# **Analysis using BEDTools**



#### 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
```

- The BED format
- BEDTools overview
  - BEDTools versions
  - Input format considerations
  - About strandedness
- About GFF/GTF annotation files
  - Filter annotations based on desired feature type
  - Convert GFF/GTF format to BED with ID in the name field
- Exercises
  - Use bedtools merge to collapse overlapping annotations
  - Use bedtools multicov to count feature overlaps
  - Use bedtools genomecov to create a signal track

## The BED format

**BED** (Browser Extensible Data) format is a simple *text format for location-oriented data* (genomic regions) developed to support UCSC Genome Browser tracks. Standard **BED** files have 3 to 6 Tab-separated columns, although up to 12 columns are defined. (Read more about the UCSC Genome Browser's official BED format.)



#### Memorize the 6 main BED fields

These 6 BED fields are so important that you should memorize them. Keep repeating "chrom, start, end, name, score, strand" until the words trip off your tongue :

- 1. chrom (required) string naming the chromosome or other contig
- 2. start (required) the 0-based start position of the region
- 3. **end** (required) the **1-based** end position of the region
- 4. name (optional) an arbitrary string describing the region
  - for BED files loaded as UCSC Genome Browser tracks, this text is displayed above the region
- 5. score (optional) an integer score for the region
  - for BED files to be loaded as UCSC Genome Browser tracks, this should be a number between 0 and 1000, higher = "better"
  - for non-GenBrowse BED files, this can be any integer value (e.g. the length of the region)
- 6. strand (optional) a single character describing the region's strand
  - + plus strand (Watson strand) region
  - -- minus strand (Crick strand) region
  - . no strand the region is not associated with a strand (e.g. a transcription factor binding region)

# Important rules for BED format:

- The number of fields per line must be consistent throughout any single BED file
  - o e.g. they must all have 3 fields or all have 6 fields
- The first base on a contig is numbered 0
  - o versus 1 for BAM file positions
  - o so the a **BED** start of 99 is actually the 100th base on the contig
  - but *end* positions are 1-based
    - so a **BED** end of 200 is the 200th base on the contig
  - o the length of a BED region is end start
    - not end start + 1, as it would be if both coordinates with 0-based or both 1-based
  - o this difference is the single greatest source of errors dealing with **BED** files!

Note that the UCSC Genome Browser also defines many BED-like data formats (e.g. **bedGraph**, **narrowPeak**, **tagAlign** and various RNA element formats). See supported UCSC Genome Browser data formats for more information and examples.

In addition to standard-format **BED** files, one can create *custom* **BED** files that have at least 3 of the standard fields (*chrom*, *start*, *end*), followed by any number of custom fields. For example:

- A BED3+ file contains the 3 required BED fields, followed by some number of user-defined columns (all records with the same number)
- A BED6+ file contains the 3 required BED fields, 3 additional standard BED fields (name, score, strand), followed by some number of user-defined columns

As we will see, BEDTools functions require BED3+ input files, or BED6+ if strand-specific operations are requested.

# **BEDTools** overview

The **BEDTools** suite is a set of utilities for manipulating **BED** and **BAM** files. We call it the "Swiss army knife" for genomic region analyses because its subcommands are so numerous and versatile. Some of the most common **bedtools** operations perform set-theory functions on regions: intersection (**intersect**), union (**merge**), set difference (**subtract**) – but there are many others. The table below lists some of the most useful sub-commands along with applicable use cases.

Sub- command	Description	Use case(s)
bamtobed	Convert <b>BAM</b> files to <b>BED</b> format.	You want to have the <i>contig</i> , <i>start</i> , <i>end</i> , and <i>strand</i> information for each mapped alignment record in separate fields. Recall that the strand is encoded in a <b>BAM</b> flag (0x10) and the exact end coordinate requires parsing the CIGAR string.
bamtofastq	Extract <b>FASTQ</b> sequences from <b>BAM</b> alignment records.	You have downloaded a <b>BAM</b> file from a public database, but it was not aligned against the reference version you want to use (e.g. it is <b>hg19</b> and you want an <b>hg38</b> alignment). To re-process, you need to start with the original <b>FASTQ</b> sequences.
getfasta	Get <b>FASTA</b> entries corresponding to regions.	You want to run <i>motif analysis</i> , which requires the original <b>FASTA</b> sequences, on a set of regions of interest. In addition to the <b>BAM</b> file, you must provide <b>FASTA</b> file(s) for the genome/reference used for alignment (e.g. the <b>FASTA</b> file used to build the aligner index).
genomecov	Generate <i>per-base</i> genome-wide <i>signal</i> tr ace	<ul> <li>Produce a per-base genome-wide signal (in bedGraph format), for example for a ChIP-seq or ATAC-seq experiment. After conversion to binary bigWig format, such tracks can be visualized in the Broad's IGV (Integrative Genome Browser) application, or configured in the UCSC Genome Browser as custom tracks.</li> </ul>
coverage	Compute <i>coverage</i> of your regions	<ul> <li>You have performed a WGS (whole genome sequencing) experiment and want to know if has resulted in the desired coverage depth.</li> <li>Calculate what proportion of the (known) transcriptome is covered by your RNA-seq alignments. Provide the transcript regions as a BED or GFF/GTF file.</li> </ul>
multicov	Count <b>overlaps</b> between one or more <b>BAM</b> files and a set of regions of interest.	<ul> <li>Count RNA-seq alignments that overlap a set of genes of interest. While this task is usually done with a specialized RNA-seq quantification tool (e.g. featureCounts or HTSeq), bedtools multicov can provide a quick estimate, e.g. for QC purposes.</li> </ul>
merge	Combine a set of possibly- overlapping regions into a single set of non- overlapping regions.	Collapse overlapping gene annotations into per-strand non-overlapping regions before counting (e.g with featureCounts or HTSeq). If this is not done, the source regions will potentially be counted multiple times, once for each (overlapping) target region it intersects.
subtract	Remove unwanted regions.	Remove rRNA gene regions from a merged gene annotations file before counting.
intersect	Determine the <b>overlap</b> betw een two sets of regions.	Similar to <b>multicov</b> , but can also report the overlapping regions, not just count them.
closest	Find the genomic features nearest to a set of regions.	For a set of significant ChIP-seq transcription factor (TF) binding regions ("peaks") that have been identified, determine nearby genes that may be targets of TF regulation.

We will explore a few of these functions in our exercises.

# **BEDTools versions**

**BEDTools** is under active development and is always being refined and extended. Unfortunately, sometimes changes are made that are incompatible with previous **BEDTools** versions. For example, a major change to the way **bedtool merge** functions was made after **bedtools v2.17.0**.

So it is important to know which version of **BEDTools** you are using, and read the documentation carefully to see if changes have been made since your version.

Login to Is6, start and idev session, then load the BioContainers bedtools module, and check its version.

#### Start an idev session

```
idev -m 120 -N 1 -A OTH21164 -r CoreNGS-Fri
# or
idev -m 90 -N 1 -A OTH21164 -p development

module load biocontainers
module load bedtools
bedtools --version # should be bedtools v2.27.1
```

# Input format considerations

- Most BEDTools functions now accept either BAM or BED files as input.
  - BED format files must be BED3+, or BED6+ if strand-specific operations are requested.
- When comparing against a set of regions, those regions are usually supplied in either BED or GTF/GFF.
- All text-format input files (BED, GTF/GFF, VCF) should use Unix line endings (linefeed only).

The most important thing to remember about comparing regions using **BEDTools**, is that *all input files must share the same set of contig names* and be based on the same reference! For example, if an alignment was performed against a human **GRCh38** reference genome from **Gencode**, use annotations from the corresponding **GFF/GTF** annotations.

#### **About strandedness**

By default many **bedtools** utilities that perform overlapping, consider reads overlapping the feature on **either strand**, but can be made **strand-specific** with the **-s** or **-S** option. This **strandedness** options for **bedtools** utilities refers the orientation of the **R1** read with respect to the feature's (gene's) strand.

- -s says the R1 read is sense stranded (on the same strand as the gene).
- -S says the R1 read is antisense stranded (the opposite strand as the gene).

RNA-seq libraries can be constructed with 3 types of strandedness:

- 1. sense stranded the R1 read should be on the same strand as the gene.
- 2. antisense stranded the R1 read should be on the opposite strand as the gene.
- 3. unstranded the R1 could be on either strand

Which type of RNA-seq library you have depends on the library preparation method – so ask your sequencing center! Our yeast RNA-seq library is **sense stranded** (note that most RNA-seq libraries prepared by GSAF are **antisense stranded**).

If you have a stranded RNA-seq library, you should use either -s or -S to avoid false counting against a gene on the wrong strand.

#### About GFF/GTF annotation files

Annotation files that you retrieve from public databases are often in GTF (Gene Transfer Format) or one of the in GFF (General Feature Format) formats (usually GFF3 these days).

Unfortunately, both formats are obscure and hard to work with directly. While **bedtools** does accept annotation files in **GFF/GTF** format, you will not like the results. This is because the most useful information in a **GFF/GTF** file is in a loosely-structured *attributes* field.

Also unfortunately, there are a number of variations of both annotation formats However both GTF and GFF share the first 8 Tab-separated fields:

- 1. segname The name of the chromosome or contig.
- 2. source Name of the program that generated this feature, or other data source (e.g. database)
- 3. feature\_type Type of the feature, for example:
  - CDS (coding sequence), exon
  - gene, transcript
  - start\_codon, stop\_codon
- 4. start Start position of the feature, with sequence numbering starting at 1.
- 5. end End position of the feature, with sequence numbering starting at 1.
- 6. score A numeric value. Often but not always an integer.
- 7. strand Defined as + (forward), (reverse), or . (no relevant strand)
- 8. frame For a CDS, one of 0, 1 or 2, specifying the reading frame of the first base; otherwise '.'

The **Tab**-separated columns will care about are (1) **seqname**, (3) **feature\_type** and (4,5) **start**, **end**. The reason we care is that when working with annotations, we usually only want to look at annotations of a particular type, most commonly **gene**, but also **transcript** or **exon**.

So where is the real annotation information, such as the unique gene ID or gene name? Both formats also have a **9th** field, which is usually populated by a set of *name/value* pair *attributes*, and that's where the useful information is (e.g. the unique feature identifier, name, and so forth).

Take a quick look at a yeast annotation file, sacCer\_R64-1-1\_20110208.gff using less.

# Start an idev session idev -m 120 -N 1 -A OTH21164 -r CoreNGS-Fri # or idev -m 90 -N 1 -A OTH21164 -p development module load biocontainers

```
Look at GFF annotation entries with less
```

bedtools --version # should be bedtools v2.27.1

module load bedtools

```
mkdir -p $SCRATCH/core_ngs/bedtools
cd $SCRATCH/core_ngs/bedtools
cp $CORENGS/yeast_rnaseq/yeast_mrna.sort.filt.bam* .
cp $CORENGS/catchup/references/gff/sacCer3.R64-1-1_20110208.gff .

# Use the less pager to look at multiple lines
less sacCer3.R64-1-1_20110208.gff

# Look at just the most-important Tab-separated columns
cat sacCer3.R64-1-1_20110208.gff | grep -v '#' | cut -f 1,3-5 | head -20

# Include the ugly 9th column where attributes are stored
cat sacCer3.R64-1-1_20110208.gff | grep -v '#' | cut -f 1,3,9 | head
```

In addition to comment lines (starting with #), you can see the **chrl** contig names in column 1 and various feature types in column 3. You see also see tags like **Name=YAL067C**;gene=**SEO1**; among the *attributes* on some records, but in general the *attributes* column information is *really ugly*.

To summarize, we have two problems to solve:

- 1. We only care about a subset of feature types (here genes), and
- 2. We want the important annotation information gene names and IDs to appear as regular columns instead of weird name/value pairs.

# Filter annotations based on desired feature type

One of the first things you want to know about your annotation file is what gene features it contains. Here's how to find that: (Read more about what's going on here at piping a histogram)

```
mkdir -p $SCRATCH/core_ngs/bedtools
cd $SCRATCH/core_ngs/bedtools
cp $CORENGS/catchup/references/gff/sacCer3.R64-1-1_20110208.gff .
```

```
Create a histogram of all the feature types in a GFF
```

```
cd $SCRATCH/core_ngs/bedtools
cat sacCer3.R64-1-1_20110208.gff | grep -v '^#' | cut -f 3 | \
sort | uniq -c | sort -k1,1nr | more
```

You should see something like this.

#### Histogram of yeast annotation features

```
7077 CDS
6607 gene
 480 noncoding_exon
 383 long_terminal_repeat
 376 intron
337 ARS
 299 t.RNA
 190 region
129 repeat_region
 102 nucleotide_match
 89 transposable_element_gene
 77 snoRNA
 50 LTR_retrotransposon
 32 telomere
  31 binding_site
  27 rRNA
  24 five_prime_UTR_intron
  21 pseudogene
 17 chromosome
 16 centromere
  15 ncRNA
   8 external_transcribed_spacer_region
  8 internal_transcribed_spacer_region
   6 snRNA
   3 gene_cassette
   2 insertion
```

Let's create a file that contains only the 6607 gene entries:

```
Filter GFF gene feature with awk

cat sacCer3.R64-1-1_20110208.gff | grep -v '#' | \
   awk 'BEGIN{FS=0FS="\t"}{ if($3=="gene"){print} }' \
   > sc_genes.gff
wc -1 sc_genes.gff
```

The line count of sc\_genes.gff should be 6607 – one for each gene entry.

# Convert GFF/GTF format to BED with ID in the name field

Our sc\_genes.gff annotation subset now contains only the 6607 genes in the Saccharomyces cerevisiae genome. This addresses our first problem, but entries in this file still have the important information – the gene ID and name – in the loosely-structured 9th attributes field.

If we want to associate reads with features, we need to have the feature names where they are easy to extract!

What most folks to is find some way to convert their **GFF/GTF** file to a **BED** file, parsing out some (or all) of the name/value attribute pairs into **BED** file columns after the standard 6. You can find such conversion programs on the web – or write one yourself. Or you could use the BiolTeam conversion script, **/work/projects/BiolTeam/common/script/gtf\_to\_bed.pl**. While it will not work 100% of the time, it manages to do a decent job on most **GFF/GTF** files. And it's pretty easy to run.



# Let Anna know if you run into problems

If this script doesn't work on your annotation file, please let Anna know. She is always looking for cases where the conversion fails, and will try to fix it.

Here we just give the script the GFF file to convert, plus a 1 that tells it to URL decode weird looking text (e.g. our Note attribute values).

```
mkdir -p $SCRATCH/core_ngs/bedtools
cd $SCRATCH/core_ngs/bedtools
cp $CORENGS/catchup/references/gff/sacCer3.R64-1-1_20110208.gff .
```

#### Convert GFF to BED with BiolTeam script

The program reads the input file twice – once to gather all the attribute names, and then a second time to write the attribute values in well-defined columns. You'll see output like this:

To find out what the resulting columns are, look at the header line out the output BED file:

```
head -1 sc_genes.converted.bed
```

For me the resulting 16 attributes are as follows (they may have a different order for you). I've numbered them below for convenience.

#### Converted BED attributes

```
1. chrom 2. start 3. end 4. featureType 5. length 6. strand
7. source 8. frame 9. Alias 10. ID 11. Name 12. Note
13. Ontology_term 14. dbxref 15. gene 16. orf_classification
```

The final transformation is to do a bit of re-ordering, dropping some fields. We'll do this with **awk**, because **cut** can't re-order fields. While this is not strictly required, it can be helpful to have the critical fields (including the gene ID) in the 1st 6 columns. We do this separately for the header line and the rest of the file so that the BED file we give **bedtools** does not have a header (but we know what those fields are). We would normally preserve valuable annotation information such as **Ontology\_term**, **dbxref** and **Note**, but drop them here for simplicity.

```
mkdir -p $SCRATCH/core_ngs/bedtools
cd $SCRATCH/core_ngs/bedtools
cp $CORENGS/catchup/bedtools_merge/*.gff .
cp $CORENGS/catchup/bedtools_merge/sc_genes.converted.bed
```

#### Re-order the final BED fields

```
head -1 sc_genes.converted.bed | sed 's/\r//' | awk '
BEGIN{FS=0FS="\t"}{print $1,$2,$3,$10,$5,$6,$15,$16}
' > sc_genes.bed.hdr

tail -n +2 sc_genes.converted.bed | sed 's/\r//' | awk '
BEGIN{FS=0FS="\t"}
{ if($15 == "") {$15 = $10} # make sure gene name is populated
    print $1,$2,$3,$10,$5,$6,$15,$16}
' > sc_genes.bed
```

One final detail. Annotation files you download may have non-Unix (*linefeed*-only) line endings. Specifically, they may use Windows line endings (*carriage return + linefeed*). (Read about Line ending nightmares.) The expression sed 's\rangle return with the context of the sed (substitution editor) tool to replace carriage return characters (\rangle r) with nothing, removing them from the output.

Finally, the 8 re-ordered attributes are:

#### Re-ordered BED attributes

```
1. chrom 2. start 3. end 4. ID 5. length 6. strand 7. gene 8. orf_classification
```

\*\*Whew\*\*! That was a lot of work. Welcome to the world of annotation wrangling – it's never pretty! But at least the result is much nicer looking. Examine the results using more or less or head:

#### **Examine our BED-format annotations**

```
cat sc_genes.bed | head -20
```

Doesn't this look better? (I've tidied up the output a bit below.)

```
315 + YAL069W Dubious
255 + YAL068W-A Dubious
363 - PAU8 Verified
228 + YAL067W-A Uncharacterized
1782 - SEO1 Verified
309 + YAL066W Dubious
387 - YAL065C Uncharacterized
381 + YAL064W-B Uncharacterized
381 - YAL064C-A Uncharacterized
285 + YAL064W Verified
291 - YAL063C-A Uncharacterized
3969 - FLO9 Verified
1374 + GDH3 Verified
1254 + BDH2 Uncharacterized
1149 + BDH1 Verified
423 - YAL059C-A Dubious
639 + ECM1 Verified
1509 + CNE1 Verified
351 - YAL056C-A Dubious
                 649
chrI
          334
                               YAL069W
chrI
          537
                    792
                               YAL068W-A
          1806 2169 YAL068C
chrI
chrI 2479 2707 YAL067W-A
chrI 7234 9016 YAL067C
chrI 10090 10399 YAL066W
chrI
         11564 11951 YAL065C
          12045 12426
13362 13743
chrI
                               YAL064W-B
                               YAL064C-A
chrI
        21565 21850 YAL064W
chrI
chrI 22394 22685 YAL063C-A
chrI 23999 27968 YAL063C
chrI 31566 32940 YAL062W
chrI
         33447 34701 YAL061W
chrI
          35154
                    36303
                               YAL060W
          36495 36918 YAL059C-A
chrI
         36508 37147 YAL059W
chrI
chrI 37463 38972 YAL058W
                                                             - YAL056C-A Dubious
chrI 38695 39046 YAL056C-A
                                                  351
          39258 41901 YAL056W
                                                                                       Verified
chrT
                                                    2643 +
                                                                         GPB2
```

Note that value in the 8th column. In the yeast annotations from **SGD** there are 3 gene classifications: *Verified*, *Uncharacterized* and *Dubious*. The *Dubious* ones have no experimental evidence so are generally excluded.

```
mkdir -p $SCRATCH/core_ngs/bedtools
cd $SCRATCH/core_ngs/bedtools
cp $CORENGS/catchup/bedtools_merge/*.gff .
cp $CORENGS/catchup/bedtools_merge/sc_genes* .
```

## Exercise: How many genes in our sc\_genes.bed file are in each category?

Use cut to isolate that field, sort to sort the resulting values into blocks, then uniq -c to count the members of each block.

```
cut -f 8 sc_genes.bed | sort | uniq -c
```

#### You should see this:

```
810 Dubious
897 Uncharacterized
4896 Verified
4 Verified|silenced_gene
```

If you want to further order this output listing the most abundant category first, add another sort statement:

```
cut -f 8 sc_genes.bed | sort | uniq -c | sort -k1,lnr
```

The -k 1,1nr options says to sort on the 1st field (*whitespace* delimited) of input, using numeric sorting, in reverse order (i.e., largest first). Which produces:

```
4896 Verified
897 Uncharacterized
809 Dubious
4 Verified|silenced_gene
```

# **Exercises**

# Use bedtools merge to collapse overlapping annotations

One issue that often arises when dealing with **BED** regions is that they can overlap one another. For example, on the yeast genome, which has very few non-coding areas, there are some overlapping **ORFs** (**Open Reading Frames**), especially *Dubious* ORFs that overlap *Verified* or *Uncharacterized* ones. When **bedtools** looks for overlaps, it will count a read that overlaps *any* of those overlapping **ORFs** – so some reads can be counted twice.

One way to avoid this double-counting is to collapse the overlapping regions into a *merged* set of *non-overlapping regions* – and that's what the **bedtool s merge** utility does (http://bedtools.readthedocs.io/en/latest/content/tools/merge.html).

Here we're going to use bedtools merge to collapse our gene annotations into a non-overlapping set, first for all genes, then for only non-Dubious genes.

The output from bedtools merge always starts with 3 columns: chrom, start and end of the merged region only.

Using the -c (column) and -o (operation) options, you can have information added in subsequent fields. Each comma-separated column number following -c specifies a column to operate on, and the corresponding comma-separated function name following the -o specifies the operation to perform on that column in order to produce an additional output field.

For example, our sc\_genes.bed file has a gene name in column 4, and for each (possibly merged) gene region, we want to know the *number* of gene regions that were collapsed into the region, and also *which* gene names were collapsed.

We can do this with -c 6,4,4 -o distinct,count,collapse, which says that three custom output columns should be added:

- the 1st custom column should result from collapsing distinct (unique) values of gene file column 6 (the strand, + or -)
   since we will ask for stranded merging, the merged regions will always be on the same strand, so this value will always be + or -
- the 2nd custom output column should result from counting the gene names in column 4 for all genes that were merged, and
- the 3rd custom output should be a comma-separated collapsed list of those same column 4 gene names

bedtools merge also requires that the input BED file be sorted by locus (chrom + start), so we do that first, then we request a strand-specific merge (-s):

```
mkdir -p $SCRATCH/core_ngs/bedtools
cd $SCRATCH/core_ngs/bedtools
cp $CORENGS/yeast_rnaseq/*.gff .
cp $CORENGS/yeast_rnaseq/sc_genes.bed* .
cp $CORENGS/yeast_rnaseq/yeast_mrna.sort.filt.bam* .
module load biocontainers
module load bedtools
```

#### Use bedtools merge to collapse overlapping gene annotations

```
cd $SCRATCH/core_ngs/bedtools
sort -k1,1 -k2,2n sc_genes.bed > sc_genes.sorted.bed
bedtools merge -i sc_genes.sorted.bed -s -c 6,4,4 -o distinct,count,collapse > merged.sc_genes.txt
```

The first few lines of the merged.sc\_genes.txt file look like this (I've tidied it up a bit):

```
2-micron
         251 1523 + 1
                                  R0010W
2-micron
          1886 3008
                           1
                                  R0020C
           3270
                 3816
                            1
                                  R0030W
2-micron
           5307
                 6198
2-micron
                                  R0040C
                            2
chrT
           334
                 792
                       +
                                  YAL069W,YAL068W-A
                           1
           1806 2169 -
                                 YAL068C
chrI
           2479 2707 +
chrI
                           1
                                 YAL067W-A
chrI
          7234 9016
                           1
                                 YAL067C
           10090 10399 +
                            1
                                  YAI.066W
chrT
chrI
           11564
                 11951
                                  YAT-065C
```

Output column 4 has the region's strand. Column 5 is the count of merged regions, and column 6 is a comma-separated list of the merged gene names.

Exercise: Compare the number of regions in the merged and before-merge gene files.

```
wc -1 sc_genes.bed merged.sc_genes.txt
```

There were 6607 genes before merging and 6485 after.

Exercise: How many regions represent only 1 gene, 2 genes, or more?

Output column 5 has the gene count.

```
cut -f 5 merged.sc_genes.txt | sort | uniq -c | sort -k2,2n
```

Produces this histogram:

```
6374 1
105 2
4 3
1 4
1 7
```

There are 111 regions (105 + 4 + 1 + 1) where more than one gene contributed.

Exercise: Repeat the steps above, but first create a good.sc\_genes.bed file that does not include Dubious ORFs.

```
cd $SCRATCH/core_ngs/bedtools
grep -v 'Dubious' sc_genes.bed > good.sc_genes.bed

sort -k1,1 -k2,2n good.sc_genes.bed > good.sc_genes.sorted.bed
bedtools merge -i good.sc_genes.sorted.bed -s \
   -c 6,4,4 -o distinct,count,collapse > merged.good.sc_genes.txt

wc -l good.sc_genes.bed merged.good.sc_genes.txt
```

There were 5797 "good" (non-Dubious) genes before merging and 5770 after.

```
cut -f 5 merged.good.sc_genes.txt | sort | uniq -c | sort -k2,2n
```

Produces this histogram:

```
5750 1
18 2
1 4
1 7
```

Now there are only 20 regions where more than one gene was collapsed. Clearly eliminating the *Dubious* ORFs helped.

So there's one more thing we need to do to create a valid **BED** format file. Our merged.good.sc\_genes.txt columns are *chrom*, *start*, *end*, *strand*, *merged\_region\_count*, *merged\_region\_s*), but the **BED6** specification is: *chrom*, *start*, *end*, *name*, *score*, *strand*.

To make a valid **BED6** file, we'll include the first 3 output columns of merged.good.sc\_genes.txt (*chrom, start, end*), but if *strand* is to be included, it should be in column 6. Column 4 should be *name* (we'll put the collapsed gene name list there), and column 5 a *score* (we'll put the region count there).

We can use awk to re-order the fields:

```
cat merged.good.sc_genes.txt | awk '
BEGIN{FS=OFS="\t"}
{print $1,$2,$3,$6,$5,$4}' > merged.good.sc_genes.bed
```

We're now (finally!) actually going to do some gene-based analyses of a yeast RNA-seq dataset using **bedtools** and the **BED**-formatted, merged yeast gene annotation file we created above.

In this section we'll use **bedtools multicov** to count RNA-seq reads that overlap our gene features. The **bedtools multicov** command (http://bedtools.readthedocs.io/en/latest/content/tools/multicov.html) takes a feature file (**GFF/BED/VCF**) and counts how many reads from one or more input **BAM** files overlap those feature. The input **BAM** file(s) must be position-sorted and indexed.

Make sure you're in an idev session, since we will be doing some significant computation, and make bedtools and samtools available.

```
Start an idev session

idev -m 120 -N 1 -A OTH21164 -r CoreNGS-Fri
# or
idev -m 90 -N 1 -A OTH21164 -p development
```

Copy over the yeast RNA-seq files we'll need (also copy the GFF gene annotation file if you didn't make one).

```
# Get the merged yeast genes bed file if you didn't create one
mkdir -p $SCRATCH/core_ngs/bedtools_multicov
cd $SCRATCH/core_ngs/bedtools_multicov
cp $CORENGS/catchup/bedtools_merge/merged*bed .

# Copy the BAM file
cd $SCRATCH/core_ngs/bedtools_multicov
cp $CORENGS/yeast_rnaseq/yeast_mrna.sort.filt.bam* .
```

Exercises: How many reads are represented in the yeast\_mrna.sort.filt.bam file? How many mapped? How many proper pairs? How many duplicates? What is the distribution of mapping qualities? What is the average mapping quality?

samtools flagstat for the different read counts.

samtools view + cut + sort + uniq -c for mapping quality distribution

samtools view + awk for average mapping quality

```
cd $SCRATCH/core_ngs/bedtools_multicov
samtools flagstat yeast_mrna.sort.filt.bam | tee yeast_mrna.flagstat.txt
```

```
samtools flagstat output
```

```
3347559 + 0 in total (QC-passed reads + QC-failed reads)
24317 + 0 secondary
0 + 0 supplementary
922114 + 0 duplicates
3347559 + 0 mapped (100.00% : N/A)
3323242 + 0 paired in sequencing
1661699 + 0 read1
1661543 + 0 read2
3323242 + 0 properly paired (100.00% : N/A)
3323242 + 0 with itself and mate mapped
0 + 0 singletons (0.00% : N/A)
0 + 0 with mate mapped to a different chr
0 + 0 with mate mapped to a different chr (mapQ>=5)
```

There are 3323242 total reads, all mapped and all properly paired. So this must be a quality-filtered BAM. There are 922114 duplicates, or about 28%.

To get the distribution of mapping qualities:

```
samtools view yeast_mrna.sort.filt.bam | cut -f 5 | sort | uniq -c
```

```
distribution of mapping qualities
   498 20
   6504 21
  1012 22
   355 23
  1054 24
  2800 25
   495 26
 14133 27
   282 28
   358 29
   954 30
  1244 31
   358 32
  6143 33
   256 34
   265 35
  1112 36
   905 37
   309 38
  4845 39
  5706 40
   427 41
  1946 42
  1552 43
  1771 44
  6140 45
  1771 46
  3049 47
  3881 48
  3264 49
  4475 50
 15692 51
 25378 52
 16659 53
 18305 54
  7108 55
  2705 56
 59867 57
  2884 58
  2392 59
3118705 60
```

To compute average mapping quality:

```
samtools view yeast_mrna.sort.filt.bam | awk '
BEGIN{FS="\t"; sum=0; tot=0}
{sum = sum + $5; tot = tot + 1}
END{printf("mapping quality average: %.1f for %d reads\n", sum/tot,tot) }'
```

Mapping qualities range from 20 to 60 – excellent quality! Because the majority reads have mapping quality 60, the average is 59. So again, there must have been quality filtering performed on upstream alignment records.

Here's how to run bedtools multicov in stranded mode, directing the standard output to a file:

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

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

#### Run bedtools multicov to count BAM alignments overlapping a set of genes

```
cd $SCRATCH/core_ngs/bedtools_multicov
bedtools multicov -s -bams yeast_mrna.sort.filt.bam \
  -bed merged.good.sc_genes.bed > yeast_mrna_gene_counts.bed
```

#### Exercise: How may records of output were written? Where is the count of overlaps per output record?

```
wc -1 yeast_mrna_gene_counts.bed
```

6485 records were written, one for each feature in the merged.sc\_genes.bed file.

The overlap count was added as the last field in each output record (here field 7, since the input annotation file had 6 columns).

#### Exercise: How many features have non-zero overlap counts?

```
cut -f 7 yeast_mrna_gene_counts.bed | grep -v '^0' | wc -l
# or
cat yeast_mrna_gene_counts.bed | \
  awk '{if ($7 > 0) print $7}' | wc -l
```

Most of the genes (6141/6485) have non-zero read overlap counts.

#### Exercise: What is the total count of reads mapping to gene features?

```
cat yeast_mrna_gene_counts.bed | awk '
BEGIN{FS="\t":sum=0;tot=0}
{if($7 > 0) { sum = sum + $7; tot = tot + 1 }}
END{printf("%d overlapping reads in %d genes\n", sum, tot) }'
```

There are 1,152,831 overlapping reads in 6,141 non-0 gene annotations.

# Use bedtools genomecov 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 that shows the *count of overlapping bases* for the 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/igy/download) or in the UCSC Genome Browser (https://genome.ucsc.edu/).

- Go to the UCSC Genome Browser: https://genome.ucsc.edu/
- Select Genomes from the top menu bar
- Select Human from POPULAR SPECIES
  - o under Human Assembly select Feb 2009 (GrCh37/hg19)
  - select GO
- In the hg19 browser page, the Layered H3K27Ac track is a signal track
  - the x-axis is the genome position
  - the y-axis represents the count of ChIP-seq reads that overlap each position
    - where the ChIP'd protein is H3K27AC (histone H3, acetylated on the Lysine at amino acid position 27)

The **bedtools genomecov** function (https://bedtools.readthedocs.io/en/latest/content/tools/coverage.html), with the **-bg** (bedgraph) option produces output in **bedGraph** format. 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 CoreNGS-day5
# or
idev -m 90 -N 1 -A OTH21164 -p development

module load biocontainers
module load bedtools

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

Then calling bedtools genomecov is easy. The -bg option says to report the depth in bedGraph format.

```
cd $SCRATCH/core_ngs/bedtools_genomecov
bedtools genomecov -bg -ibam yeast_mrna.sort.filt.bam > yeast_mrna.genomecov.bedGraph
wc -l yeast_mrna.genomecov.bedGraph # 1519274 lines
```

The **bedGraph (BED3+)** format has only 4 columns: **chrom start end value** and does not need to include positions with 0 reads. Here the count is the number of reads covering each base in the region given by **chrom start end**, as you can see looking at the first few lines with **head**:

```
4390
chrI
        4348
                         2
chrI
        4390
                4391
                         1
chrI
        4745
                4798
                         2
                4799
chrI
        4798
                         1
chrI
        4949
                4957
                         2
chrI
        4957
                 4984
                         4
        4984
                4997
chrI
                         6
        4997
                4998
chrI
                         5
        4998
                5005
chrI
chrI
        5005
                5044
                         2
                5045
        5044
                         1
chrI
chrI
        6211
                6268
                         2
chrI
        6268
                6269
                         1
                7257
        7250
                         3
chrI
                7271
chrI
        7257
                         4
chrI
        7271
                7274
                         6
                7278
chrI
        7274
                         7
chrI
        7278
                7310
                         8
chrI
        7310
                 7315
                         6
chrI
        7315
                 7317
                         5
```

Because this **bedGraph** file is for the small-ish (12Mb) yeast genome, and for reads that cover only part of that genome, it is not too big – only ~34M. But depending on the species and read depth, **bedGraph** files can get very large, so there is a coresponding binary format called **bigWig** (see https://genome. ucsc.edu/goldenpath/help/bigWig.html). The program to covert a **bedGraph** file to **bigWig** format is part of the **UCSC Tools** suite of programs. Look for it with **module spider**, and note that you can get information about all the tools in it using module spider with a specific container version:

Looking at the help for **bedGraphToBigWig**, we'll need a file of chromosome sizes. We can create one from our **BAM** header, using a **PerI** substitution script, which I prefer to **sed** (see Tips and tricks#perlpatternsubstitution):

```
module load ucsc_tools
cd $SCRATCH/core_ngs/bedtools_genomecov
bedGraphToBigWig # look at its usage
# create the needed chromosome sizes file from our BAM header
module load samtools
samtools view -H yeast_mrna.sort.filt.bam | grep -P 'SN[:]' | \
 perl -pe 's/.*SN[:]//' | perl -pe 's/LN[:]//' > sc_chrom_sizes.txt
cat sc chrom sizes.txt
# displays:
chrI 230218
chrII 813184
chrIII 316620
chrIV 1531933
chrV
       576874
chrVI 270161
chrVII 1090940
chrVIII 562643
chrIX 439888
       745751
chrX
       666816
chrXI
chrXII 1078177
chrXIII 924431
chrXIV 784333
chrXV 1091291
chrXVI 948066
       85779
chrM
```

Finally, call **bedGraphToBigWig** after sorting the **bedGraph** file again using the **sort** format **bedGraphToBigWig** likes. (You can try calling **bedGraphToBigWig** without sorting to see the error).

```
cd $SCRATCH/core_ngs/bedtools_genomecov
export LC_COLLATE=C
sort -k1,1 -k2,2n yeast_mrna.genomecov.bedGraph > yeast_mrna.genomecov.sorted.bedGraph
bedGraphToBigWig yeast_mrna.genomecov.sorted.bedGraph sc_chrom_sizes.txt yeast_mrna.genomecov.bw
```

See the size difference between the bedGraph and the bigWig files. The bigWig (9.7M) is less that 1/3 the size of the bedGraph (34M).

```
cd $SCRATCH/core_ngs/bedtools_genomecov
ls -lh yeast_mrna.genome*
```

Since the **bigWig** file is binary, not text, you can't use commands like **cat**, **head**, **tail** on them directly and get meaningful output. Instead, just as **zcat** converts **gzip**'d files to text, and **samtools view** convets binary **BAM** files to text, the **bigWigToBedGraph** program can convert binary **bigWig** format to text. That's a different **BioContainers** module (**ucsc-bigwigtobedgraph**) and the default container version doesn't work, so we'll specifically load one that does:

```
# The default version of is broken, so load this specific biocontainers version
module load ucsc-bigwigtobedgraph/ctr-357--1

# see usage for bigWigToBedGraph:
bigWigToBedGraph

cd $SCRATCH/core_ngs/bedtools_genomecov
# use the program to view a few lines of the binary bigWig file
bigWigToBedGraph yeast_mrna.genomecov.bw stdout | head
```