

Class19

Daniel Xu

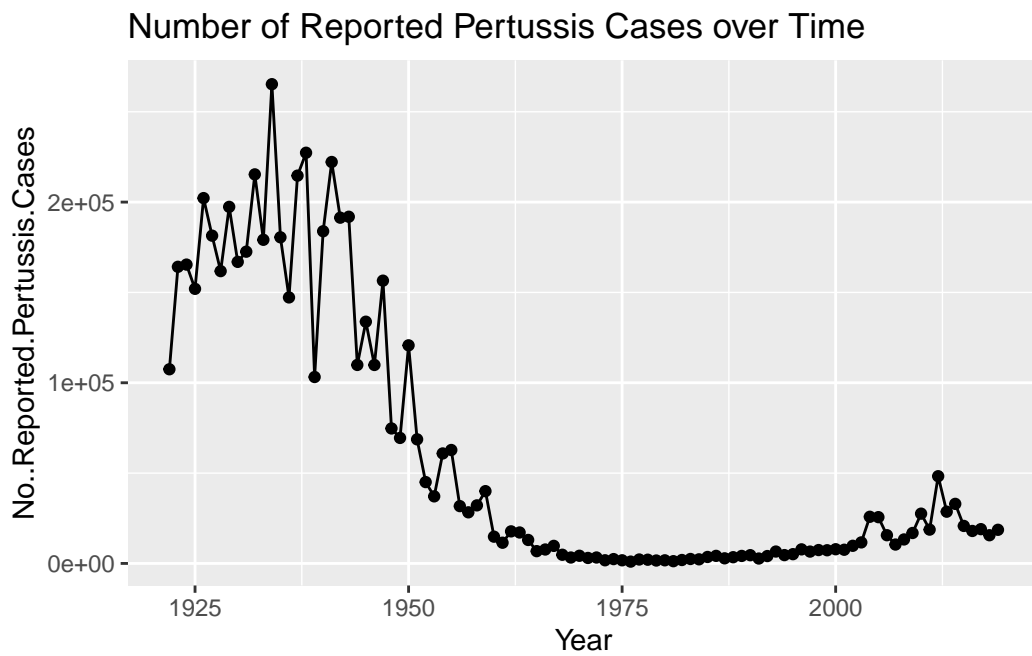
Q1)

```
cdc<-data.frame(
  Year = c(1922L,
    1923L, 1924L, 1925L, 1926L, 1927L, 1928L,
    1929L, 1930L, 1931L, 1932L, 1933L, 1934L, 1935L,
    1936L, 1937L, 1938L, 1939L, 1940L, 1941L,
    1942L, 1943L, 1944L, 1945L, 1946L, 1947L, 1948L,
    1949L, 1950L, 1951L, 1952L, 1953L, 1954L,
    1955L, 1956L, 1957L, 1958L, 1959L, 1960L,
    1961L, 1962L, 1963L, 1964L, 1965L, 1966L, 1967L,
    1968L, 1969L, 1970L, 1971L, 1972L, 1973L,
    1974L, 1975L, 1976L, 1977L, 1978L, 1979L, 1980L,
    1981L, 1982L, 1983L, 1984L, 1985L, 1986L,
    1987L, 1988L, 1989L, 1990L, 1991L, 1992L, 1993L,
    1994L, 1995L, 1996L, 1997L, 1998L, 1999L,
    2000L, 2001L, 2002L, 2003L, 2004L, 2005L,
    2006L, 2007L, 2008L, 2009L, 2010L, 2011L, 2012L,
    2013L, 2014L, 2015L, 2016L, 2017L, 2018L,
    2019L),
  No..Reported.Pertussis.Cases = c(107473,
    164191, 165418, 152003, 202210, 181411,
    161799, 197371, 166914, 172559, 215343, 179135,
    265269, 180518, 147237, 214652, 227319, 103188,
    183866, 222202, 191383, 191890, 109873,
    133792, 109860, 156517, 74715, 69479, 120718,
    68687, 45030, 37129, 60886, 62786, 31732, 28295,
    32148, 40005, 14809, 11468, 17749, 17135,
    13005, 6799, 7717, 9718, 4810, 3285, 4249,
    3036, 3287, 1759, 2402, 1738, 1010, 2177, 2063,
    1623, 1730, 1248, 1895, 2463, 2276, 3589,
```

```
4195,2823,3450,4157,4570,2719,4083,6586,
4617,5137,7796,6564,7405,7298,7867,
7580,9771,11647,25827,25616,15632,10454,
13278,16858,27550,18719,48277,28639,
32971,20762,17972,18975,15609,18617)
```

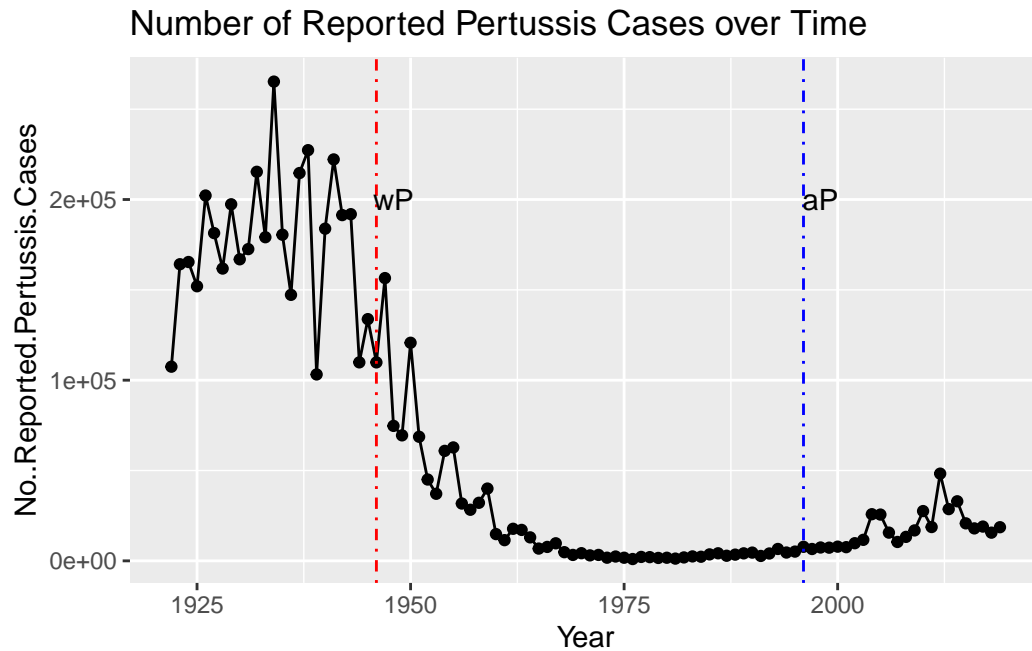
```
)
```

```
library(ggplot2)
ggplot(cdc,aes(Year,No..Reported.Pertussis.Cases))+geom_point()+geom_line()+labs(title="Nu
```



```
oplot<-ggplot(cdc,aes(Year,No..Reported.Pertussis.Cases))+geom_point()+geom_line()+labs(ti

oplot+geom_vline(xintercept=1946,color="red",linetype=4)+geom_vline(xintercept=1996,color=
```



Q3) Pertussis has increased in frequency in comparison to before the aP vaccine. This could be explained by aP being less effective than wP.

```
library(jsonlite)
```

Warning: package 'jsonlite' was built under R version 4.2.2

```
subject <- read_json("https://www.cmi-pb.org/api/subject", simplifyVector = TRUE)
```

```
head(subject, 3)
```

	subject_id	infancy_vac	biological_sex	ethnicity	race
1	1	wP	Female Not	Hispanic or Latino	White
2	2	wP	Female Not	Hispanic or Latino	White
3	3	wP	Female	Unknown	White

	year_of_birth	date_of_boost	dataset
1	1986-01-01	2016-09-12	2020_dataset
2	1968-01-01	2019-01-28	2020_dataset
3	1983-01-01	2016-10-10	2020_dataset

Q4)96 Q5) 66 Female, 30 Male

```
table(subject$biological_sex)
```

Female	Male
66	30

```
table(subject$biological_sex,subject$race)
```

	American Indian/Alaska Native	Asian	Black or African American
Female	0	18	2
Male	1	9	0

	More Than One Race	Native Hawaiian or Other Pacific Islander
Female	8	1
Male	2	1

	Unknown or Not Reported	White
Female	10	27
Male	4	13

Native,Asian,Black,Multiple,other,not reported,white Female: 0,18,2,8,1,10,27 Male:1,9,0,2,1,4,13

```
library(lubridate)
```

Warning: package 'lubridate' was built under R version 4.2.2

Loading required package: timechange

Warning: package 'timechange' was built under R version 4.2.2

Attaching package: 'lubridate'

The following objects are masked from 'package:base':

date, intersect, setdiff, union

```
subject$age <- today() - ymd(subject$year_of_birth)
library(dplyr)
```

Attaching package: 'dplyr'

The following objects are masked from 'package:stats':

filter, lag

The following objects are masked from 'package:base':

intersect, setdiff, setequal, union

```
ap <- subject %>% filter(infancy_vac == "aP")
round( summary( time_length( ap$age, "years" ) ) )
```

Min.	1st Qu.	Median	Mean	3rd Qu.	Max.
23	25	26	25	26	27

```
# wP
wp <- subject %>% filter(infancy_vac == "wP")
round( summary( time_length( wp$age, "years" ) ) )
```

Min.	1st Qu.	Median	Mean	3rd Qu.	Max.
28	32	35	36	40	55

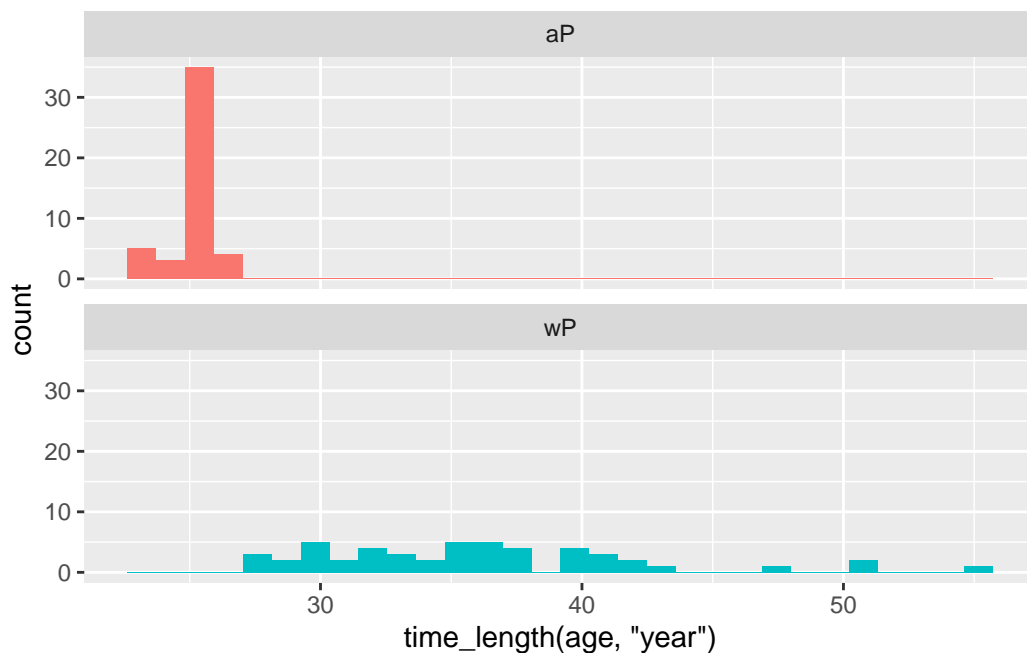
Q7) i) 36,ii)25,iii) yes they are significantly different Q8)

```
int <- ymd(subject$date_of_boost) - ymd(subject$year_of_birth)
age_at_boost <- time_length(int, "year")
head(age_at_boost)
```

```
[1] 30.69678 51.07461 33.77413 28.65982 25.65914 28.77481
```

```
ggplot(subject) +
  aes(time_length(age, "year"),
      fill=as.factor(infancy_vac)) +
  geom_histogram(show.legend=FALSE) +
  facet_wrap(vars(infancy_vac), nrow=2)
```

`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.



Q9) yes they are significantly different

```
specimen <- read_json("https://www.cmi-pb.org/api/specimen", simplifyVector = TRUE)
titer <- read_json("https://www.cmi-pb.org/api/ab_titer", simplifyVector = TRUE)
```

Q10)

```
meta <- inner_join(specimen, subject)
```

Joining, by = "subject_id"

```
dim(meta)
```

```
[1] 729 14
```

```
head(meta)
```

```
specimen_id subject_id actual_day_relative_to_boost
1           1           1                        -3
2           2           1                       736
3           3           1                        1
4           4           1                        3
5           5           1                        7
6           6           1                       11
planned_day_relative_to_boost specimen_type visit infancy_vac biological_sex
1                             0         Blood      1         wP         Female
2                            736         Blood     10         wP         Female
3                             1         Blood      2         wP         Female
4                             3         Blood      3         wP         Female
5                             7         Blood      4         wP         Female
6                            14         Blood      5         wP         Female
ethnicity race year_of_birth date_of_boost dataset
1 Not Hispanic or Latino White 1986-01-01 2016-09-12 2020_dataset
2 Not Hispanic or Latino White 1986-01-01 2016-09-12 2020_dataset
3 Not Hispanic or Latino White 1986-01-01 2016-09-12 2020_dataset
4 Not Hispanic or Latino White 1986-01-01 2016-09-12 2020_dataset
5 Not Hispanic or Latino White 1986-01-01 2016-09-12 2020_dataset
6 Not Hispanic or Latino White 1986-01-01 2016-09-12 2020_dataset
age
1 13481 days
2 13481 days
3 13481 days
4 13481 days
5 13481 days
6 13481 days
```

```
abdata <- inner_join(titer, meta)
```

```
Joining, by = "specimen_id"
```

```
dim(abdata)
```

```
[1] 32675    21
```

Q11)

```
IgE IgG IgG1 IgG2 IgG3 IgG4 6698 1413 6141 6141 6141 6141
```

```
table(abdata$isotype)
```

```
 IgE  IgG  IgG1  IgG2  IgG3  IgG4  
6698 1413 6141 6141 6141 6141
```

```
table(abdata$visit)
```

```
 1    2    3    4    5    6    7    8  
5795 4640 4640 4640 4640 4320 3920   80
```

There is significantly less people

```
ig1 <- abdata %>% filter(isotype == "IgG1", visit!=8)  
head(ig1)
```

	specimen_id	isotype	is_antigen_specific	antigen	MFI	MFI_normalised
1	1	IgG1	TRUE	ACT	274.355068	0.6928058
2	1	IgG1	TRUE	LOS	10.974026	2.1645083
3	1	IgG1	TRUE	FELD1	1.448796	0.8080941
4	1	IgG1	TRUE	BETV1	0.100000	1.0000000
5	1	IgG1	TRUE	LOLP1	0.100000	1.0000000
6	1	IgG1	TRUE	Measles	36.277417	1.6638332

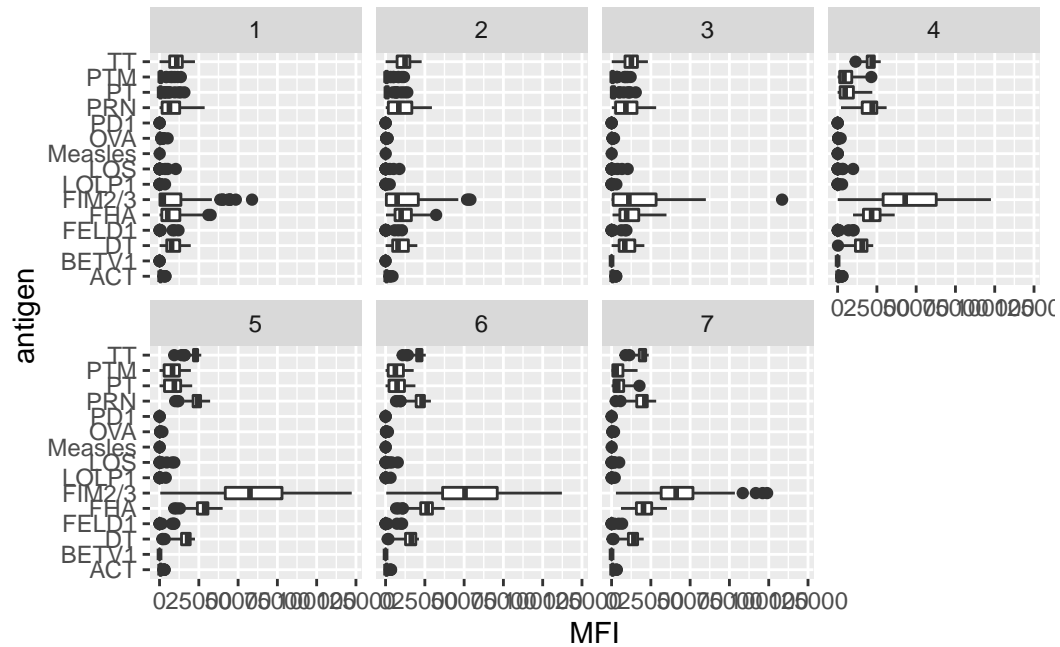
	unit	lower_limit_of_detection	subject_id	actual_day_relative_to_boost
1	IU/ML	3.848750	1	-3
2	IU/ML	4.357917	1	-3
3	IU/ML	2.699944	1	-3
4	IU/ML	1.734784	1	-3
5	IU/ML	2.550606	1	-3

6	IU/ML	4.438966	1	-3
	planned_day_relative_to_boost	specimen_type	visit	infancy_vac biological_sex
1	0	Blood	1	wP Female
2	0	Blood	1	wP Female
3	0	Blood	1	wP Female
4	0	Blood	1	wP Female
5	0	Blood	1	wP Female
6	0	Blood	1	wP Female

	ethnicity	race	year_of_birth	date_of_boost	dataset
1	Not Hispanic or Latino	White	1986-01-01	2016-09-12	2020_dataset
2	Not Hispanic or Latino	White	1986-01-01	2016-09-12	2020_dataset
3	Not Hispanic or Latino	White	1986-01-01	2016-09-12	2020_dataset
4	Not Hispanic or Latino	White	1986-01-01	2016-09-12	2020_dataset
5	Not Hispanic or Latino	White	1986-01-01	2016-09-12	2020_dataset
6	Not Hispanic or Latino	White	1986-01-01	2016-09-12	2020_dataset

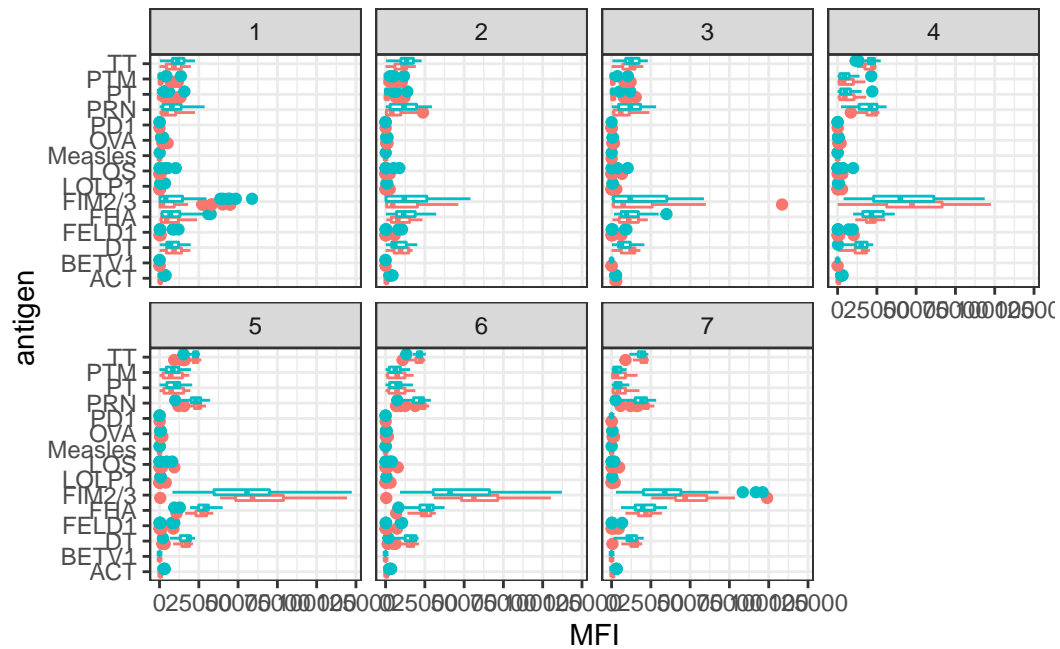
	age
1	13481 days
2	13481 days
3	13481 days
4	13481 days
5	13481 days
6	13481 days

```
ggplot(ig1) +
  aes(MFI, antigen) +
  geom_boxplot() +
  facet_wrap(vars(visit), nrow=2)
```



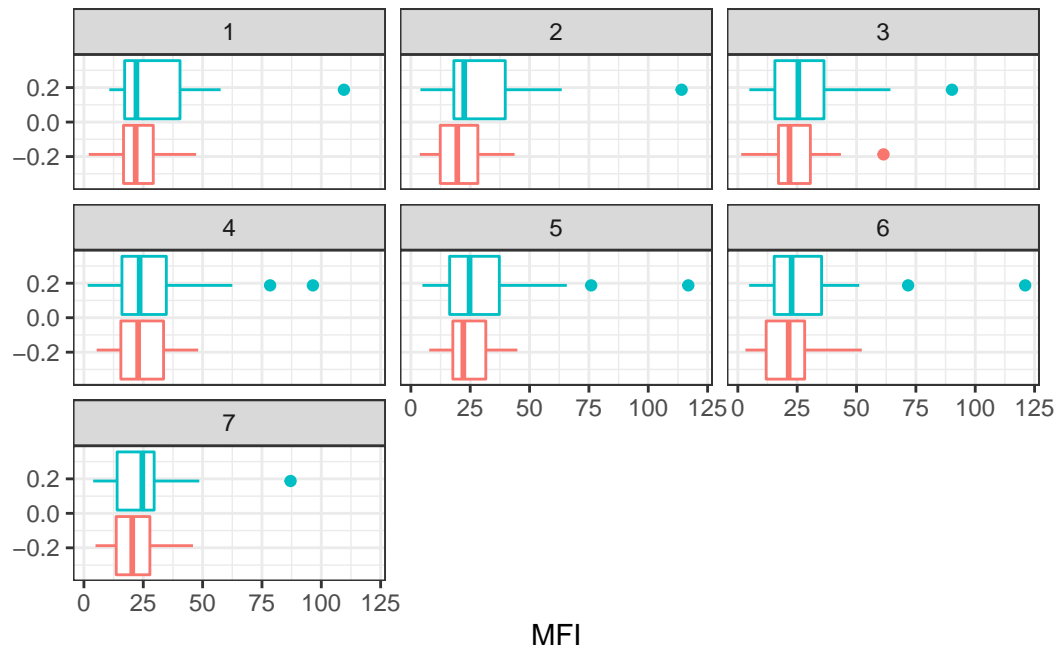
Q14) FIM2/3 Antigen seems to be responding. Maybe this antigen is what corresponds to pertussis

```
ggplot(ig1) +
  aes(MFI, antigen, col=infancy_vac ) +
  geom_boxplot(show.legend = FALSE) +
  facet_wrap(vars(visit), nrow=2) +
  theme_bw()
```

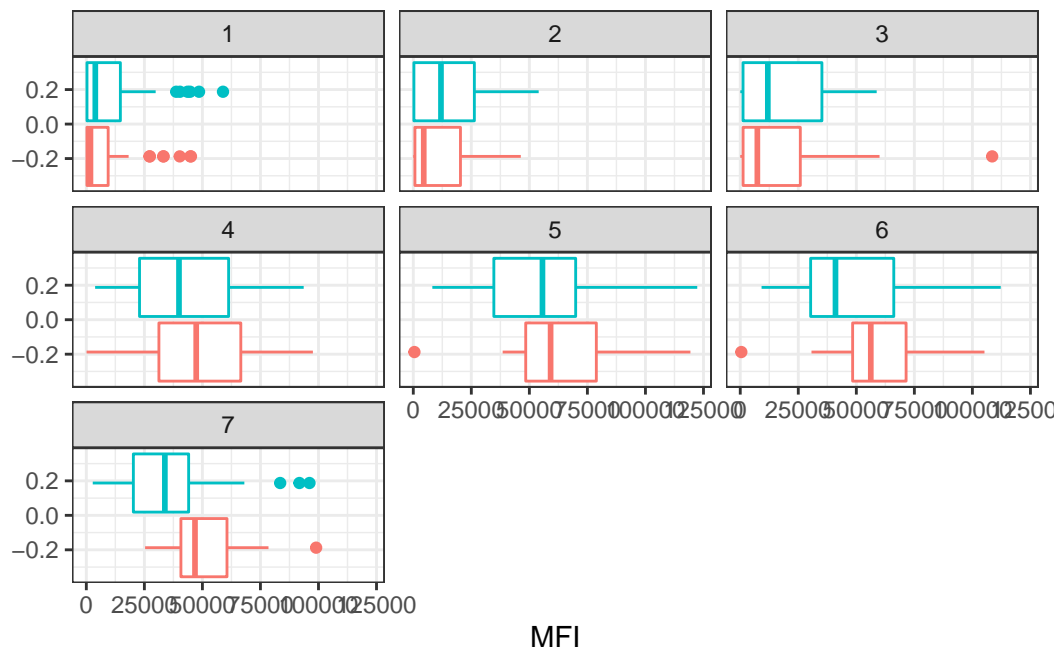


Q15)

```
filter(ig1, antigen=="Measles") %>%
  ggplot() +
  aes(MFI, col=infancy_vac) +
  geom_boxplot(show.legend = FALSE) +
  facet_wrap(vars(visit)) +
  theme_bw()
```



```
filter(ig1, antigen=="FIM2/3") %>%
  ggplot() +
  aes(MFI, col=infancy_vac) +
  geom_boxplot(show.legend = FALSE) +
  facet_wrap(vars(visit)) +
  theme_bw()
```



Q16) over time the antigen levels of the aP are significantly growing over time compared to the wP. In both the antigen level increases over time

Q17) aP has higher antigen level over time compared to wP

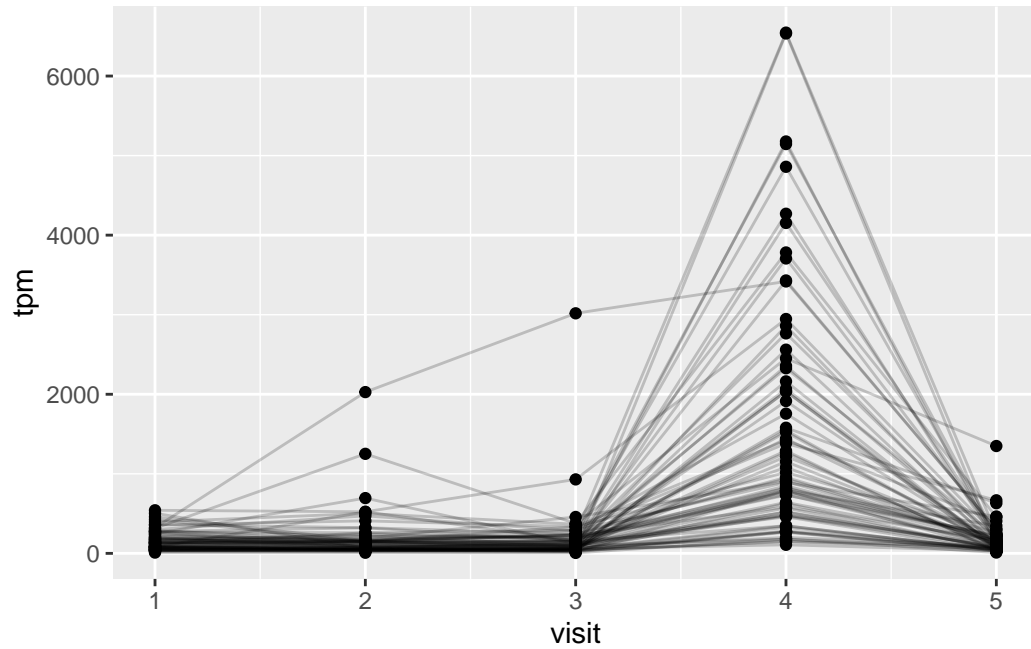
```
url <- "https://www.cmi-pb.org/api/v2/rnaseq?versioned_ensembl_gene_id=eq.ENSOG00000211896."

rna <- read_json(url, simplifyVector = TRUE)
ssrna <- inner_join(rna, meta)
```

Joining, by = "specimen_id"

Q18)

```
ggplot(ssrna) +
  aes(visit, tpm, group=subject_id) +
  geom_point() +
  geom_line(alpha=0.2)
```



Q19)

It is at maximum level at visit 4

Q20)

No the ab titer data shows continuous increase instead of the peak there is at visit 4 Maybe maximum expression occurs at visit 4 and the consequences of this peak expression is seen over time by the gradual increase in ab titer.