R for data analysis	
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3 steps to	
Basic data analysis	
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1. Reading in data	
• read.table()	
<ul><li>read.csv(), read.delim()</li></ul>	
2. Analysis	
Manipulating & reshaping the data	
Any maths you like	
<ul> <li>Plotting the outcome</li> <li>High level plotting functions (covered tomorrow)</li> </ul>	
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3. Writing out results	
write.table()	
• write.csv()	
A simple walkthrough	
Exemplifies 3 steps to R analysis	
Exemplines 3 steps to K analysis	
50 neuroblastoma patients were tested for NMYC gene copy number	
by interphase nuclei FISH	-
<ul> <li>Amplification of NMYC correlates with worse prognosis</li> </ul>	
We have count data	
<ul> <li>Numbers of cells per patient assayed</li> <li>For each we have NMYC copy number relative to base ploidy</li> </ul>	
<ul> <li>We need to determine which patients have amplifications</li> </ul>	
<ul> <li>(i.e &gt;33% of cells show NMYC amplification)</li> </ul>	

## Step 1. Read in the data

Patient	Nuclei	NB_Amp	NB_Nor	NB_Del
1	42	0	34	8
2	40	3	30	7
3	56	6	50	0
4	42	5	37	0
5	32	1	30	1
6	70	10	53	7
7	65	3	58	4
8	40	4	31	5
9	60	0	54	6
10	61	0	57	4
11	43	13	29	1

This data is a tab delimited text file Each row is a record, each column is a field Columns are separated by tabs in the text.

We need to read in the results table and assign it to an object (rawData)

If the data had been comma separated values, then sep="," read.csv("08\_NBcountData.csv")
?read.table for a full list of arguments

08\_NBcountData.R (script commands)

08\_NBcountData.txt (data file)

## Step 2. Analysis (reshaping data & maths)

- Our analysis involves identifying patients with > 33% NB amplification
  - $\label{lem:prop <- rawData$NB\_Amp / rawData$Nuclei $\#$ create an index of results}$
  - $\texttt{amp} \leftarrow \texttt{which(prop} > 0.33) \ \# \ \texttt{Get} \ \texttt{sample} \ \texttt{names} \ \texttt{of} \ \texttt{amplified} \ \texttt{patients}$
- We can plot a simple chart of the % NB amplification
  - plot(prop, ylim=c(0,1.2))abline(h=0.33,lwd=1.5,lty=2)



These 2 samples are amplified (11 & 23)

## Step 3. Outputting the results

- We write out a data frame of results (patients > 33% NB amplification) as a 'comma separated values' text file
  - write.csv(rawData[amp,],file="selectedSamples.csv") # Export table, file name = selectedSamples.csv
    - Files are directly readable by Excel and Calc
- ${\mbox{\ensuremath{\bullet}}}$  Its often helpful to double check where the data has been saved
  - Use get working directory function
    - getwd() # print working directory

Data analysis exercise: Which samples are near normal?	
Patients are near normal if:	
(NB_Amp/Nuclei <0.33 & NB_Del ==0)	
<ul> <li>Modify the condition in our previous code to find these patients</li> </ul>	
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<ul> <li>Write out a results file of the samples that match these criteria, and open it in a spreadsheet program</li> </ul>	
08_NBcountData.R (script commands)	
Solution to NB normality test  Basic data analysis	
<pre>&gt; norm &lt;- which( prop &lt; 0.33 &amp; rawData\$NB_Del==0) &gt; norm</pre>	
[1] 3 4 7 15 20 24 36 37 42 47	
> write.csv(rawData[norm,],"My_NB_output.csv")	