UCSC GENOME BROWSER AND HOMOLOGENE

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UCSC Genome Browser

- The University of California, Santa Cruz (UCSC) Genome Browser (http://genome.ucsc.edu) contains a variety of tools
- The Genome Browser allows for interactive visualization of genomic data, which is visualized based on tracks:
 - NCBI RefSeq information
 - OMIM alleles
 - Clinical variants and SNPs, etc
- The Table Browser allows you to download sequence or feature data from all (selected) genes
 - Can download promoter regions, introns, exons, etc

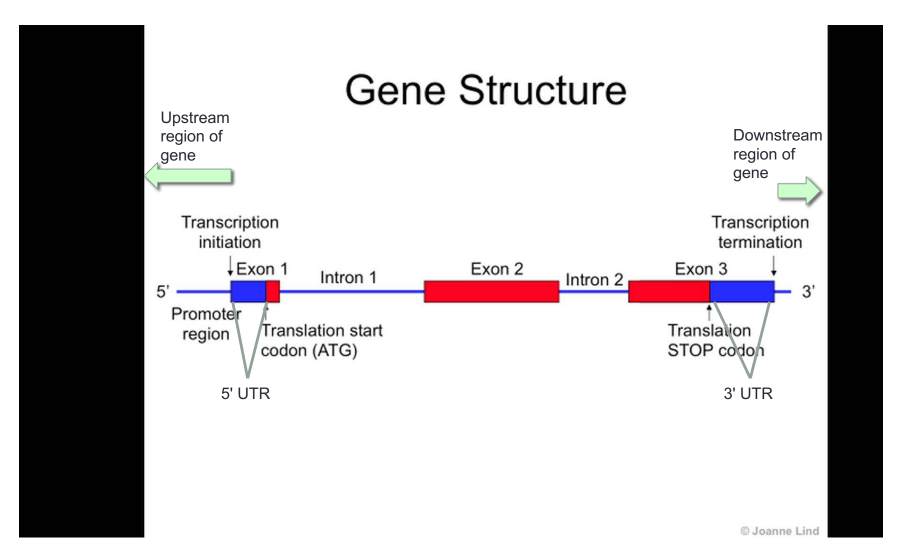
UCSC Genome Browser examples

- Using the Genome Browser, let's look at the gene MAOA
 - Let's hide all tracks, then show the following (use pack view):
 - Genes and Gene Predictions:
 - GenCode v44
 - NCBI RefSeq
 - mRNA and ESTs
 - Human mRNAs
 - Phenotype and Literature
 - OMIM genes (display as "full")
 - Variation
 - dbSNP 155
 - Let's zoom in on CYS266PHE

UCSC Table Browser examples

- Using the Table Browser to get selected information about genes/proteins
- Let's look at all NCBI RefSeq genes in humans
 - Select clade (organism type) and
 - For humans, select "Mammal" for clade and "Human" for genome
 - For group, select "Genes and Gene Predictions"
 - For track we will use "NCBI RefSeq"
 - For region we will use "genome"
 - We can enter specific genes (identifiers), if we want
 - For output format, select "selected fields from .."
 - We will select name, chrom, exonCount, name2
 - Click "Get output" and select features

Overview of Gene Structure



Original image and video: https://www.youtube.com/watch?v=h5HI2OqOJA0

UCSC Table Browser examples

- Using the Table Browser to get sequence information
 - Specify organism, database, and genes as before
 - For output format, select "sequence" and click "Get output"
 - You will then need to specify what parts of the sequence to get
- Can you get the following?
 - Get the CDS exon and intron sequences for BRCA2
 - Get the 5' UTR exon sequences for all human genes

HomoloGene

- HomoloGene is an automated system for detecting homologs from genomic data, and the accompanying database
 - Link: https://www.ncbi.nlm.nih.gov/homologene
 - Two genes are homologs if they share a common ancestral gene
 - Orthologs are homologs as a result of speciation
 - Paralogs are homologs resulting from gene duplication

Ortholog example (true story)

- I have received data from a mouse experiment from a collaborator
- High expression of the following genes (MGI symbols) were associated with resistance to radiation treatment in mice with bladder cancer:
 - Arl4a
 - Birc6
 - Car2
 - Oog3
 - Zfp110
- The next step is to evaluate these genes in human bladder cancer patients – what are the orthologs of these genes?