IGV Practical Session.

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This session will give you short introduction into using IGV. We will be using externally hosted data available from Encode including ChIP-seq and RNA-seq data.

Please feel free to try other datasets once you have run through the tasks but remember IGV is hosted on your machine so less powerful computers will feel the memory load of large numbers of datasets.

ChIP-seq data

This section looks at how to load and manipulate some ChIP-seq data. The data is from a lymphoblastoma cell line and contains ChIP-seq samples for several histone types.

Loading IGV and moving around the gene of interest.

- First, load IGV and select the hg19 genome.
- ❖ Navigate to chr12:6,641,585-6,649,537.
- Navigate to Gapdh.
- ❖ Zoom out to see the surrounding genes.
- ❖ Select the "RefSeq Genes" track and expand to see all transcript isoforms.
- Scroll to the left to discover the non-coding RNA SCRNA2.
- ❖ Click on SCRNA to bring up a window with further information.
- ❖ Double click on Gapdh to recentre window on the gene.
- ❖ Add a region of interest at the Gapdh locus using the IGV menu Regions -> Region Navigator -> add
- Right click the "Region of interest" and edit description to show "Active gene"

Loading data

- ❖ Go to the IGV menu -> File -> load from server
- From the menu follow drop-down

Tutorials -> UI Basics (Encode)

GM12878 CFCF

GM12878 H3K27ac

GM12878 H3K27me3

GM12878 H3K36me3

GM12878 H3K4me3

GM12878 H3K9ac

Controlling IGV display

- Select all tracks and then select "autoscale."
- ❖ At the Gapdh locus, investigate the enrichment of signal over gene body.
- ❖ Set all the tracks to maintain current data ranges (*deselect autoscale*)
- Once set, navigate to PIANP.

- ❖ Note the difference in enrichment of ChIP-seq.
- ❖ Add as "Region of interest" and edit description to show "Inactive gene"
- ❖ Zoom out to compare enrichment across neighbouring genes.
- ❖ Go to Regions -> Gene Lists.. -> "proneural dev genes"
- ❖ Inspect the signal across genes to determine their expression state.
- * Return to main view by selecting all from chromosome dropdown.
- Colour all track by a unique colour. Good idea to make K27me3 the most distinct colour to rest.
- Autoscale all tracks.
- Select all tracks and create an overlay,

select tracks -> Create Overlay Track

Change track height to 100

Select tracks -> Change Track Height

Revisit Gapdh, PIANP and gene list. Scan across genome to identify silent and active gene expression.

RNA-seq data

Start a new session

❖ Clear data by going to IGV menu -> File -> New Session...

Loading the data

- ❖ Go to the IGV menu -> File -> load from server
- ❖ From the menu follow drop-down

Tutorials -> RNA-seq (Body Map)

Heart

Lung

❖ Go to IGV menu -> Views -> Preferences -> Alignments

Tick "show junction track"

Set "visibility range threshold' to 15kb

- ❖ Go to SLC25A3 gene.
- Collapse reads track (named Heart/Liver)

Inspect RNA-seq data

- Select tracks -> Colour Alignments by -> Read Strand to identify strand of transcript.
- ❖ Expand "Features track" to identify alternative exons/transcripts.
- ❖ Inspect coverage tracks to discover areas of coverage unique to Heart or Liver sample.
- Inspect junction tracks to evaluate alternative splicing of transcripts between tissue
- ❖ Select junction tracks -> Expand
- ❖ Compare major (first) transcript variant in Heart and Tissue
- Click on junctions to identify start and end of spans.

Another way to inspect splicing

❖ Select tracks -> Shashmi plot Select Gene Track -> Refseq Genes Select Alignment Tracks -> Heart + Liver

❖ Save image and compare junctions across tissue.