

Introduction to Biological Big Data Exercises

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# Introduction

In these exercises we are going to explore a number of different online resources to see what we can find out about one specific gene. This will give you the chance to play around in some of the most useful repositories of biological information and see how they might be able to help you in your future work.

The gene we are going to use as our example is human TEC – a protein tyrosine kinase.

# Exercise 1: Genome Exploration

We are going to see what is known about the location, transcripts and structure of the TEC gene using two of the biggest genome browser sites.

## Ensembl

Go to [www.ensembl.org](http://www.ensembl.org) and select the human genome

Use the search function to find the gene page for TEC (it should be ENSG00000135605)

* How many different transcripts are listed for TEC?
* How many encode a protein?
* Of the protein coding transcripts how many are likely to be removed by nonsense mediated decay?
* How long in both transcript base pairs and protein amino acids is the main splice form?

Click on the  tab at the top to take you to the genome browser view of this gene. This will display a track based view of a region of the genome. Many of the tracks shown will be present twice since there will be a separate track for the top and bottom genome strands. Tracks at the top, above the central blue line are top strand (running left to right), those under the blue line are bottom strand (running right to left). Tracks with no direction are drawn right at the bottom.

In the chromosome view you can use the controls to zoom in/out and move along the chromosome



When moving along a chromosome it’s often useful to click on the double arrow to go into drag mode, and then drag in the main view to move it along.

* How close is TEC to the centromere of chr 4?
* Which genes flank TEC on either side? Answer this for both conventional protein coding genes and for small RNA genes.
* Look at the structure of the TEC transcripts. From which end (5’ or 3’) are most of the splice variants most truncated?

You can click on  to select the tracks you want to see in the display. You’ll get a main list of “Active Tracks” initially, and then you can select from a large number of tracks on the left which you can add.

Turn off some of the default tracks you’re not using at the moment. Turning off unneeded tracks will make the loading of new views *much* quicker.

Add a track of CpG islands from the “Simple Features” section.

* Is there a CpG island in the promoter of TEC?

Move back to the TEC gene tab, and we can have a look at the range of species where a homolog of TEC is found. Click on the Gene Tree from the menu on the left and have a look at the species where TEC has been found. You can see the species on the tree, and on the right a summary of the exon structure.

* In which placental mammals has TEC been observed? You will need to click on the placental group and expand the sub-tree to see the individual animals.
* Which phylogenetic group has a large deletion at the start of the gene (blank area on the exon structure picture)? Is this region missing from all animals in this group?

Click on the Orthologs item in the main left menu to get a table of all of the TEC orthologs. Find the ortholog for the crocodile (*Crocodylus porosus*)

* What type of ortholog is this (1-to-1 or 1-to-many)?
* How conserved is the DNA sequence (percentage identity)

Click on the Compare Regions link to open up a multi genome view of the two species around the TEC gene.

* Can you see conserved blocks (the light green shading) covering all of the exons of human TEC?

Expand the view to include the flanking genes in both species (drag a box in the Region Comparison section covering the genes you want to see and select to realign). You need to do this separately in both species.

* Do all of the surrounding genes have orthologs in the two species?
* Are there any obvious changes in structure between the transcripts in this region?

Go back to the main TEC gene tab and then select the page for the primary transcript (ENST00000381501.8)

* How many exons does this transcript have?
* How many of them form part of the coding sequence?

From the left hand menu select the Exons sequence view. Configure the page and change the options to turn off the highlighting of variants.

Find the start of the translated region (blue letters, should start with ATG).

* The Kozak consensus sequence is gcc Rcc ATG g (where the ATG is the first translated base). How well does the sequence at the start of the TEC CDS match this consensus?
* Does the coding sequence run through into the last exon?
* Which is the longest intron, and how long is it?

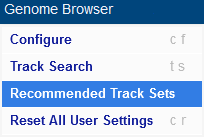
Move to the cDNA sequence section of the Transcript tab

* Download a fastA file of the TEC-201 transcript (turn off all of the other sequence types it offers to give you alongside the cDNA sequence)

## UCSC Browser

Go to <http://genome.ucsc.edu/> and opt to connect to their genome browser. You may be prompted to connect to a local mirror site – it’s fine (good indeed) to use the closest mirror.

Make sure you’re using the hg38 human genome, and find the TEC gene and have a look at the default information you get. You can be sure to get the default tracks by going to

 then 

The basic controls you need to know are:

1. You can click and drag in the main view to move left/right
2. You can hold control whilst dragging to select a region to zoom in to
3. You can zoom in out using the buttons at the top 

Zoom out 10X from the initial view of TEC

* Which genes can you see?
* How many splice variants do they have?

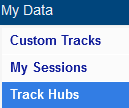
Look at the tracks showing conservation across species

* Which part of which gene has the highest overall conservation?
* Do you observe a loss of conservation in intergenic regions (the bits between genes)?

You can right-click on a track to get some options for how it is displayed and to turn it off completely.

* Hide the RefSeq curated gene track
* Change the ENCODE cCRE track to being a ‘squish’ presentation

One of the biggest benefits to using UCSC is the amount of additional data you can incorporate from other sources. Tracks from other groups come from “track hubs” which you can connect to to see the additional information.

Press on  to see the selection of available track hubs. Have a read down the list to see what sorts of information you can get at and add to your UCSC browser.

* Find the DNA Methylation hub and connect to it. Your session will restart, so re-find the TEC gene and you should see a load of new methylation tracks.
* Zoom out from TEC and see if you can find a region near the gene where the methylation level drops substantially
* Is the extent of the unmethylated region the same in all of the tissues which are shown?

## BioMart

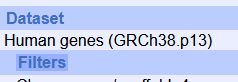
Finally in this section we are going to have a look at the BioMart interface which allows you to do large scale queries against genome annotation information. Although there is a stand-alone site for this at <http://www.biomart.org/>, practically speaking the easiest way to use it is to use the implementation at Ensembl, <http://ensembl.org/biomart/martview/>.

In this section of the exercise we’re going to get you to export a lot of information around all of the human genes on chromosome 4 (the chromosome the TEC gene is on).

To begin, go to the URL above and select that you wish to analyse Ensembl Genes, and that you want to work with the latest human genome.

### Filters

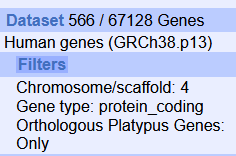
The next step will be to pick the genes about which you wish to find information. To do this you will need to apply filters. You should automatically be taken to the filter section of BioMart as soon as you select the genome, but you can get back to this part at any time by clicking on the links on the left of the page.



The filters we would like you to apply are:

* We only want to use genes on chr 4 (under the REGION section)
* We only want to see protein coding genes (under the GENE section)
* We only want to see genes which have an ortholog in the platypus (under the MULTI SPECIES COMPARISON section)

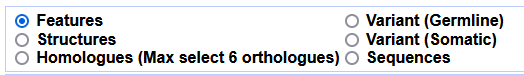
After you’ve made these choices you can click on the “Count” button at the top of the left hand menu to see how many genes remain after applying the filter.



Note that the exact number you see here will change with every Ensembl release as either the human or platypus gene sets get updated. Don’t worry if the number you see doesn’t match exactly as long as it’s similar and the filter descriptions look right.

### Attributes

Next you need to say what it is that you want to know about the genes you have selected. Again there are many different options for this and you can see them by clicking on the attributes link on the left.



You can export information about:

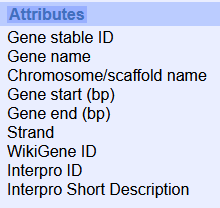
* Annotation features – any of the pieces of annotation for a gene or transcript
* The Structure of a gene – how it breaks down into transcripts, exons and CDS regions
* Homologs – the equivalent Ensembl identifier for homologous genes in other species
* Variants – the details and properties for geneline or somatic variants
* Sequences – the underlying sequence (genome, transcript, CDS exons etc)

We are going to export only feature annotation information so you can choose Features under the attributes options.

You can then see three additional groups of pieces of information you can add. You can pick as many of these as you like.

For our search the information we want to know is:

* The Ensembl Gene ID (Gene Stable ID)
* The Gene Name
* The gene’s location (chromosome, start, end, strand)
* The WikiGene ID (under EXTERNAL)
* The Interpro ID and short description (under Proteins)



Once you’ve selected those you need to click on the Results button which is at the top left of the interface.



That should then show you a short preview of the data you’ve collected so you can check that it looks right. To download the full set of results you need to use the option at the top to save the results to a file.



If you save to a TSV or CSV file then you should be able to open the file directly in Excel to see all of the contents.

### Sequence Query

Once you’ve completed the example above see if you can use BioMart to export out all of the spliced transcript (cDNA) sequences for the TEC gene. The sequences should be in fastA format and should just have the Ensembl transcript ID in their header. ie:

>ENST00000381501

ACTCTGGGGCGCTAGGCTCCGGACTCCGCGGCGCAGACTGCACCTCGCAGTCTCCCCAGG

TCCGCCCAGCAGCCGCGCTTCAGCCAGAATACTGGGATCTTCAGTGGCAGGAGGAGTAAT

CAGAAGACGGAGATGAATTTTAACACTATTTTGGAGGAGATTCTTATTAAAAGGTCACAG

CAGAAAAAGAAGACATCGCCCTTAAACTACAAAGAGAGACTTTTTGTACTTACAAAGTCC

…

>ENST00000505452

AATACTGGGATCTTCAGTGGCAGGAGGAGTAATCAGAAGACGGAGATGAATTTTAACACT

ATTTTGGAGGAGATTCTTATTAAAAGGTCACAGCAGAAAAAGAAGACATCGCCCTTAAAC

TACAAAGAGAGACTTTTTGTACTTACAAAGTCCATGCTAACCTACTATGAGGGTCGAGCA

GAGAAGAAATACAGAAAGGGGTTTATTGATGTTTCAAAAATCAAGTGTGTGGAAATAGTG

AAGAATGATGATGGTGTCATTCCCTGTCAAAATAAGTATCCATTTCAGTCCACAAAGCAG

GGACCTGTGGGTGAAGAAGTTAAAAGAAGAAATAAAGAACAACAATAATATTATGATTAA

ATATCATCCTAAATTCTGGACAGATGGAAGTTATCAGTGTTGTAGACAAACTGAAAAATT

etc.

# Exercise 2: Proteins, domains, structures

## Protein Domains

Start by going to the SMART website <http://smart.embl-heidelberg.de> and look up the information for the TEC gene. For this you’ll need to know the identifier for the TEC protein sequence. For TEC the protein access is P42680. You would have seen this in the Ensembl information for the TEC protein.



Put the accession code into the SMART search and see what domains are found for TEC. How many domains are there? Are they all globular domains or are there any disordered or low complexity linkers present?

For each domain read the information about it. Try to see whether it’s a binding or catalytic domain, and if it’s a binding domain see what it binds to (DNA, Protein, Cofactors etc).

On the domain plot you should see some vertical lines running down the image. What are they, and what is signified by their colour?

Do the same search on Pfam and compare the list of domains annotated. Are there any differences between the results from the two sites?

Now try to find Interpro domains which may contain details of specific binding or active site pockets. The interpro search is at <https://www.ebi.ac.uk/interpro/> but you’ll need the TEC protein sequence to do the search, so go back to Ensembl to get this (it’s only one sequence so you can do it from the main Ensembl web site not through BioMart). The interpro searches can take a little while to run so feel free to move on to the next part whilst this is running and come back to look at the results later.

Once you have the results have a look at the summary figure on the Overview and see whether you can see both large domains (such as you got from SMART and Pfam), but also smaller hits within those large domains.

To get more information click on the Entries tab at the top of the results to get a tabular view of the hits.

See if you can use these results to find where the active site of the Tyrosine kinase domain is and see what details are provided about it.

## Protein Structures

### Experimental Structures

We know that TEC is a multi-domain protein so it’s unlikely that there will be a single structure covering all of the protein, but there may be structures for some of the individual domains.

Go to <https://www.rcsb.org/> and see what you can find. Search initially with the protein accession code for human TEC and see if you get any hits. If you do, which of the domain(s) do they cover.

Next you can try to be less specific and search for the TEC gene name to see if anything else comes up from other species. Just be aware that when searching you need to scroll down quite a long way in the search options to say that you’re providing a gene name. Do you get any additional hits? Do they cover more domains than you saw before?

Have a look at the structures you’ve found. As well as the tabular and image information you can also select the 3D view of the protein and get an interactive view of the structure to play with.

You should have seen that there are still some domains for which you’ve seen no structural information. As a final search go to the “Advanced Search” option and use the sequence of the TEC protein to search against RCSB. Do you now get any additional hits. Why did these not come up before? Do they cover any domains which you didn’t previously see?

Can you see any examples of structures which are complexed with either substrates or inhibitors to the catalytic active site. If so, see if the small molecule sits in the same position as you found the InterPro active site motif in the sequence.

### Predicted Structures

In the experimental structures we found structures for some of the TEC domains, but always as isolated components, and not all of the sequence was covered.

Go to <https://alphafold.ebi.ac.uk> and look for the human TEC protein. Do you find a predicted structure for the whole protein? If so which parts are confidently predicted and which parts are more uncertain? Do these relate to the domain structure of the protein? You can make selections in the heatmap below the structure to highlight the relevant regions in both the 3D structure and the linear sequence.

# Exercise 3: Reactions, Pathways, Functions

## Biochemical Reactions for TEC

Before we get into more complex pathways or functional groups we can look at the reaction which is catalysed by TEC. This will be stored in the Rhea database, but it’s not easy to find catalytic proteins there, only reactants and products. To find the correct reaction you either need to find the Enzyme Classification (EC) number of the protein, or go to a page which annotates the reaction directly.

One of the best resources for protein information is UniProt which collates information about all proteins and links out to a large number of other databases, including Rhea. If you look in the “General Identifiers” section of the Ensembl gene page you should find links to lots of other data sources, including UniProt.

Find the uniprot entry for TEC and you should see the reaction for its catalytic activity and a link to the relevant reaction in Rhea. Have a look and see what reactants and products are in the reaction which is catalysed.

You should also be able to find the EC number for the reaction.

Take a look at the Rhea reaction and see whether there are proteins listed which catalyse the reverse reaction to the TEC kinase. If you can’t see any then look at the reactions involving O-phospho-L-tyrosyl-[protein] and see if you can find a corresponding reverse reaction that way.

## Pathways

### Reactome

Got to the Reactome database at [www.reactome.org](http://www.reactome.org) and search for TEC. You’ll get many hits but you should be able to select just the human protein.

Expand the tree of hits to the pathway browser and see the range of pathways that this protein is involved in. You should see entries under both immune system and signal transduction.

Have a look at the pathway browser for the three major pathway branches which mention TEC and try to determine the following:

* Which protein complexes with TEC after IL3 stimulation. Which domain in the protein mediates this interaction?
* In the innate immune response which lipid is involved in the reaction involving TEC? What is the endpoint of the chain of reactions that this is part of?
* Under the signal transduction tree, which membrane complex does TEC (as well as several other tyrosine kinases) associate with?

### KEGG

We can also look in KEGG to see which of its pathways TEC turns up in. From the main KEGG page at <https://www.genome.jp/kegg/> go to KEGG GENES and search for TEC, find it on the list of hits (it’s not the top hit – it should be entry 7006).

From the gene page you should be able to link through to the two KEGG pathways which involve TEC. See which these are, and whether you can see the correspondence to the pathways you saw previously in reactome.

## Gene Ontology

Finally, in this section we’re going to take a look at the Gene Ontology and see what information it holds on TEC.

Go to the main Gene Ontology web site at <http://geneontology.org/> and search for TEC. You’ll get a lot of hits, so add a filter to only look at the hits from humans. You should see a set of around 39 Gene Ontology hits.

From the list see if you can see a group which covers the main catalytic activity of TEC. Find the most specific catalytic category and then open the page for that category. Have a look at the Inferred Tree View to get a text representation of the hierarchical view from that category. Also try going to Graph Views and construct a figure in QuickGO so you can see the structure of the annotations from that category all the way back to the root of that part of the GO tree.

On the ontology hits you can filter by Ontology (aspect) – this allows you to see hits in the three branches of the tree

* P – Biological Process
* F – Molecular Function
* C – Cellular Component

Have a look to see which process categories are listed and see if they seem familiar from the pathway analysis you did before.

Look at the cellular component hits to see where TEC is found and acts. Try selecting some of the references so you can see the sources of information on which the assignments are based.

# Exercise 4: Regulation and Interactions

## Transcription Factors

### UCSC Browser

Go back to UCSC browser, find TEC and then adjust the zoom to look more closely at the promoter. In the set of available tracks look under the regulation section and turn on the ENCODE regulation track. This should give you two potentially useful views, firstly there should be a track of DNase accessible sites, showing where the chromatin is unwound and would therefore be accessible to the binding of transcription factors. Secondly you should also be able to see a track of ChIP-Seq peak positions for a large number of factors so you can see where different factors were observed to have bound.

This sort of prediction can be useful, but is also prone to over-predicting, so just because you see a bound factor in an open region doesn’t necessarily mean that you can be sure that that factor is actually regulating the gene.

To help focus in on likely candidates you can change the display mode of the transcription factor track to “squish” to get less detail, but see more features. You can also look at the colouring of the features – the darker the feature the stronger the peak / motif hit, so you can look initially at only the strongest hits.

In the TEC promoter, which features in the Transcription Factor track have the strongest hits? Are they all actually transcription factors or is anything else in there.

### JASPAR

One of the factors with a strong hit in TEC is Runx3. Take a look in the Jaspar database <http://jaspar.genereg.net> and find the entry for this transcription factor and see what the binding motif looks like.

### Hocomoco

Find Runx3 in Hocomoco (<https://hocomoco12.autosome.ru/>) and see what the motif there looks like. Is it the same as in Jaspar? If not, why not? Have a look at the subfamily of factors to which Runx3 belongs – how many species has this been detected in? How many other members of this family are there?

### Factorbook

Find Runx3 in factorbook (<https://www.factorbook.org>). Have a look at the description they provide for it. One of the other datasets they provide is a graph to show in which tissues each factor is expressed. Have a look where Runx3 is highly expressed. Does this make sense with what you saw before about the pathways where TEC is important?

In ReMap2022 (<https://remap.univ-amu.fr>) find Runx3, and see how many experimental datasets they show which have measured this genome wide. Which cell type(s) were used. How many peaks were detected for this factor?

## Interactions

Now we can look at some of the sources of information which detail the interactions which TEC has with other proteins or factors.

### BioGRID

We can start with BioGRID (<https://thebiogrid.org>). Find TEC in their search and have a look at the list of interactions they have. You want to look at the evidence column on the right hand side of the list. Ideally you want to see low throughput studies (the lighter colours), and preferably for an interaction to have been confirmed by multiple studies.

Based on this how many interactions have solid evidence and what are they to. Do any of them fit with what you saw in the pathways section? Are there any proteins which you’d have expected to see which either don’t appear or have less evidence than you might have expected?

If you have a look at the details of some of the interactions (click on the evidence count box) you will see that for some there are explanations of the function of the interaction. Have a look and see what some of these are.

### String

String (<https://string-db.org>) has one of the nicest collections and presentation of interactions. Find human TEC and then look at the network of interactions they show. The number of lines connecting the points indicates the types of evidence used to link different proteins. In general the more lines the better!

Have a look at the set of proteins and click on them to see some information about them. Do you think that this set fits better with what you saw in the pathway databases for TEC than the results from BioGRID?

As well as the interaction network view you can also click on “Viewers” and look at the co-expression matrix for TEC and the proteins with which it is predicted to interact. How well do the occurrences of these proteins match? Is the co-occurrence better if you just look in human, or if you expand to other species?

# Exercise 5: Variant Exploration

## Variants in Ensembl

As always, the easiest place to start looking for information is Ensembl. From the TEC gene page select the Variant Table view to bring up a list of all of the variants which have been recorded for this gene.

You will often find that the initial view of variants isn’t particularly useful since the size of the table can be huge. There are some options above the table to filter the list to get to the subset of variants which are actually of interest to you.



The main filters you’d want to apply would firstly be on the consequences of the SNPs. Let’s assume that we are only interested in SNPs which affect the protein sequence, either by changing or truncating it. To do this select the “Consequences” button and select “PTV or Missense”.

Secondly, we can filter by the impact scores. There are two scores recorded by Ensembl, SIFT and PolyPhen. Both of these are measures calculated on the basis of the chemical change in the amino acids (ie a change from uncharged to acidic would be more important than between two uncharged amino acids), and also on the degree of conservation of the position being mutated – more conserved positions are likely to be more functionally important. You will find that these can be useful to get to the more interesting variants, but that some variants may not yet have a score calculated so will have a blank measure.

To cut down the full list apply filters to only see variants where both the SIFT and PolyPhen scores are above 0.8. This should get you a fairly small list.

For the variants which remain which position is mutated and what are the amino acid changes? If you look back at the results from the domain analysis can you see which of the domains are affected? Do any of them affect the catalytic site of the kinase?

## dbSNP

The most interesting SNP in TEC is rs749636109. Find this variant in dbSNP (<https://www.ncbi.nlm.nih.gov/snp/>) and then click through from the search results to the SNP page.

Are there any reported clinical consequences for this SNP?

Look at the frequency with which the SNP has been observed. How prevalent is it in the population? In which geographic region(s) has it been observed?

## ClinVar

The ClinVar database reports variants with clinical relevance. We know from our investigations above that the most significant SNPs in TEC are not associated with a clinical outcome. We can see if there are other variants which are more relevant though.

Go to <https://www.ncbi.nlm.nih.gov/clinvar/> and search for TEC. Are there any clinical variants which affect this gene? If so what sort of variants are they, and how specifically are they tied to TEC?

## OMIM

On a slightly wider scale we can look to see what is known about the phenotypes which are associated with TEC by looking in the OMIM database (<https://www.omim.org>). Look up TEC and see what is known.

Are there any drugs which are known to target TEC? If so, what diseases are they used to treat?

Is there an knockout animal model for TEC? If so, which species and is it only TEC which is affected? What phenotypic effects are seen in the knockout?

Are there any papers which have described the structure and function of TEC in detail?

## COSMIC

Rather than looking at naturally occurring variants within the general population we can instead look at somatic variants which appear in cancer cases. Go to <https://cancer.sanger.ac.uk> and look up TEC.

In which domain of the gene are the largest number of somatic mutations seen?

Which tissue types are the highest frequencies of mutation seen?

Which types of mutation (functional consequence) are most frequent?

Are there any specific publications associated with the role of TEC, or similar kinase in cancer?

# Final Summary Exercise

Take a look at the entry for TEC in both Uniprot (<https://www.uniprot.org/>) WikiGenes (<https://www.wikigenes.org/>) and Gene Cards (<https://www.genecards.org/>). How much of the information we’ve seen through the previous exercises is collated within these summaries, or the gene page of sites such as Ensembl?

# Exercise 6: Finding data in GEO

Although you can find data directly in GEO, in most cases the route to getting to the data is finding a paper which describes an interesting piece of biology you want to pursue. We can search for interesting papers in the pubmed database (<https://pubmed.ncbi.nlm.nih.gov/>)

We are going to look for information relating to the Prox1 gene, specifically we’d like to find out what effect knocking this gene out in embryonic tissue has.

In pubmed search for “prox1 embryonic knockout transcriptome” and find a paper which is obviously based around RNA-Seq data (it may not be the top hit), and which includes all of these terms.

Follow the link to the paper and see if you can get the full text (probably as a PDF). See if they give a GEO accession (GSEXXXX) for the data they use. If they do is it data they created, or public data they re-analysed?

Search for the accession code you find in GEO (<https://www.ncbi.nlm.nih.gov/geo/>) and find the details for the dataset. Is the paper you found the original paper for this dataset? If not which paper first published it?

How many samples are included in this dataset? What experimental conditions do they represent and how many replicates of each condition are there?

Look at one of the Sample pages for this study in GEO. As well as raw data the authors are required to deposit quantitated data. What quantitation did they perform in this instance and what data have they provided?

Which type of sequencing platform was used to generate this sequence data?

From the main GEO study page follow the link through to the SRA run selector and look at the details of the files which have been deposited with this entry. How many runs (sequencing lanes) were produced for each sample (experiment)? Have a look at the first run in the list. How many reads (spots) are in this sample? How long is the read length?

From the GEO entry find the SRA database accession for this data. Take this and search for it in sra explorer <https://sra-explorer.info>. Can you see all of the runs you saw in the SRA run selector? Add the runs to your basket and then generate a list of download URLs where you could get the data if you wanted to download it all.

Finally put the GSE accession number into the text search (not the accession search) of <https://www.ebi.ac.uk/ena/>. Find the relevant study page and check that you can see the samples. See that you could click on the links to the individual fastq files to download them (but don’t actually download them).

# Exercise 7: Exploring FlowRepository

Here we will look at the data available in FlowRepository and download one of the available files and view it.

Go to <https://flowrepository.org>

Search for “PBMC” which is a type of standard blood sample (peripheral blood mononuclear cells), and includes T, B and NK cells.

From the original list of hits further filter for CD4 and find an experiment which describes the measurement of CD4 and IFN-g in monocytes. Look at the details for this submission.

Who submitted the sample?

What is the publication associated with this entry?

How many experimental conditions are in this dataset and what do they represent?

How complete is the metadata for this study (look at the MiFlowCyt score)?

Go to the downloads for this study and see which FCS files are available. We only want to download the two experimental datasets, we can leave out the no treatment control (NTC) samples. Once the zip file is created you can directly open it in the xarchiver program, and then extract the FCS files to the “Big\_Data” folder which should be in your home directory.

From the desktop start the terminal program (the black square in the toolbar at the bottom of the screen). Once that’s open you can launch fcsalyzer, which is a free viewer for fcs files. To do this just type fcsalyzer into the terminal and press return.

To view the files select the density plot button  from the toolbar (the one on the right of the two buttons which look similar). After selecting it draw a square covering about half the width of the page. After you’ve drawn the square you should see a file selector, select the Unstim FCS file.

Once you’ve done this draw a second plot to the right of the first, this time containing the Stim FCS file. Look at the patterning of the two markers in the two conditions and see how the expression of the surface markers has changed between the two conditions.

# Exercise 8: Mass Spec Repositories

## Protein Data in ProteomeExchange

Go to <http://www.proteomexchange.org/> and select the option to Access public data. You should see an interactive plot which shows you how many datasets have been deposited from different species, and using different types of spectrometer.

How many studies are there from Rat?

Use the search to find datasets coming from pituitary and find a dataset from human which profiled this in 2019.

Which of the underlying hosting databases contains the full dataset for this study?

Find the entry for this data in the underlying repository.

What type of Mass Spectrometer generated this data?

Which publication is associated with the data?

Have a look at the samples which were submitted as part of the study and see what information was recorded about each of them.

Click through to the FTP site associated with this data and check how many files you can see and what format they are in.

## Metabolomic data in Metabolomics Workbench

Go to the main metabolomics workbench page at <https://www.metabolomicsworkbench.org/>

In the quick search at the top search for the accession ST000899.

You should find one match – click through so you can see it.

What was the purpose of this study? What biological material was collected for it? What experimental conditions does it contain and how many samples from each condition are there?

Select the option to Show Named Metabolites to see what compounds were detected in the study, remembering that these will be metabolites rather than proteins.

Note that the list of molecules is divided into sections based on the type of Ionisation which was used to detect them. Different molecules are efficiently detected using different types of ionisation (positive and negative) in different runs of the spectrometer.

From the list of metabolites find adipoylcarnitine.

Select it then press the button at the top to show the values for this metabolite. Then draw a bar graph of the levels in the different samples.  Remembering that the samples are in groups of 20, does it look like something interesting might be happening with this metabolite?

Go back and draw a boxplot by factor.  Which of the conditions does this metabolite seem to be downregulated in?

We can do a larger scale analysis of the differential abundance of all metabolites between different conditions. Go back to the main study page and select 

We will construct a volcano plot of the results of a pairwise comparison of two conditions. This plots the p-value (y-axis) against the magnitude of change (x-axis). Select the volcano tool, and choose the Control as Group1 and Crohn disease as group 2 and run the analysis with the default options.

Have a look at the results, there is a table of results at the bottom of the page and you can click through to see the volcano plot and other summaries of the results. You should be able to see the adipoylcarnitine as an outlier along with several other metabolites.

Repeat the analysis on the same groups using the negative ion mode data instead of positive ion mode. Note how you get a different set of hits. In negative ion mode you should see that sucrose is a strong positive hit. Go back to the original full list of metabolites and find sucrose on the list and draw the barplot for it and check that you can see a strong increase in the crohn disease samples (21-40)

# Exercise 9: Imaging Resources

## BioImage Archive

First we’re going to look at the more general data in the BioImage Archive.

Go to <https://www.ebi.ac.uk/bioimage-archive/>

### Gel Images

Do a search for studies about Crispr deposited in 2020 (there should only be one).

Have a look at the gel images which are attached to this study.

### Microscopy Movies

Do a search for “locomotor” and find the dataset which observes movement in wild type and mutant *C.elegans* worms.

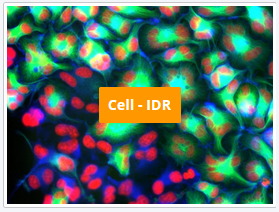
Look in the list of files, you should see a series of AVI movie files. Pick one (preferably a smallish one!), download it and see if you can view the movie.

## Image Data Resource

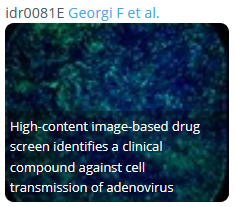
We’re going to look at the information and data we can get from studies in the image data resource.

Go to <https://idr.openmicroscopy.org/>

We’re going to look at the cell level data since this is where most of the higher throughput data generally comes from.



Under Infection Studies look for study idr0081E. This is a study which looked at the ability of 1280 small molecule compounds to inhibit the activity of Human Adenovirus infection. It has high throughput data visualising the starting human cells and then the activity of the virus after treatment with each of the compounds.



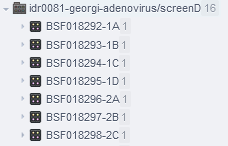
If you select the study and open the main data browser opened at the main study information.

On the left of the interface you have a list of all of the datasets in IDR. For this study there are several folders of information. At the top level you have the main idr0081 folder containing the overall information about the study. As you select that you should see some high level information appear in the sidebar on the right, but no image data.

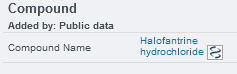


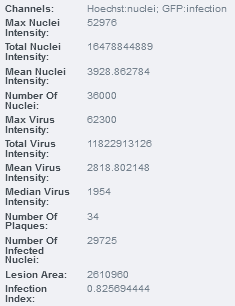
The image data was collected in 5 distinct screens and these are listed as separate folders. You can select them in turn and look at the screen details on the right to see what they represent. You should see that screens A, B and C are all control experiments with either no real treatment or an artificial positive control. The actual screen data starts in screen D, so you should select this.

Under the screen D folder you should then see a set of 16 384-well plates. As you click on a plate you’ll see a thumbnail of the images obtained from each well. Each image shows a small collection of human cells infected with Adenovirus and stained with two stains, Hoechst in blue which stains the human nuclei and GFP in green which represents the activity of the virus in these cells.



If you click on a well you will see the details of both the treatment in that well, ie which drug was added, and also the quantitated values which were extracted. You can see a measure for the mean and median virus intensity, and for the number of viral plaques (colonies).





You should be able to glance over the plate and see some wells are mostly blue, indicating heathy human cells with little viral activity, some are green, indicating strong viral infection, and some are blank indicating that the treatment was likely toxic to the cells.

Have a look through the data and see if you can identify compounds which did both well and badly at preventing viral infection and check that the extracted values from the well agree with your assessment.

For any specific well you can also look at the image in more detail. Select an image from the plate and then at the top of the right hand sidebar there is an option to open the “Full Viewer”. Try this and you can see an expanded version of the image. In the viewer you can zoom in more to see more details, and you can adjust the intensities of the two different stains to get a more balanced view of the image. You should be able to see distinct blue nuclei along with specific bright green viral plaques.

# Exercise 10: Looking at variant data with IGV

In this exercise we’re going to get you to use the IGV program to take a look at some data from the 1000genomes project. We’re going to look at an individual BAM file of aligned reads so we can see the comparison of the sequence to the reference genome to identify SNPs, and then we’re going to look at a VCF file where the collective information for SNPs across hundreds of genomes from different parts of the world has been collated so we can see what can be done with larger scale datasets.

## Navigation

Open IGV. Select the human genome hg19 from the dropdown box on the top left.

The chromosome names are shown on the ruler at the top and the genes at the bottom.

Select a chromosome from the dropdown box next to the genome dropdown.

Zoom in to see a gene by highlighting a region on the chromosome ruler, double clicking in the data tracks or using the zoom bar at the top right.

Click and drag to find a gene on the forward strand. Zoom right in to see the individual nucleotides.

Click on the DNA sequence to see the 3 frame amino acid translation.

Zoom out using Alt + click or the zoom bar and find a gene on the reverse strand.

Zoom in until you can see the DNA sequence.

Click on the arrow to the left of the bases to switch to viewing reverse stranded bases. Click again to return to forward strand view.

Enter the name of your favourite gene in the search bar, view splice variants by right clicking and selecting “expanded” or “squished”.

## Viewing Aligned Reads

Make sure you have the genome set as Human hg19. This isn’t using the latest human genome, but the data that we’re using has been aligned to hg19.

Go to **File -> load from server -> 1000 Genomes -> Alignments -> GBR -> exome ->** Select **HG00101** and **HG00103**

Set the option to show the centre line by going to **View -> preferences ->**  **Alignments ->** scroll down and check the box for **Show center line**

Go to chr12:7,945,539-7,945,578

In the two data tracks **Right click -> color alignments by -> read strand**

Right click -> sort alignments by -> base

Right click -> Squished to fit more reads on the screen

Which sample looks like it’s heterozygous for the SNP and which is homozygous?

Single click the exome coverage track above the reads to view more information – this should show the read counts for each nucleotide at this position, along with percentages for each base.

The data sets that you have loaded show all the individual reads that were mapped for these samples. Now we’ll have a look at a much larger set of samples (including the 2 samples for which we have the exome data).

## Variant Exploration

### Load the data

**File -> Load from Server -> 1000 Genomes ->** Select **Phase 3 genotypes** and click Ok

If you’ve navigated away from the SNP located at chr12:7,945,539-7,945,578 move back there.

light blue means that sample is homozygous for the SNP at that location, dark blue means heterozygous. Find the samples in the variant data that match with the exome samples and check that the SNP call matches between the two.

There are over 1000 samples in the variant data set so remove the exome tracks in order to fit more of the variant data on the screen. **Right click -> remove track.** Do this for the exome coverage tracks too.

### Load sample information

**File -> Load from Server -> 1000 Genomes -> Sample information**

To make use of the sample information go to

**View -> Select attributes to show** select pop and super\_pop

Pop is the country of the individual, super\_pop are groups of countries, mainly continents.

Change the view so more samples can be seen on the screen by

**right click -> Display mode -> squished**

Group by super\_pop

**right click -> Group by super\_pop**

Zoom out and browse the data to see whether there seem to be many differences between the super populations

Have a look at these SNPs and see whether there are differences between the populations

The SNP references can be pasted into the search bar e.g. rs917997, then IGV will move to the location if it finds a match. The view should be centred on the locus, zoom in to view more information. Single click on the SNP in the phase 3 genotypes track to check it’s the right SNP – the rs number should be in the ID field

Right click on the SNP to sort by genotype in order to see the proportion of homozygous and heterozygous samples in each super population.

Rs917997 – associated with inflammatory bowel disease

Rs334 – sickle cell anaemia, this isn’t present in most of the samples, and might be a bit tricky to find, it’s at position 5,248,232. Scroll down to the African group to see the SNPs.

rs2040410 type 1 diabetes, IGV doesn’t seem to find this from the reference, so the location can be used instead chr6:32602698-32602699

# Exercise 11: Looking at RNA-Seq data with SeqMonk

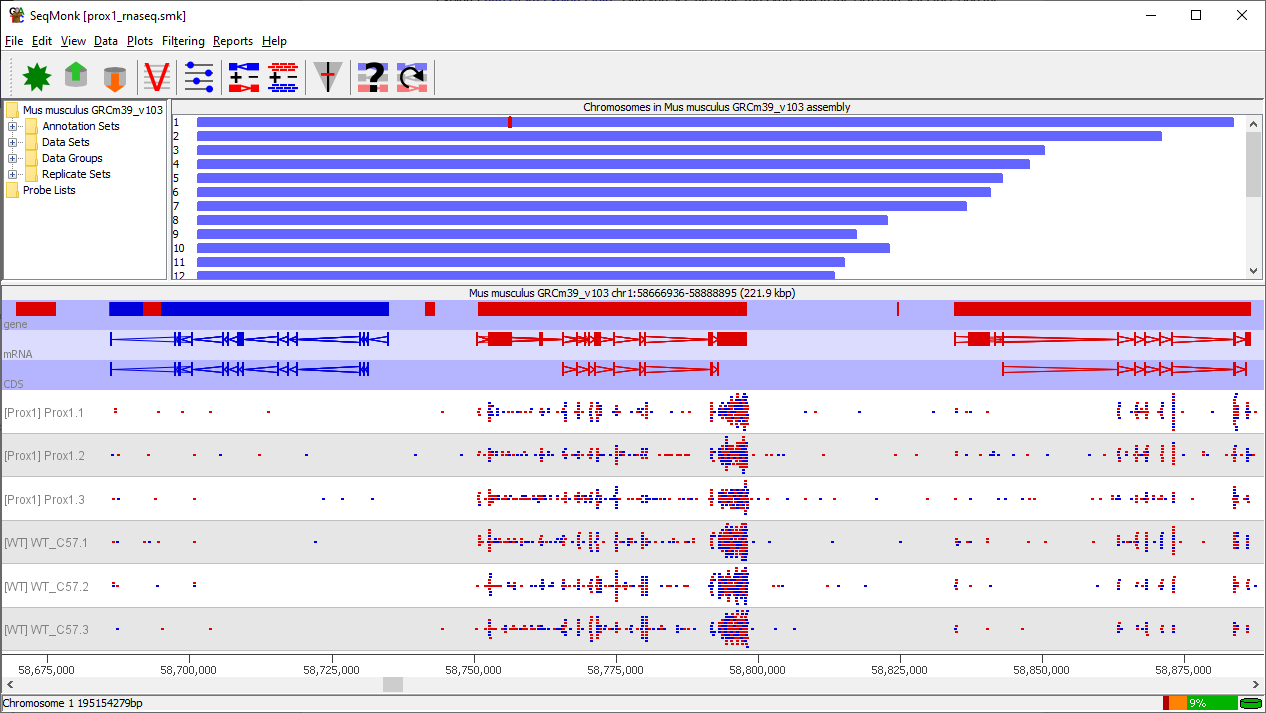
SeqMonk is a graphical program for the exploration and analysis of mapped data – ie sequence data which has been converted to positions in a reference genome.

In this exercise you’re going to use SeqMonk to analyse the RNA-Seq data you found in the GEO exercise. We’ve downloaded and processed this and loaded it into a SeqMonk project so that you can see some of the capabilities of the program. This exercise IS NOT intended to be a guide for the analysis of RNA-Seq data – come on our RNA-Seq course if you want to find out about that – but more an illustration of the software which can help you with this analysis and the sort of results you can find.

## Loading and Exploring the raw data

Launch seqmonk by opening a terminal and typing seqmonk and pressing return.

From within the program select File > Open Project and then select the prox1\_rnaseq.smk project file. You should see the program open and show you a view of the genome along with some annotation and data tracks for part of one chromosome.



Have a look around the genome. You can move left and right along a chromosome using the scroll wheel on your mouse or trackpad, or the left/right arrow keys on your keyboard. You can zoom in by dragging a box over part of the bottom panel, and you can zoom out by clicking the right mouse button (control + click on a mac).

The colour of both features and reads represents the genomic strand. Red = top strand (left to right), Blue = bottom strand (right to left).

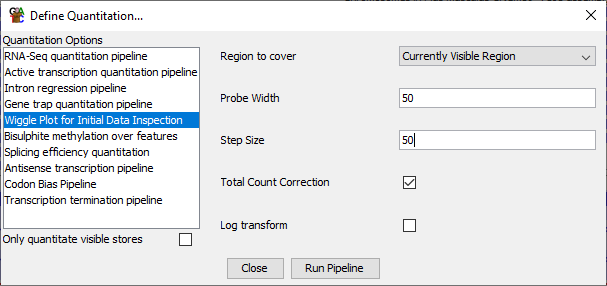
Can you see reads aligning to the exons of transcripts? Can you see examples of genes with both high and low levels of expression.

From looking at the reads over a given gene can you say whether this is a directional, or non-directional library?

This experiment is supposed to be a conditional knockout of the Prox1 gene so we can look at that to check. Press the  button in the toolbar to open the search and search for prox1. You should see two hits, the main prox1 gene, and another gene which transcribes from the same promoter in the opposite direction. Double click on the prox1 gene to move the view to see it.

Do you still see any data for Prox1 in the Prox1 KO? Does the pattern of reads look different in any way?

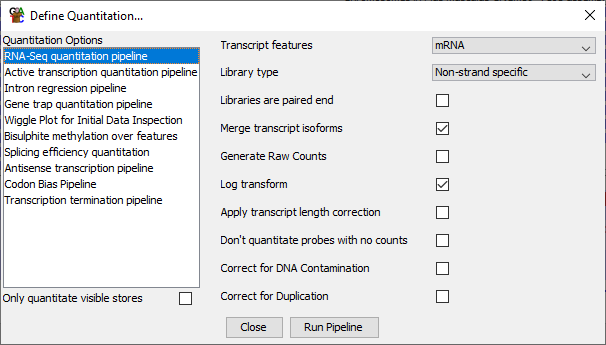
We can quantitate the expression over this region to get a clearer view of what’s going on. Select Data > Quantitation Pipelines > Wiggle Plot for initial data inspection and then change the Probe Size and Step Size to both by 50bp and press Run Pipeline



You should now see a quantitative view of the Prox1 gene. Use the  button on the toolbar to open a control which allows you to change the y-axis scaling and see if you can see a quantitative difference between the WT and Prox1KO samples over Prox1. How is the expression affected in both exons and introns.

## Quantitation and exploration

Now we can quantitate the gene level expression. To do this select Data > Quantitation Pipelines > RNA-Seq Quantitation Pipeline. Run the pipeline with its default settings.



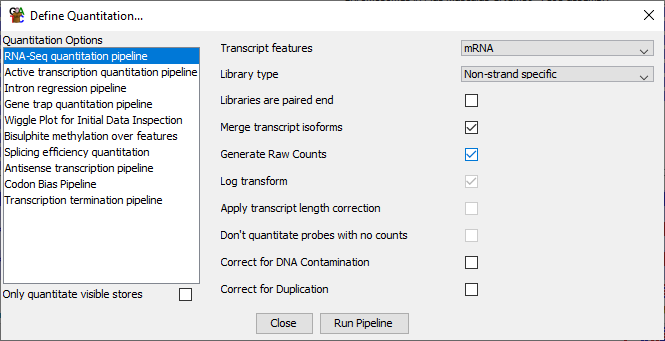
You’ll get a warning about your probe list being replaced. This is fine so just say Yes when prompted.

You now have a gene level quantitation of expression. You can therefore look at some other properties of this data. Firstly you can see whether the WT and KO samples look consistently different to each other. To do this you can select Plots > Data Store Similarity > Data Store Tree. This draws a clustered tree for all of your samples based on the similarity of their data. Do you see a clear separation of the WT and Prox1KO samples?

You can also look at the overall expression comparison of the two conditions. Select Plots > Scatterplot and then plot Prox1 vs WT. Do you see obvious differences on the scatterplot? Double click on some of the genes with the largest difference (those furthest from the diagonal) to look at their raw data in the main view. Can you see differences in the mapped read distributions. Are they consistent between the replicates of the two conditions?

## Statistical Analysis

We’re going to use the DESeq2 R package to select significantly differentially expressed genes in this data. This requires that we generate a set of raw read counts for each gene (not the normalised counts we currently have). To do this go back to Data > Quantitation Pipelines > RNA-Seq Quantitation Pipeline and turn on the option to “Generate Raw Counts” and rerun the pipeline.

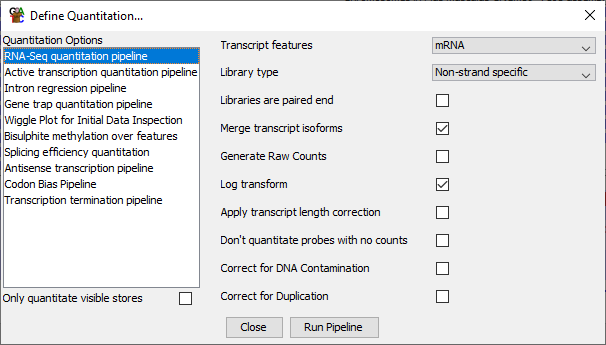


We can then run DESeq from within SeqMonk. Select Filters > Filter by Statistical Test > Count based Statistics > Replicated Data > DESeq2.

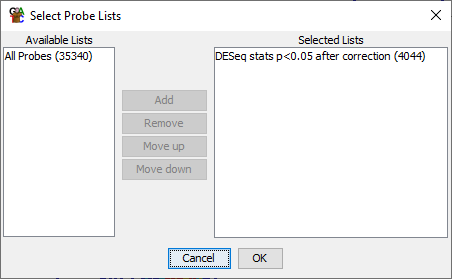
You can press “Run Test” using the default options. You will see an R session open and run a bunch of code and then you should get a hit list back. You can use the default name for the list.

## Viewing Statistical Results

Now that we have our statistical hits we need to regenerate normalised quantitation values. We can do this by going back to Data > Quantitation Pipelines > RNA-Seq Quantitation Pipeline and turning OFF the option to “Generate Raw Counts” and rerun the pipeline.



Now you can redraw the scatterplot to look at the hits. Select Plots > Scatterplot and plot Prox1 vs WT again, but now select the option to “Highlight Sublists” and select the DESeq list you just created and add it to the plot.



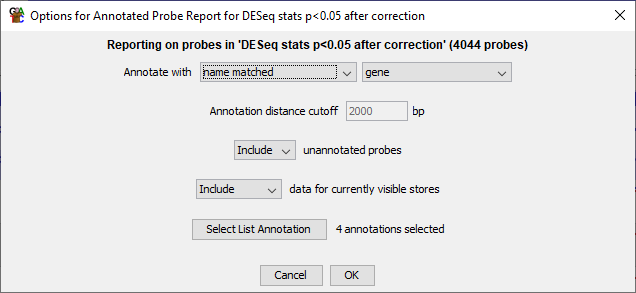
You should now be able to see that the hits are the points on the outside of the cloud of genes.

## Reporting and Biological Themes

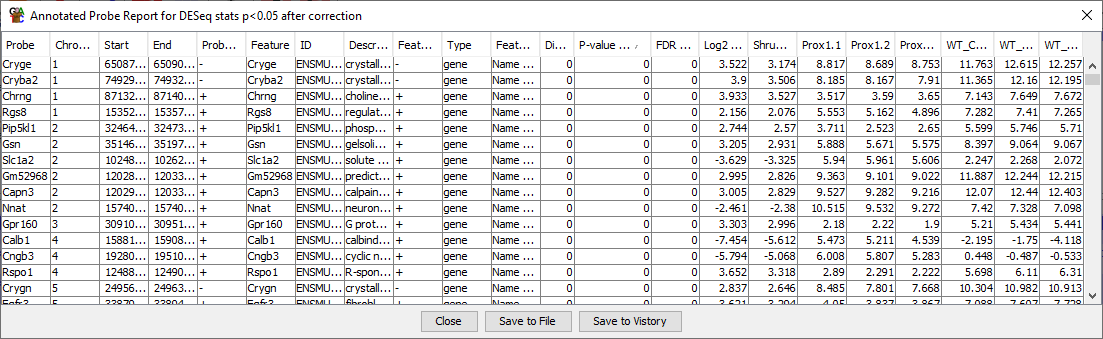
Finally we’re going to generate a table of results and use this to do a Gene Ontology search for relevant biological processes in our hits. To do this we’re initially going to need to select our DESeq list in our data view (the top left window). Use the icon next to “All Probes” to expand the tree so you can see the DESeq list, and select it.



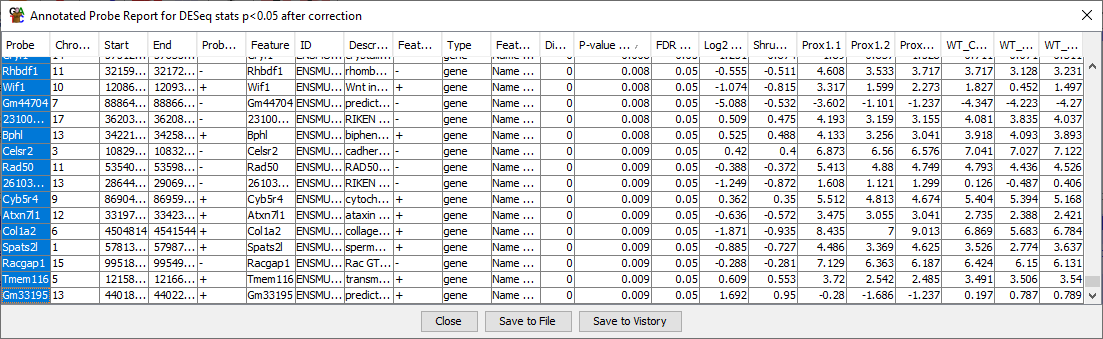
Now select Reports > Annotated Probe Report and press OK to generate a report of the hits. You should see a table appear.



We need to sort the hits by the p-value of the statistical test. You can click on the P-value column header to do this (make sure the lowest values are at the top of the list)



Finally, we’re going to copy the sorted list of gene names for the hits so we can put this into a Gene Ontology search. Select the first Gene Name from the top of the list then scroll to the bottom, hold shift, and click on the last gene to select all of the gene names. Press Control + C to copy the list of gene names.



Finally go to <http://cbl-gorilla.cs.technion.ac.il/> and paste the list of gene names into the search box. Change the species to “Mus musculus” and press “Search Enriched GO Terms”. You will see a graphical representation of the Gene Ontology with enriched regions coloured in yellow – orange. Look at the terms which come up (you’ll need to scroll across to see them all). Given that this was a study to look at the effects of knockouts in the lens of the eye, do you see terms which seem sensible in this context?

# Using the Linux Command Line

## Exercise 12: Runing Programs in Bash

The figlet command draws pretty graphical representations of text you supply, something like this:

\_\_\_ \_ \_ \_

|\_ \_| | | \_\_\_\_\_ \_\_\_\_\_ | | (\_)\_ \_\_ \_ \_\_\_ \_\_

| | | |/ \_ \ \ / / \_ \ | | | | '\_ \| | | \ \/ /

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Read the man page for figlet to work out how to use it

Write your name (if you put spaces in you need to put your name into quotes)

Get your name centred in the terminal

xcowsay is a graphical program which makes a cow say something

Run xcowsay -t 0 “I am a graphical program”

* + Note that you can’t enter more commands in the terminal until you click on the cow to make it go away
  + Read the man page to find out what the -t 0 means

## Exercise 13: Using the file system

In your home directory use mkdir create a folder called compare

Move into the seqmonk\_genomes/Saccharomyces cerevisiae directory in your home directory. Make sure you use tab completion to write the folder name.

* + Note that there is a space in the second folder name. How does the command line completion deal with this?

Try the following commands and note the differences

* + ls
  + ls -l
  + ls \*
  + ls -ld \*

Using ls list the contents of directories containing a 4 in their name (\*4\*)

Use the head command to simultaneously show the first line only of all of the I.dat files in any of the subdirectories (\*/I.dat)

* + Are the chrI sequences all the same length?

Use less to look at Mito.dat in the EF4 directory.

* + See if you can find the first rRNA gene
  + What is its position?

Using cp copy Mito.dat into the compare directory in your home directory

* + Use nano to edit the file
  + Change Mito to Mitochondrion in the ID and AC header lines
  + Save the file and exit nano
  + Using mv rename the file from Mito.dat to Mitochondrion.txt

Copy the original Mito.dat file (the one at ~/seqmonk\_genomes/Saccharomyces\ cerevisiae/EF4/Mito.dat) to the same filename in your compare directory

Run diff on Mitochrondrion.txt and Mito.dat to see that it finds the edits you made to the file.

## Exercise 14: Automation in Bash

Go into the FastQ\_Data directory and look at one of the fastq files using less

* + Less is clever enough to realise that the file needs to be decompressed so you can just pass the file to less directly
  + Now validate that one of the files can be successfully decompressed
    - Run zcat on the file, but…
    - Throw away the STDOUT output (using > /dev/null) so that you just see errors or warnings

Calculate the signatures of all of the fastq files using the sha1sum program

* + Start by running sha1sum on one fastq file to see how it works
  + Now run it on the entire contents of FastQ\_Data using a wildcard (rather than a loop)
    - Write the results (STDOUT) to a file in your home directory using >~/signatures.txt

Use nohup to run the fastqc program on all of the fastq.gz files (\*fastq.gz)

* + Check the nohup.out file to see that it has finished.

Once the fastqc jobs have finished, run multiqc . (note the dot to specify it should run in the current directory) to assemble the fastqc output into a single report

## Optional Exercise: Installing Software with BioConda

We have pre-installed the fastqc program for you on the systems you’re working on, but if we hadn’t you could have installed it from bioconda

Start by finding where fastqc is currently installed by running the following Bash command:

which fastqc

You should see that it’s currently being accessed in /usr/local/bin.

Now you can install BioConda. You can follow the instructions at <https://bioconda.github.io/user/install.html>

You’ll need to accept the license agreement, and you can allow it to activate the conda environment automatically. Once you’ve installed it you’ll need to close the shell you had open and start another one for the changes to take effect. If it’s worked you should see your command prompt to change to:

**(base)** student@ip-172-31-40-53:~$

..where the (base) at the start shows that conda is active. After setting up the channels as instructed on the bioconda install page you can install fastqc by running:

conda install fastqc

You will see that it installs a ton of other things, this is the downside to conda, it’s self contained and therefore duplicates lots of your system software to be sure it will work.

After the install has completed run:

which fastqc

..again, and you should see that you now have a new version of the program installed in your conda directory. If you want you can try reprocessing some of the fastq files to check the new version works.

# Exercise 15: Interactive Data Analysis in R Notebooks

In this exercise we’re going to give you a flavour of the sort of general data analysis which you can perform in an R notebook using the Tidyverse packages in R.

We’re going to explore and analyse a small dataset where neutrophils were treated with different inhibitors and then their ability to invade cells was measured. The dataset has quantitative measures of invasion for multiple replicates of different treatments.

The structure of the input data file is:

|  |  |
| --- | --- |
| **Treatment** | **Invasion** |
| DMSO | 144.4393 |
| DMSO | 135.7167 |
| DMSO | 57.88828 |
| TGX.221 | 99.61073 |
| TGX.221 | 115.3576 |
| etc. | etc. |

# Setting up the Notebook

Before we get into the actual analysis we’re going to set up the template for the notebook and write a description of what we’re going to do using simple markdown syntax.

Open rstudio (just type rstudio in a shell) and select File > New File > R Notebook from the menu. You will see a template notebook structure open.

You can immediately save the notebook in a file called neutrophils.Rmd in the Big\_Data folder, which is where the data we’re going to import is located.

Our analysis is going to be in four parts:

1. Loading in and looking at the data
2. Plotting our data
3. Calculating some summarised values from the data
4. Performing a statistical analysis

## Adding a basic markdown structure

We’re going to start by simply describing what we’re going to do in this analysis. Delete everything below the header in the template notebook which opens (from “This is an [R Markdown] …”)

Use what you were shown about markdown syntax to make level 1 headers (underlined with equals signs) for the four sections we showed above and write a little bit of text to say what you’re going to do in each one (just make it up!). In the loading section you can describe the conditions of the experiment and format them as a numbered list. The conditions are:

1. DMSO
2. TGX221
3. PI103
4. Akt1

Once you’ve added your markdown, save the document and compile it by selecting “Knit to HTML” from the drop down menu immediately above the editor. Check to see that the headings and list have rendered as you’d expect.

If you like you can modify the header of your document to add in the option to include a floating table of contents.

---

title: "Neutrophil Analysis"

output:

html\_document:

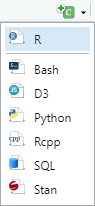
df\_print: paged

toc: true

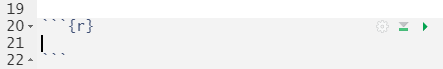
toc\_float: true

---

Save and re-render the document and check the table of contents shows up.

Now we have our basic document structure we can start adding in code. For each code block we want to add you need to put your cursor into the document at the point where you want to insert the code block and then use the menu above the editor to insert an R block.

You will see a block like this insert into the document.



You need to put your code between the upper and lower delimiters. When you want to run your code press the green play button at the top right. The output from the code will embed into the document just below the code block.

Before getting into the analysis we need to load the tidyverse package which we’re going to be using. We’re also going to load rstatix which is a library we’ll later use for statistical testing. Create an initial code block just after the header section and write:

library(tidyverse)

library(rstatix)

You should see something like the following appear in the document:

Registered S3 methods overwritten by 'dbplyr':

method from

print.tbl\_lazy

print.tbl\_sql

-- Attaching packages --------------------------------------------------------------------------------------------------------- tidyverse 1.3.1 --

v ggplot2 3.3.3 v purrr 0.3.4

v tibble 3.1.2 v dplyr 1.0.6

v tidyr 1.1.3 v stringr 1.4.0

v readr 1.4.0 v forcats 0.5.1

-- Conflicts ------------------------------------------------------------------------------------------------------------ tidyverse\_conflicts() --

x dplyr::filter() masks stats::filter()

x dplyr::lag() masks stats::lag()

Registered S3 method overwritten by 'data.table':

method from

print.data.table

Attaching package: “rstatix”

The following object is masked from “package:stats”:

filter

## Loading and Viewing Raw data

The data is contained in a file called neutrophil\_invasion.csv (a comma separated value file). We’re going to load this in using the read\_csv() function from tidyverse.

Put a code block into your loading section containing:

read\_csv("neutrophil\_invasion.csv") -> data

data

This will load in the data and save it under the name ‘data’. Putting data on its own prints out the contents of data so you see the table of results we’re working with. You can use the controls under the table to see the remaining rows.

We can also try using a filtering option to have a look at all of the data we have for just one of the conditions.

Open a new code block and run:

data %>%

filter(Treatment == "Akt1") %>%

arrange(Invasion)

To see all of the Akt1 data and sort the rows from the lowest to highest Invasion values.

## Plotting the data

Now we want to plot out the data to see what we’re working with and what might be happening in the different conditions. To do this we’re going to use the ggplot2 package (which is part of tidyverse) and is the most common plotting system used in R.

To create a basic stripchart view of the values for each condition create a new code block in the plotting section and run:

data %>%

ggplot(aes(x=Treatment, y=Invasion)) +

geom\_jitter()

You might find that it’s not very easy to tell which points come from which treatment, so we can help by adding colouring to the points, and making them slightly bigger.

data %>%

ggplot(aes(x=Treatment, y=Invasion, colour=Treatment)) +

geom\_jitter(size=4)

See that ggplot automatically assigns colours and creates a colour legend next to the plot for us.

Have a look at the data and write into the document what you think you can see in the data.

We can also use a summarised representation of the data to visulalise it. In a new code block we can draw a boxplot of the data.

data %>%

ggplot(aes(x=Treatment, y=Invasion)) +

geom\_boxplot(fill="yellow")

See if this makes the trends in the data any clearer.

## Calculating Summarised Values

We can see that there appear to be differences in the mean values for invasion in the different conditions, however we can also see that the spread of points is also somewhat different and it would be good to quantify this too.

We are therefore going to summarise the data by splitting it into groups based on the Treatment, and then calculating a mean and standard deviation for each subgroup.

To do this create a new code block in the summarisation section and run the following:

data %>%

group\_by(Treatment) %>%

summarise(

mean=mean(Invasion),

stdev=sd(Invasion)

) -> summarised\_data

summarised\_data

This will calculate and store the mean and stdev for the invasion scores for each treatment and then will display it for us to see.

Which sample(s) show a marked shift in their mean value? How close are the StDev values for the treatment(s) which change?

We can also plot this summarisation out as a barplot. To do that we create a new block and use:

summarised\_data %>%

ggplot(aes(x=Treatment, y=mean, ymin=mean-stdev, ymax=mean+stdev)) +

geom\_col(fill="orange") +

geom\_errorbar(size=1, width=0.4)

This will use our summarised values, where the height of the bars is the calculated mean, and the errorbars are the mean +/- the standard deviation.

## Statistical Analysis

Finally we can do a statistical analysis of the different conditions. We’re going to use an ANOVA test to compare all of the Treatments\* and then a Tukey pos-hoc test to tell us which combinations of Treatments were the most significant.

\* This is slightly shaky from a statistical point of view since the ANOVA assumes that all of your samples will have similar standard deviation, and ours shows a bit of a shift, but for the purposes of demonstration we’re going to apply the test anyway. In real life we’d go and talk to a statistician to see whether the difference in StDev is big enough that it might invalidate the test results.

Firstly we’re going to use an ANOVA to compare all of the conditions to see if there is any evidence of a difference in any of the means.

In your statistics section create a new block and run:

data %>%

anova\_test(Invasion~Treatment) %>%

as\_tibble()

The Invasion~Treatment means that what we’re testing is the ability of the test to predict the Invasion level based on knowledge of the Treatment.

Look at the value for the p column. If this is below 0.05 then somewhere in the data there is a significant difference in the mean values. Does the value you see agree with what you’d expect having looked at the data?

We can finally go on to do a set of pairwise comparisons between all of the Treatments to determine which of them is going to be the most significant. Create a new block and run;

data %>%

tukey\_hsd(Invasion~Treatment) %>%

arrange(p.adj)

Which pairs of samples have significant results (p.adj < 0.05). What do they have in common?

Once you’ve finished this you can re-render your notebook to create an HTML output.

# Exercise 16: Python Application Development

In this section we’re going to build a small python application. This is going to be a program which asks the user for the name of a human gene and then queries the Ensembl API for this name, retrieves the Ensembl Gene ID, and then uses this to get the cDNA sequence of all of the transcripts for that gene.

To do this we will be using code examples from both Ensembl’s API documentation, but also using the BioPython project to parse the sequence files for us.

This is NOT intended to turn you into a python programmer in an hour or so, but should help illustrate how you can automate access to these kinds of data sources.

## Overall Structure

We’re going to develop this application in 4 stages

1. Collect a gene name from the user
2. Turn this gene name into an Ensembl ID
3. Get the cDNA sequences for the Ensembl ID
4. Parse the sequences and collect the names and lengths of the transcripts

Before we get into these, at the top of the script we need to load in the helper packages we’re going to use later on:

import requests, sys

from Bio import SeqIO

import io

The first is used for the Ensembl API requests and the other two are used when parsing the fasta sequence data. After we’ve put these in you can save the file under the name gene\_stats.py.

## Asking the user for a gene name

To ask the user for a piece of data we can use the input function. This allows us to ask a question, wait for their answer and then store it so we can use it later.

query = input("Which gene? ")

When the query comes back it will have a newline character on the end (because they pressed return to submit it), and we need to remove that, which we can do with a method called strip.

query = query.strip()

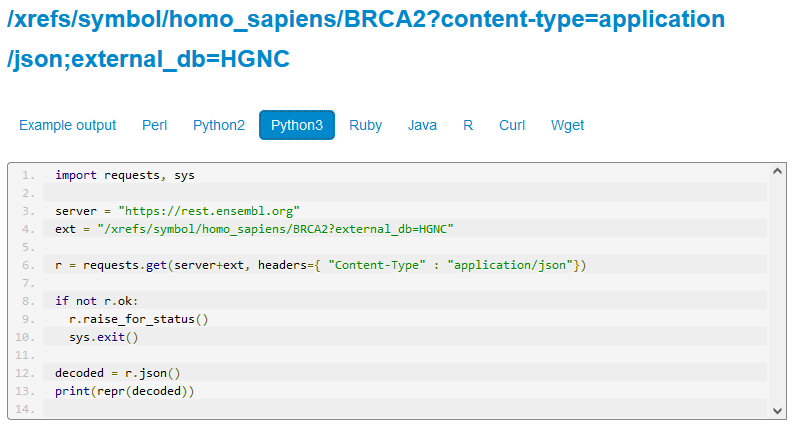
If you want to check what they got you can print out any piece of data using the print function. You can pass multiple pieces of data to this separating them with commas.

print("They asked for",query)

Check that the script works this far and you can see the gene name they asked for.

## Converting a gene name into an Ensembl ID

That was fairly simple, but now we get a little more complicated. We’re going to use the Ensembl API web resource to query the Ensembl database for that gene and get back the gene (or possibly genes) which use that name. To do this we’re going to adapt the instructions which Ensembl provide on their website (<https://rest.ensembl.org/documentation/info/xref_external>), they even give snippets of code in various languages to help you implement this for yourself. One of their examples is pretty much exactly what we want so we’re going to adapt that:



In our case we need to change this slightly, firstly to insert the gene name we collected from the user, and secondly to extract the Ensembl gene id directly from the response rather than just printing the whole thing.

We start by putting in the server details:

server = "https://rest.ensembl.org"

..but then for the request we need to join our gene name into the query string:

ext = "/xrefs/symbol/homo\_sapiens/"+query+"?external\_db=HGNC"

Now we can make the request exactly as they suggest and check that it worked.

r = requests.get(server+ext, headers={"Content-Type":"application/json"})

if not r.ok:

r.raise\_for\_status()

sys.exit()

The editor should automatically indent your code after the if statement. When you get to the end of the block you can use the backspace key to align your next statement with the left side of the screen.

Assuming it did work then we can get the decoded data and extract the first gene id. The code where we retrieve information from within decoded gets the first entry in the list (entry 0), and then gets the id value from that entry. This code needs to be back against the left side of the editor.ex

decoded = r.json()

gene\_id = decoded[0]['id']

Finally we can print out the result to check it’s working:

print("Query=",query,"ID=",gene\_id)

If you query with BRCA2 you should get ENSG00000139618 back as the gene id.

## Getting all of the cDNA sequences from the query gene

Now that we have an Ensembl gene ID we can use a different API to get all of the transcript sequences associated with that gene. For this we’re going to use a different API, this one: <https://rest.ensembl.org/documentation/info/sequence_id>

This time none of the examples are exactly what we want, so we’re going to have to build a query using the set of parameters we have at the top. The things we need to set are:

1. We’re querying with a gene id
2. The type of sequence we want is cdna
3. We want to retrieve multiple sequences (one gene can have several transcripts)
4. We want fasta format data back

We’re using the same server as before so we don’t need to set that up again, but we’re using a different query so we need to build that using the Ensembl ID we got back from the last query.

ext = "/sequence/id/"+gene\_id+"?type=cdna&multiple\_sequences=1"

So here we input our Ensembl ID and say that we want cdna sequence, and that we are happy to have multiple sequences returned.

We can now run the query, and say we want the output in fasta format. We can check that it worked in the same way as before:

r = requests.get(server+ext, headers={ "Content-Type" : "text/x-fasta"})

if not r.ok:

  r.raise\_for\_status()

  sys.exit()

If you want a quick check that this worked then you can print out the full response, but you probably don’t want to leave this in your code once you know that it’s working as you’ll see that the full sequences can be pretty big! This code again needs to be out of the if block and back against the left of the editor.

print(r.text)

See if you can get this far and see the sequences coming back from Ensembl.

## Parsing the fasta files and getting sequence details

For the final part we’re going to use a sequence parser which is part of the biopython project (<https://biopython.org/>). Specifically we’re going to use a package called SeqIO which deals with reading and writing sequences (<https://biopython.org/docs/1.75/api/Bio.SeqIO.html>).

The SeqIO parser is designed to read data from a file, and we’ve actually got it as a piece of text, so before we can parse it we need to use a little helper called io.String which makes a piece of text look like a file so SeqIO will read it.

s = io.StringIO(r.text)

Now we can use SeqIO to parse the data.

fasta = SeqIO.parse(s,"fasta")

Finally we can iterate through the sequences, pulling out the name and length and printing the details.

for f in fasta:

    name = f.id

    length = len(f)

    print("Found transcript",name,"with length",length)

This should get us our final program. Try running the program and trying out a few gene names to check it works.

Which gene? BRCA2

They asked for BRCA2

Query= BRCA2 ID= ENSG00000139618

Found transcript ENST00000544455.6 with length 11854

Found transcript ENST00000530893.6 with length 2011

Found transcript ENST00000380152.8 with length 11954

Found transcript ENST00000680887.1 with length 11880

Found transcript ENST00000614259.2 with length 11763

Found transcript ENST00000665585.1 with length 2598

Found transcript ENST00000528762.1 with length 495

Found transcript ENST00000470094.1 with length 842

Found transcript ENST00000666593.1 with length 523

Found transcript ENST00000533776.1 with length 523

Some example genes to try would be:

Sox2

Nanog

Fos

..but feel free to try your own favourites too.