

Variant Data Comparison Between Websites with Machine Learning Analysis

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Contents

1	Abstract	3
2	Background	3
3	Data sources	3
4	Algorithms and Data Sets	3
4.1	Data Sets	3
4.2	Algorithms	4
4.2.1	Types of Clustering	4
4.2.1.1	Partitioning-based clustering	4
4.2.1.1.1	K-means	4
4.2.1.1.2	K-medoids (PAM)	4
4.2.1.2	Hierarchical clustering	5
4.2.1.2.1	Agglomerative hierarchical clustering	5
4.2.1.2.2	Divisive hierarchical clustering	5
5	Scraping Websites	5
5.1	Scraping the ExAC Website	5
5.2	Scraping the HGMD Website	5
6	Coding with R	10
6.1	Scraping Websites with RSelenium	18
6.2	Shaping the Data	25
6.2.1	Remove Duplicates	25
6.2.1.1	Verifying Results	32
6.3	Analyzing the Data	32
6.3.1	Cluster HGMD Data	33
6.3.1.1	Load Data.	33
6.3.1.2	Formating the data	33
6.3.2	Clustering	36
6.3.2.1	Number of Clusters	36
6.3.2.1.1	Averaged Silhouette Width and Gap Statistic	36
6.3.2.1.2	Hartigan's rule	38
6.3.2.1.3	Number of Clusters Results	39
6.3.2.2	Partitioning-based clustering	39
6.3.2.2.1	k-means	39
6.3.2.2.2	K-medoids clustering in R	45
6.3.3	Gap statistic	47
6.3.4	Hierarchical clustering in R	50
6.3.4.1	Plotting to determinine the cluster level.	56
7	Evaluation	57
7.1	System Time	57
7.2	Precision and Recall	57
8	Conclusion	58
9	References	58

1 Abstract

The final project for the class DA5030 Introduction to Data Mining and Machine Learning is a data mining project that scrapes data from the HGMD website and compares it to the ExAC website and reports the duplicate and unique variants. Then the unique data is analyzed using unsupervised learning algorithms.

2 Background

There is a website called, HGMD, which has a list of all published mutations in every gene. The user enters the specific gene and the website shows a list of all of the published mutations, which contains the variant and position. However, some of these publications are from 20 years ago where they might not be considered mutations today, now that we have more information. Genetic information is constantly being updated as more research is being conducted. To check if the published mutation is still considered a mutation, there is another website, ExAC, which has a list of all known variants for all known genes that are not mutations. The user can enter the specific gene and get a list of all the variants and the position. Then the variants and positions are compared from each website, if the two match, then the variant is no longer considered a mutation. Currently, geneticists need to perform this analysis manually by visually comparing the websites. The reason I want to do this project is because my wife and her colleges need to do this manually and her bioinformatics department is too busy at this time to find a solution. My hope is to find a solution and present it to her team and eventually get a job in their bioinformatics group.

To perform this project I plan to have the user enter the URL's with the gene info into R. Then R will scrape the websites and put the data into data frames. I will compare the data frames and print the differences. I am not sure of the details now, I will need to research the best method to do this (use R and compare or make a database with the info and use SQL). I want to also see if it is possible to enter the gene into R, as a text input, and have R search the database for the gene info then scrape the website, I am not sure if it is possible and I will need to research it.

This project is dependent on if it is possible to use R to scrape the information from the websites, I am not sure if it is possible with R and I am currently trying to figure it out. Also, I will need permission to use the HGMD website, it is very expensive to get a license and I will ask the company for a 2 month trial for a graduate research project. If they give me a trial then I can do it, if not, it is not feasible to pursue this project.

3 Data sources

Variant data used will be scraped from ExAC and HGMD websites.

4 Algorithms and Data Sets

4.1 Data Sets

The R packages used for this assignment are:

For scraping and shaping the data

- `rvest`[1]
- `RSelenium`[2]
- `plyr`[3]

For analysis

- ggplot2[4]
- cluster[5]
- amap[6]
- useful[7]

4.2 Algorithms

For analysis I will use unsupervised learning algorithms.

The goal for unsupervised learning is to model the underlying structure or distribution in the data in order to learn more about the data. Algorithms are left to their own devices to discover and present the interesting structure in the data[8].

4.2.1 Types of Clustering

Clustering (e.g., k-means, mixture models, hierarchical clustering). Cluster analysis or clustering is the task of grouping a set of objects in such a way that objects in the same group (called a cluster) are more similar (in some sense or another) to each other than to those in other groups (clusters)[9]. There are various types of cluster analysis.

- Partitioning-based clustering (K-means and its variants)
- Hierarchical clustering

4.2.1.1 Partitioning-based clustering

4.2.1.1.1 K-means

k-means clustering is a method of vector quantization, originally from signal processing, that is popular for cluster analysis in data mining. k-means clustering aims to partition n observations into k clusters in which each observation belongs to the cluster with the nearest mean, serving as a prototype of the cluster. This results in a partitioning of the data space into Voronoi cells[10].

4.2.1.1.2 K-medoids (PAM)

The K-medoids or Partitioning Around Medoids (PAM) algorithm (Kaufman & Rousseeuw'87) is related to the k-means algorithm and but uses medoid shifts rather than reassigning points based on Euclidean distance. Each cluster is represented by one of the objects (i.e. points) in the cluster A medoid is a point in a cluster whose dissimilarity to all the points in the cluster is minimal. Medoids are similar in concept to means or centroids, but medoids are always members of the data set. That is, in 2D Cartesian space a centroid can be any valid x,y coordinate. Whereas a medoid must be one of the data points[11].

Pseudocode for the k-medoid clustering (Partitioning Around Medoids (PAM)) algorithm:

Initialize: randomly select[citation needed] (without replacement) k of the n data points as the medoids.

Associate each data point to the closest medoid.

While the cost of the configuration decreases:

For each medoid m , for each non-medoid data point o :
Swap m and o , recompute the cost

If the total cost of the configuration increased in the previous step, undo the swap.

4.2.1.2 Hierarchical clustering

In hierarchical clustering the idea is to group data objects (i.e. points) into a tree of clusters. That is, hierarchical clustering is a method of cluster analysis which seeks to build a hierarchy of clusters[12].

These trees (hierarchies) generally fall into two types:

4.2.1.2.1 Agglomerative hierarchical clustering

Initially each data object (i.e. point) is in its own cluster. Iteratively the clusters are merged together from the “bottom-up.” The two most similar/closest objects are aggregated into the same cluster/data object. Then the next two, until there is just one cluster/data object. This agglomerative approach results in straggly (long and thin) clusters due to a chaining effect. It is also sensitive to noise[11].

4.2.1.2.2 Divisive hierarchical clustering

In divisive hierarchical clustering all data objects (i.e. points) are initially in one cluster. These clusters are successively divided recursively in a “top-down” manner. The cluster is broken into two clusters that are most dissimilar. Then each of those clusters is broken into two clusters that are most dissimilar. This continues until each cluster is a single data object (i.e. point)[11].

5 Scraping Websites

Below is a walkthrough without showing the code. I did this because I needed to use if statements and the first chunk of code is over 250 lines. This section will show what the program does and the next section will show the code.

In order to scrape the ExAC website I need to use RSelenium and rvest. This is because the table was constructed using javascript, specifically jQuery. When a website makes use of JavaScript to display data, the rvest and XML packages miss the required functionality.

Selenium is a web automation tool that literally “drives” the browser, so it can see anything you see when you right click and inspect element in Chrome or Firefox, making it possible to scrape the information with rvest. This vastly widens the universe of content that can be extracted from automation, but can be slow as all content must be rendered in the browser.

The great thing about RSelenium is that the user does not need to download and install a Selenium server, RSelenium will download it and run it automatically. Below is the R code.

5.1 Scraping the ExAC Website

Once RSelenium is up and running, the website is opened through RSelenium as defined in the code, figure 1 shows an example of what it looks like.

Then the gene name is automatically entered (this is an input in the code) and RSelenium searches for the gene and scrapes the data and puts it into a data frame. Figure 2 shows an image of the website.

5.2 Scraping the HGMD Website

Now that the ExAC data has been collected it is time to move to HGMD. I began by opening the website, see figure 3 as reference.

Then the username and password were entered automatically, which are defined in the code.

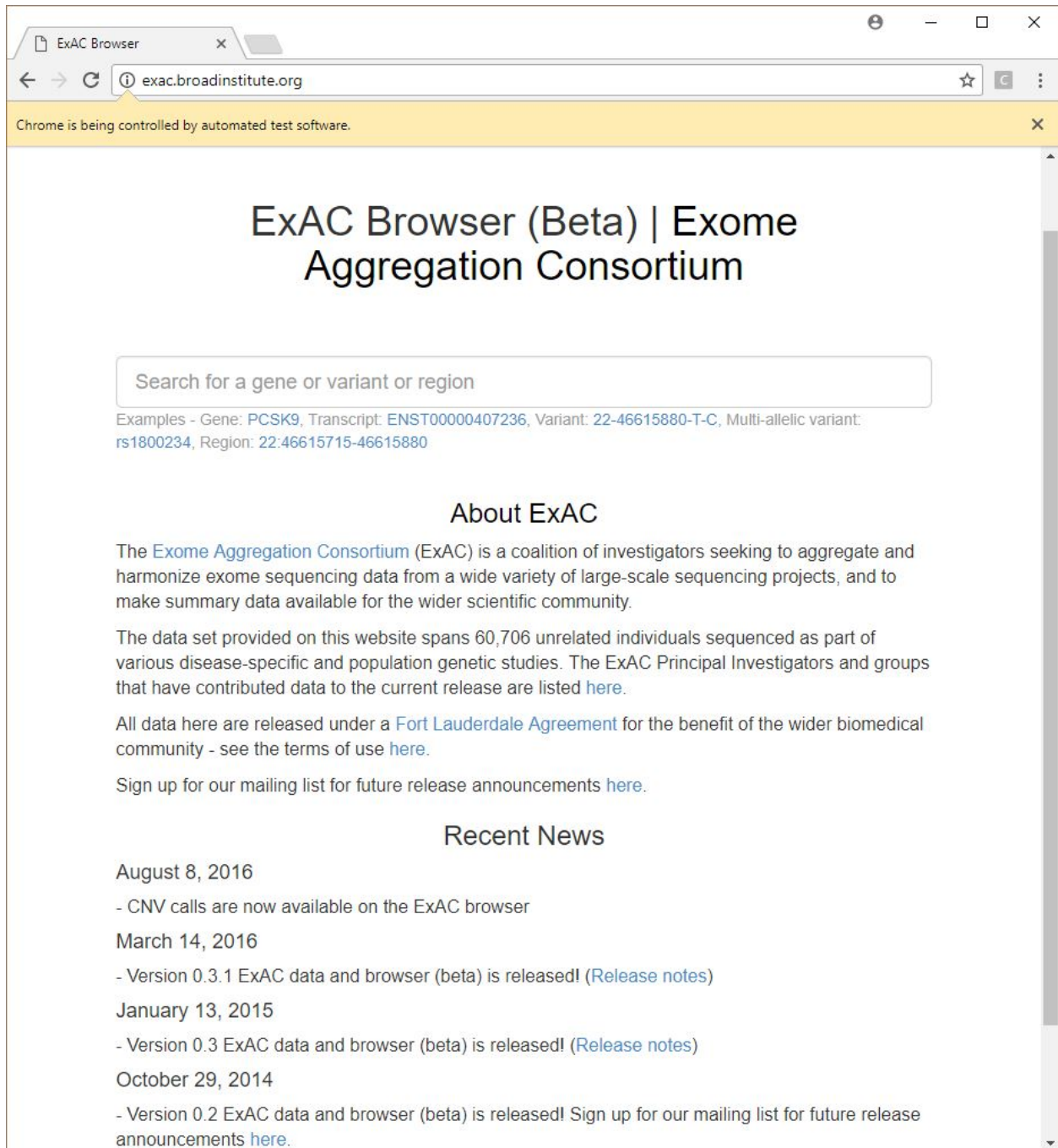


Figure 1: ExAC Webpage

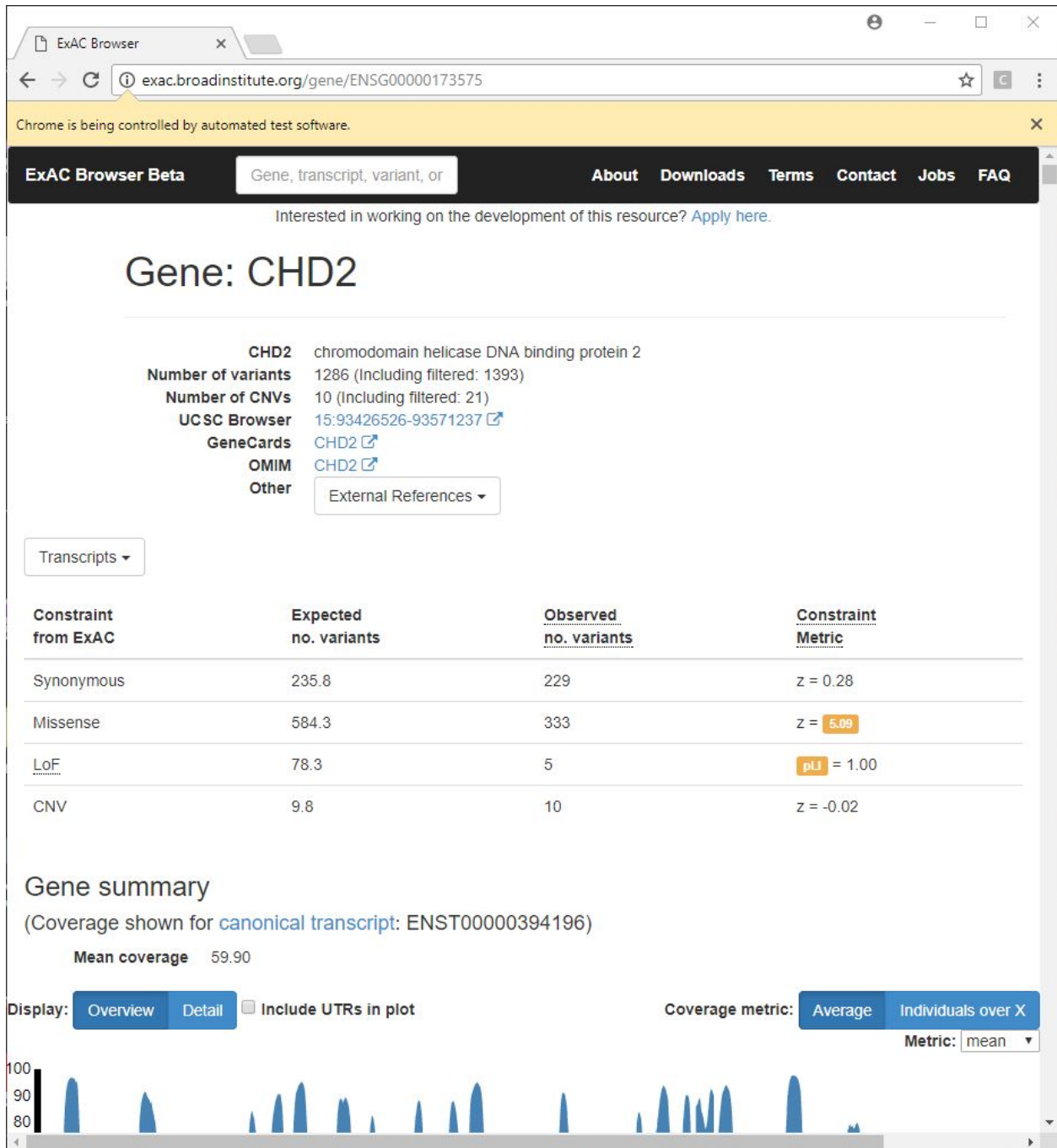


Figure 2: ExAC CHD2 Summary

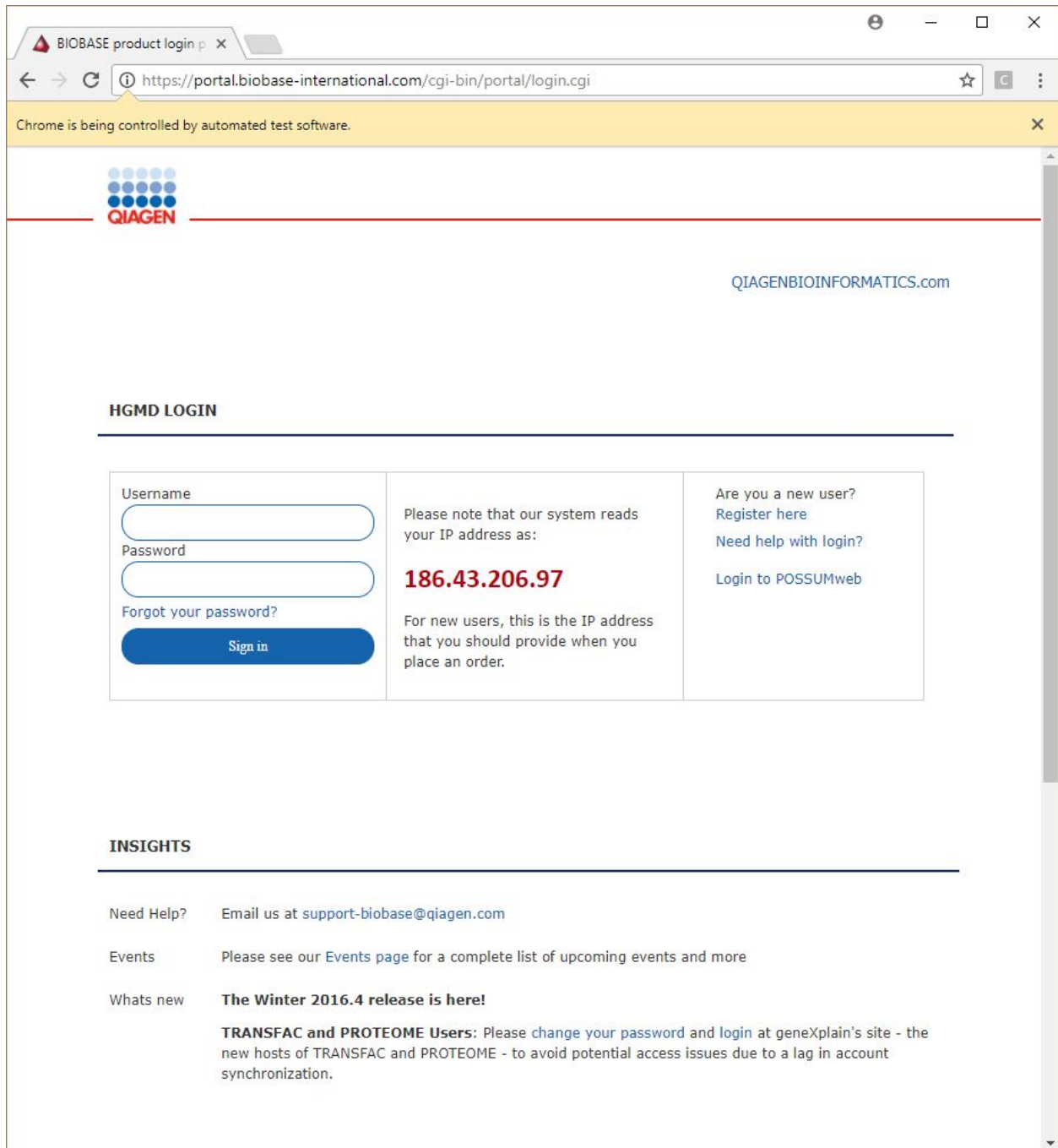


Figure 3: HGMD Webpage

The screenshot shows a web browser window with the address bar displaying `https://portal.biobase-international.com/cgi-bin/portal/login.cgi`. A yellow notification bar at the top states "Chrome is being controlled by automated test software." The page header includes the QIAGEN logo and the text "QIAGENBIOINFORMATICS.com" and "BIOBASE BIOLOGICAL DATABASES A QIAGEN® Company".

The main content area features a promotional banner for ANNOVAR with HGMD data, stating "Pair ANNOVAR with HGMD data for efficient annotation of genomes" and a "Learn more" button. Below this, a diagram illustrates the workflow: "INPUT" (a DNA sequence) is processed by "ANNOVAR" and "HGMD" (represented as gears) to produce "OUTPUT" (a disease annotation).

A "Subscription Renewal" section indicates that the user's subscription expires in "05 days" and advises them to "Contact your account manager to renew today".

A "Need Help?" section provides contact information for support.

A "Release Updates" section mentions that "The 2016.3 release is now available."

At the bottom, a "You are currently subscribed to the following product(s):" section lists "HGMD® trial" with a "Logout" button.

Figure 4: HGMD Webpage After Password

Figure 4 is what the website looks like after the username and password are entered.

Now the link needs to be “clicked”. Figure 5 shows the website after the linked has been “clicked”.

Then an authentication popup window appears where the username and password need to be entered. RSe-
lenium deals with this by putting the username and password into the URL. The code chunk is shown below:

```
# An authentication window pops up, to get around this
# I need to switch to the main window
# get all windows
windows <- remDr$getWindowHandles()

# Swith to main window
remDr$switchToWindow(windows[[1]])

# This enters the username and password into the url
HGMD_url2 <-
  paste(
    "http://",
    username,
    ":",
    password,
    "@hgmdtrial.biobase-international.com/hgmd/pro/start.php",
    sep = ""
  )

# This goes to the HGMD website
remDr$navigate(HGMD_url2)

# Swith to other window
remDr$switchToWindow(windows[[2]])

# Close the window
remDr$closeWindow()

# Swith to main window
remDr$switchToWindow(windows[[1]])
```

Figure 6 is what the website looks like after the authentication information is entered.

Next, RSelenium “clicks” the gene tab in the upper left corner to search for the gene. Figure 7 shows the result and below id the code.

Next, the gene is entered in the textbox and “enter” is pressed. Then a webpage is displayed with a summary of the gene, as shown in figure 8.

Next, the “Get all mutations” link is pressed and this opens a webpage with all of the mutations in four different categories: Missense, splicing, deletions, and insertion. These are the four tables that are needed to be scraped. The webpage are shown in the images below.

6 Coding with R

R studio was configured with the following parameters before beginning the project:

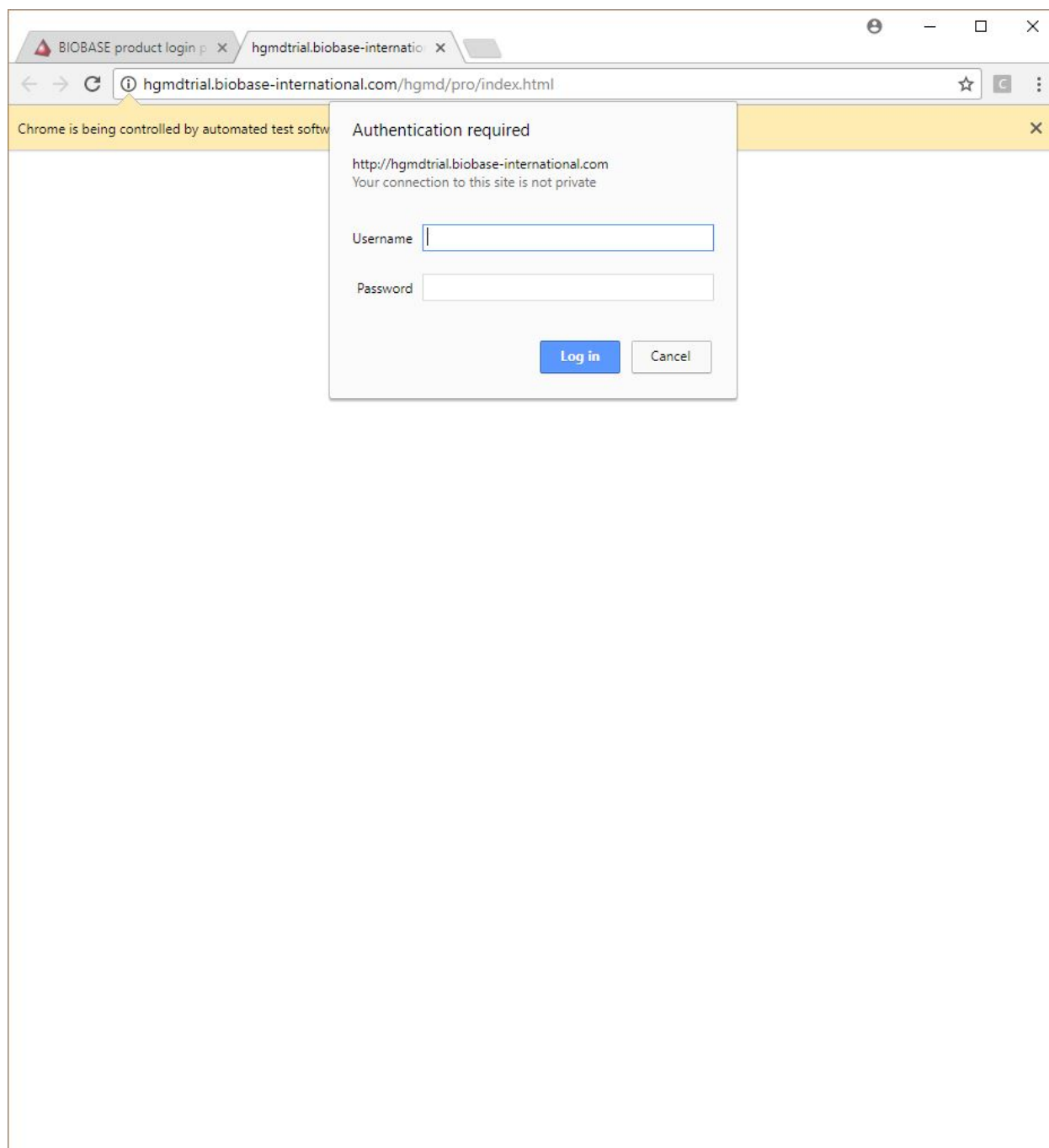


Figure 5: HGMD Webpage After Click

hgmdtrial.biobase-international.com/hgmd/pro/start.php

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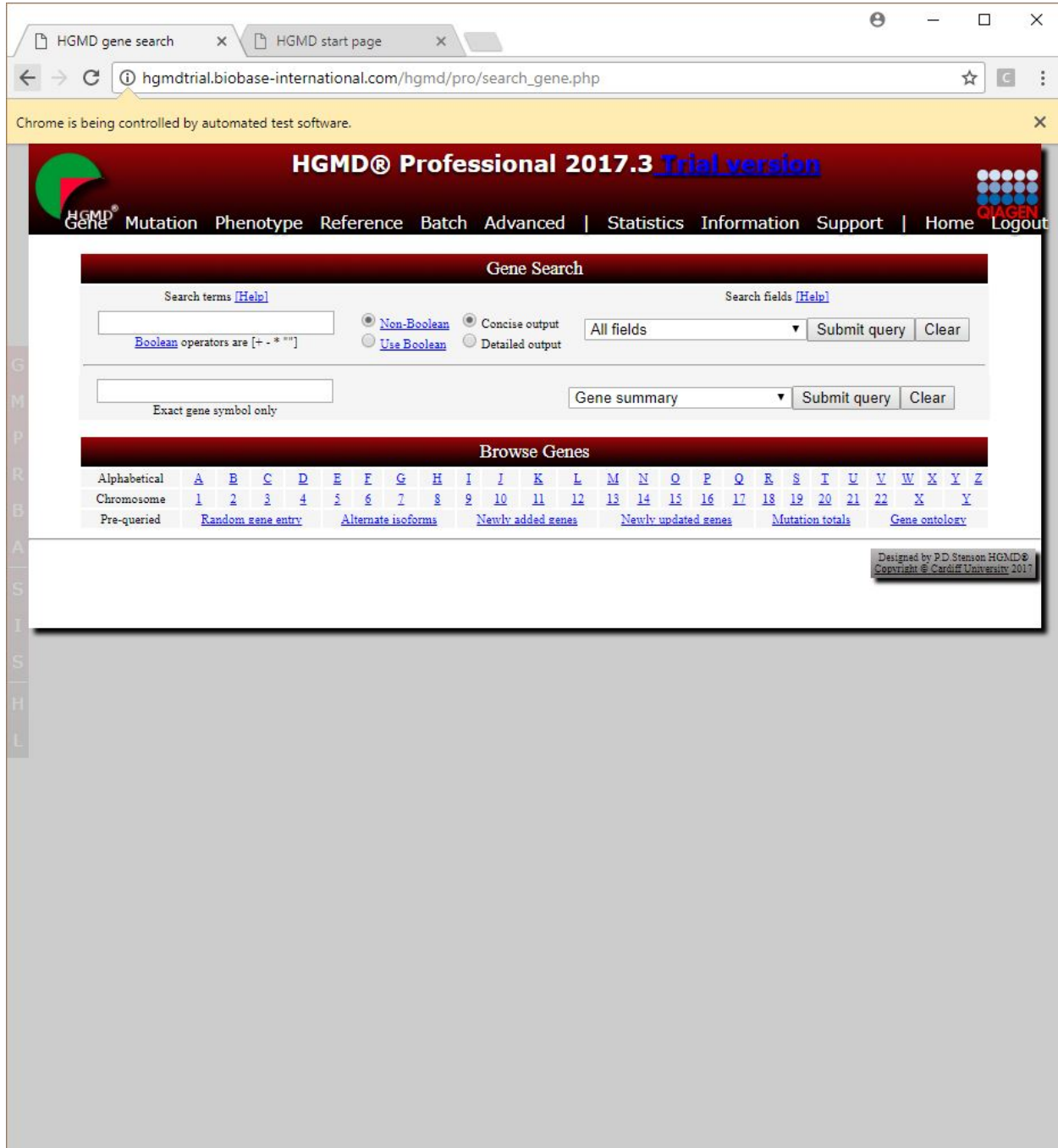
Gene Mutation Phenotype Reference Batch Advanced | Statistics Information Support | Home Logout

HGMD 2017.3 comprises the following core tables:

Data type:	Description:	Entries:
Missense/nonsense	Single base-pair substitutions in coding regions are represented as a triplet codon change.	120901
Splicing	Substitutions with consequences for mRNA splicing are presented in brief with information specifying the relative position of the lesion with respect to a numbered intron donor or acceptor splice site. Positions given as positive integers refer to a 3' (downstream) location, negative integers refer to a 5' (upstream) location.	19247
Regulatory	Substitutions causing regulatory abnormalities are logged in with thirty nucleotides flanking the site of the mutation on both sides. The location of the mutation relative to the transcriptional initiation site, initiation codon, polyadenylation site or termination codon is given.	3925
Small deletions	Micro-deletions (20 bp or less) are presented in terms of the deleted bases in lower case plus, in upper case, 10 bp DNA sequence flanking both sides of the lesion. The numbered codon is preceded in the given sequence by the caret character (^).	31571
Small insertions/duplications	Micro-insertions (20 bp or less) are presented in terms of the inserted bases in lower case plus, in upper case, 10 bp DNA sequence flanking both sides of the lesion. The numbered codon is preceded in the given sequence by the caret character (^).	13167
Small indels	Micro-indels (20 bp or less) are presented in terms of the deleted/inserted bases in lower case plus, in upper case, 10 bp DNA sequence flanking both sides of the lesion. The numbered codon is preceded in the given sequence by the caret character (^).	2970
Gross deletions	Information regarding the nature and location of each lesion is logged in narrative form because of the extremely variable quality of the original data reported.	15999
Gross insertions/duplications	Information regarding the nature and location of each lesion is logged in narrative form because of the extremely variable quality of the original data reported.	3942
Complex rearrangements	Information regarding the nature and location of each lesion is logged in narrative form because of the extremely variable quality of the original data reported.	1925
Repeat variations	Information regarding the nature and location of each lesion is logged in narrative form because of the extremely variable quality of the original data reported.	511
Mutation total		214158
With chromosomal coordinates (GRCh38/hg38)		189578
With HGVS descriptions		190134

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Figure 6: HGMD Webpage Database



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Gene Search

Search terms [\[Help\]](#) Search fields [\[Help\]](#)

☒ Non-Boolean ☐ Concise output ☐ Detailed output All fields

[Boolean](#) operators are [+ - * **]

☐ Use Boolean ☐ Detailed output

Exact gene symbol only Gene summary

Browse Genes

Alphabetical	A	B	C	D	E	F	G	H	I	J	K	L	M	N	O	P	Q	R	S	T	U	V	W	X	Y	Z
Chromosome	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	X	Y		
Pre-queried	Random gene entry		Alternate isoforms		Newly added genes		Newly updated genes		Mutation totals		Gene ontology															

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Figure 7: HGMD Webpage for Genes

hgmdtrial.biobase-international.com/hgmd/pro/gene.php?gene=CHD2

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Gene Symbol	Location	Gene description	cDNA sequence	Extended cDNA	RefSeqGene	cDNA viewer	Viewer (DEPRECATED)
CHD2 (Aliases: EEOC)	15q26	Chromodomain helicase DNA binding protein 2 (Aliases: ATP-dependent helicase CHD2; CHD-2)	NM_001271.3	Extended cDNA	NG_012826.1	CDS mutations	View mutations

Mutation type	Total number of mutations	Mutation data sorted by location
Missense/nonsense	19	Get missense/nonsense
Splicing substitutions	4	Get splicing
Regulatory substitutions	0	No mutations
Small deletions	7	Get small deletions
Small insertions/duplications	4	Get small insertions
Small indels	0	No mutations
Gross deletions	8	Get gross deletions
Gross insertions/duplications	0	No mutations
Complex rearrangements	1	Get complex rearrangements
Repeat variations	0	No mutations
TOTAL	43	Get all mutations

Variant class	Number of mutations	Mutation data by class
DM	24	Get all DM
DM?	19	Get all DM?

Disease/phenotype	Number of mutations	Mutation data by disease/phenotype
Autism spectrum disorder	7	Phenotype ID 1816339378
Epileptic encephalopathy	6	Phenotype ID 2012727453
Autism	4	Phenotype ID 1086878777
Developmental delay, intellectual disability & epilepsy	4	Phenotype ID 1121202983
Dravet syndrome-like myoclonic epileptic encephalopathy	3	Phenotype ID 114269671
Lennox-Gastaut syndrome	3	Phenotype ID 1009838363
Epilepsy, photosensitive	2	Phenotype ID 1138588393
Intellectual disability	2	Phenotype ID 84606745

Figure 8: HGMD Webpage for Gene Summary

HGMD Mutation Result x HGMD start page x

hgmdtrial.biobase-international.com/hgmd/pro/all.php

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[missense/nonsense](#) [splicing](#) [small deletions](#) [small insertions](#) [gross deletions](#) [complex rearrangements](#)

Missense/nonsense : 19 mutations [\[back to top\]](#)

HGMD accession	HGMD codon change	HGMD amino acid change	HGVS (nucleotide)	HGVS (protein)	Variant class	Reported phenotype	Reference	Extra information
CM1410645	TCA-TGA	Ser112Term	c.335C>G	p.S112*	DM	Intellectual disability	Hamdan (2014) <i>PLoS Genet</i> 10, e1004772	hg38 hg19 dbSNP
CM1314729	CGA-TGA	Arg121Term	c.361C>T	p.R121*	DM	Epileptic encephalopathy	Carvill (2013) <i>Nat Genet</i> 45, 825	hg38 hg19 CpG dbSNP
CM153783	CCC-CTC	Pro218Leu	c.653C>T	p.P218L	DM?	Eyelid myoclonia with absences	Galizia (2015) <i>Brain</i> 138, 1198	hg38 hg19 COM
CM1311116	CGA-TGA	Arg466Term	c.1396C>T	p.R466*	DM	Dravet syndrome-like myoclonic epileptic encephalopathy	Suls (2013) <i>Am J Hum Genet</i> 93, 967	hg38 hg19 CpG dbSNP
CM1314730	CTG-CCG	Leu823Pro	c.2468T>C	p.L823P	DM	Epileptic encephalopathy	Carvill (2013) <i>Nat Genet</i> 45, 825	hg38 hg19 dbSNP
CM125579	GAT-GGT	Asp856Gly	c.2567A>G	p.D856G	DM?	Autism	Neale (2012) <i>Nature</i> 485, 242 Uddin (2014) <i>Nat Genet</i> 46, 742 [Additional report]	hg38 hg19 dbSNP
CM177698	CAA-TAA	Gln906Term	c.2716C>T	p.Q906*	DM?	Autism spectrum disorder	Yuen (2017) <i>Nat Neurosci</i> 20, 602	hg38 hg19 COM
CM1617343	CGG-CAG	Arg1000Gln	c.2999G>A	p.R1000Q	DM?	Autism spectrum disorder	Wang (2016) <i>Nat Commun</i> 7, 13316	hg38 hg19 COM CpG
CM1617344	GGT-GAT	Gly1174Asp	c.3521G>A	p.G1174D	DM?	Autism spectrum disorder	Wang (2016) <i>Nat Commun</i> 7, 13316	hg38 hg19 COM
CM153791	CGA-GGA	Arg1313Gly	c.3937C>G	p.R1313G	DM	Epilepsy, photosensitive	Galizia (2015) <i>Brain</i> 138, 1198	hg38 hg19
CM153778	CGG-CAG	Arg1345Gln	c.4034G>A	p.R1345Q	DM	Epilepsy, juvenile myoclonic	Galizia (2015) <i>Brain</i> 138, 1198	hg38 hg19 CpG
CM177497	AGT-GGT	Ser1406Gly	c.4216A>G	p.S1406G	DM?	Autism spectrum disorder	Yuen (2017) <i>Nat Neurosci</i> 20, 602	hg38 hg19 COM
CM153792	GTG-ATG	Val1479Met	c.4435G>A	p.V1479M	DM	Photoparoxysmal response with febrile seizures	Galizia (2015) <i>Brain</i> 138, 1198	hg38 hg19 CpG dbSNP
CM162166	TGG-TGT	Trp1534Cys	c.4602G>T	p.W1534C	DM?	Epileptic encephalopathy, early onset	Helbig (2016) <i>Genet Med Epub, Epub</i>	hg38 hg19 COM
CM153777	CGA-TGA	Arg1637Term	c.4909C>T	p.R1637*	DM	Epilepsy, idiopathic photosensitive occipital	Galizia (2015) <i>Brain</i> 138, 1198 O'Roak (2014) <i>Nat Commun</i> 5, 5595 [Additional phenotype]	hg38 hg19 CpG dbSNP
CM1414846	CAG-TAG	Gln1641Term	c.4921C>T	p.Q1641*	DM?	Autism	O'Roak (2014) <i>Nat Commun</i> 5, 5595	hg38 hg19 dbSNP

Figure 9: HGMD Webpage for Missense

hgmdtrial.biobase-international.com/hgmd/pro/all.php

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Splicing : 4 mutations [\[back to top\]](#)

HGMD accession	HGMD splicing mutation	HGVS (nucleotide)	Variant class	Reported phenotype	Reference	Extra information
CS149377	IVS3 as C-T +9	c.390C>T	DM?	Lennox-Gastaut syndrome	Appenzeller (2014) Am J Hum Genet 95, 360	hg38 hg19 COM CpG
CS1314621	IVS12 ds G-A +1	c.1502+1G>A	DM	Lennox-Gastaut syndrome	Allen (2013) Nature 501, 217	hg38 hg19 COM dbSNP
CS153788	IVS13 ds G-A +5	c.1719+5G>A	DM	Epilepsy, photosensitive	Galizia (2015) Brain 138, 1198	hg38 hg19
CS1311117	IVS14 as A-C -2	c.1810-2A>C	DM	Dravet syndrome-like myoclonic epileptic encephalopathy	Suls (2013) Am J Hum Genet 93, 967	hg38 hg19 dbSNP

Small deletions : 7 mutations [\[back to top\]](#)

HGMD accession	HGMD deletion	HGVS (nucleotide)	Variant class	Reported phenotype	Reference	Extra information
CD177189	GAAAG ^{Δ16} ACCATcCAGACCATAT	c.1552delC	DM?	Moderate intellectual disability, autism spectrum disorder and obsessive compulsive disorder	Bowling (2017) Genome Med 9,	hg38 hg19 COM
CD129351	GAAAGAT ^{Δ603} AAG_E15I15_gTGTGTAATTA	c.1809+1delG	DM?	Intellectual disability, nonsyndromic	Rauch (2012) Lancet 380, 1674 O'Roak (2014) Nat Commun 5, 5595 [Additional phenotype]	hg38 hg19 COM dbSNP
CD1613710	AATGAT ^{Δ626} GACTctttATTGTATAAA	c.1880_1883delCTTT	DM	Epileptic encephalopathy	Moller (2016) Mol Syndromol 7, 210	hg38 hg19
CD1414848	CTTTT ^{Δ964} AATAAagaaGAGCTGACAG	c.2895_2898delAGAA	DM?	Autism	O'Roak (2014) Nat Commun 5, 5595 Bi (2012)	hg38 hg19 COM

Figure 10: HGMD Webpage for Splicing and Deletions

hgmdtrial.biobase-international.com/hgmd/pro/all.php

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Small insertions : 4 mutations [\[back to top\]](#)

HGMD accession	HGMD insertion	HGVS (nucleotide)	Variant class	Reported phenotype	Reference	Extra information
CI177155	GAGTGG ¹²⁶² GGGGgTGAAGATGA	c.3787dupG	DM?	Moderate intellectual disability, seizures and speech delay	Bowling (2017) Genome Med 9,	h38 h39 COM
CI143035	GAAAAAA ¹³⁹¹ AAAaCAGAAGAAGA	c.4173dupA	DM	Lennox-Gastaut syndrome	Lund (2014) Epilepsy Behav 33C, 18 Galante (2015) Brain 138: 1198 [Additional phenotype] Pinto (2016) Brain Dev : [Additional phenotype]	h38 h39
CI1414847	ACTAT ¹⁶⁴⁹ GGTGGgTGGCAACAAC	c.4949dupG	DM?	Autism	O'Roak (2014) Nat Commun 5, 5595	h38 h39
CI170083	CAGAAAG ¹⁶⁹⁸ AGGgCCTTATGACC	c.5094dupG	DM	Intellectual disability	Gauthier-Vassero (2017) Am J Med Genet A 173, 62	h38 h39 COM

Gross deletions : 8 mutations [\[back to top\]](#)

HGMD accession	DNA level	Description	Variant class	Reported phenotype	Reference	Extra information
CG145323	gDNA	~191 kb incl. entire gene	DM	Developmental delay, intellectual disability & epilepsy	Chénier (2014) J Neurodev Disord 6, 9	
CG145324	gDNA	~210 kb incl. entire gene	DM	Developmental delay, intellectual disability & epilepsy	Chénier (2014) J Neurodev Disord 6, 9	
CG145326	gDNA	~237 kb incl. entire gene & RGMA	DM	Developmental delay, intellectual disability & epilepsy	Chénier (2014) J Neurodev Disord 6, 9	
CG145325	gDNA	~78 kb incl. part of gene	DM	Developmental delay, intellectual disability & epilepsy	Chénier (2014) J Neurodev Disord 6, 9	
CG145057	gDNA	~83 kb partial gene	DM?	Autism spectrum disorder	Pinto (2014) Am J Hum Genet 94, 677	COM
CG149829	gDNA	106kb incl entire gene & RGMA	DM	Intellectual disability, epilepsy & truncal obesity	Courage (2014) Eur J Med Genet 57, 520	
CG149828	gDNA	415kb incl entire gene & RGMA	DM	Intellectual disability, epilepsy & truncal obesity	Courage (2014) Eur J Med Genet 57, 520	
CG121687	gDNA	511kb incl entire gene & RMGA	DM	Epilepsy, developmental delay, autism & facial dysmorphism	Capelli (2012) Eur J Med Genet 55, 132	COM

Complex rearrangements : 1 mutation [\[back to top\]](#)

HGMD accession	Description	Variant class	Reported phenotype	Reference	Extra information
----------------	-------------	---------------	--------------------	-----------	-------------------

Figure 11: HGMD Webpage for Insertions

```

# clears the console in RStudio
cat("\014")

# clears environment
rm(list = ls())

# Set working directory
setwd("C:/R/DA5030/final project")

# Load required packages
# For scraping and shaping the data
library(rvest)
library(RSelenium)
library(plyr)

# For analysis
require(ggplot2)
require(cluster)
require(ama)
require(useful)

```

6.1 Scraping Websites with RSelenium

This section shows the code that was used to scrape the websites. Unfortunately, I only had a week trial to HGMD website so my username and password will no longer work. To work around this, I scraped and saved the HGMD tables as CSV files and this code will open those up instead of scraping the website.

To start, the user needs to enter a gene to search, since I no longer have access to HGMD, this code will work with genes CHD2 and HCN2 with the CSV files provide:

```

# Enter the gene to search
gene <- "CHD2"

```

Once the gene is entered, RSelenium will start. I want to add that there are a couple of ‘if’ statements that are in the code. The first checks if the gene entered is an actual gene, if it is not it lets the user know that it is not a valid gene and stops the program. The next if statement checks to see if the user name and password are valid for the HGMD website, if they are not, the code will load csv files that are provided with the code, if the username and password are good, RSelenium will work as intended.

```

# From https://stackoverflow.com/questions/42468831/how-to-set-up-rselenium-for-r
# This runs a selenium server with chrome browser, wait for necessary files to
# download
rD <- rsDriver()

```

```
## checking Selenium Server versions:
```

```
## BEGIN: PREDOWNLOAD
```

```
## BEGIN: DOWNLOAD
```

```
## BEGIN: POSTDOWNLOAD
```

```
## checking chromedriver versions:
```

```
## BEGIN: PREDOWNLOAD
```

```
## BEGIN: DOWNLOAD
```

```
## BEGIN: POSTDOWNLOAD
## checking geckodriver versions:
## BEGIN: PREDOWNLOAD
## BEGIN: DOWNLOAD
## BEGIN: POSTDOWNLOAD
## checking phantomjs versions:
## BEGIN: PREDOWNLOAD
## BEGIN: DOWNLOAD
## BEGIN: POSTDOWNLOAD
## [1] "Connecting to remote server"
## $applicationCacheEnabled
## [1] FALSE
##
## $rotatable
## [1] FALSE
##
## $mobileEmulationEnabled
## [1] FALSE
##
## $networkConnectionEnabled
## [1] FALSE
##
## $chrome
## $chrome$chromedriverVersion
## [1] "2.33.506120 (e3e53437346286c0bc2d2dc9aa4915ba81d9023f)"
##
## $chrome$userDataDir
## [1] "C:\\Users\\Josh\\AppData\\Local\\Temp\\scoped_dir16188_23106"
##
##
## $takesHeapSnapshot
## [1] TRUE
##
## $pageLoadStrategy
## [1] "normal"
##
## $databaseEnabled
## [1] FALSE
##
## $handlesAlerts
## [1] TRUE
##
## $hasTouchScreen
## [1] FALSE
##
## $version
## [1] "62.0.3202.94"
##
## $platform
```

```
## [1] "Windows NT"
##
## $browserConnectionEnabled
## [1] FALSE
##
## $nativeEvents
## [1] TRUE
##
## $acceptSslCerts
## [1] TRUE
##
## $locationContextEnabled
## [1] TRUE
##
## $webStorageEnabled
## [1] TRUE
##
## $browserName
## [1] "chrome"
##
## $takesScreenshot
## [1] TRUE
##
## $javascriptEnabled
## [1] TRUE
##
## $cssSelectorsEnabled
## [1] TRUE
##
## $setWindowRect
## [1] TRUE
##
## $unexpectedAlertBehaviour
## [1] ""
##
## $id
## [1] "0f09fa9b620ad036bbbe9e0f881ed46d"

# puts a variable to the session
remDr <- rD$client

# Search ExAC

# URL for ExAC
exac_url <- "http://exac.broadinstitute.org/"

# This goes to the exac website
remDr$navigate(exac_url)

# This enters the gene and hits enter. This takes the user to the gene site
# \uE007 simulates "enter".
webElem <-
  remDr$findElement(using = 'css selector', "#home-searchbox-input")
webElem$sendKeysToElement(list(gene, "\uE007"))
```

```

# This extracts the contents of the website
exac_html <- read_html(remDr$getPageSource()[[1]])

# This uses rvest to get the table from Selenium
exac <- exac_html %>%
  html_nodes("#variant_table") %>%
  html_table
dfExac = as.data.frame(exac)

# This makes sure a valid entry was made using an if statement.
# If a valid gene is not entered it ends the program and lets the
# user know that it was not a valid gene.
if (length(dfExac) == 0) {
  # If a valid name was not entered then the program stops with this message.
  paste("This is not a valid gene, please try again")

  # close Selenium
  remDr$close()
} else {
  # If a valid gene was entered, then the program will move to the HGMD website

  # Search HGMD

  # Username for HGMD
  username <- "joshua_husk"

  # Password for HGMD
  password <- "trial"

  # URL for HGMD
  HGMD_url <-
    "https://portal.biobase-international.com/cgi-bin/portal/login.cgi"

  # This goes to the HGMD website
  remDr$navigate(HGMD_url)

  # This enters the Username
  webElem <-
    remDr$findElement(
      using = 'css selector',
      "#login_form > table:nth-child(1) > tbody:nth-child(1) > tr:nth-child(2) > td:nth-child(1) > label"
    )
  webElem$sendKeysToElement(list(username))

  # This enters the password and hits enter
  webElem <-
    remDr$findElement(
      using = 'css selector',
      "#login_form > table:nth-child(1) > tbody:nth-child(1) > tr:nth-child(4) > td:nth-child(1) > label"
    )
  webElem$sendKeysToElement(list(password, "\uE007"))
}

```

```

exac_html <- read_html(remDr$pageSource()[[1]])

# This section of the code checks to see if the username and password are correct

# This will give 'Error! No subscription' if the subscription is expired
checkOutput <- exac_html %>%
  html_nodes("#login_form > table > tbody > tr:nth-child(8) > td > span") %>%
  html_text()

# This will give 'Error: A wrong username or password was entered.' Or 'Error:
# Subscription has expired.' if the wrong info was entered
checkOutput2 <- exac_html %>%
  html_nodes("#login_form > table > tbody > tr:nth-child(8) > td > span > font") %>%
  html_text()

# If password and user name are bad this will load the excel files
# If the password and user name are good, then it will scrape the website

if (checkOutput == "Error! No subscription" |
    checkOutput2 == "Error: A wrong username or password was entered. " |
    checkOutput2 == "Error: Subscription has expired.") {
  # close Selenium
  remDr$close()

  # This will open the CSV files after it checks if it is empty, if the files
  # are empty then it sets it to zero

  if (file.info(paste("dfHGMDMissense_", gene, ".csv", sep = ""))$size > 5) {
    dfHGMDMissense <- read.csv2(
      paste("dfHGMDMissense_", gene, ".csv", sep = ""),
      sep = ",",
      stringsAsFactors = FALSE
    )
  } else {
    dfHGMDMissense <- 0
  }

  if (file.info(paste("dfHGMDSplice_", gene, ".csv", sep = ""))$size > 5) {
    dfHGMDSplice <- read.csv2(
      paste("dfHGMDSplice_", gene, ".csv", sep = ""),
      sep = ",",
      stringsAsFactors = FALSE
    )
  } else {
    dfHGMDSplice <- 0
  }

  if (file.info(paste("dfHGMDDelete_", gene, ".csv", sep = ""))$size > 5) {
    dfHGMDDelete <- read.csv2(
      paste("dfHGMDDelete_", gene, ".csv", sep = ""),
      sep = ",",

```

```

    stringsAsFactors = FALSE
  )
} else {
  dfHGMDDelete <- 0
}

if (file.info(paste("dfHGMDInsert_", gene, ".csv", sep = ""))$size > 5) {
  dfHGMDInsert <- read.csv2(
    paste("dfHGMDInsert_", gene, ".csv", sep = ""),
    sep = ",",
    stringsAsFactors = FALSE
  )
} else {
  dfHGMDInsert <- 0
}

} else {
  # This will scrape the website
  # This finds the link to the database
  webElem <-
    remDr$findElement(
      using = 'css selector',
      "#form-template > tbody > tr:nth-child(2) > td > div > table > tbody > tr:nth-child(2) > td > a"
    )
  # click the search link
  webElem$clickElement()

  # An authentication window pops up, to get around this
  # I need to switch to the main window
  # get all windows
  windows <- remDr$getWindowHandles()

  # Switch to main window
  remDr$switchToWindow(windows[[1]])

  # This enters the username and password into the url
  HGMD_url2 <-
    paste(
      "http://",
      username,
      ":",
      password,
      "@hgmdtrial.biobase-international.com/hgmd/pro/start.php",
      sep = ""
    )

  # This goes to the HGMD website
  remDr$navigate(HGMD_url2)

  # Switch to other window
  remDr$switchToWindow(windows[[2]])

```

```

# Close the window
remDr$closeWindow()

# Swith to main window
remDr$switchToWindow(windows[[1]])

# This finds the gene link
webElem <-
  remDr$findElement(using = 'css selector',
    "body > div.top > div.links > a:nth-child(1)")

# click the search link
webElem$clickElement()

# This enters the gene and hits enter
webElem <-
  remDr$findElement(
    using = 'css selector',
    'body > div.content > form:nth-child(2) > table > tbody > tr > td:nth-child(1) > input[type="text"]'
  )
webElem$sendKeysToElement(list(gene, "\uE007"))

# This finds the mutation link
webElem <-
  remDr$findElement(
    using = 'css selector',
    'body > div.content > form > table:nth-child(5) > tbody > tr:nth-child(12) > td:nth-child(3) > a'
  )

# click the search link
webElem$clickElement()

# This extracts the contents of the website
HGMD_html <- read_html(remDr$getPageSource()[[1]])

# This uses rvest to get the missense table from Selenium
HGMDMissense <- HGMD_html %>%
  html_nodes("body > div.content > table:nth-child(6)") %>%
  html_table
dfHGMDMissense = as.data.frame(HGMDMissense)
colnames(dfHGMDMissense)[3] <- "Consequence"

# This uses rvest to get thesplicing table from Selenium
HGMDSplice <- HGMD_html %>%
  html_nodes("body > div.content > table:nth-child(9)") %>%
  html_table
dfHGMDSplice = as.data.frame(HGMDSplice)
colnames(dfHGMDSplice)[3] <- "Consequence"

# This uses rvest to get thesplicing table from Selenium
HGMDDelete <- HGMD_html %>%
  html_nodes("body > div.content > table:nth-child(12)") %>%
  html_table

```



```

dfHGMDDelete = as.data.frame(HGMDDelete)
colnames(dfHGMDDelete)[3] <- "Consequence"

# This uses rvest to get the inserts table from Selenium
HGMDInsert <- HGMD_html %>%
  html_nodes("body > div.content > table:nth-child(15)") %>%
  html_table
dfHGMDInsert = as.data.frame(HGMDInsert)
colnames(dfHGMDInsert)[3] <- "Consequence"

# close Selenium
remDr$close()
}
}

```

6.2 Shaping the Data

Now that the data has been collected, it is time to compare the two and remove the duplicates and keep what is remaining of the HGMD data. I also removed some columns that are not needed that does not contain pertinent information

6.2.1 Remove Duplicates

Before removing the duplicates, the ExAC data needs to be reformatted so that the variants are easier to read and the Consequence column will match the Consequence column in the HGMD data:

```

# This section removes duplicates and prints out the unique file,
# This also prints out the duplicates in a separate file for reference

# format the ExAC data to make it easier to read and compatible with HGMD

# The headers were stored in row one, this makes row one the header and then deletes
# the row
colnames(dfExac) <- dfExac[1, ]
dfExac <- dfExac[-1, ]

# This reformats the Variant data, there was a bunch of junk that was scrapped
# with it.
# Remove whitespace
dfExac$Variant <- gsub(" ", "", dfExac$Variant, fixed = TRUE)

# Remove extra characters
dfExac$Variant <- gsub("\n", "", dfExac$Variant, fixed = TRUE)

# This makes the consequence column compatible with Exac by removing the p.
dfExac$Consequence <-
  gsub("p.", "", dfExac$Consequence, fixed = TRUE)

# We only need data that has an entry in the Consequence column, any blank data
# can be removed, to do this, NA is added to the Consequence column and then all
# NA's are removed.

```

```
dfExac$Consequence[dfExac$Consequence == ""] <- NA
dfExac_sub <- na.omit(dfExac)
```

Once the ExAC data is formatted, then the data can be compared. This code will check to make sure there is a dataframe and then compare the data and remove all duplicates then print it to a CSV file. It also searches the reference info and pulls out the first publish date of the mutation and puts it in a new column called year.

I also made it so that the data can tell if it is a missense or nonsense, splice, or frameshift for the CHD2 gene, and if it causes loss of function. This will be used to analyze the data.

This code also prints out the duplicate information in a separate file too.

```
# Compare the data

# This makes sure there is data in the dataframe before analyzing,
# If there is no data it prints a statement that there is no data
# and moves to the next data set.
if (exists("dfHGMDMissense") &&
    is.data.frame(get("dfHGMDMissense")) == 'FALSE') {
  paste("There is no data to analyze for HGMD Missense")
} else {
  # Removing duplicates for Missense

  # This combines the data frames by using plyr
  df1_mis <- join(dfHGMDMissense, dfExac_sub, type = "full")

  # This removes the duplicates
  df1_mis <-
    df1_mis[!(
      duplicated(df1_mis$Consequence) |
      duplicated(df1_mis$Consequence, fromLast = TRUE)
    ), ]

  # This keeps the rows needed
  keep <-
    c(
      "Consequence",
      "HGMD.codonchange",
      "HGVS.nucleotide.",
      "HGVS.protein.",
      "Variantclass",
      "Reported.phenotype",
      "Reference"
    )
  df2_mis <- df1_mis[keep]

  # This omits all na's and gives the final value with no duplicates
  dfHGMDMissense_final <- na.omit(df2_mis)

  # Add columns for additional analysis
  # Extract publish year from reference and make new column
  dfHGMDMissense_final$Year <-
    as.numeric(sub(
      "\\D*\\((\\d\\d\\d\\d\\d\\d)\\).*",
```

```

    "\\1",
    dfHGMDMissense_final$Reference
  ))

# Create mutation type column
# If the ending in Consequence is Term then it is a Nonsense
# Define Nonsense
nonsense <-
  grep(".+Term$",
    dfHGMDMissense_final$Consequence,
    perl = TRUE,
    value = TRUE)

# Make new column
dfHGMDMissense_final$Mutation.type <-
  ifelse(dfHGMDMissense_final$Consequence == nonsense,
    "Nonsense",
    "Missense")

# Make overall Consequence column
dfHGMDMissense_final$Overall.consequence <-
  ifelse(dfHGMDMissense_final$Mutation.type == "Missense",
    "nLoF",
    "LoF")

# For reference, this will get all of the duplicates
# This collects the duplicates
dfHGMDMissense_Duplicate <-
  merge(dfExac_sub, dfHGMDMissense, by = "Consequence", all = FALSE)
}

```

```
## Joining by: Consequence, Reference
```

```

# This makes sure there is data in the dataframe before anlaying,
# If there is no data it prints a statem that there is no data
# and moves to the next data set.
if (exists("dfHGMDSplice") &&
  is.data.frame(get("dfHGMDSplice")) == 'FALSE') {
  paste("There is no data to analyze for HGMD Splices")
} else {
  # Removing duplicates for Splice

  # This combines the data frames
  df1_spl <- join(dfHGMDSplice, dfExac_sub, type = "full")

  # This removes the duplicates
  df1_spl <-
    df1_spl[!(
      duplicated(df1_spl$Consequence) |
      duplicated(df1_spl$Consequence, fromLast = TRUE)
    ),]

  # This keeps the rows needed
  keep <-

```

```

c(
  "Consequence",
  "HGMD.splicing.mutation",
  "Variantclass",
  "Reported.phenotype",
  "Reference"
)
df2_spl <- df1_spl[keep]

# This omits all na's and gives the final value with no duplicates
dfHGMDSplice_final <- na.omit(df2_spl)

# Add columns for additional analysis
# Extract publish year from reference and make new column
dfHGMDSplice_final$Year <-
  as.numeric(sub(
    "\\D*\\((\\d\\d\\d\\d\\d\\d)\\).*",
    "\\1",
    dfHGMDSplice_final$Reference
  ))

# Create mutation type column
dfHGMDSplice_final$Mutation.type <-
  rep("splice", nrow(dfHGMDSplice_final))

# Make overall Consequence column
dfHGMDSplice_final$Overall.consequence <-
  ifelse(dfHGMDSplice_final$Mutation.type == "Missense",
    "nLoF",
    "LoF")

# For reference, this will get all of the duplicates
# This collects the duplicates
dfHGMDSplice_Duplicate <-
  merge(dfExac_sub, dfHGMDSplice, by = "Consequence", all = FALSE)
}

```

```
## Joining by: Consequence, Reference
```

```

# This makes sure there is data in the dataframe before analyzing,
# If there is no data it prints a statem that there is no data
# and moves to the next data set.
if (exists("dfHGMDDelete") &&
  is.data.frame(get("dfHGMDDelete")) == 'FALSE') {
  paste("There is no data to analyze for HGMD Deletions")
} else {
  # Removing duplicates for Deletions
  # This combines the data frames
  df1_del <- join(dfHGMDDelete, dfExac_sub, type = "full")

  # This removes the duplicates
  df1_del <-
    df1_del[!(
      duplicated(df1_del$Consequence) |

```

```

    duplicated(df1_del$Consequence, fromLast = TRUE)
  ),]

# This keeps the rows needed
keep <-
  c(
    "Consequence",
    "HGMD.deletion",
    "Variantclass",
    "Reported.phenotype",
    "Reference"
  )
df2_del <- df1_del[keep]

# This omits all na's and gives the final value with no duplicates
dfHGMDDelete_final <- na.omit(df2_del)

# Add columns for additional analysis
# Extract publish year from reference and make new column
dfHGMDDelete_final$Year <-
  as.numeric(sub(
    "\\D*\\((\\d\\d\\d\\d\\d\\d)\\).*",
    "\\1",
    dfHGMDDelete_final$Reference
  ))

# Create mutation type column
# If the ending in Consequence is Term then it is a Nonsense
# Define Nonsense
dfHGMDDelete_final$Mutation.type <- dfHGMDDelete_final$Consequence

# If it has a plus or minus it is a splice
dfHGMDDelete_final$Mutation.type <-
  gsub(pattern = ".+\\+|.+.+\\-|.+",
        dfHGMDDelete_final$Mutation.type,
        replacement = "splice")

# Everything else should be a frame shif
dfHGMDDelete_final$Mutation.type <-
  gsub(pattern = "c\\.+",
        dfHGMDDelete_final$Mutation.type,
        replacement = "frameshift")

# Make overall Consequence column
dfHGMDDelete_final$Overall.consequence <-
  ifelse(dfHGMDDelete_final$Mutation.type == "Missense",
        "nLoF",
        "LoF")

# For reference, this will get all of the duplicates
# This collects the duplicates
dfHGMDDelete_Duplicate <-
  merge(dfExac_sub, dfHGMDDelete, by = "Consequence", all = FALSE)

```

```

}

## Joining by: Consequence, Reference
# This makes sure there is data in the dataframe before anlaying,
# If there is no data it prints a statem that there is no data
# and moves to the next data set.
if (exists("dfHGMDInsert") &&
    is.data.frame(get("dfHGMDInsert")) == 'FALSE') {
  paste("There is no data to analyze for HGMD Insertions")
} else {
  # Removing duplicates for Inserts

  # This combines the data frames
  df1_ins <- join(dfHGMDInsert, dfExac_sub, type = "full")

  # This removes the duplicates
  df1_ins <-
    df1_ins[!(
      duplicated(df1_ins$Consequence) |
      duplicated(df1_ins$Consequence, fromLast = TRUE)
    ),]

  # This keeps the rows needed
  keep <-
    c(
      "Consequence",
      "HGMD.insertion",
      "Variantclass",
      "Reported.phenotype",
      "Reference"
    )
  df2_ins <- df1_ins[keep]

  # This omits all na's and gives the final value with no duplicates
  dfHGMDInsert_final <- na.omit(df2_ins)

  # Add columns for additional analysis
  # Extract publish year from reference and make new column
  dfHGMDInsert_final$Year <-
    as.numeric(sub(
      "\\D*\\((\\d\\d\\d\\d\\d\\d)\\).*",
      "\\1",
      dfHGMDInsert_final$Reference
    ))

  # Create mutation type column
  # If the ending in Consequence is Term then it is a Nonsense
  # Define Nonsense
  dfHGMDInsert_final$Mutation.type <- dfHGMDInsert_final$Consequence

  # If it has a plus or minus it is a splice
  dfHGMDInsert_final$Mutation.type <-
    gsub(pattern = "\\+|\\-|\\.", "",

```

```

    dfHGMDInsert_final$Mutation.type,
    replacement = "splice")

# Everything else should be a frame shif
dfHGMDInsert_final$Mutation.type <-
  gsub(pattern = "c\\..+",
    dfHGMDInsert_final$Mutation.type,
    replacement = "frameshift")

# Make overall Consequence column
dfHGMDInsert_final$Overall.consequence <-
  ifelse(dfHGMDInsert_final$Mutation.type == "Missense",
    "nLoF",
    "LoF")

# For reference, this will get all of the duplicates
# This collects the duplicates
dfHGMDInsert_Duplicate <-
  merge(dfExac_sub, dfHGMDInsert, by = "Consequence", all = FALSE)
}

## Joining by: Consequence, Reference
# This merges all non-duplicate data
dfHGMD_final <-
  join(dfHGMDMissense_final, dfHGMDSplice_final, type = "full")

## Joining by: Consequence, Variantclass, Reported.phenotype, Reference, Year, Mutation.type, Overall.c
dfHGMD_final <-
  join(dfHGMD_final, dfHGMDDelete_final, type = "full")

## Joining by: Consequence, Variantclass, Reported.phenotype, Reference, Year, Mutation.type, Overall.c
dfHGMD_final <-
  join(dfHGMD_final, dfHGMDInsert_final, type = "full")

## Joining by: Consequence, Variantclass, Reported.phenotype, Reference, Year, Mutation.type, Overall.c
# This checks to see if there are non-duplicates to write to a file
rowIns_final <- nrow(dfHGMD_final)

# This prints to a csv file if there are non-duplicate.
if (rowIns_final > 0) {
  write.csv(
    dfHGMD_final,
    file = paste("HGMD_final_", gene, ".csv", sep = ""),
    row.names = FALSE
  )
}

# This merges all duplicate data
dfHGMD_Duplicate <-
  join(dfHGMDMissense_Duplicate, dfHGMDSplice_Duplicate, type = "full")

```

```
## Joining by: Consequence, Variant, Chrom, Position, RSID, Reference.x, Alternate, Protein Consequence
dfHGMD_Duplicate <-
  join(dfHGMD_Duplicate, dfHGMDDelete_Duplicate, type = "full")

## Joining by: Consequence, Variant, Chrom, Position, RSID, Reference.x, Alternate, Protein Consequence
dfHGMD_Duplicate <-
  join(dfHGMD_Duplicate, dfHGMDInsert_Duplicate, type = "full")

## Joining by: Consequence, Variant, Chrom, Position, RSID, Reference.x, Alternate, Protein Consequence
# This checks to see if there are duplicates to write to a file
rowIns_Duplicate <- nrow(dfHGMD_Duplicate)

# This prints to a csv file if there are duplicates.
if (rowIns_Duplicate > 0) {
  write.csv(
    dfHGMD_Duplicate,
    file = paste("HGMD_Duplicate_", gene, ".csv", sep = ""),
    row.names = FALSE
  )
}
```

6.2.1.1 Verifying Results

The missense data was the only table with redundant information. To verify the results, I will use the merge function to find what is similar. Then compare before and after to make sure what was the same is now removed.

```
comparison <- merge(dfExac_sub, dfHGMDMissense, by = "Consequence", all=FALSE)
comparison$Consequence
```

```
## [1] "Arg1000Gln" "Arg1685His" "Val1479Met"
```

```
# This is the original and it contains the values
```

```
apply(dfHGMDMissense, 1, function(r)
  any (r %in% c("Arg1000Gln", "Arg1685His", "Val1479Met")))
```

```
## [1] FALSE FALSE FALSE FALSE FALSE FALSE FALSE TRUE FALSE FALSE FALSE
```

```
## [12] FALSE TRUE FALSE FALSE FALSE FALSE FALSE TRUE
```

```
# This is the final and it does not contain the values
```

```
apply(dfHGMDMissense_final, 1, function(r)
  any (r %in% c("Arg1000Gln", "Arg1685His", "Val1479Met")))
```

```
##      1      2      3      4      5      6      7      9     10     11     12     14
## FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE
##     15     16     17     18
## FALSE FALSE FALSE FALSE
```

6.3 Analyzing the Data

I will perform clustering to look how the data is grouped.

6.3.1 Cluster HGMD Data

I cleared the previous data and configured R with the following parameters before beginning the analysis:

```
# clears the console in RStudio
cat("\014")

# clears environment
rm(list = ls())
```

6.3.1.1 Load Data.

I opened the variant data from the gene CHD2 containing all known mutations.

To format the data, the data is separated by ',', stringsAsFactors = FALSE so that the strings in a data frame will be treated as plain strings and not as factor variables. I set na strings for missing data.

Below is my R code:

```
# Some csv files are really big and take a while to open. This command checks to
# see if it is already opened, if it is, it does not open it again.
# I also omitted the first column
if (!exists("dfHGMD_final")) {
  dfHGMD_final <-
    read.csv2(
      "HGMD_final_CHD2.csv",
      sep = ",",
      stringsAsFactors = FALSE,
      na.strings = c("", "NA")
    )
}
```

6.3.1.2 Formating the data

Inorder to cluster the data I need numeric data. I will make new columns with Consequence, Variantclass, Reported.phenotype, Mutation.type, Overall.consequence, and year converted to numeric.

For Consequence the data will change to: Create column and assign a Consequence with a num Ser112Term = 1, Arg121Term = 2, Pro218Leu = 3, Arg466Term = 4, Leu823Pro = 5, Asp856Gly = 6, Gln906Term = 7, Gly1174Asp = 8, Arg1313Gly = 9, Arg1345Gln = 10, Ser1406Gly = 11, Trp1534Cys = 12, Arg1637Term = 13, Gln1641Term = 14, Trp1657Term = 15, Arg1679Term = 16, c.390C>T = 17, c.1502+1G>A = 18, c.1719+5G>A = 19, c.1810-2A>C = 20, c.1552delC = 21, c.1809+1delG = 22, c.1880_1883delCTTT = 23, c.2895_2898delAGAA = 24, c.3734delA = 25, c.4233_4236delAGAA = 26, c.4256_4274del19 = 27, c.3787dupG = 28, c.4173dupA = 29, c.4949dupG = 30, c.5094dupG = 31

```
# Make Consequence numeric
dfHGMD_final$Consequence.num <- 1:nrow(dfHGMD_final)
```

For Reported Phenotype the data will change to: Autism = 1 Epilepsy = 2 Eyelid myoclonia = 3 Intellectual disability = 4

```
# Make Reported.phenotype numeric
# Assign number to Reported.phenotype
# Create column
dfHGMD_final$Reported.phenotype.num <-
  dfHGMD_final$Reported.phenotype

# Use regex to make data numeric
```

```

# Autism = 1
dfHGMD_final$Reported.phenotype.num <-
  gsub("^Autism.*",
    dfHGMD_final$Reported.phenotype.num,
    replacement = "1")

# Epilepsy = 2
dfHGMD_final$Reported.phenotype.num <-
  gsub("^Epilep.*|^Dravet.*|^Lennox-Gastaut.*",
    dfHGMD_final$Reported.phenotype.num,
    replacement = "2")

# Eyelid myoclonia = 3
dfHGMD_final$Reported.phenotype.num <-
  gsub("^Eyelid.*",
    dfHGMD_final$Reported.phenotype.num,
    replacement = "3")

# Intellectual disability = 4
dfHGMD_final$Reported.phenotype.num <-
  gsub("^Intellectual.*|^intellectual disability.*",
    dfHGMD_final$Reported.phenotype.num,
    replacement = "4")

```

For Variant Class the data will change to: DM = 1 DM? = 2

```

# Make Variantclass numeric
# Assign number to Variantclass.num
dfHGMD_final$Variantclass.num<-dfHGMD_final$Variantclass
dfHGMD_final$Variantclass.num<-ifelse(dfHGMD_final$Variantclass=="DM", 1, 2)

```

For Overall.consequence the data will change to: LoF = 1 nLoF = 2

```

# Make Overall.consequence numeric
# Assign number to Overall.consequence.num
dfHGMD_final$Overall.consequence.num <-
dfHGMD_final$Overall.consequence
dfHGMD_final$Overall.consequence.num <-
ifelse(dfHGMD_final$Overall.consequence == "LoF", 1, 2)

```

For Mutation.type the data will change to: frameshift = 1 Missense = 2 Nonsense = 3 splice = 4

```

# Make Reported.phenotype numeric
# Assign number to Reported.phenotype
# Create column
dfHGMD_final$Mutation.type.num <-
  dfHGMD_final$Mutation.type

# Use regex to make data numeric
# frameshift = 1
dfHGMD_final$Mutation.type.num <-
  gsub("^frameshift",
    dfHGMD_final$Mutation.type.num,
    replacement = "1")

# Missense = 2

```

```
dfHGMD_final$Mutation.type.num <-
  gsub("^Missense",
    dfHGMD_final$Mutation.type.num,
    replacement = "2")

# Nonsense = 3
dfHGMD_final$Mutation.type.num <-
  gsub("^Nonsense",
    dfHGMD_final$Mutation.type.num,
    replacement = "3")

# Splice = 4
dfHGMD_final$Mutation.type.num <-
  gsub("^splice",
    dfHGMD_final$Mutation.type.num,
    replacement = "4")
```

Change data to numeric and make new dataframe:

```
# Convert to numeric
dfHGMD_final[8] <- sapply(dfHGMD_final[8], as.numeric)
dfHGMD_final[14:18] <- sapply(dfHGMD_final[14:18], as.numeric)

# Columns to keep
keep <-
  c(
    "Consequence.num",
    "Reported.phenotype.num",
    "Variantclass.num",
    "Overall.consequence.num",
    "Mutation.type.num",
    "Year")

# Make new dataframe with keep data
dfHGMDCluster <- dfHGMD_final[keep]

# Check dataframe
str(dfHGMDCluster)

## 'data.frame': 31 obs. of 6 variables:
## $ Consequence.num : num 1 2 3 4 5 6 7 8 9 10 ...
## $ Reported.phenotype.num : num 4 2 3 2 2 1 1 1 2 2 ...
## $ Variantclass.num : num 1 1 2 1 1 2 2 2 1 1 ...
## $ Overall.consequence.num: num 1 1 2 2 2 2 2 2 2 2 ...
## $ Mutation.type.num : num 3 3 2 2 2 2 2 2 2 2 ...
## $ Year : num [1:31, 1] 2014 2013 2015 2013 2013 ...
## ..- attr(*, "dimnames")=List of 2
## .. ..$ : NULL
## .. ..$ : chr "Year"

summary(dfHGMDCluster)

## Consequence.num Reported.phenotype.num Variantclass.num
## Min. : 1.0 Min. :1.000 Min. :1.000
```

```
## 1st Qu.: 8.5      1st Qu.:1.500      1st Qu.:1.000
## Median :16.0     Median :2.000      Median :1.000
## Mean   :16.0     Mean   :2.097      Mean   :1.484
## 3rd Qu.:23.5     3rd Qu.:2.000      3rd Qu.:2.000
## Max.    :31.0     Max.    :4.000      Max.    :2.000
## Overall.consequence.num Mutation.type.num      Year.Year
## Min.     :1.000      Min.     :1.000      Min.     :2012.0000
## 1st Qu.:1.000      1st Qu.:1.000      1st Qu.:2013.0000
## Median :1.000      Median :2.000      Median :2015.0000
## Mean    :1.323      Mean    :2.194      Mean    :2014.5806
## 3rd Qu.:2.000      3rd Qu.:3.000      3rd Qu.:2016.0000
## Max.    :2.000      Max.    :4.000      Max.    :2017.0000
```

6.3.2 Clustering

Clustering is grouping like with like such that:

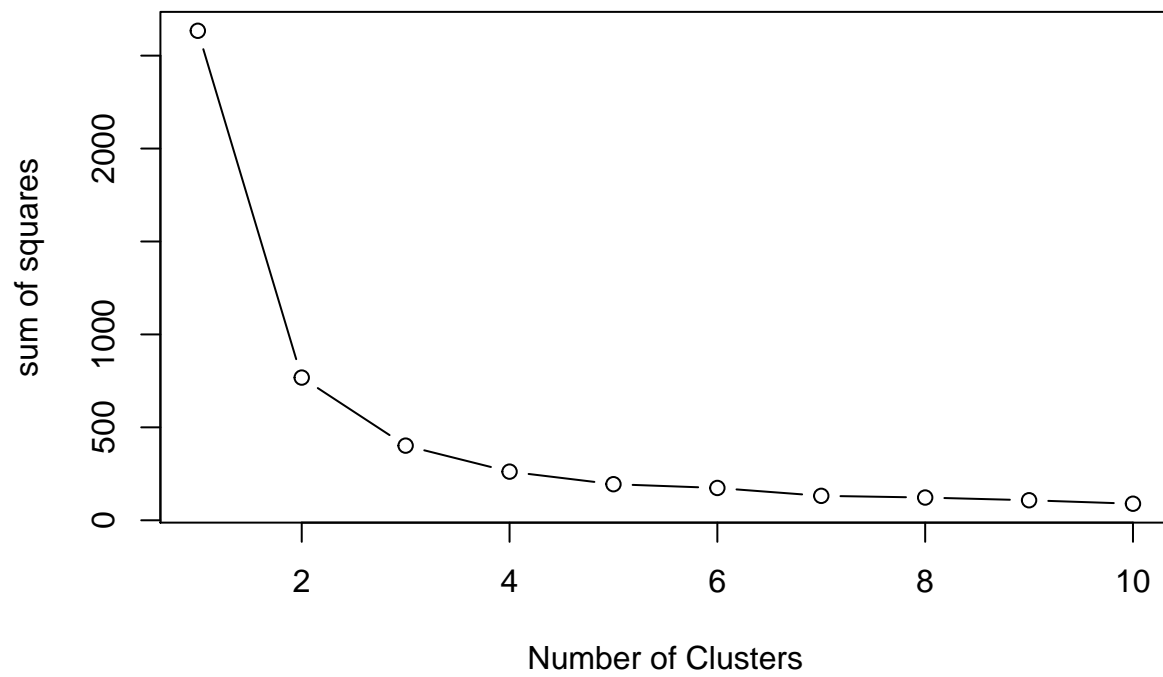
1. Similar objects are close to one another within the same cluster.
2. Dissimilar to the objects in other clusters.

6.3.2.1 Number of Clusters

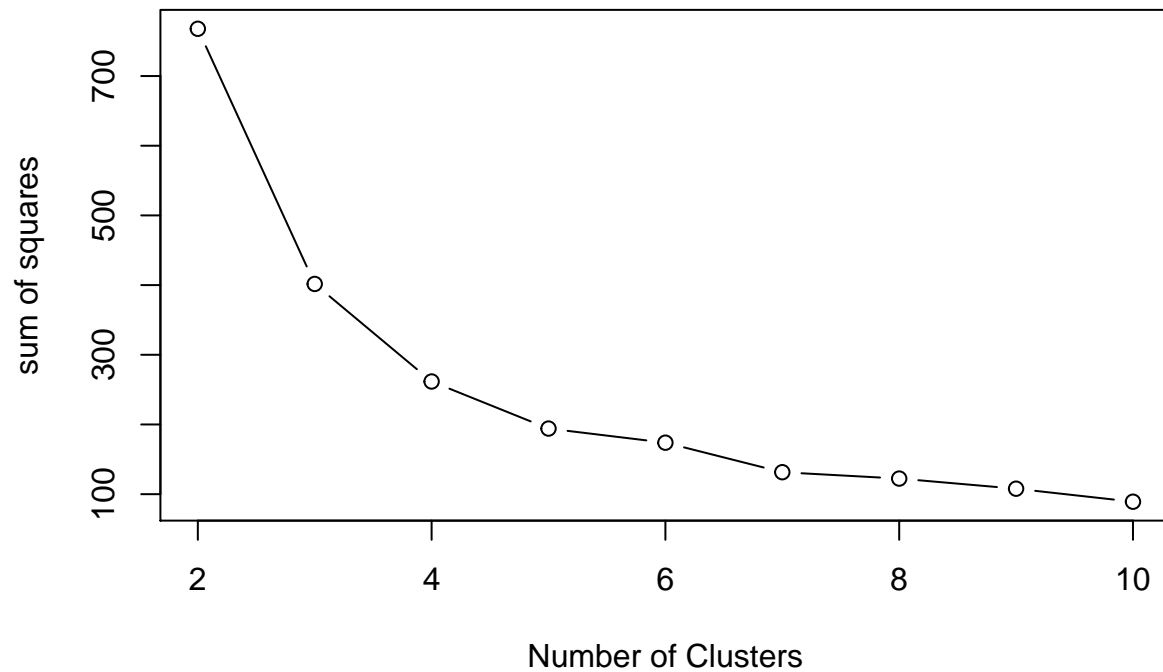
Before I can begin clustering analysis, I need to determine the number of clusters. For determining “the right number of clusters”, for this analysis I will use the averaged Silhouette width and Gap statistic and Hartigan’s rule.

6.3.2.1.1 Averaged Silhouette Width and Gap Statistic

```
# Determining number of clusters
sos <- (nrow(dfHGMDCluster) - 1) * sum(apply(dfHGMDCluster, 2, var))
for (i in 2:10)
  sos[i] <- sum(kmeans(dfHGMDCluster, centers = i)$withinss)
plot(1:10,
     sos,
     type = "b",
     xlab = "Number of Clusters",
     ylab = "sum of squares")
```

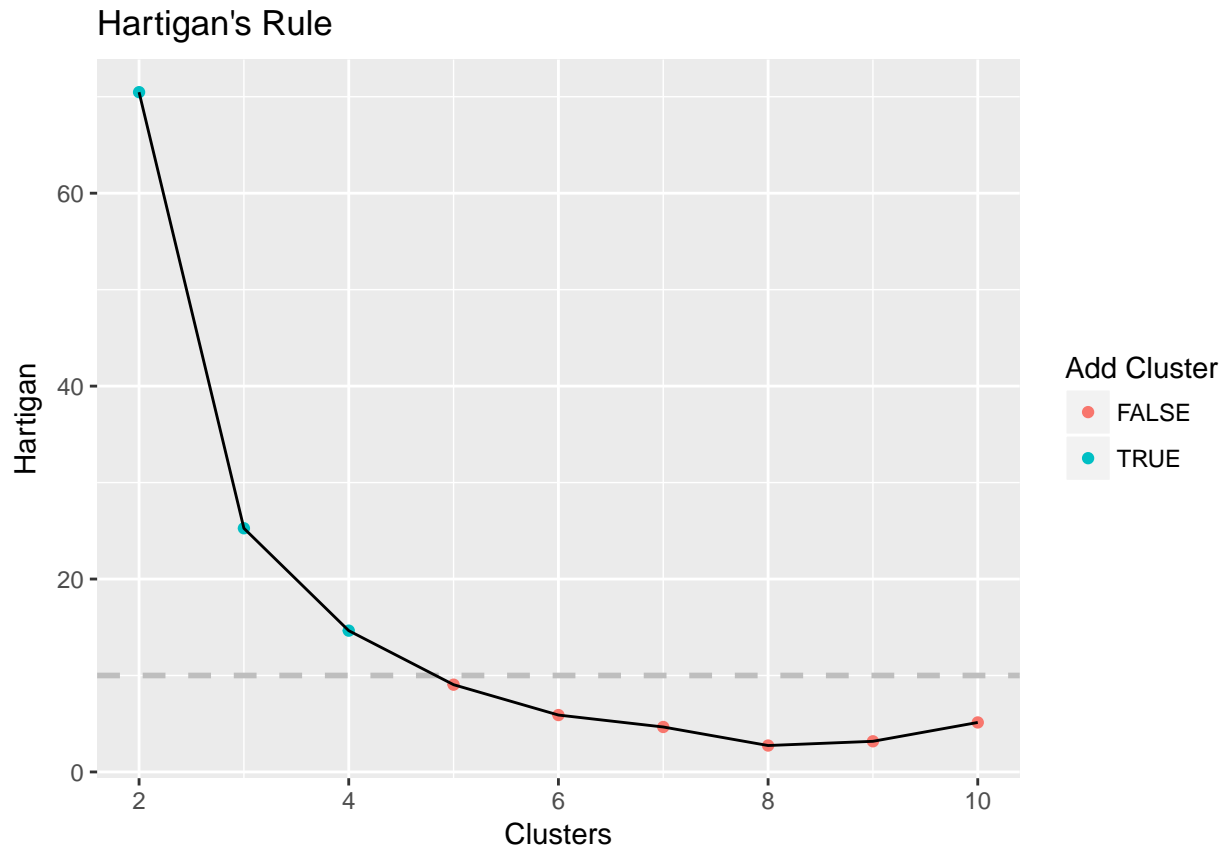


```
plot(2:10,  
     sos[c(2:10)],  
     type = "b",  
     xlab = "Number of Clusters",  
     ylab = "sum of squares")
```



6.3.2.1.2 Hartigan's rule

```
# Hartigans's rule FitKMean (similarity)
# require(useful)
best<-FitKMeans(dfHGMDCluster,max.clusters=10, seed=111)
PlotHartigan(best)
```



6.3.2.1.3 Number of Clusters Results

Based off of the graphs above it looks like I should perform clustering with 2 to 4 clusters. The analysis looks better with 2 after some trial and error.

6.3.2.2 Partitioning-based clustering

Partitioning algorithms construct various partitions and then evaluate them by some criterion. Hierarchy algorithms, two examples are k-means and k-medoids algorithms.

6.3.2.2.1 k-means

k-means clustering is a method of vector quantization, originally from signal processing. k-means clustering aims to partition n observations into k clusters in which each observation belongs to the cluster with the nearest mean, serving as a prototype of the cluster. This results in a partitioning of the data space into Voronoi cells.

Below I apply the k-means algorithm with a calculation that makes the k-means calculation more stable, it performs this analysis 1000 times and takes the ones with the least error:

```
# Clustering with 2 clusters
k <- 2
trials <- 1000
dfHGMDCluster.2.cluster <- kmeans(dfHGMDCluster, k, nstart = trials)
dfHGMDCluster.2.cluster
```

```
## K-means clustering with 2 clusters of sizes 16, 15
##
## Cluster means:
##   Consequence.num Reported.phenotype.num Variantclass.num
## 1             8.5             1.875000             1.500000
## 2            24.0             2.333333             1.466667
##   Overall.consequence.num Mutation.type.num      Year
## 1                 1.625             2.375 2014.625
## 2                 1.000             2.000 2014.533
##
## Clustering vector:
## [1] 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2
##
## Within cluster sum of squares by cluster:
## [1] 397.0 370.8
## (between_SS / total_SS = 70.8 %)
##
## Available components:
##
## [1] "cluster"      "centers"      "totss"        "withinss"
## [5] "tot.withinss" "betweenss"    "size"         "iter"
## [9] "ifault"
```

Evaluating model performance

```
# Evaluating model performance
# look at the size of the clusters
dfHGMDCluster.2.cluster$size
```

```
## [1] 16 15
```

```
# look at the cluster centers
dfHGMDCluster.2.cluster$centers
```

```
##   Consequence.num Reported.phenotype.num Variantclass.num
## 1             8.5             1.875000             1.500000
## 2            24.0             2.333333             1.466667
##   Overall.consequence.num Mutation.type.num      Year
## 1                 1.625             2.375 2014.625
## 2                 1.000             2.000 2014.533
```

```
names(dfHGMDCluster)
```

```
## [1] "Consequence.num"      "Reported.phenotype.num"
## [3] "Variantclass.num"    "Overall.consequence.num"
## [5] "Mutation.type.num"   "Year"
```

```
# mean of 'Consequence.num' by cluster
```

```
Consequence <-
```

```
  aggregate(data = dfHGMDCluster, Consequence.num ~ dfHGMDCluster.2.cluster$cluster, mean)
Consequence
```

```
##   dfHGMDCluster.2.cluster$cluster Consequence.num
## 1                               1             8.5
## 2                               2            24.0
```

```
# mean of 'Reported.phenotype.num' by cluster
```

```
Reported.phenotype <-
```



```

aggregate(data = dfHGMDCluster,
           Reported.phenotype.num ~ dfHGMDCluster.2.cluster$cluster,
           mean)
Reported.phenotype

##    dfHGMDCluster.2.cluster$cluster Reported.phenotype.num
## 1                                1             1.875000
## 2                                2             2.333333

# mean of 'Variantclass' by cluster
Variantclass <-
  aggregate(data = dfHGMDCluster,
            Variantclass.num ~ dfHGMDCluster.2.cluster$cluster,
            mean)
Variantclass

##    dfHGMDCluster.2.cluster$cluster Variantclass.num
## 1                                1             1.500000
## 2                                2             1.466667

# mean 'year' by cluster
year <-
  aggregate(data = dfHGMDCluster, Year ~ dfHGMDCluster.2.cluster$cluster, mean)
year

##    dfHGMDCluster.2.cluster$cluster    Year
## 1                                1 2014.625
## 2                                2 2014.533

```

Multidimensional scaling (MDS)

Multidimensional scaling (MDS) is a means of visualizing the level of similarity of individual cases of a high-dimensional dataset.

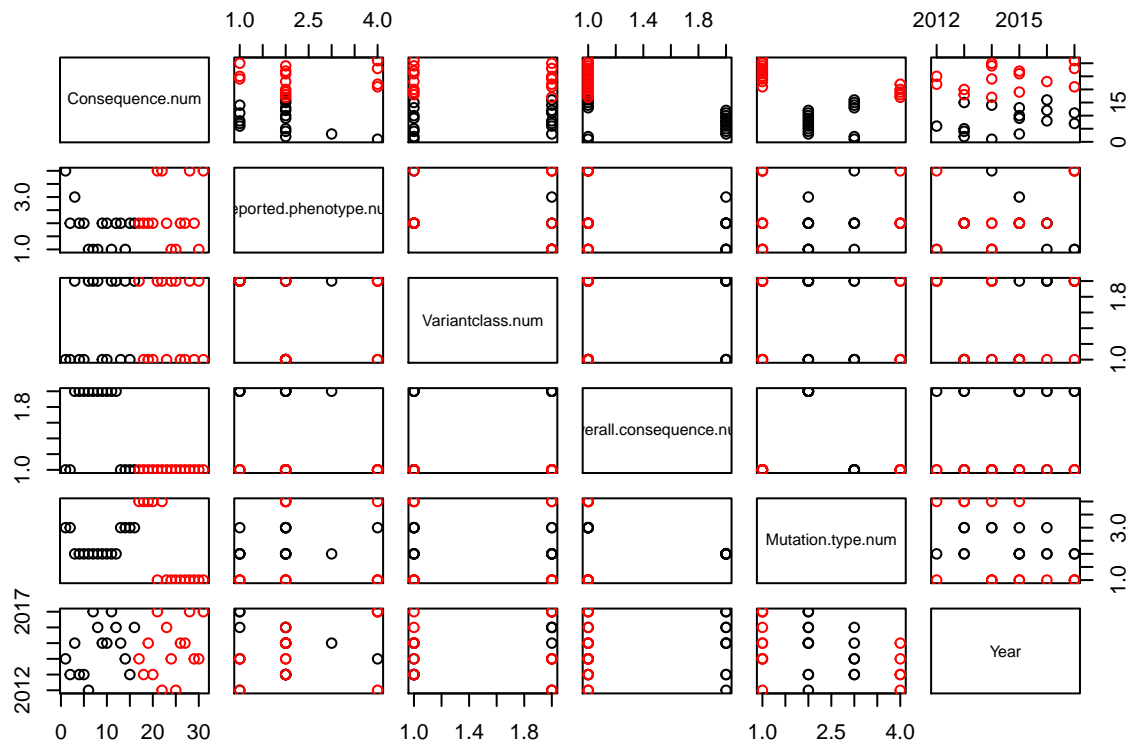
MDS attempts to find an embedding from the I objects into \mathbb{R}^N such that distances are preserved.

Below are some MDS plots:

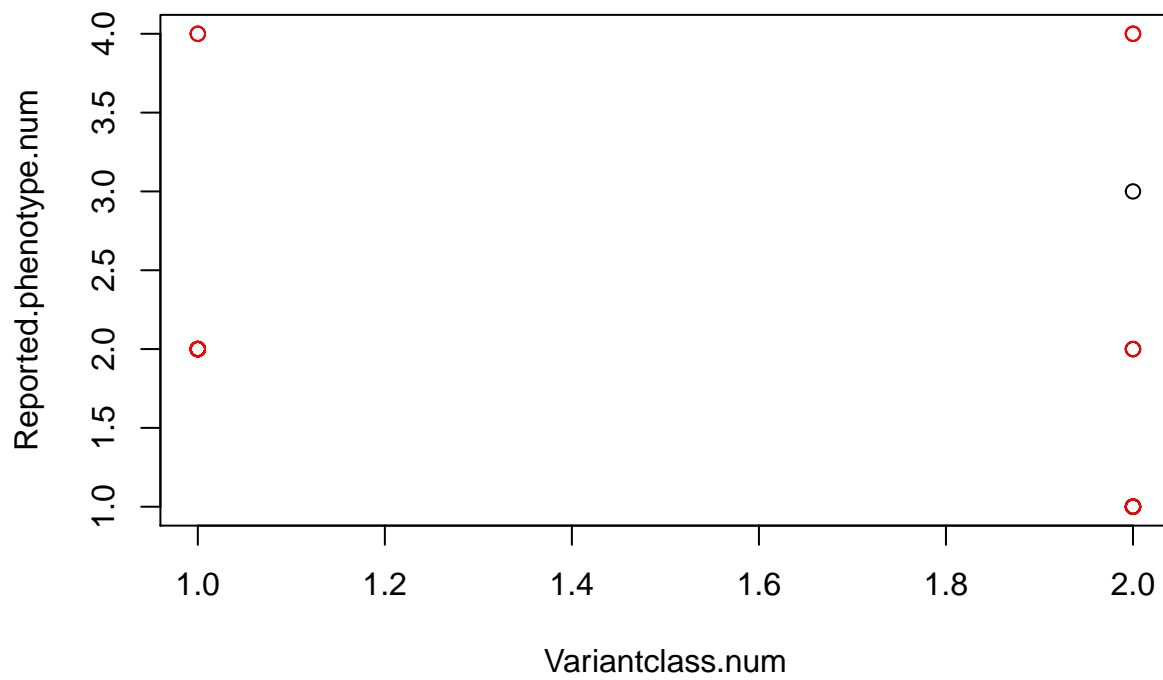
```

plot(dfHGMDCluster, col = dfHGMDCluster.2.cluster$cluster)    # Plot Clusters

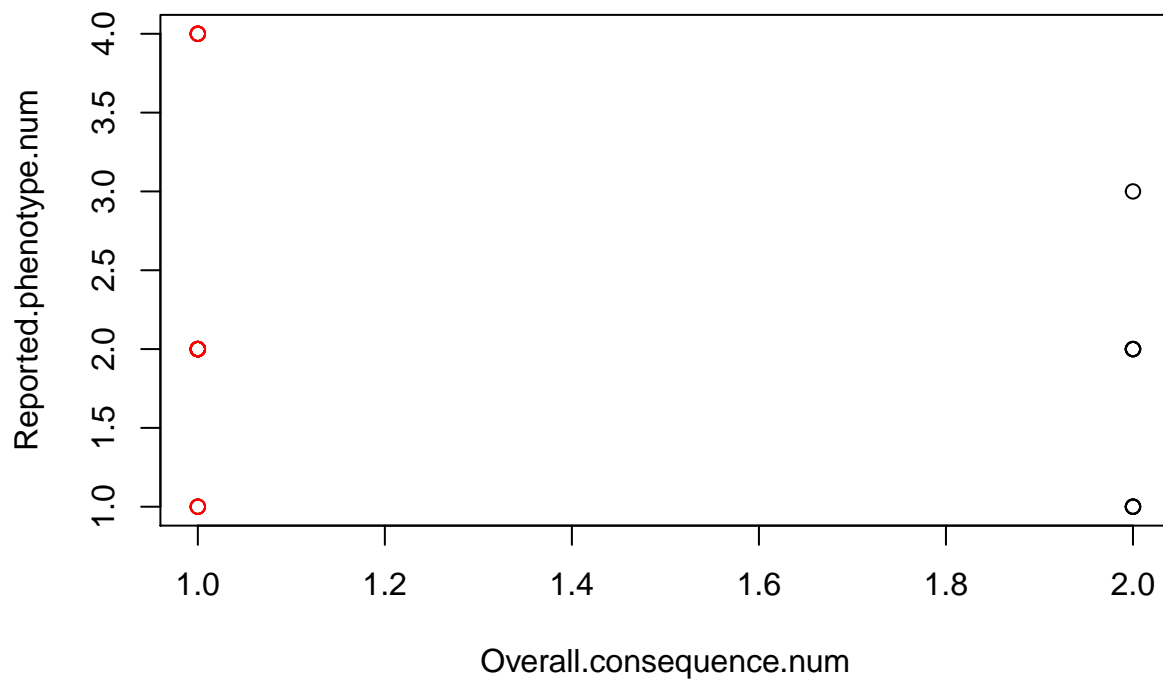
```



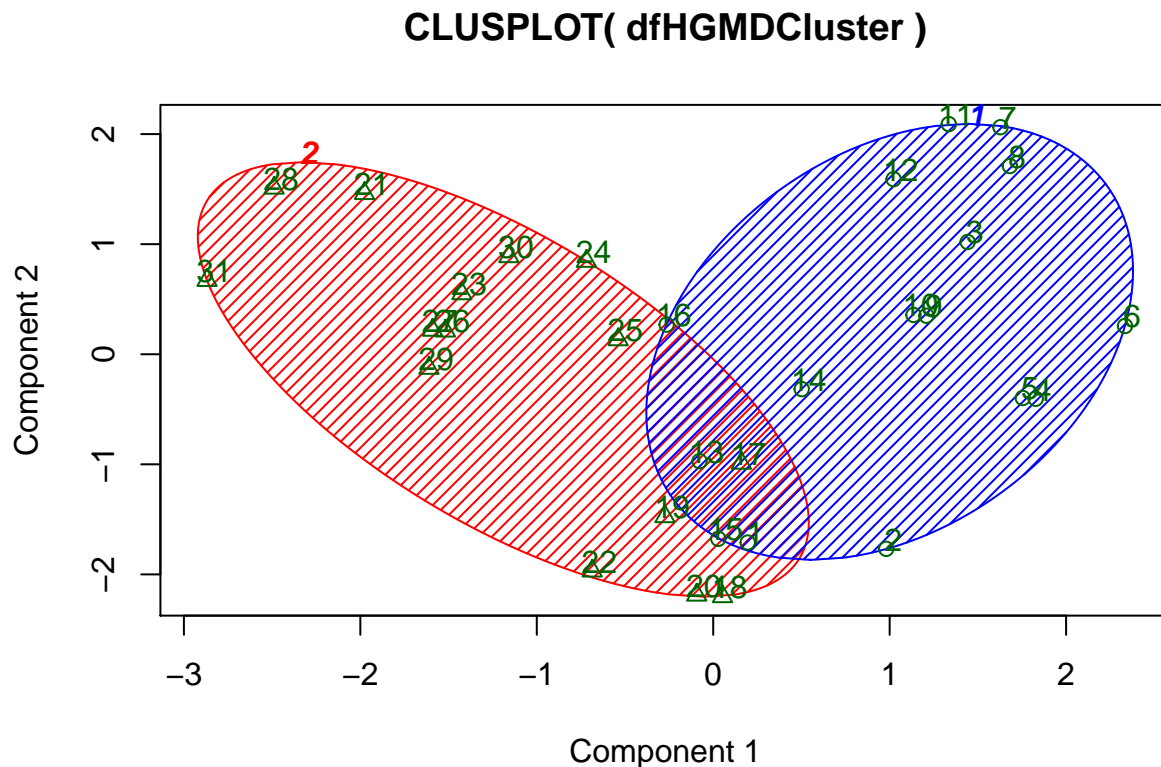
```
# Plot Consequence.num and Reported.phenotype.num
plot(dfHGMDCluster[c("Variantclass.num", "Reported.phenotype.num")], col =
  dfHGMDCluster.2.cluster$cluster)
points(dfHGMDCluster.2.cluster$centers,
  col = 1:2,
  pch = 8)
```



```
# Plot Mutation.type.num and Reported.phenotype.num
plot(dfHGMDCluster[c("Overall.consequence.num", "Reported.phenotype.num")], col =
  dfHGMDCluster.2.cluster$cluster)
points(dfHGMDCluster.2.cluster$centers,
  col = 1:2,
  pch = 8)
```



```
# Centroid Plot against 1st two discriminant functions
clusplot(
  dfHGMDCluster,
  dfHGMDCluster.2.cluster$cluster,
  color = TRUE,
  shade = TRUE,
  labels = 2,
  lines = 0
)
```



These two components explain 57.01 % of the point variability.

```
# library(fpc)
# plotcluster(dfHGMDCluster, dfHGMDCluster.2.cluster$cluster)
```

For Overall.consequence the data will change to: LoF = 1 nLoF = 2

For Variant Class the data will change to: DM = 1 DM? = 2

For Reported Phenotype the data will change to: Autism = 1 Epilepsy = 2 Eyelid myoclonia = 3 Intellectual disability = 4

The plots look OK. There is overlapping in the centroid plot. I wish this would have clustered better.

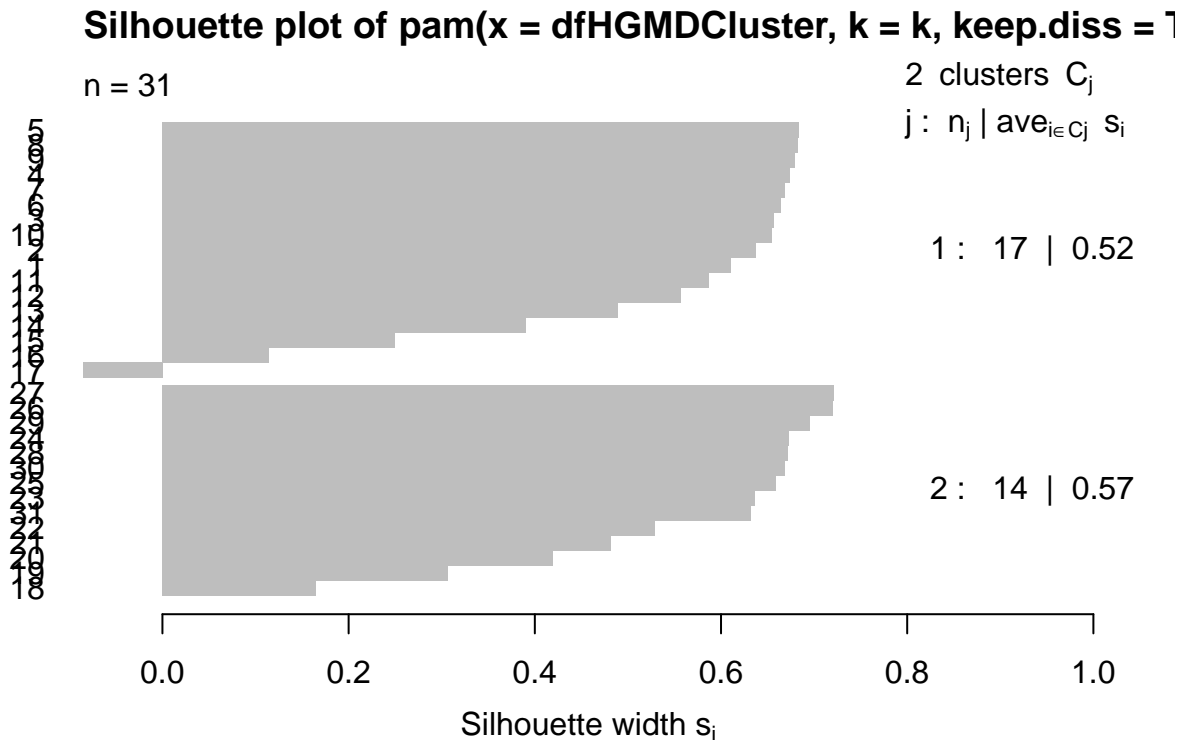
6.3.2.2.2 K-medoids clustering in R

The K-medoids or Partitioning Around Medoids (PAM) algorithm is related to the k-means algorithm and but uses medoid shifts rather than reassigning points based on Euclidean distance. Each cluster is represented by one of the objects (i.e. points) in the cluster. A medoid is a point in a cluster whose dissimilarity to all the points in the cluster is minimal. Medoids are similar in concept to means or centroids, but medoids are always members of the data set. That is, in 2D Cartesian space a centroid can be any valid x.y coordinate. Whereas a medoid must be one of the data points.

Below I use R to apply the k-medoids algorithm:

```
# PAM
k <- 2
dfHGMDCluster.pam.2.clust <-
pam(dfHGMDCluster, k, keep.diss = TRUE, keep.data = TRUE)
dfHGMDCluster.pam.2.clust
```

```
## Medoids:
##      ID Consequence.num Reported.phenotype.num Variantclass.num
## [1,]  9                9                2                1
## [2,] 26                26                2                1
##      Overall.consequence.num Mutation.type.num Year
## [1,]                2                2 2015
## [2,]                1                1 2015
## Clustering vector:
## [1] 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 2 2 2 2 2 2 2 2 2 2 2 2
## Objective function:
##      build      swap
## 5.869938 4.628850
##
## Available components:
## [1] "medoids"      "id.med"      "clustering"  "objective"  "isolation"
## [6] "clusinfo"     "silinfo"     "diss"        "call"       "data"
plot(dfHGMDCluster.pam.2.clust, which.plots = 2)
```



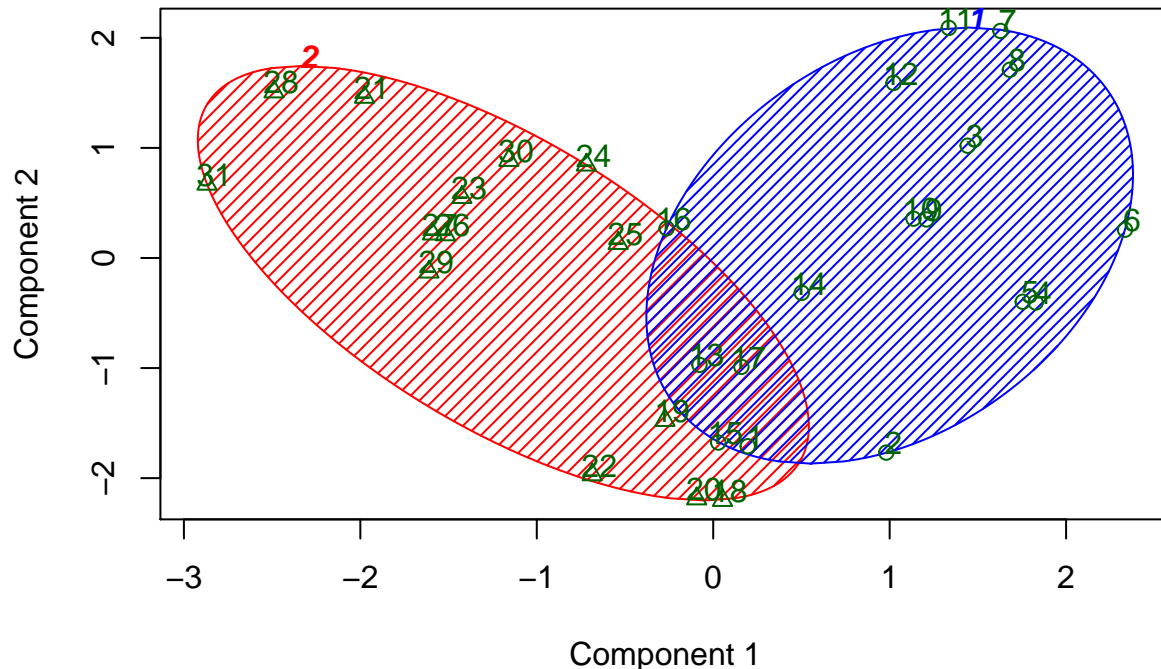
Average silhouette width : 0.55

```
# long lines good - means greater within cluster similarity

# Centroid Plot against 1st two discriminant functions
clusplot(
dfHGMDCluster.pam.2.clust,
color = TRUE,
shade = TRUE,
labels = 2,
```

```
lines = 0
)
```

```
usplot(pam(x = dfHGMDCluster, k = k, keep.diss = TRUE, keep.data = T
```



These two components explain 57.01 % of the point variability.

The silhouette plot looks good, long lines are good, short ones are not. It appears that there are more long lines.

The centroid plot looks a little better than the k-means plot, but there seems to be overlap with the clusters.

6.3.3 Gap statistic

`clusGap()` calculates a goodness of clustering measure, the \hat{a} gap statistic. For each number of clusters k , it compares $\log(W(k))$ with $E^*[\log(W(k))]$ where the latter is defined via bootstrapping, i.e. simulating from a reference distribution.

`maxSE(f, SE.f)` determines the location of the maximum of f , taking a \hat{a} 1-SE rule into account for the SE methods. The default method “firstSEmax” looks for the smallest k such that its value $f(k)$ is not more than 1 standard error away from the first local maximum. This is similar but not the same as “Tibs2001SEmax”, Tibshirani et al’s recommendation of determining the number of clusters from the gap statistics and their standard deviations.

```
gap <-
clusGap(dfHGMDCluster, FUNcluster = pam, K.max = 10) # Bootstrapping
gap$Tab
```

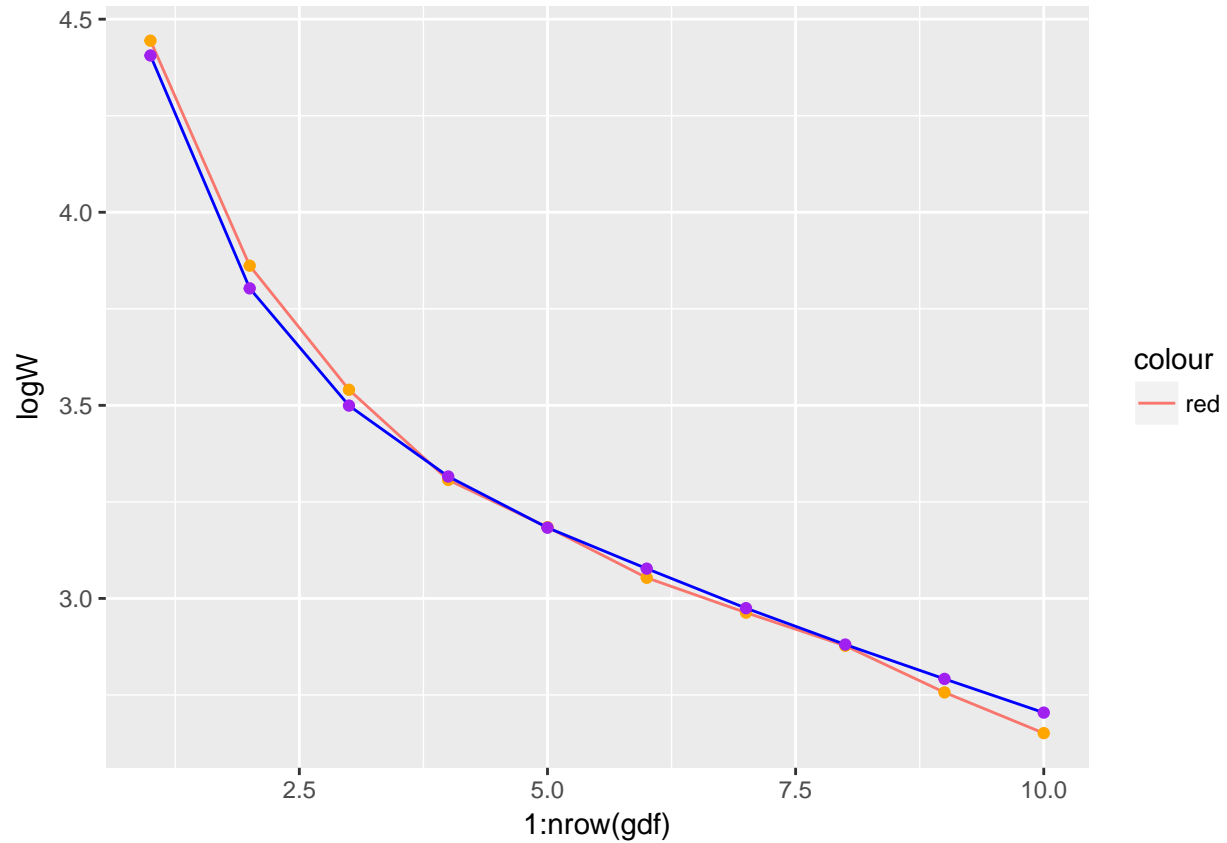
```
##          logW    E.logW          gap    SE.sim
## [1,] 4.444358 4.405934 -0.038424818 0.07974331
## [2,] 3.861596 3.802927 -0.058668961 0.06715870
```

```
## [3,] 3.540359 3.499303 -0.041056420 0.06031533
## [4,] 3.306797 3.315699 0.008901448 0.05136976
## [5,] 3.184518 3.183143 -0.001375534 0.05836193
## [6,] 3.053437 3.077105 0.023668841 0.05769219
## [7,] 2.963147 2.974743 0.011595714 0.05789634
## [8,] 2.877389 2.880581 0.003192080 0.06044070
## [9,] 2.756605 2.791579 0.034974451 0.06270258
## [10,] 2.651120 2.704217 0.053096953 0.06781101
```

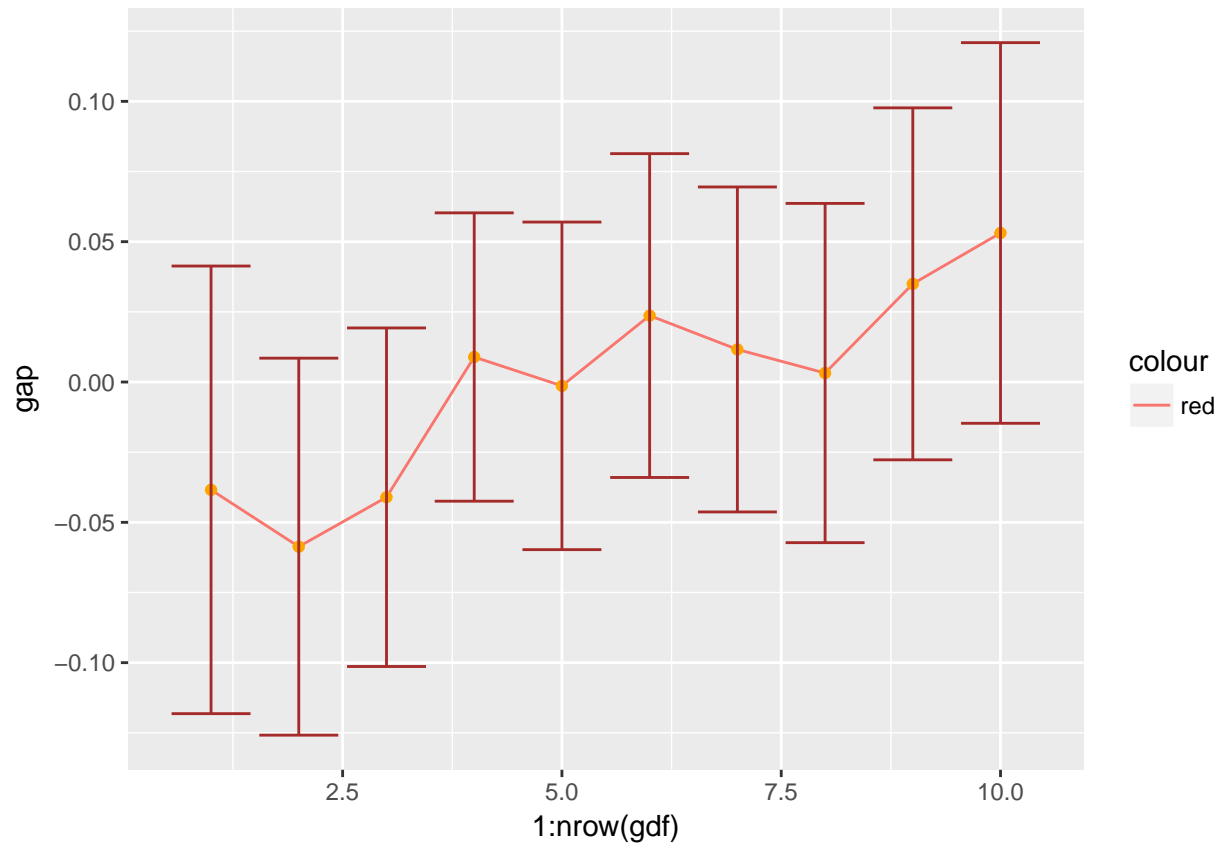
```
gdf <- as.data.frame(gap$Tab)
head(gdf)
```

```
##      logW      E.logW      gap      SE.sim
## 1 4.444358 4.405934 -0.038424818 0.07974331
## 2 3.861596 3.802927 -0.058668961 0.06715870
## 3 3.540359 3.499303 -0.041056420 0.06031533
## 4 3.306797 3.315699 0.008901448 0.05136976
## 5 3.184518 3.183143 -0.001375534 0.05836193
## 6 3.053437 3.077105 0.023668841 0.05769219
```

```
qplot(
  x = 1:nrow(gdf),
  y = logW,
  data = gdf,
  geom = "line",
  color = "red"
) +
  geom_point(aes(y = logW), color = "orange") +
  geom_line(aes(y = E.logW), color = "blue") +
  geom_point(aes(y = E.logW), color = "purple")
```

```
# Gap statistic
qplot(
  x = 1:nrow(gdf),
  y = gap,
  data = gdf,
  geom = "line",
  color = "red"
) +
  geom_point(aes(y = gap), color = "orange") +
  geom_errorbar(aes(ymin = gap - SE.sim, ymax = gap + SE.sim), color = "brown")
```



It looks like around 2, the gap increases at a higher slope, so I think a k of 2 is good.

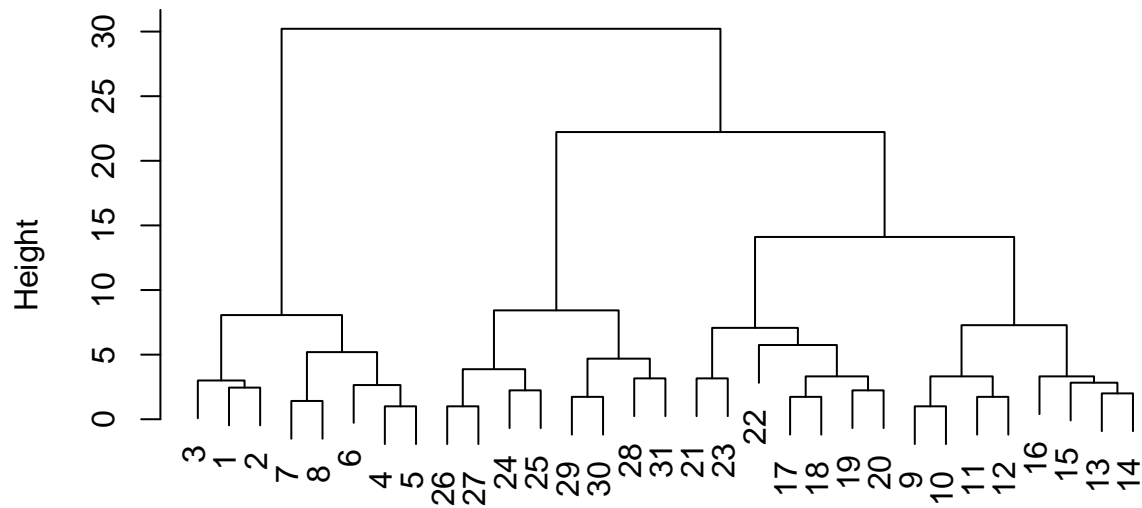
6.3.4 Hierarchical clustering in R

In hierarchical clustering the idea is to group data objects (i.e. points) into a tree of clusters. That is, hierarchical clustering is a method of cluster analysis which seeks to build a hierarchy of clusters.

Below is my analysis using Hierarchical clustering:

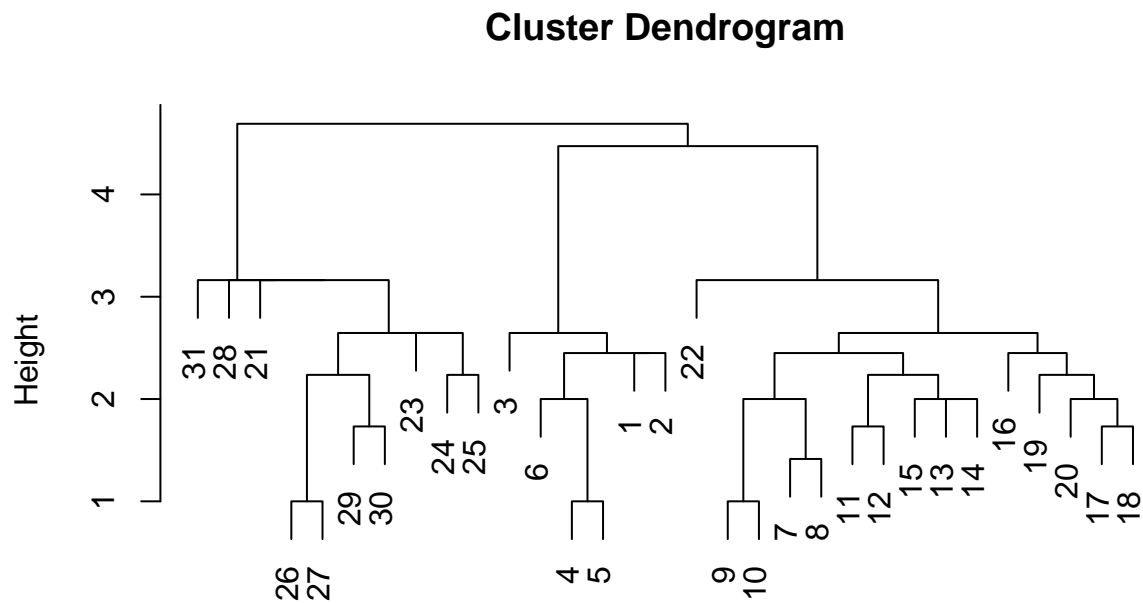
```
dfHGMDCluster.h.clust<- hclust(d=dist(dfHGMDCluster))
plot(dfHGMDCluster.h.clust)
```

Cluster Dendrogram



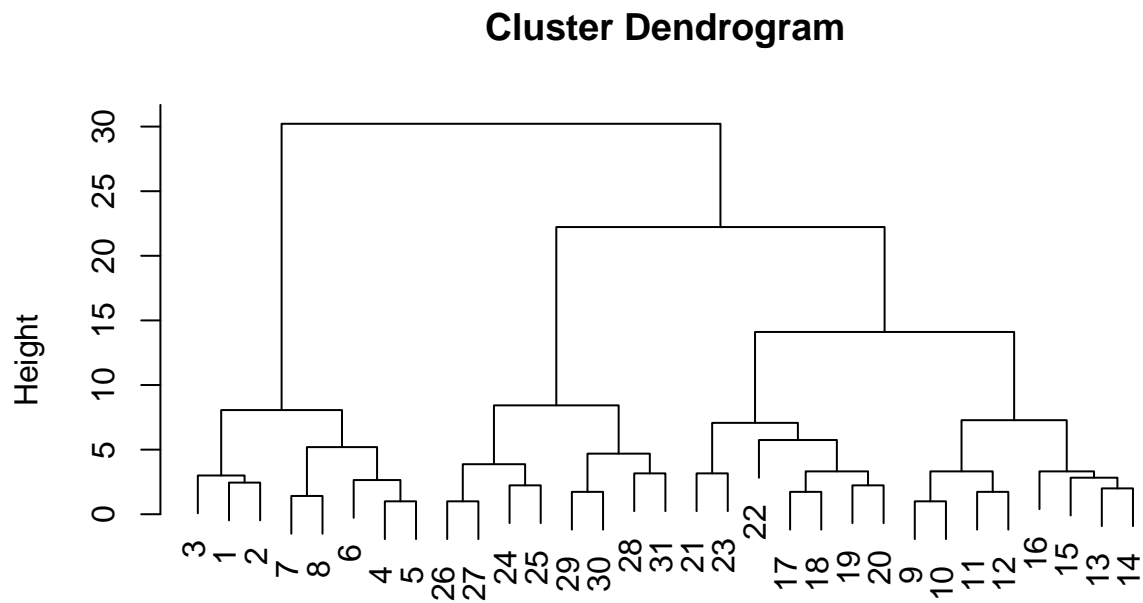
```
dist(dfHGMDCluster)
hclust (*, "complete")
```

```
dfHGMDCluster.h.clust.si<- hclust(dist(dfHGMDCluster), method = "single")
dfHGMDCluster.h.clust.co<- hclust(dist(dfHGMDCluster), method = "complete")
dfHGMDCluster.h.clust.av<- hclust(dist(dfHGMDCluster), method = "average")
dfHGMDCluster.h.clust.ce<- hclust(dist(dfHGMDCluster), method = "centroid")
dfHGMDCluster.h.clust.me<- hclust(dist(dfHGMDCluster), method = "ward.D")
plot(dfHGMDCluster.h.clust.si)
```



```
dist(dfHGMDCluster)
hclust (*, "single")
```

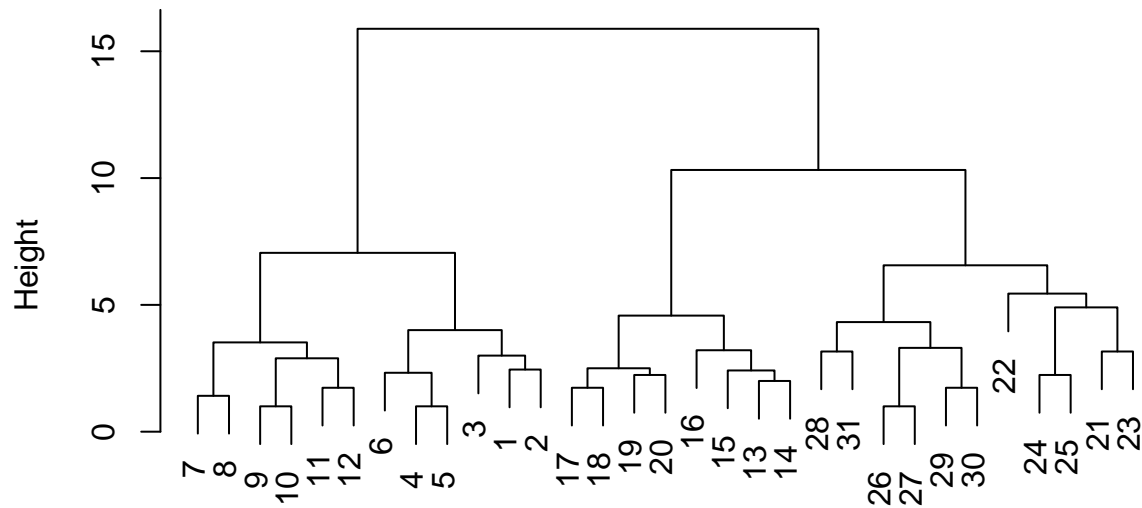
```
plot(dfHGMDCluster.h.clust.co)
```



```
dist(dfHGMDCluster)  
hclust (*, "complete")
```

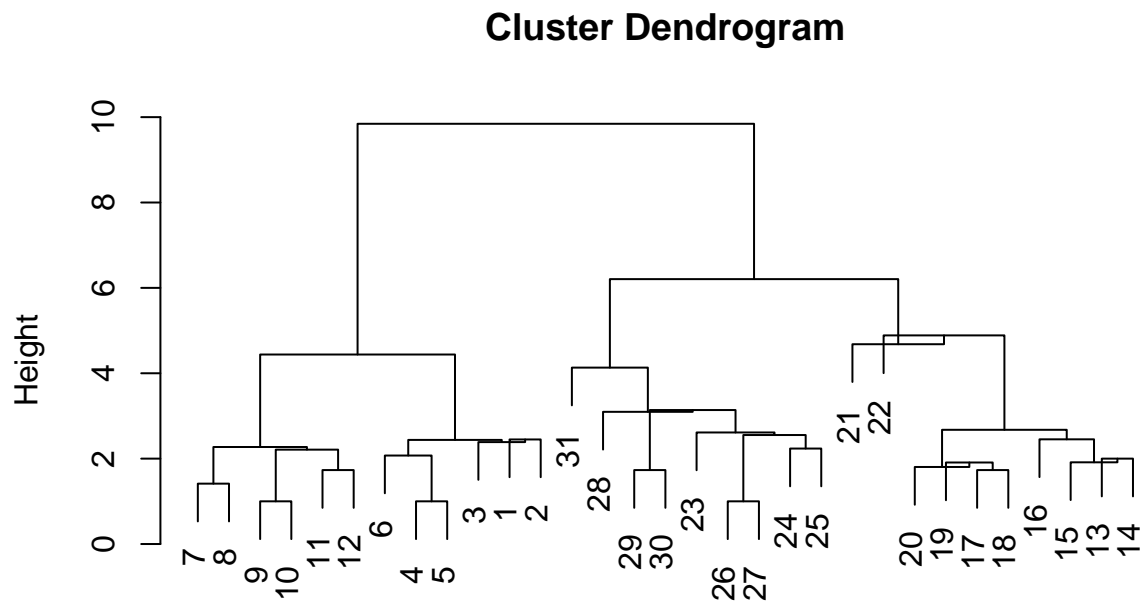
```
plot(dfHGMDCluster.h.clust.av)
```

Cluster Dendrogram



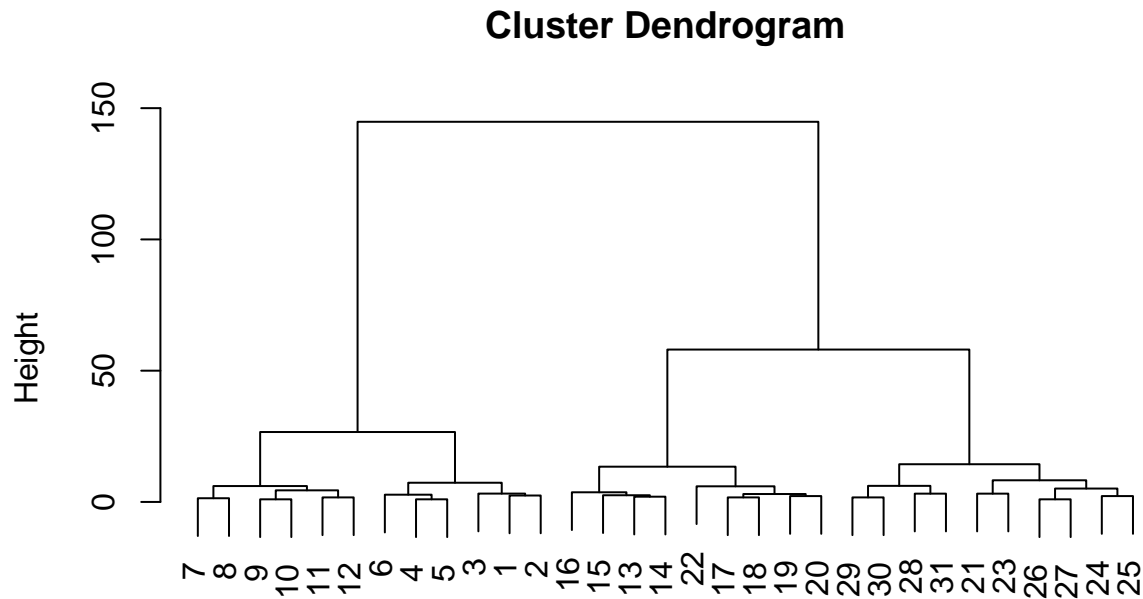
```
dist(dfHGMDCluster)
hclust (*, "average")
```

```
plot(dfHGMDCluster.h.clust.ce)
```



```
dist(dfHGMDCluster)
hclust (*, "centroid")
```

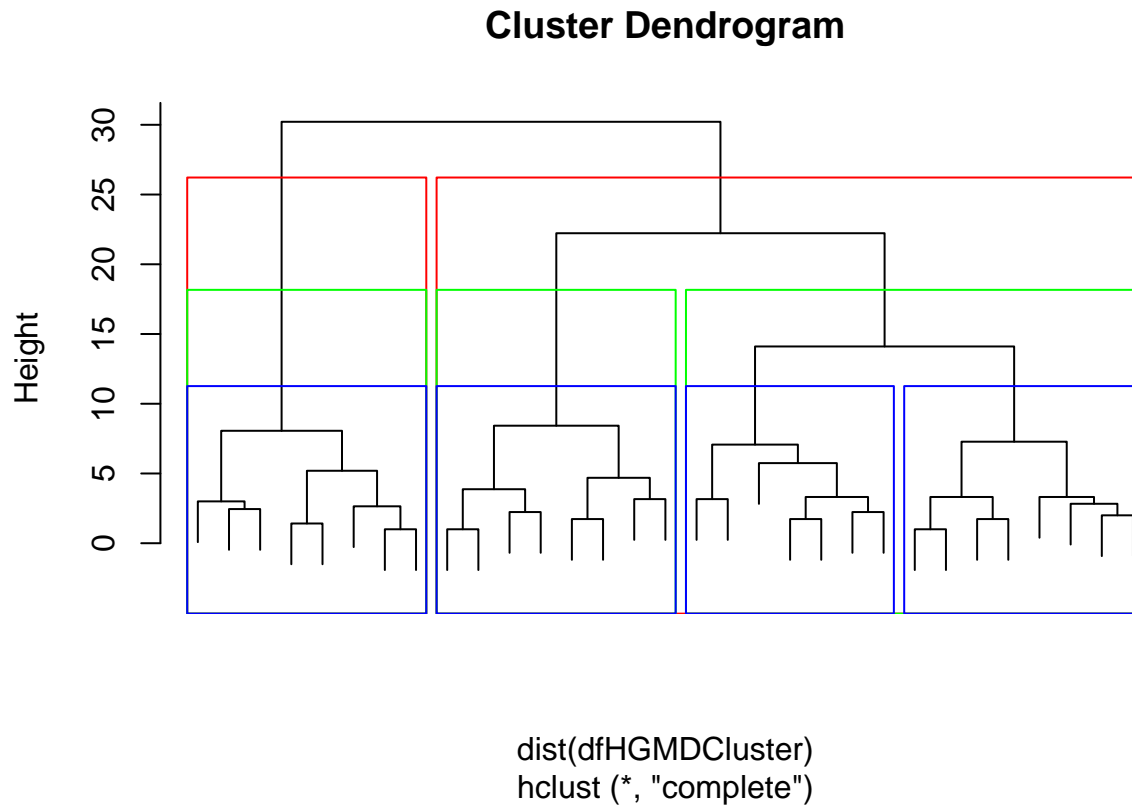
```
plot(dfHGMDCluster.h.clust.me)
```



```
dist(dfHGMDCluster)
hclust (*, "ward.D")
```

6.3.4.1 Plotting to determine the cluster level.

```
plot(dfHGMDCluster.h.clust, labels = FALSE)
rect.hclust(dfHGMDCluster.h.clust, k=2, border="red")
rect.hclust(dfHGMDCluster.h.clust, k=3, border="green")
rect.hclust(dfHGMDCluster.h.clust, k=4, border="blue")
```

This looks like $k = 3$ is a good number to cluster with.

7 Evaluation

7.1 System Time

system run time:

Table 1: Total Run Time

user	system	elapsed
74.72	2.23	132.91

7.2 Precision and Recall

Precision: fraction of retrieved docs that are relevant = relevant/retrieved
 Recall: fraction of relevant docs that are retrieved = retrieved/relevant

Table 2: Precision and Recall

Retrieved?	Relevant	Non-relevant
Retrieved	3	0
Not Retrieved	0	16

Retrieved?	Relevant	Non-relevant
------------	----------	--------------

Precision $P = 3/(3 + 0) = 1$

Recall $R = 3/(3 + 0) = 1$

8 Conclusion

9 References

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