_	20	1
chr	7234	9016	YAL067C 1	_	21	1
chrT	7221	9016	VAL.067C 1	_	21	1
chir I	7234	0016	IALUU/C I	-	22	1
CHT1	1234	ANTP	YALUO/C I	-	∠ <i>3</i>	1
chrI	7234	9016	YALU67C 1	-	24	1
chrI	7234	9016	YAL067C 1	-	25	1
chrI	7234	9016	YAL067C 1	-	26	1
chrI	7234	9016	YAL067C 1	-	27	1
chrI	7234	9016	YAL067C 1	-	28	1
chrI	7234	9016	YAL067C 1	-	29	1
chrI	7234	9016	YAL067C 1	-	30	1
chrI	7234	9016	YAL067C 1	_	31	1
chr	7234	9016	YAL067C 1	_	32	1
ahr	7234	9016	VAL 067C 1	_	22	1
apro T	7001	9010 0016	VALUOIC 1	-	22	1
CHT1	1234	ANTP	YALUO/C I	-	54	1
chrI	7234	9016	YAL067C 1	-	35	1
chrI	7234	9016	YAL067C 1	-	36	1
chrI	7234	9016	YAL067C 1	-	37	1
chrI	7234	9016	YAL067C 1	-	38	2
chrI	7234	9016	YAL067C 1	-	39	2
chrI	7234	9016	YAL067C 1	-	40	2
chrI	7234	9016	YAL067C 1	_	41	3
chrT	7234	9016	YAL067C 1	_	42	3
ahrt	7237	001C	VAL0670 1	_	72 // C	с С
chri	7234	9010	IALUO/C I	-	43	3
chrl	7234	9016	YALU67C 1	-	44	3
chrI	7234	9016	YAL067C 1	-	45	4
chrI	7234	9016	YAL067C 1	-	46	4
chrI	7234	9016	YAL067C 1	-	47	4
chrI	7234	9016	YAL067C 1	-	48	4
chrI	7234	9016	YAL067C 1	-	49	4
chrI	7234	9016	YAL067C 1	_	50	4
						-

A proper **bedGraph** file has only 4 columns: *chrom start end value* and does not need to include positions with 0 reads, so we'll convert the **bedtools coverage** output to **bedGraph** using **awk**. We re-sort the output so that plus and minus strand positions are adjacent.

```
cat yeast_mrna.gene_coverage.txt | awk '
BEGIN{FS=OFS="\t"}
{if ($8>0) {print $1,$2-1+$7,$2+$7,$8}}' | \
   sort -kl,l -k2,2n -k3,3n > yeast_mrna.gene_coverage.almost.bedGraph
wc -l yeast_mrna.gene_coverage.almost.bedGraph # 5,710,186 -- better, but still big
```

While we probably could consider this file to have **bedGraph** format, it's preferable to combine adjacent per-base coordinates with the same count into larger regions, e.g.

Щ то ото 1	haaa aaumt		
# per-	base count	- 5	
chrI	7271	7272	2
chrI	7272	7273	2
chrI	7273	7274	2
chrI	7274	7275	3
chrI	7275	7276	3
chrI	7276	7277	3
chrI	7277	7278	3
# corr	esponding	region	counts
chrI	7271	7274	6
chrI	7274	7278	12

Here's some awk to do this:

```
cat yeast_mrna.gene_coverage.almost.bedGraph | awk '
BEGIN{FS=OFS="\t"; chr=""; start=-1; end=-1; count=0}
{if (chr != $1) { # new contig; finish previous
  if (count > 0) { print chr,start,end,count }
  chr=$1; start=$2; end=$3; count=$4
} else if (($2==end || $2==end+1) && ($4==count)) { # same or adjacent position with same count
  end=$3;
} else { # new region on same contig; finish prev
  if (count > 0) { print chr,start,end,count}
   start=$2; end=$3; count=$4
}
}
END{ # finish last
 if (count > 0) { print chr,start,end,count }
}' > yeast_mrna.gene_coverage.bedGraph
wc -l yeast_mrna.gene_coverage.bedGraph # 1,048,510 -- much better!
```

#### Make sure the total counts match!

cat yeast_mrna.gene_coverage.txt   awk '	
BEGIN{tot=0}{tot=tot+\$8}END{print tot}'	# should be 86703686
cat yeast_mrna.gene_coverage.almost.bed   awk '	
BEGIN{tot=0}{tot=tot+\$4}END{print tot}'	# should also be 86703686
cat yeast_mrna.gene_coverage.bedGraph   awk '	
$BEGIN{tot=0}{tot=tot+$4*($3-$2)}END{print tot}'$	# should also be 86703686

Now our yeast\_mrna.gene\_coverage.bedGraph file is a proper bedGraph, whose first lines look like this:

chrI 7250 7271 1 7274 2 chrI 7271 chrI 7274 7278 3 chrI 7278 7310 4 7317 chrI 7310 3 7317 7349 chrI 2 chrI 7349 7353 1 7500 7556 1 chrI chrI 8851 8891 1 chrI 11919 11951 1

х

# A brief introduction to bedtools

Now that we have a BAM file with only the reads we want included, we can do some more sophisticated analysis using **bedtools**. **Bedtools** changes from version to version, and here we are using version 2.22, the newest version, and what is currently installed on **stampede**. You can check what versions of **bedtools** are installed by using the following command on **stampede**:

module spider bedtools

First, log on to the login8 node on stampede and make a directory in scratch called bedtools in your scratch folder. Then copy your filtered BAM file from the samtools section into this folder.

```
ssh user@login8.stampede.tacc.utexas.edu #if you are not already logged in!
cd $SCRATCH/core_ngs
mkdir bedtools
cd samtools
cp yeast_pairedend_sort.mapped.ql.bam ../bedtools
cd ../bedtools
```

If you were unable to make the filtered and sorted BAM file from the previous section, you can copy it over from my scratch directory:

```
cd bedtools
cp /scratch/01786/awh394/core_ngs/bedtools/yeast_pairedend_sort.mapped.ql.bam .
```

# bedtools bamtofastq: converting a BAM file to a fastq file

Sometimes, especially when working with external data, we need to go from a BAM file back to a fastq file. This can be useful for re-aligning reads using a different aligner, different settings on the original aligner used. It can also be useful for extracting the sequence of interesting regions of the genome after you have manipulated your BAM file.

For this exercise, you'll be using bamtofastq. This function takes an aligned BAM file as input and outputs a fastq format file. You can use the options if you have paired end data to output R1 and R2 reads for your fastq file. This type of function is especially useful if you need to to analyze sequences after you've compared several BAM or bed files.

bedtools bamtofastq -i input.bam -fq output.fastq

#### Exercise 1: convert BAM to fastq and look at the quality scores

# solution code module load bedtools bedtools bamtofastq -i yeast\_pairedend\_sort.mapped.ql.bam -fq yeast\_pairedend\_sort.mapped.ql.fastq #takes 1-2 minutes more yeast\_pairedend\_sort.mapped.ql.fastq

Here is an example of two sequences (and their corresponding quality scores):

#### two lines of a fastq file

@HWI-ST1097:127:COW5VACXX:5:2212:10568:79659

TACCCTCCAATTACCCATATCCAACCCACTGCCACTTACCCTACCATCCACCATGACCTACTCACCATACTGTTCTTCTACCCACCATAT

TAGGGTAAGTTTGAGATGGTATATACCCTACCATCCACCATGACCTACTCACCATACTGTTCTTCTACCCACCATATTGAAACGCTAACAAATGATCGTAA

?B@DF2ADHFHHFHJIIIGCIHIGGIJJJJGHIIIGIJEHHIGGGAHEGGFGHIECGIJIIIJIIIIJJJJJJJE>EHDHEEEBCDD?CDDBDDDDDDCDB

As we discussed earlier, the top line is the identifier for the sequence produced, the second line defines which bases were produced, the third line indicates the strand the sequence is aligned to, and the fourth line indicates the ASCII based quality scores for each character in the second line.

# bedtools bamtobed: converting a BAM file into a bed file

While it's useful to be able to look at the fastq file, many analyses will be easiest to perform in bed format. Bed format is a simple tab delimited format that designates various properties about segments of the genome, defined by the chromosome, start coordinates and end coordinates. Bedtools provides a simple utility to convert BAM files over into bed files, termed bamtobed.

```
bedtools bamtobed -i input.bam > output.bed #output to a file bedtools bamtobed -i input.bam \mid more #output to more
```

Note that the output will be piped to standard out unless you redirect to a program (head, more, less) or a file (output.bed). Now we'll put this example to use and convert our filtered BAM file from the samtools section into a bed file.

#### Exercise 2: Convert the filtered yeast paired end BAM to bed using bamtobed, look at your file in more, and find the number of lines in the file

Hint: direct the output to a file first, then use more to look at the converted file visually; use ctrl+c to quit more.

solution code
<pre>module load bedtools #if you haven't loaded it in for this session bedtools bamtobed -i yeast_pairedend_sort.mapped.ql.bam &gt; yeast_pairedend_sort.mapped.ql.bed</pre>
more yeast_pairedend_sort.mapped.ql.bed #to examine the bed file visually wc -l yeast_pairedend_sort.mapped.ql.bed #to get the number of lines in a file

Here is what my output looks like:

#### output from the code above

wc -l yeast\_pairedend\_sort.mapped.ql.bed
528976 yeast\_pairedend\_sort.mapped.ql.bed

chrI219320HWI-ST1097:127:C0W5VACXX:5:2212:10568:79659/137+chrI266344HWI-ST1097:127:C0W5VACXX:5:2115:19940:13862/229+chrI368469HWI-ST1097:127:C0W5VACXX:5:2115:19940:13862/129-chrI684785HWI-ST1097:127:C0W5VACXX:5:2212:10568:79659/237-chrI871955HWI-ST1097:127:C0W5VACXX:5:1103:4918:43976/229+chrI871948HWI-ST1097:127:C0W5VACXX:5:1104:2027:42518/229+
chrI       266       344       HWI-ST1097:127:C0W5VACXX:5:2115:19940:13862/2       29       +         chrI       368       469       HWI-ST1097:127:C0W5VACXX:5:2115:19940:13862/1       29       -         chrI       684       785       HWI-ST1097:127:C0W5VACXX:5:2115:19940:13862/1       29       -         chrI       684       785       HWI-ST1097:127:C0W5VACXX:5:2212:10568:79659/2       37       -         chrI       871       955       HWI-ST1097:127:C0W5VACXX:5:1103:4918:43976/2       29       +         chrI       871       948       HWI-ST1097:127:C0W5VACXX:5:1104:2027:42518/2       29       +
chrI       368       469       HWI-ST1097:127:C0W5VACXX:5:2115:19940:13862/1       29       -         chrI       684       785       HWI-ST1097:127:C0W5VACXX:5:2212:10568:79659/2       37       -         chrI       871       955       HWI-ST1097:127:C0W5VACXX:5:1103:4918:43976/2       29       +         chrI       871       948       HWI-ST1097:127:C0W5VACXX:5:1104:2027:42518/2       29       +
chrI       684       785       HWI-ST1097:127:C0W5VACXX:5:2212:10568:79659/2       37       -         chrI       871       955       HWI-ST1097:127:C0W5VACXX:5:1103:4918:43976/2       29       +         chrI       871       948       HWI-ST1097:127:C0W5VACXX:5:1104:2027:42518/2       29       +
chri         871         955         HWI-ST1097:127:C0W5VACXX:5:1103:4918:43976/2         29         +           chri         871         948         HWI-ST1097:127:C0W5VACXX:5:1104:2027:42518/2         29         +
chrI 871 948 HWI-ST1097:127:COW5VACXX:5:1104:2027:42518/2 29 +
chrI 871 948 HWI-ST1097:127:C0W5VACXX:5:1109:3153:38695/2 29 +
chrI 871 948 HWI-ST1097:127:C0W5VACXX:5:2109:6222:11815/2 29 +
chrI 871 948 HWI-ST1097:127:C0W5VACXX:5:2113:5002:59471/2 29 +
chrI 871 948 HWI-ST1097:127:COW5VACXX:5:2113:7803:87146/2 29 +
chrI 971 1072 HWI-ST1097:127:C0W5VACXX:5:1103:4918:43976/1 29 -
chrI 978 1079 HWI-ST1097:127:C0W5VACXX:5:1104:2027:42518/1 29 -
chrI 978 1079 HWI-ST1097:127:C0W5VACXX:5:1109:3153:38695/1 29 -
chrI 978 1079 HWI-ST1097:127:C0W5VACXX:5:2109:6222:11815/1 29 -
chrI 978 1079 HWI-ST1097:127:C0W5VACXX:5:2113:5002:59471/1 29 -
chrI 978 1079 HWI-ST1097:127:COW5VACXX:5:2113:7803:87146/1 29 -
chrI 978 1079 HWI-ST1097:127:COW5VACXX:5:2203:1231:50183/1 37 -

Note the "stacks" of reads that are occurring on similar coordinates on the same strand of the genome. We'll deal with those in the bedtools merge section.

See also: bedtools bedtobam, if you need to get back to a bam file from a bed file (some programs take bam files as input). Documentation here.

# bedtools coverage: how much of the genome does my data cover?

One way of characterizing data is to understand what percentage of the genome your data covers. What type of experiment you performed should affect the coverage of your data. A ChIP-seq experiment will cover binding sites, and a RNA-seq experiment will cover expressed transcripts. **Bedtools** coverage allows you to compare one bed file to another and compute the breadth and depth of coverage.

bedtools coverage -a experiment.bed -b reference\_file.bed

The resulting output will contain several additional columns which summarize this information:

After each interval in B, coverageBed will report:

- 1. The number of features in A that overlapped (by at least one base pair) the B interval.
- 2. The number of bases in B that had non-zero coverage from features in A.
- 3. The length of the entry in B.
- 4. The fraction of bases in B that had non-zero coverage from features in A.

For this exercise, we'll use a bed file that summarizes the S. cerevisiae genome, version 3 (aka sacCer3). For this class, I've made a bed file out of the genome, using the file sacCer3.chrom.sizes. First go and copy the file over from my scratch directory:

cd bedtools #if you aren't already there cp /scratch/01786/awh394/core\_ngs.test/bedtools/sacCer3.chrom.sizes.bed .

more sac	Cer3.	chrom.sizes.bed
chrIV	1	1531933
chrXV	1	1091291
chrVII	1	1090940
chrXII	1	1078177
chrXVI	1	948066
chrXIII	1	924431
chrII	1	813184
chrXIV	1	784333
chrX	1	745751
chrXI	1	666816
chrV	1	576874
chrVIII	1	562643
chrIX	1	439888
chrIII	1	316620
chrVI	1	270161
chrI	1	230218
chrM	1	85779

The format is bed3 - just chrom, start (which is always 1) and stop, which is always the length of the chromosome, for this type of bed file.

Now use bedtools coverage to find the coverage of the file output.bed over the sacCer3 genome and examine the output coverage.

Exercise 3: Find the coverage of your bed file over the sacCer3 genome

#### solution code

module load bedtools #again, if not already loaded bedtools coverage -a sacCer3.chrom.sizes.bed -b yeast\_pairedend\_sort.mapped.ql.bed > sacCer3coverage.bed more sacCer3coverage.bed #this file should have 17 lines, one for each chromosome

And here is what my output looks like:

more sacCer3coverage.bed								
chrIV	1	1531933	70633	1026387	1531932	0.6699951		
chrXV	1	1091291	47871	710376	1091290	0.6509507		
chrVII	1	1090940	49762	722821	1090939	0.6625677		
chrXII	1	1078177	48155	658373	1078176	0.6106359		
chrXVI	1	948066	43531	612122	948065	0.6456540		
chrXIII	1	924431	40054	618798	924430	0.6693833		
chrII	1	813184	35818	539222	813183	0.6631004		
chrXIV	1	784333	32565	513382	784332	0.6545468		
chrX	1	745751	30743	472357	745750	0.6333986		
chrXI	1	666816	27950	446567	666815	0.6697015		
chrV	1	576874	26918	381078	576873	0.6605926		
chrVIII	1	562643	23424	356421	562642	0.6334774		
chrIX	1	439888	15953	276571	439887	0.6287319		
chrIII	1	316620	13701	199553	316619	0.6302623		
chrVI	1	270161	10662	167222	270160	0.6189740		
chrI	1	230218	7972	128701	230217	0.5590421		
chrM	1	85779	3264	58599	85778	0.6831472		

It's worth noting that just computing coverage over the genome isn't the most useful thing, but you might compute coverage over a set of genes or regions of interest. Coverage is really useful coupled with intersect or subtract as well.

# bedtools merge: collapsing bookended elements (or elements within a certain distance)

When we originally examined the bed files produced from our BAM file, we can see many reads that overlap over the same interval. While this level of detail is useful, for some analyses, we can collapse each read into a single line, and indicate how many reads occurred over that genomic interval. We can accomplish this using **bedtools** merge.

bedtools merge [OPTIONS] -i experiment.bed > experiment.merge.bed

Bedtools merge also directs the output to standard out, to make sure to point the output to a file or a program. While we haven't discussed the options for each bedtools function in detail, here they are very important. Many of the options define what to do with each column (-c) of the output (-o). This defines what type of operation to perform on each column, and in what order to output the columns. Standard bed6 format is chrom, start, stop, name, score, strand and controlling column operations allows you to control what to put into each column of output. The valid operations defined by the -o operation are as follows:

- sum, min, max, absmin, absmax,
- mean, median,
- collapse (i.e., print a delimited list (duplicates allowed)),
- distinct (i.e., print a delimited list (NO duplicates allowed)),
- count
- count\_distinct (i.e., a count of the unique values in the column)

For this exercise, we'll be summing the number of reads over a region to get a score column, using distinct to choose a name, and using distinct again to keep track of the strand. For the -c options, define which columns to operate on, in the order you want the output. In this case, to keep the standard bed format, we'll list as -c 5,4,6 and -o count\_distinct,sum,distinct, to keep the proper order of name, score, strand. Strandedness can also be controlled with **m** erge, using the -s option.

#### Exercise 4: Use bedtools merge to merge an experiment, look at the output and see how many lines there are in the file.

Hint: make sure to remove whitespace between lists for the -c and -o options!

#### solution code

```
bedtools merge -s -c 4,5 -o count_distinct,sum -i yeast_pairedend_sort.mapped.ql.bed > yeast_pairedend_sort.
mapped.ql.merge.bed
more yeast_pairedend_sort.mapped.ql.merge.bed
wc -l yeast_pairedend_sort.mapped.ql.merge.bed
#without strand considered
bedtools merge -c 4,5,6 -o count_distinct,sum,distinct -i yeast_pairedend_sort.mapped.ql.bed >
yeast_pairedend_sort.noStrand.mapped.ql.merge.bed
```

wc -l yeast_pairedend_sort.noStrand.mapped.ql.merge.bed										
40	40319 yeast_pairedend_sort.noStrand.mapped.ql.merge.bed #without the -s option									
wc -l yeast_pairedend_sort.mapped.ql.merge.bed										
76	76601 yeast_pairedend_sort.mapped.ql.merge.bed #with the -s option									
mo	ore ye	ast_pair	edend_sc	ort.map	pped.q	1.merge.bed				
cł	ırI	219	344	+	2	66				
cł	ırI	368	469	-	1	29				
cł	ırI	684	785	-	1	37				
cł	ırI	871	955	+	6	174				
cł	ırI	971	1079	-	7	211				
cł	ırI	1216	1322	+	6	157				
cł	ırI	1347	1437	-	6	157				
cł	ırI	2892	2993	+	14	406				
cł	ırI	3010	3111	+	1	37				
cł	ırI	3013	3107	-	14	406				
mo	ore ye	ast_pair	edend_sc	ort.nos	Strand	l.mapped.ql.merge.bed				
cł	ırI	219	344	2	66	+				
cł	ırI	368	469	1	29	-				
cł	ırI	684	785	1	37	-				
cł	ırI	871	955	6	174	+				
cł	ırI	971	1079	7	211	-				
cł	ırI	1216	1322	6	157	+				
cł	ırI	1347	1437	6	157	-				
cł	ırI	2892	2993	14	406	+				
cł	ırI	3010	3111	15	443	+,-				

Note the change in column order in the first set of commands. We can use awk like this to change the column order, either piped in the original command or after the fact:

### using awk for column reordering

```
#after the creation of the first file
cat yeast_pairedend_sort.mapped.ql.merge.bed | awk '{print $1 "\t" $2 "\t" $3 "\t" $5 "\t" $6 "\t" $4}' >
yeast_pairedend_sort.mapped.ql.merge.reorder.bed
```

#piped in-line bedtools merge -s -c 4,5 -o count\_distinct,sum -i yeast\_pairedend\_sort.mapped.ql.bed | awk '{print \$1 "\t" \$2 "\t" \$3 "\t" \$5 "\t" \$6 "\t" \$4}' > yeast\_pairedend\_sort.mapped.ql.merge.bed

# bedtools intersect: identifying where two experiments overlap (or don't overlap)

One useful way to compare two experiments (especially biological replicates, or similar experiments in two yeast strains/cell lines/mouse strains) is to compare where reads in one experiment overlap with reads in another experiment. Bedtools offers a simple way to do this using the intersect function.

bedtools intersect options	
bedtools intersect [OPTIONS] -a <file> \</file>	-b <file1, file2,,="" filen=""></file1,>

The intersect function has many options that control how to report the intersection. We'll be focusing on just a few of these options, listed below.

-a and -b indicate what files to intersect. in -b, you can specify one, or several files to intersect with the file specified in -a.

- wa: Write the original entry in A for each overlap.
- wb: Write the original entry in B for each overlap. Useful for knowing what A overlaps. Restricted by -f and -r.
- loj: Perform a "left outer join". That is, for each feature in A report each overlap with B. If no overlaps are found, report a NULL feature for B.
  wo: Write the original A and B entries plus the number of base pairs of overlap between the two features. Only A features with overlap are reported. Restricted by -f and -r.
- wao: Write the original A and B entries plus the number of base pairs of overlap between the two features. However, A features w/o overlap are also reported with a NULL B feature and overlap = 0. Restricted by -f and -r.
- f: Minimum overlap required as a fraction of A. Default is 1E-9 (i.e. 1bp).

- v: Only report those entries in A that have no overlap in B. Restricted by -f and -r. Useful to report what doesn't overlap, the inverse of typical usage.
- names: When using multiple databases (-b), provide an alias for each that will appear instead of a file Id when also printing the DB record.

In this section, we'll intersect two human experiments - one from sequencing RNA, and one from sequencing micro RNA. Copy these files over to your directory:

#### copy some files over to intersect

```
cd $SCRATCH/core_ngs/
mkdir intersect
cd intersect
cp /corral-repl/utexas/BioITeam/core_ngs_tools/alignment/bam/human_mirnaseq_hg19.bam .
cp /corral-repl/utexas/BioITeam/core_ngs_tools/alignment/bam/human_rnaseq_bwa.bam .
ls -lah
```

```
-rwxrwxr-x 1 awh394 G-801021 19M May 22 18:57 human_mirnaseq_hg19.bam
-rwxrwxr-x 1 awh394 G-801021 6.6M May 22 18:57 human_rnaseq_bwa.bam
```

Before we can intersect these files, we need to perform the pipeline we used in **samtools** to **index**, **sort** and **filter** the files, and **bedtools** to convert from BAM over to bed, then collapse down the files using **merge**. Below is a little workflow to help you through it on the files you just copied above.

My output (for length of bed files) is in the comments.

```
a samtools/bedtools workflow
module load samtools #if you haven't loaded it up this session
#sort both files
samtools sort human_mirnaseq_hq19.bam human_mirnaseq_hq19_sort # will take 1-2 minutes
samtools sort human_rnaseq_bwa.bam human_rnaseq_bwa_sort # will take 1-2 minutes
#index the new files
samtools index human_mirnaseq_hg19_sort.bam
samtools index human_rnaseq_bwa_sort.bam
#filter the sorted files, reindex the new filtered files
samtools view -b -F 0x04 -q 1 -o human_mirnaseq_hg19_sort.mapped.q1.bam human_mirnaseq_hg19_sort.bam
samtools view -b -F 0x04 -q 1 -o human_rnaseq_bwa_sort.mapped.ql.bam human_rnaseq_bwa_sort.bam
samtools index human_mirnaseq_hg19_sort.mapped.q1.bam
samtools index human_rnaseq_bwa_sort.mapped.ql.bam
#convert filtered bam files to bed format
module load bedtools #if you haven't loaded it in for this session
bedtools bamtobed -i human_mirnaseq_hg19_sort.mapped.q1.bam > human_mirnaseq_hg19_sort.mapped.q1.bed
bedtools bamtobed -i human_rnaseq_bwa_sort.mapped.ql.bam > human_rnaseq_bwa_sort.mapped.ql.bed
#check the length of the files:
wc -l *.bed
# 164806 human_mirnaseq_hg19_sort.mapped.q1.bed
   22538 human_rnaseq_bwa_sort.mapped.ql.bed
#merge the bed files, check the length again
bedtools merge -s -c 4,5 -o count_distinct,sum -i human_mirnaseq_hg19_sort.mapped.ql.bed | awk '{print $1 "\t"
$2 "\t" $3 "\t" $5 "\t" $6 "\t" $4}' > human_mirnaseq_hq19_sort.mapped.q1.merge.bed
bedtools merge -s -c 4,5 -o count_distinct,sum -i human_rnaseq_bwa_sort.mapped.ql.bed | awk '{print $1 "\t" $2
"\t" $3 "\t" $5 "\t" $6 "\t" $4}' > human_rnaseq_bwa_sort.mapped.q1.merge.bed
wc -l *.merge.bed
# 14794 human_mirnaseq_hg19_sort.mapped.ql.merge.bed
# 7134 human_rnaseq_bwa_sort.mapped.ql.merge.bed
```

If we run low on time, you can copy the merged bed files over from my directory on scratch:

cds

cd intersect

cp /scratch/01786/awh394/core\_ngs/intersect/human\_mirnaseq\_hg19\_sort.mapped.ql.merge.bed .

 $\texttt{cp /scratch/01786/awh394/core_ngs/intersect/human\_rnaseq\_bwa\_sort.mapped.ql.merge.bed .}$ 

#### Exercise 5: Intersect two experiments using intersect and examine the output

My output is commented in this code block.

cd inter module i bedtools bed > he	rsect load bedtool: s intersect g19_rnaseq_m	s #if you ha -wo -a human irnaseq_inte	ven't _rnas rsect	loade eq_bwa .bed	d it ı _sort	up yet t .mapped.	his session ql.merge.bed	-b human_mi	rnase	q_hg19	_sort	.mapped.ql.merge.
wc -l hg #38 hg19	g19_rnaseq_m 9_rnaseq_mirn	irnaseq_inte naseq_inters	rsect ect.b	.bed ed								
more hg	19_rnaseq_mi:	rnaseq_inter	sect.	bed								
#chr1	20987370	20987471	1	37	-	chr1	20987402	20987430	1	12	-	28
#chr1	25555557	25555616	1	37	+	chr1	25555612	25555636	1	2	-	4
#chr1	25555557	25555617	1	37	-	chr1	25555612	25555636	1	2	-	5
#chr1	28906396	28906497	1	37	+	chr1	28906368	28906405	6	246	-	9
#chr1	33245783	33245884	1	37	+	chr1	33245880	33245908	1	24	-	4

Using the options we've specified (-wo) the resulting file will have entries for file A, file B and the number of base pairs overlap between the feature in A and the features in B, but **we'll only retain lines where there is an overlap between A and B**. We could also use the -v option to only contain areas with NO intersection, or control the intersections with -f and -r options. Bedtools intersect is a powerful tool, and it's always a good idea to ask "what is this code going to do?" while you're testing analysis workflows. It can be very useful to pipe your output to more when you are unsure of the output of a command, as such:

#### pipe-ing output to more

bedtools intersect -wo -a human\_rnaseq\_bwa\_sort.mapped.ql.merge.bed -b human\_mirnaseq\_hg19\_sort.mapped.ql.merge. bed | more

## bedtools closest: when you want to know how far your regions are from a test set

The manual page for **bedtools closest** has a really nice image of how closest behaves with overlapping options. Bedtools closest first looks for any overlaps of B with A, if it finds an overlap, the overlap in B with the highest proportional overlap with A is reported. If there are no overlaps, then it looks for the closest genomic feature proximal to A (using distance from the start or end of A to do this).

#### bedtools intersect options

bedtools closest [OPTIONS] -a <FILE> \
 -b <FILE1, FILE2, ..., FILEN>

Much like **bedtools intersect**, **bedtools closest** takes an A file and a series of B files. So if you wanted to determine the distance of your regions of interest to several different classes of genes, **bedtools closest** would be a useful tool for that analysis.

- s: Require same strandedness. That is, find the closest feature in B that overlaps A on the \_same\_ strand. By default, overlaps are reported without respect to strand.
- S: Require opposite strandedness. That is, find the closest feature B that overlaps A on the \_opposite\_ strand. By default, overlaps are reported without respect to strand.
- d: In addition to the closest feature in B, report its distance to A as an extra column. The reported distance for overlapping features will be 0.
- D: Like -d, report the closest feature in B, and its distance to A as an extra column. However unlike -d, use negative distances to report upstream features.
  - ref Report distance with respect to the reference genome. B features with a lower (start, stop) are upstream.
  - a Report distance with respect to A. When A is on the strand, "upstream" means B has a higher (start, stop).
  - b Report distance with respect to B. When B is on the strand, "upstream" means A has a higher (start, stop).
- io: Ignore features in B that overlap A. That is, we want close, yet not touching features only.
- iu: Ignore features in B that are upstream of features in A. This option requires -D and follows its orientation rules for determining what is "upstream".
- id: Ignore features in B that are downstream of features in A. This option requires -D and follows its orientation rules for determining what is "downstream"
- names: When using multiple databases (-b), provide an alias for each that will appear instead of a file Id when also printing the DB record.

In this section, we'll intersect the human\_rnaseq\_bwa\_sort.mapped.q1.merge.bed file with some protein coding genes from Gencode (hg19). First go copy a couple files from my scratch directory:

copy some gencode files over	
<pre>cd \$SCRATCH/core_ngs mkdir closest cd closest cd closest cp /scratch/01786/awh394/core_ngs.test/closest/gencode.v19.proteincoding.genes.sort.merge.final . cp/intersect/human_rnaseq_bwa_sort.mapped.ql.merge.bed . #or: cp /scratch/01786/awh394/core_ngs/closest/human_rnaseq_bwa_sort.mapped.ql.merge.bed .</pre>	

-rwxrwxr-x 1 awh394 G-801021 646K May 22 20:41 gencode.v19.proteincoding.genes.sort.merge.final

Exercise 6: Identify the closest protein coding genes (on the same strand) for the human rnaseq file using closest, then sort by the distance column (largest, then smallest distance first).

My output is commented in this code block.

<pre>cd closest module load bedtools #if you haven't loaded it up yet this session sort -k1,1 -k2,2n human_rnaseq_bwa_sort.mapped.ql.merge.bed &gt; human_rnaseq_bwa.mapped.ql.merge.sort.bed #need to sort both files to the same order bedtools closest -s -d -a human_rnaseq_bwa.mapped.ql.merge.sort.bed -b gencode.v19.proteincoding.genes.sort. merge.final &gt; hg19_rnaseq_protcode_closest.bed</pre>					
wc -l hg19_rnaseq_protcode_closest.bed					
#7134 hg19_rnaseq_protcode_closest.bed #same length as the original file					
more hg19_rnaseq_protcode_closest.bed					
#chr1 880458 880529 1 37 + chr1 860260 879955 SAMD11 . + 504					
#chrl 881549 881650 1 37 - chrl 879584 894689 NOC2L 0					
#chrl 887884 887985 1 37 + chrl 860260 879955 SAMD11 . + 7930					
#chrl 892309 892410 1 37 - chrl 879584 894689 NOC2L 0					
#chr1 892475 892576 1 23 + chr1 895967 901095 KLHL17 . + 3392					
<pre>#sort by the distance to a gene, longest distances first sort -k13,13nr hg19_rnaseq_protcode_closest.bed   more #sort by the distance to a gene, shortest distances first sort -k13,13n hg19_rnaseq_protcode_closest.bed   more</pre>					

This is a nice way to examine your reads over annotated protein-coding genes. Note the strand specificity - only reads on the correct strand will be reported when there is a + strand gene and a - strand gene over the same location.

# bedtools subtract: removing features from your bed file

Bedtools subtract takes an A file and a B file, then searches for features in B that overlap A. When/if these features are identified, the overlapping portion is removed from A and the remaining portion of A is reported. If a feature in B overlaps all of a feature in A, that feature will not be reported.

bedtools subtract options

bedtools subtract [OPTIONS] -a <BED/GFF/VCF> -b <BED/GFF/VCF>

Note that **bedtools subtract** is performed on two files, and unlike some of the other utilities we've used, you can't use multiple B features here. However, you can use **cat** to join together features you'd like to subtract from your A file.

- f: Minimum overlap required as a fraction of A. Default is 1E-9 (i.e. 1bp).
- F: Minimum overlap required as a fraction of B. Default is 1E-9 (i.e., 1bp).
- r: Require that the fraction of overlap be reciprocal for A and B. In other words, if -f is 0.90 and -r is used, this requires that B overlap at least 90% of A and that A also overlaps at least 90% of B.

- e: Require that the minimum fraction be satisfied for A \_OR\_ B. In other words, if -e is used with -f 0.90 and -F 0.10 this requires that either 90% of A is covered OR 10% of B is covered. Without -e, both fractions would have to be satisfied.\*\*-s\*\* Force "strandedness". That is, only report hits in B that overlap A on the same strand. By default, overlaps are reported without respect to strand.
- S: Require different strandedness. That is, only report hits in B that overlap A on the \_opposite\_ strand. By default, overlaps are reported without respect to strand
- A: Remove entire feature if any overlap. That is, by default, only subtract the portion of A that overlaps B. Here, if any overlap is found (or -f amount), the entire feature is removed.
- N: Same as -A except when used with -f, the amount is the sum of all features (not any single feature)

Let's do a little set-up for the next exercise:

#### copy some gencode files over

```
cd $SCRATCH/core_ngs
mkdir subtract
cd subtract
cp /scratch/01786/awh394/core_ngs.test/closest/gencode.v19.proteincoding.genes.sort.merge.final .
cp /scratch/01786/awh394/core_ngs.test/closest/gencode.v19.genes.sort.merge.final .
```

# Exercise 7: remove the protein-coding genes from a gencode list of genes using subtract, then give a count of the non-protein-coding gene entries

This allows you to identify which gene regions are not protein coding, and are likely pseudogenes, but could also be miRNAs, snRNAs or other genes that aren't translated into a peptide sequence.

My output is commented in this code block.

```
cd subtract
module load bedtools #if you haven't loaded it up yet this session
bedtools subtract -a gencode.v19.genes.sort.merge.final -b gencode.v19.proteincoding.genes.sort.merge.final >
gencode.v19.not.proteincoding.genes.bed
wc -l gencode.v19.not.proteincoding.genes.bed
#23483 gencode.v19.not.proteincoding.genes.bed
more gencode.v19.not.proteincoding.genes.bed
#chr1 11869 14412 DDX11L1
                                 .
                                        +
        14363
                29806
                        WASH7P
#chr1
                31109
        29554
                        MIR1302-11
#chr1
        34554 36081
#chr1
                        FAM138A
        52473 54936
                        OR4G4P
#chr1
#chr1
        62948
                63887
                         OR4G11P
```

While the above example is not super useful in all cases, one might use the above workflow to remove genes that aren't of interest from a larger set.

# A little bit of filtering, using awk

As a final note, yesterday we taught you about using a lot of unix utilities, including **uniq**, **sort** and **cut**. One last utility I'd like to add, that is very useful for manipulating these types of tab delimited files, is **awk**. **Awk** isn't a command, but rather a little text manipulation language in it's own right (which we briefly used above to rearrange the columns in a file). While **awk** can be used to do many different things, here we'll primarily use it to sort tab delimited files based on the values present in those files. That is useful to filter your files for entries on a given chromosome, or greater than/less than a given score. If your dataset is large, this type of filtering can be invaluable! Below is an example of a simple **awk** script:

```
a simple awk script
cat file.bed | awk 'BEGIN{FS="\t";}{if ($6 == '+'){print}}' > file.plusStrand.bed
```

- 1. In the first section, we open the bed file of interest. Then we pipe that filestream to the awk program.
- 2. The section: BEGIN{FS="\t";OFS="\t";} tells awk to begin a filter, the input file is tab delimited, and the output file is also tab delimited.
- a. Generally, you can leave this section constant (if you are working with tab delimited files).
- 3. The second section: {if (\$6 == +){print}} is our selection and printing criteria.
- a. "\$6" indicates column 6, and == indicates "equals" or "matches".
- 4. The {print} command tells awk to print the whole line if the statement for column 6 evaluates to true.
- 5. Thus, the output file only contains those lines which satisfy the criteria in the selection statement.

We can do this filtering on the hg19\_rnaseq\_mirnaseq\_intersect.bed file we just created using bedtools intersect.

# You could also insist on columns 6 and 12 both being the plus strand as such:

```
cd $SCRATCH/core_ngs/intersect/
cat hg19_rnaseq_mirnaseq_intersect.bed | awk 'BEGIN{FS="\t";OFS="\t";}{if ($6 == "+" && $12 == "+"){print}}' |
more
```