

# Exercise 3

## Instructions –

- Create a folder called “exercise3” in your WEXAC-mounted classNN folder (testing\classNN\exercise3).
- Create a file with your answers called “Exercise\_3\_2019\_answers.docx”.
- Read the exercise instructions carefully and answer the questions below
- In this exercise we will use the IGV application. Install IGV according to the instructions in the lecture (<https://software.broadinstitute.org/software/igv/download>). If you are working in a Feinberg Class B computer, IGV is installed.
- This exercise is part of part of a Broad workshop from April 2017: [http://www.igv.org/workshops/BroadApril2017/IGV\\_Exercises.pdf](http://www.igv.org/workshops/BroadApril2017/IGV_Exercises.pdf)
- The exercise includes ChiP-seq and RNA-seq data. Chip-seq is a method used to analyze protein interactions with DNA. Chip-seq combines chromatin immunoprecipitation (ChIP) with massively parallel DNA sequencing to identify the binding sites of DNA-associated proteins. RNA-seq can be used to determine mRNA expression levels.

## IGV Hands-on Exercise

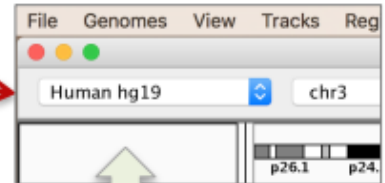
## IGV Basics

### 1. Launch IGV

### 2. Select reference genome.

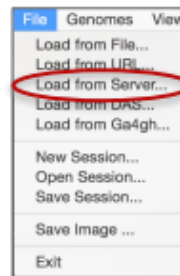
- Click on *Human hg19* in the genome drop-down menu in the upper left corner.

If you only see *Human hg18* in the menu, it's ok to select that instead



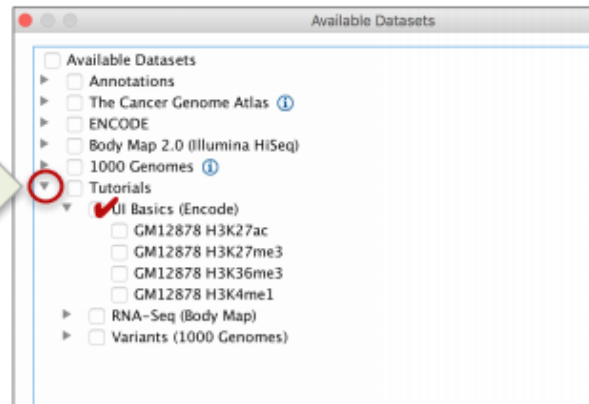
### 3. Load data from the IGV hosted server.

- Select *File > Load from Server...*
- Open the *Tutorials* menu (Use **⏪** on Mac, and **+** on Windows) and click on the *UI Basics* checkbox.



If this is the first time you run IGV, there may be **only one** entry in the menu. More about that later...

Make sure you only **open** the *Tutorials* menu. Do **not** check the box next to *Tutorials*. That will select everything under *Tutorials*, but we only want *UI Basics* for this exercise.



Four tracks are loaded: ENCODE project ChIP-seq data representing histone modifications. Each track is displayed as a bar chart of signal intensities.

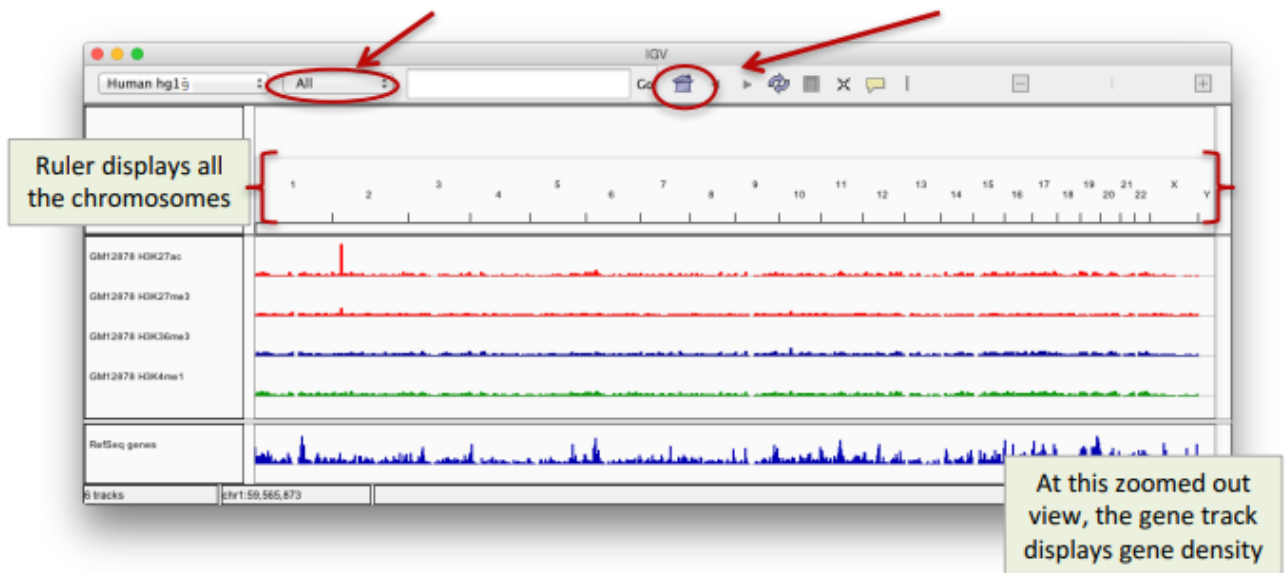


This type of data is perfect for a UI basics exercise because the tracks are visually simple – but the navigation basics are the same no matter what type of data you load.

4. **Navigate** across different genomic loci and at different zoom levels, from whole genome view and down to base-pair resolution.

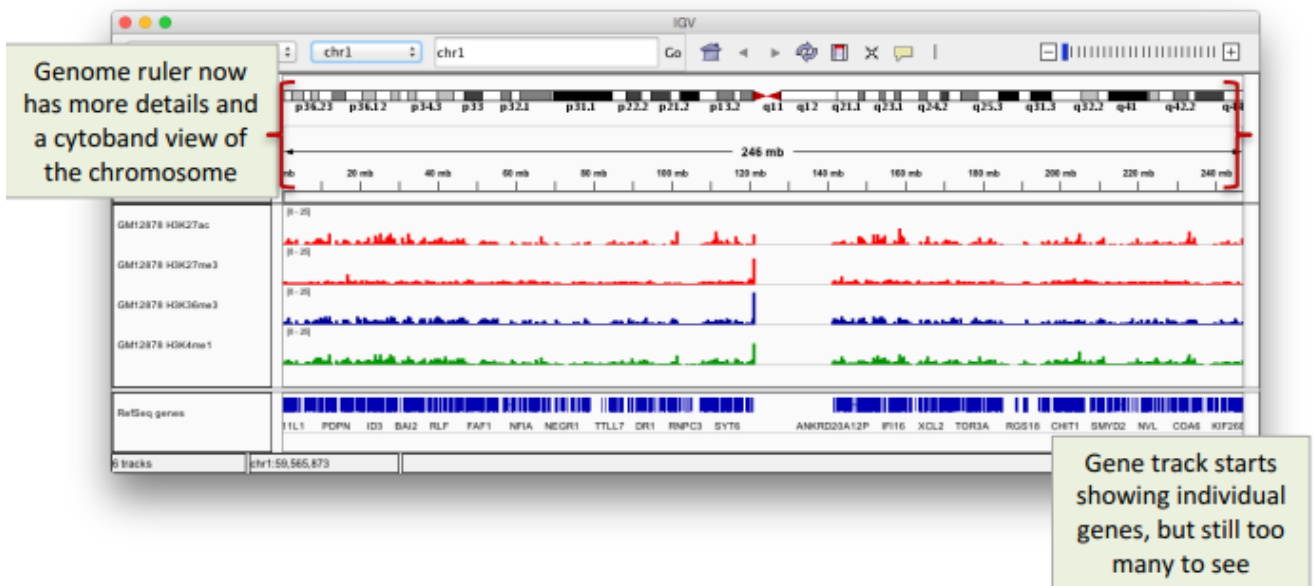
4a. Start at **whole genome view**:

- Select *All* from the chromosome drop-down menu –OR– Click the *Home* button.



4b. Zoom in to view **one whole chromosome**:

- Select *Chr1* from the chromosome drop-down menu –OR– Click the *1* in the genome ruler.



## Questions

1. Go to gene BRCA2. How many isoforms has it (Be sure the RefSeq track is expanded)?
2. Go to gene N4BP2L1.

A. How many isoforms has it?

B. What are the differences between isoforms 2 and 3 (from top to bottom) as they appear in the IGV?

Use exon numbers to answer the following questions. Note that exon 1 is the closest to the 5' of the transcript.

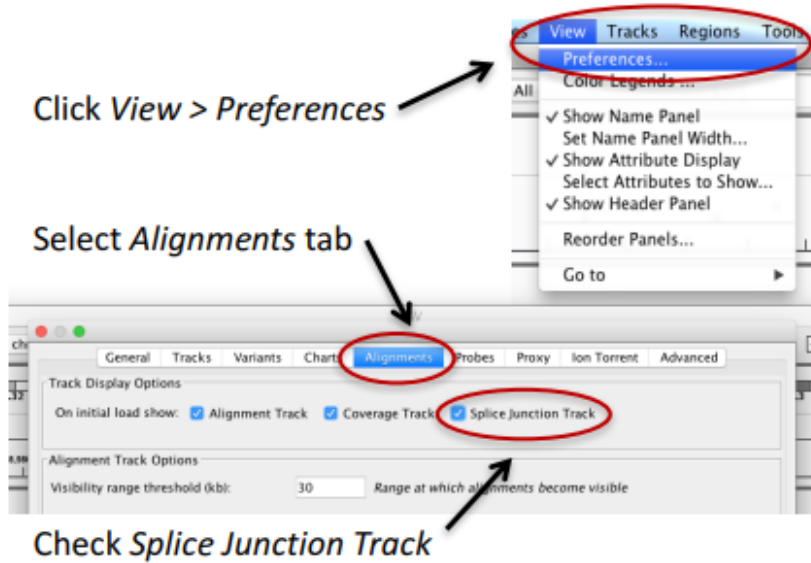
C. Which amino acids are translated from the first exon (from 5' to 3')?

D. What is the size of the first intron?

E. In which track do you see the biggest peak near the promoter?



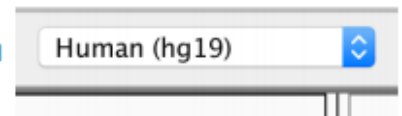
1. Set preferences for viewing RNA-seq data





In the newer version instead of "Splice Junction Track", it is written "Show junction track".

2. Load data

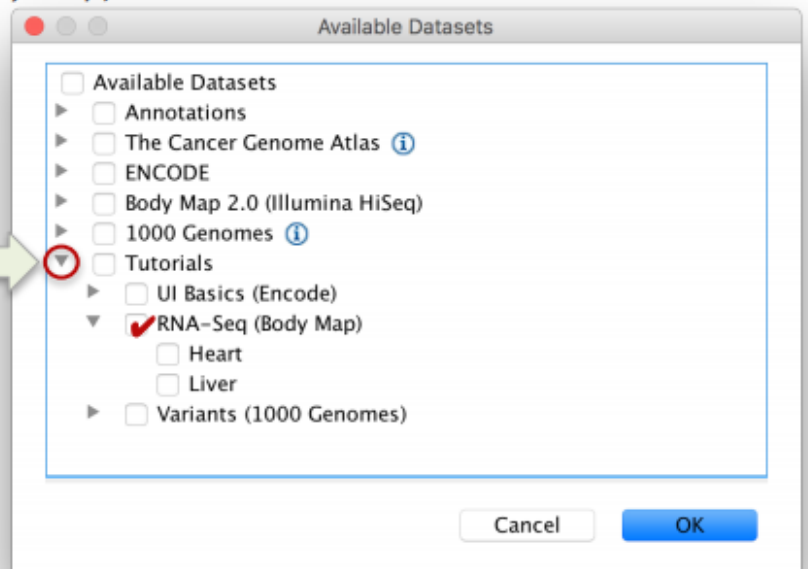
Select *Human hg19* from the genome dropdown menu



Click *File > Load from Server*

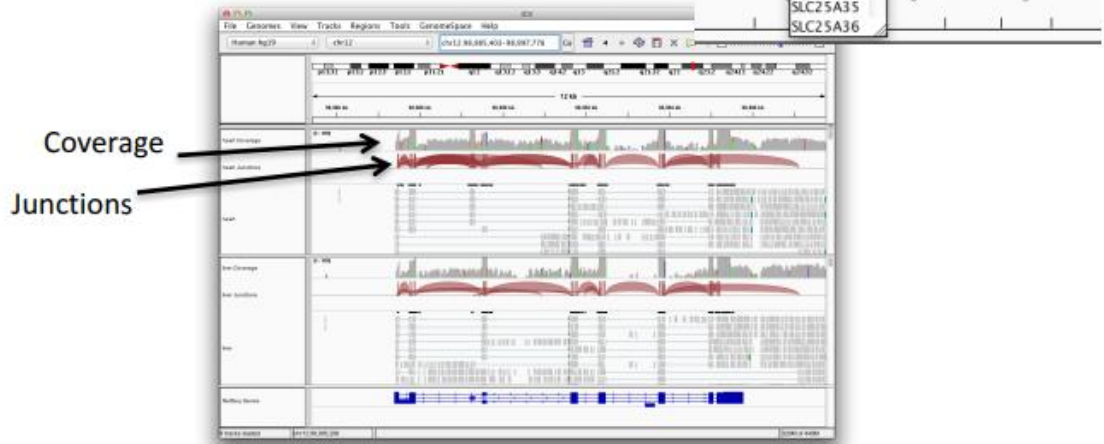
Open the *Tutorials* menu (Use  on Mac, and  on Windows) and click on *RNA-Seq (Body Map)* and then click on *OK*

Make sure you only **open** the *Tutorials* menu. Do **not** check the box next to *Tutorials*. That will select everything under *Tutorials*, but we only want *RNA-Seq* for this exercise.



### 3. Jump to gene SLC25A3

Type *SLC25A3* in the search box and click **Go**



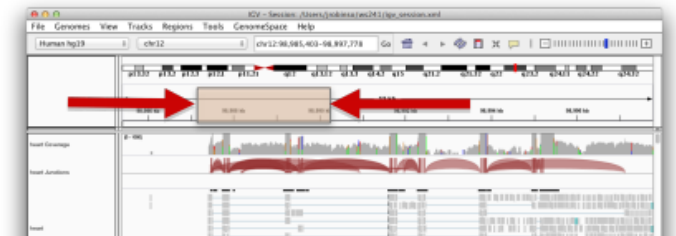
### 4. Expand gene track to see isoforms

Right-click over the *RefSeq Genes* track, and select **Squished**



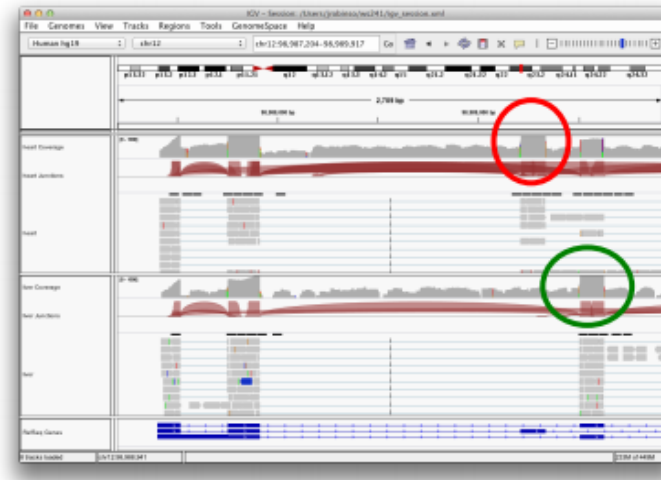
### 5. Zoom in on first 3 exons

Click and drag in ruler region over area shown

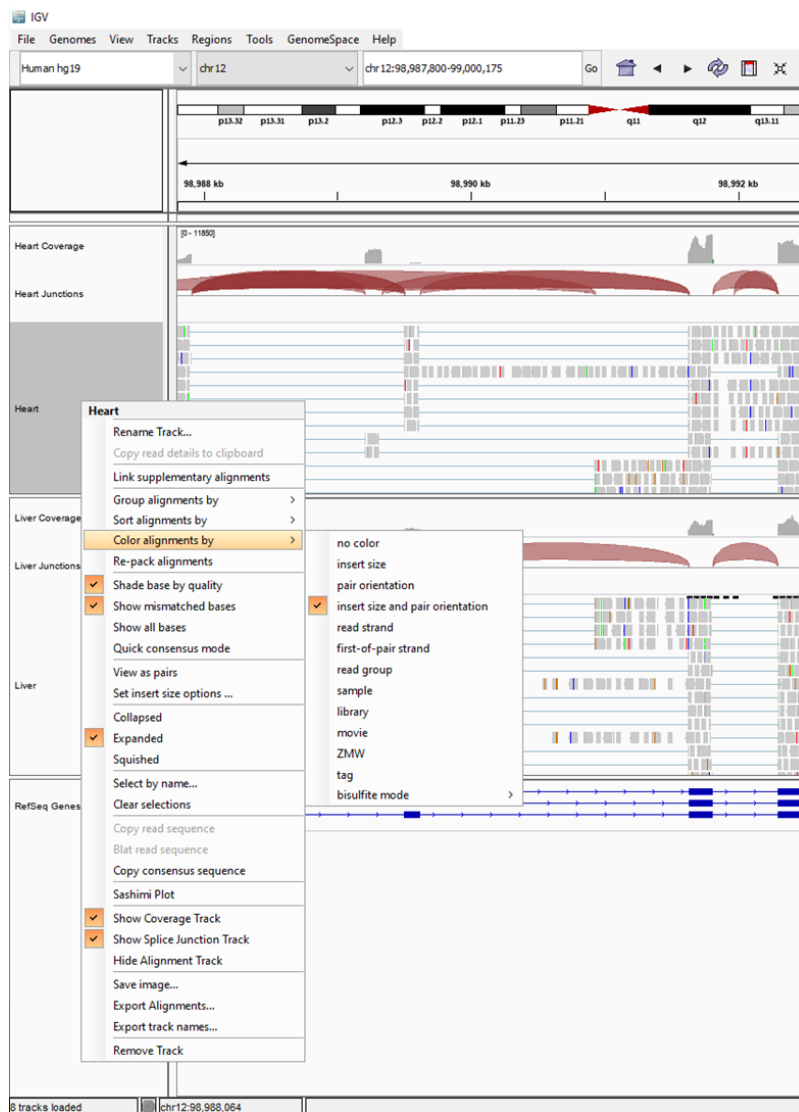


## 6. Note evidence of alternative splicing.

Observe which isoforms in the RefSeq track are expressed in each tissue.



Be sure that the Heart and Liver track alignments are colored by insert size and orientation according to the picture below (right click on the track).





## Questions

3. What is the most probable format for the files loaded in this part of the tutorial?
4. What are the gray rectangles you see in the browser?
5. Which isoforms of the SLC25A3 gene are most likely to be expressed in each tissue according to the data you see in IGV? Count the isoforms from top to bottom.

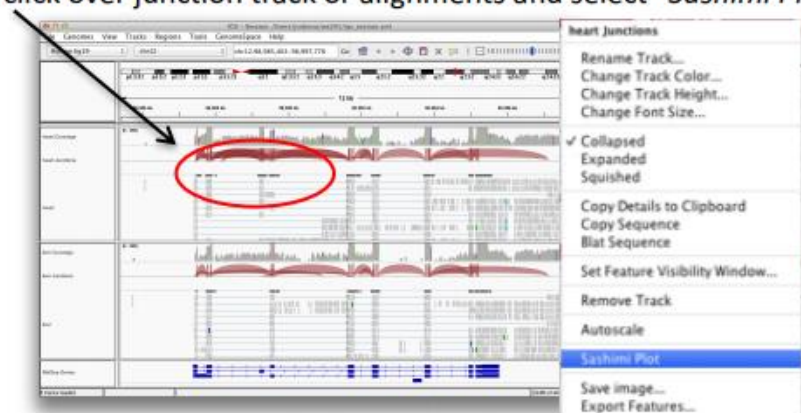
### 7. Zoom back out to view whole gene

Click the back button in the command bar to zoom out to previous view

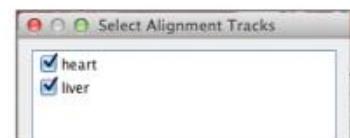


### 8. Open Sashimi plot

Right-click over junction track or alignments and select "Sashimi Plot"



Verify both *heart* and *liver* are checked, and click OK

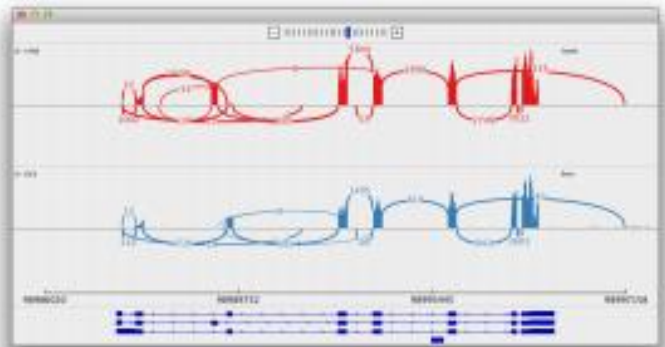


**Sashimi Plot.** Sashimi plots visualize splice junctions for multiple samples from their alignment data alongside genomic coordinates and a user-specified annotation track. IGV displays the Sashimi plot in a separate window and allows for more manipulations of the plots than the junctions track.

## 9. Examine Sashimi plot

Note:

- Arcs represent reads spanning exon junctions
- Peaks represent exon coverage



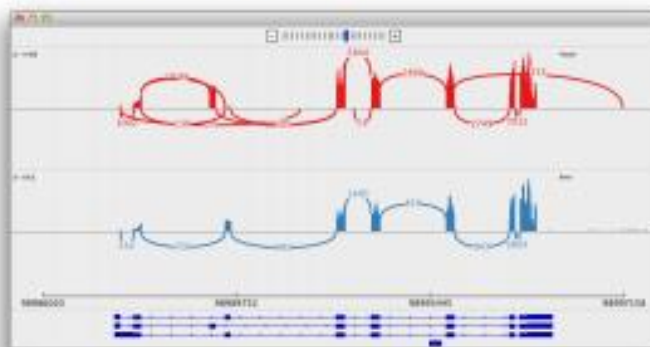
## 10. Filter out low-count splicing events

Right click over red (heart) track and select **Set Junction Coverage Min**. Enter **50** and click **OK**.

Repeat for blue (liver) track.



## 11. Compare with non-filtered view



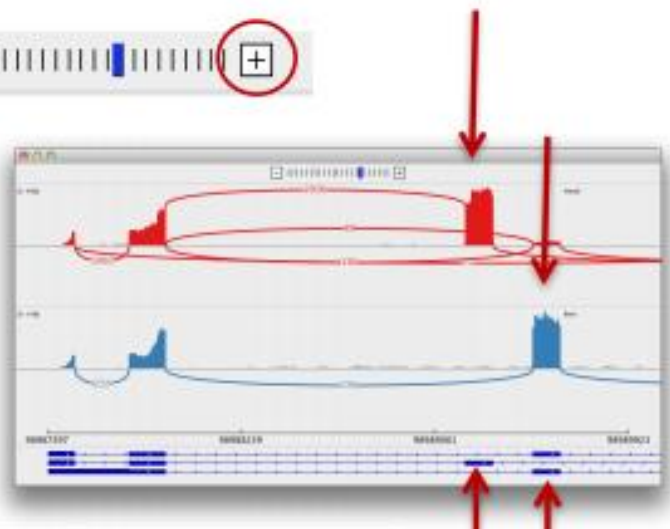
## 12. Zoom in on 5' end



Click "+" button 2 times

Click-and-drag tracks to the right to bring the first 3 exons in view.

## 13. Observe the alternative splicing of the 3<sup>rd</sup> exon



Close the Sashimi plot.

## Questions

6. A. In the liver sample, in the top read of the first exon, there is a green column, what does it mean?  
B. What is the reference genome base at the same location (coordinate)?
  
7. Go to chr21:33,979,695-34,159,621.  
A. Expand the RefSeq Genes track. Find 2 genes that are located head to head (5' UTR of both genes very close to each other). Zoom into this region.  
Which 2 genes are these? Do you think their promoters overlap?  
B. Which 2 genes have overlapping sequences in different orientations?  
C. Are there aligned reads in this region?
  
8. Go to chr12:99,041,218-99,051,312. Look at the blue and green reads in the intron of APAF1 in the liver sample. Explain their meaning.
  
9. Look at the Coverage tracks. Are the reads evenly distributed along the exons?

THE END