[Tutorial] SCN2A mutations in neurodevelopmental disorders

1. Introduction

1.1 Background

SCN2A is a voltage-gated sodium channel gene that encodes the neuronal sodium channel NaV1.2 and plays a critical role in action potential initiation during early neurodevelopment. The latest study demonstrated that it is loss of function mutations that in SCN2A that lead to autism spectrum disorders (ASD), in contrast to gain of function, which leads to infantile seizures (Ben-Shalom 2018).

In this tutorial, we will handle genetic data for SCN2A mutations identified in latest genomic studies, and then explore the data format to describe genetic mutations using R basic functions. Our tutorial will utilize the summary data from Sanders et al. (2018).

1.2 Aims

What we will do with this dataset,

- Understand the dataset from a scientific journal
- Apply some functions you have learnt from the Chapter 2 and 3

2. Explore your data

2.1. Unboxing your dataset

Here we obtain the list of mutations in the **Supplementary Table 1** from Sanders et al. (2018).

Using the rio package, reading the excel file from the file link into your workspace. If you don't have the rio package in your system, please install as following:

```
# install.packages("rio")
```

Now you can read the file from the website. This will create the d object.

```
d <- rio::import('https://www.ncbi.nlm.nih.gov/pmc/articles/PMC6015533/bin/NIHMS957592-supplement-1.xls
```

Let's explore the object you just loaded. How would you check the class of the object d?

```
class(d)
```

```
## [1] "data.frame"
```

It shows that the d object is data.frame.

Then, let's overview the data frame. We will use head function to print out first few lines.

head(d)

##		PatientII) Pati	entSex	Patie	ntAgeAt <i>A</i>	Assessment	Chr	Pos_l	ng19	Ref
##	1			М		J	7у		_	_	166384278
##	2			F	•		Зу		163706	6754	166506754
##	3	Patient2	2	F	•		18m	2	16387	5903	166478766
##	4	Case1,280269)	F	•		4y	2	165798	3270	166304847
##				M	[3y	2	166019	9786	166249879
##	6			F	•		25y	2	166060	0054	166153823
##			Alt	Туре		Effect	c.DNA p.1		ein Ir	nheri	itence
##	1	12Mb Duplio			Duplic	ationCNV				Novol	Mosaic
##	2	2.8Mb Del		CNV	Del	etionCNV				Ι	DeNovo
##	3	2.6Mb Duplio	ation	CNV	Duplic	ationCNV				Inhe	erited
##	4	507kb Duplio	ation	CNV	Duplic	ationCNV				Inhe	erited
##	5	230kb Del	etion	CNV	Del	.etionCNV				Ur	nknown
##	6	94kb Del	etion	CNV	Del	.etionCNV				Ι	DeNovo
##			Sou	rce So	urcePM	IID Ben-S	Shalom2017	Woli	ff2017	AnyF	Recurrence
##	1	Vecchi et	al 20	011	218934	19	Y		N		1
##	2	Chen et	al 20	010	203464	23	Y		N		1
##	3	Yoshitomi et	al 20	015	258432	248	Y		N		1
##	4	Thuresson et	al 20	016	271533	34	Y		N		1
##	5	Celle et	al 20	013	240804	82	Y		N		1
##	6	Bartnik et	al 20	011	208072	23	Y		N		1
##		UniqueSample	e/Fami	ly Tru	eRecur	rence Se	eizures Se	izure	eOnsetI	Days	SeizureType
##	1			Y		1	Y			90	Unknown
##	2			Y		1	N				None
##	3			Y		1	Y			3	Unknown
##	4			Y		1	Y			3	Unknown
##	5			Y		1	N				None
##	6			Y		1	Y			365	Unknown
##		ASD DD/1		ID sev	•						OtherFeatures
##	1	N	Y		Mild						clumsiness
##	2	Y	Y	S	evere		d;	ysmoı	rphia,	imma	ature myelination
##		Unknown	Y	S	evere						cerebral atrophy
##	_	N	N								•
##	5	Y	Y		evere						microcephaly
##	6	N	Y	Mod	lerate	hypotoni	la, bipola	r dis	sorder	, bel	navioral problems
##		Classificat									
		IEE_Mild/Ata									
##		ASI)/DD								
##			IEE								
##			BIS								
##)/DD								
##	6	ASI)/DD								

When you execute code in a notebook chunk, an output will be visible immediately beneath the input. From this, you can see several rows and columns in the data frame.

Let's look at the first column PatientID and check which class it is.

class(d\$PatientID)

[1] "character"

Character

Cool. Now you can see the TrueRecurrence column. What is the class of the column TrueRecurrence?

```
class(d$TrueRecurrence)
```

```
## [1] "numeric"
```

Numeric

To check the class of columns, you don't need to type an individual column. We can overview the summary of the dataset using summary function. Which column has the character class?

summary(d)

## ## ## ## ##	PatientID Length:293 Class:character Mode:character	PatientSex Length:293 Class :character Mode :character	PatientAgeAtAssess: Length:293 Class:character Mode:character	Min. :2 1st Qu.:2 Median :2 Mean :2 3rd Qu.:2 Max. :2			
##	Pos_hg19	Ref	Alt	Туре			
##	Length:293	Length:293	Length: 293	Length:293			
##	Class :character	Class :character	Class :character	Class :character			
##	Mode :character	Mode :character	Mode :character	Mode :character			
##							
##							
##							
##	Effect	c.DNA	p.Protein	Inheritence			
##	Length: 293	Length:293	Length:293	Length:293			
##	Class :character	Class :character	Class :character	Class :character			
##	Mode :character	Mode :character	Mode :character	Mode :character			
##							
##							
##	~	a 51/75	D 61 2 0045				
##	Source	SourcePMID	Ben-Shalom2017	Wolff2017			
##	Length: 293	Length: 293	Length: 293	Length: 293			
##	Class :character	Class :character	Class :character	Class :character			
##	Mode :character	Mode :character	Mode :character	Mode :character			
##							
##							
## ##	Angra	UniqueComple/Femily	TrueRecurrence Seizures				
##	AnyRecurrence Min. : 1.000	UniqueSample/Family					
##		Length: 293	'	gth:293 ss :character			
	1st Qu.: 1.000	Class :character	· · · · · · · · · · · · · · · · · · ·				
##	Median : 1.000	Mode :character	Median:1.000 Mod	e :character			

```
##
    Mean
           : 2.119
                                          Mean
                                                 :1.768
##
    3rd Qu.: 2.000
                                          3rd Qu.:1.000
                                          Max.
##
  Max.
           :10.000
                                                 :7.000
                                                                  DD/ID
  SeizureOnsetDays
                       SeizureType
                                               ASD
##
##
   Length:293
                       Length:293
                                           Length:293
                                                              Length:293
   Class : character
                       Class : character
                                           Class : character
                                                              Class : character
##
   Mode :character
                       Mode :character
                                           Mode :character
                                                              Mode : character
##
##
##
##
   DD/ID severity
                       OtherFeatures
                                           Classification
   Length:293
                       Length: 293
                                           Length: 293
##
##
    Class :character
                       Class : character
                                           Class : character
   Mode :character
                       Mode :character
                                           Mode :character
##
##
##
##
# Every column except 'Chr', 'AnyRecurrene', 'TrueRecurrence'.
```

2.2 Difference between data frame and matrix

Here we will convert the data frame into a matrix, and compare which part will be different in this. To convert a data frame into a matrix, you can use the command called as matrix.

```
m=as.matrix(d)
```

Let's overview the matrix object. Can you tell difference with data frame?

```
head(m[,'TrueRecurrence'])

## 1 2 3 4 5 6

## "1" "1" "1" "1" "1"
```

I can see the difference between matrix and data frame! Matrices can only contain a single class of d

2.3. Subset and Sort

Some patients who have the SCN2A mutation (hereafter called "SCN2A patient") often have seizures. So we want to know when the seizure occurs in development.

Let's check the class first.

```
class(d$SeizureOnsetDays)
```

```
## [1] "character"
```

Why this column contains character? Let's head the first few lines.

head(d\$SeizureOnsetDays)

```
## [1] "90" "." "3" "3" "." "365"
```

t seems that some rows contain samples who do not have seizure or unknown information. It's represented by ".", and also recorded in another column called Seizures.

head(d\$Seizures)

```
## [1] "Y" "N" "Y" "Y" "N" "Y"
```

So we want to subset rows where the seizure phenotype is available.

```
d1<- d[d$Seizures == "Y",]</pre>
```

Let's see how many samples have seizure phenotypes? Then, you can ask when is the earliest days for the representation of seizure phenotype? How can we check this? The fisrt, as seen previously the SeizureOnsetDays column is character so we cannot apply functions for numeric.

head(d1\$SeizureOnsetDays)

```
## [1] "90" "3" "365" "30" "1825"
```

So we have to convert this into numeric first.

```
d1$SeizureOnsetDays2 <- as.numeric(d1$SeizureOnsetDays)</pre>
```

Warning: NAs introduced by coercion

Hmm. There's an warning for NA introduction. This is because some rows do not have character that we can properly convert from character to numeric. So possible solutions are either you can bear with this in your downstream analyses or 2) convert character into an appropriate form of numeric conversion.

Then, the question is how can we find the rows with NA? We will ask whether the rows contains NA or not using is na function. This will return boolean as to NA presence.

is.na(d1\$SeizureOnsetDays2)

```
[1] FALSE FALSE
##
                                               [13]
##
                                                                                                       TRUE FALSE FALSE FALSE FALSE FALSE FALSE FALSE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              TRUE FALSE FALSE
##
                                               [25] FALSE FALSE
                                               [37] FALSE F
                                               [49] FALSE FALSE FALSE FALSE FALSE FALSE FALSE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 TRUE FALSE FALSE
                                               [61] FALSE F
##
                                               [73] FALSE FALSE
                                                                                                                                                                                                                                              TRUE FALSE FALSE FALSE TRUE FALSE FALSE FALSE FALSE
                                               [85] FALSE TRUE FALSE FA
##
                                              [97] FALSE F
## [109] FALSE FAL
## [121] FALSE FALSE
## [133] FALSE FALSE
```

```
## [145] FALSE FAL
```

See we can find some rows with NA. One of them is the 7th row. Let's see how it looks like.

```
d1$SeizureOnsetDays[13]
```

```
## [1] "<365"
```

```
# In my data 'd1', 13th row have NA not 7th.
```

Here you have < (angle bracket) in the character so it won't properly converted to numeric information. Did you find more of these cases?

```
d1[is.na(d1$SeizureOnsetDays2),]$SeizureOnsetDays
```

```
## [1] "<365" "<365" "<365" "<28" "<30" "<365" "<365" "<365" "<365" "<365" "<365"
```

Our NAs all contains <, which prevent converting a character into a numeric. We would fix for downstream analyses. For example, we can convert <365 into 365. One function we can try is gsub. This replace your string into a format that you may not get NA. For example,

```
# gsub('pattern in your character', 'new character you want to replace', vectors for your character)
d1$SeizureOnsetDays3 <- gsub('<', '', d1$SeizureOnsetDays)
head(d1$SeizureOnsetDays3)</pre>
```

```
## [1] "90" "3" "365" "30" "1825"
```

Let's convert them into numerics.

```
d1$SeizureOnsetDays3<-as.numeric(d1$SeizureOnsetDays3)
```

Did you get warning for this? Now we can ask our initial question. When is the earliest day for having seizure?

```
min(d1$SeizureOnsetDays3)
```

[1] 0

3. Exercise

The dataset contains more details for genetic mutations in SCN2A patients. From this information, what can we analyze further?

Here I list up few questions you can examine further.

- Finding the position of the genetic mutations within SCN2A. Which information you would use? If you are not familiar with positional information on genetic variants (or mutations), please find the **Figure 1** or the slides for Mutation (BSMS205 Session 3-1).
- Counting the recurrent mutations at the same protein position (in other words, the same mutations seen across different patients), and examine whether the patients have similar phenotype.
- Finding the position where different consequences mutations occur. Please note that "consequences" are loss-of-function (Nonsense, Frameshift) or missense.
- Sketch a plot to visualize your analysis.

Exon 1

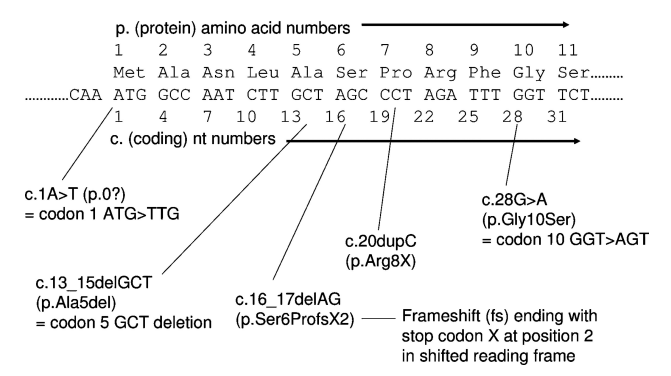


Figure 1: **Figure 1. Standard mutation nomenclature** (Ogino et al. 2007). According to this guideline, genetic mutations can be represented for coding DNA reference sequences (c. prefix) and protein-level amino acid sequences (p. prefix).

3.1 For-loops and Vectorization

Here we examine more details of genetic mutations as to their functional consequence and position of SCN2A mutations. In the dataset includes, there are two columns called c.DNA and p.Protein, containing the cDNA or protein position for the genetic mutations.

During these exercises, we will look at the concept of for-loop and vectorization, which you learn from the Chapter 3.4. Let's look at the column **p.Protein**. It contains protein positions from each patient. What would you check at the first place?

- Task 1. First, I want to overview this column using 'head()'
- Task 2. Overview this column using 'tail()'
- Task 3. Which class is it?
- Task 4. How many observations(samples)

- Task 5. How many samples with p.Protein information?
- ...
- Task N

library(tidyverse)

Let's write down your code to explore this column.

```
## -- Attaching packages ------ tidyverse 1.3.1 --
## v ggplot2 3.3.5
                    v purrr
                             0.3.4
## v tibble 3.1.3
                    v dplyr
                             1.0.7
## v tidyr
          1.1.3
                    v stringr 1.4.0
## v readr
           2.0.1
                    v forcats 0.5.1
## -- Conflicts ----- tidyverse conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                  masks stats::lag()
head(d$p.Protein)
## [1] "." "." "." "." "." "."
tail(d$p.Protein)
## [1] "p.R1902C" "p.R1902C" "p.Q1904E" "p.R1918H" "p.K1933M" "p.S1974L"
class(d$p.Protein)
## [1] "character"
nrow(d)
## [1] 293
d[d$p.Protein!=".",]%>%nrow()
```

[1] 285

If you need more information on SCN2A, please visit the Uniprot description for SCN2A. The Uniprot database contains description for protein domains.

Then, I would remove the characters from the string so we can have only numerics for positions. Here I use gsub function to extract numbers from string. Let's remove non-numeric characters from the string.

```
gsub('[^0-9]', '', 'p.R102X')
```

[1] "102"

In the dataset, we have many rows for protein positions. One way we might try is to set up for-loop to process each row.

```
for(i in 1:293){
   a <- gsub('[^0-9]','',d[i,'p.Protein'])
}</pre>
```

Do you think this is an effective approach? As we have done in your assignment, for-loop is not a good choice to process vectors because R can do vectorization for this process with a shorter and clearer code. So this mean you can apply gsub on vector and return your output to another column (could be new assign).

```
d$p.Protein1 <- gsub('[^0-9]','',d$p.Protein)
# I made new column named 'p.Protein1' in which outputs are saved.</pre>
```

3.2. Counting the recurrent mutations

Recurrent mutations are the ones that the same genetic mutations occur in multiple individuals. Recurrent mutations can be common 1) when the mutation does not affect on natural selection, 2) when the mutation is beneficial, 3) in the hotspot for a disease or strongly associated with trait. However, given we are dealing with the genetic mutations from rare disorders, the mutations in the dataset are supposed to be uniquely present in general population. Otherwise, the recurrent mutations can indicate strong association with the phenotype.

To assess the recurrent mutations, the first thing we can try is to examine whether the same mutations occur in multiple individuals. Since the dataset contains individual patients for each row, we can simply check the frequency using:

```
c <- as.data.frame(table(d$p.Protein))</pre>
```

or we can check the number of unique variants in the dataset by:

```
nrow(c)
```

[1] 203

```
# There are 203 unique variants in the dataset.
```

How many unique variants you can find? and which variants are occurred in multiple times?

```
filter(c,Freq>1 )%>%
  nrow()
```

[1] 40

```
# Among 203 unique variants, 40 are occurred in multiple times.
# But I just find out that one of them is undefined variants.
# In other words, 39 variants are occurred in multiple individuals.
```

Then, you can use other columns to check frequency for different groups. Which columns you would use for more grouping?

If you find that, please check the recurrent mutations for each group.

```
table()%>%
  as.data.frame%>%
  filter(Freq>1) # I got 40 recurrent mutations group using nrow() from here. But two groups are undefi
##
                  Freq
              ???
## 1
                     2
                     8
## 2
                     3
## 3
        c.106A>G
                     2
## 4
       c.1136G>A
## 5
       c.1267G>C
                     2
## 6
       c.2558G>A
                     9
                     2
##
  7
       c.2645G>A
## 8
       c.2695G>A
                     2
## 9
       c.2715G>C
                     2
       c.2783T>G
                     2
## 10
## 11
       c.2809C>T
                     2
## 12
       c.2932T>C
                     2
## 13
       c.2995G>A
                     6
                     2
## 14
        c.304C>T
##
  15 c.3057AA>A
                     2
## 16
       c.3631G>A
                     3
       c.3844G>T
                     2
## 17
                     2
## 18
      c.386+2T>C
## 19
       c.3956G>A
                     6
                     2
## 20
       c.4007C>A
## 21
       c.4025T>C
                     4
##
  22
        c.408G>T
                     2
## 23
       c.4303C>T
                     3
## 24
       c.4436A>C
                     2
## 25
       c.4565G>C
                     2
                     2
## 26
       c.4591C>A
      c.476+1G>A
                     2
## 27
## 28
       c.4777G>A
                     2
## 29
       c.4886G>A
                     2
##
  30
       c.5318C>T
                     3
                     2
## 31
        c.562C>T
## 32
       c.5644C>G
                     2
                     6
## 33
       c.5645G>A
##
  34
       c.5704C>T
                     2
##
  35
        c.632G>A
                     2
                     2
##
  36
        c.638T>A
                     2
##
   37
        c.710T>A
                     2
##
   38
        c.781G>A
## 39
        c.788C>T
                     8
## 40
        c.982T>G
                     2
```

I can use 'c.DNA' column to check frequency for different groups.

d\$c.DNA%>%

3.3 What is the proportion of diagnosis for SCN2A patient?

SCN2A mutation can have multiple different consequences for disease phenotypes. It can cause ASD but also other neurodevelopmental conditions. In total cases, how many phenotypes occur in SCN2A patients. Then, calculate the proportion of the phenotypes among total cases.

```
# I used column 'Classification' in which each patients are classified into different phenotypes.
d$Classification%>%
  table()%>%
  as.data.frame()%>%
 mutate(proportion=Freq/293)
##
                   . Freq proportion
## 1
              ASD/DD 92 0.31399317
## 2
                       36 0.12286689
                 BIS
## 3
                 IEE 111 0.37883959
## 4 IEE_Mild/Ataxia
                       7 0.02389078
## 5
               Other
                        3 0.01023891
## 6
                        5 0.01706485
       Schizophrenia
## 7
             Unclear 39 0.13310580
Then, you might be intrigued to whether females and males have different occurrence in each disorder. Let's
#Calculate total number of Female
Female_total <- d%>%
  filter(PatientSex=="F")%>%
  nrow()
Female_total #114
## [1] 114
#Calculate total number of Male
Male_total <- d%>%
  filter(PatientSex=="M")%>%
  nrow()
Male_total #119
## [1] 119
#Male and Female have difference occurences in each disorder. Check this below.
d%>%
  filter(PatientSex %in% c("F","M"))%>%
  group_by(PatientSex,Classification)%>%
  count()%>%
 mutate(proportion=ifelse(PatientSex=="F",n/Female_total,n/Male_total))
## # A tibble: 13 x 4
               PatientSex, Classification [13]
## # Groups:
##
      PatientSex Classification
                                     n proportion
                                             <dbl>
##
                 <chr>
      <chr>
                                 <int>
##
  1 F
                 ASD/DD
                                    26
                                           0.228
## 2 F
                 BIS
                                          0.0965
                                    11
## 3 F
                 IEE
                                    57
                                          0.5
## 4 F
                 IEE_Mild/Ataxia
                                     3
                                          0.0263
## 5 F
                 Other
                                     2
                                          0.0175
                                          0.00877
## 6 F
                 Schizophrenia
                                     1
```

```
7 F
                  Unclear
                                      14
                                            0.123
##
    8 M
                  ASD/DD
                                      44
                                            0.370
##
   9 M
                 BIS
                                      12
                                            0.101
                  IEE
                                            0.361
## 10 M
                                      43
## 11 M
                  IEE_Mild/Ataxia
                                       4
                                            0.0336
## 12 M
                  Other
                                       1
                                            0.00840
## 13 M
                  Unclear
                                      15
                                            0.126
```

Another question you can ask is whether different mutation consequences occur in each phenotype. Let's find out how many mutation consequences are observed in each phenotype.

```
group_by(Classification,Effect)%>%
  count()
## # A tibble: 25 x 3
               Classification, Effect [25]
## # Groups:
      Classification Effect
##
##
      <chr>
                      <chr>>
                                               <int>
##
   1 ASD/DD
                      DeletionCNV
                                                   3
##
   2 ASD/DD
                     DuplicationCNV
                                                   1
   3 ASD/DD
                      Frameshift
                                                  17
   4 ASD/DD
##
                     Missense
                                                  40
##
    5 ASD/DD
                      Nonsense
                                                  16
##
    6 ASD/DD
                      PopulationVariantInExAC
                                                   3
   7 ASD/DD
                      SpliceSite
                                                  12
##
   8 BIS
                      DuplicationCNV
                                                   1
##
   9 BIS
                      Frameshift
                                                   1
## 10 BIS
                      Missense
                                                  34
```

3.4. Find the position where different consequences of mutations occur

If you checked the recurrent mutations, you might want to find a locus where two or more variants occur. Such loci might indicate functionally important position of the gene and you might find some insight as to a cause of disease.

```
r_locus <- d%>%
  group_by(c.DNA)%>%
  count()%>%
  filter(n>1)
r_locus$loci <- gsub('[^0-9]','',r_locus$c.DNA)%>%
  as.numeric()
r_locus
```

```
## # A tibble: 40 x 3
## # Groups:
               c.DNA [40]
      c.DNA
##
                    n loci
##
      <chr>
                <int> <dbl>
  1 ???
##
                    2
                         NA
  2 .
                         NA
  3 c.106A>G
                    3
                        106
##
```

... with 15 more rows

```
4 c.1136G>A
                       1136
##
    5 c.1267G>C
                       1267
    6 c.2558G>A
                       2558
                       2645
    7 c.2645G>A
                    2
    8 c.2695G>A
                       2695
##
    9 c.2715G>C
                    2
                       2715
## 10 c.2783T>G
                       2783
## # ... with 30 more rows
```

3.5. Sketch a plot to visualize your analysis

When you examine the dataset, you would draw something to show your output. Though we haven't learnt how to plot data yet, we can have a quick sketch for the dataset. There's no restriction on your suggestion. Please submit your hand-drawing for the plot you would like to show from this dataset.

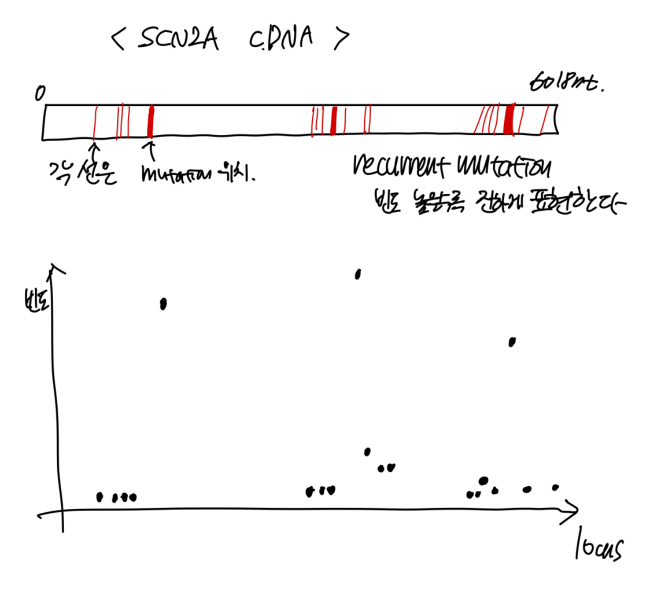


Figure 2: plot