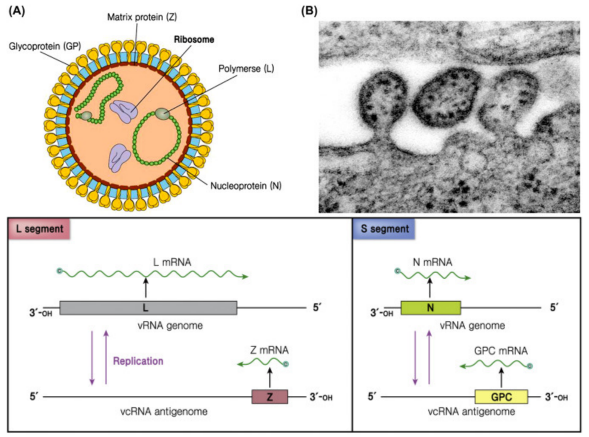
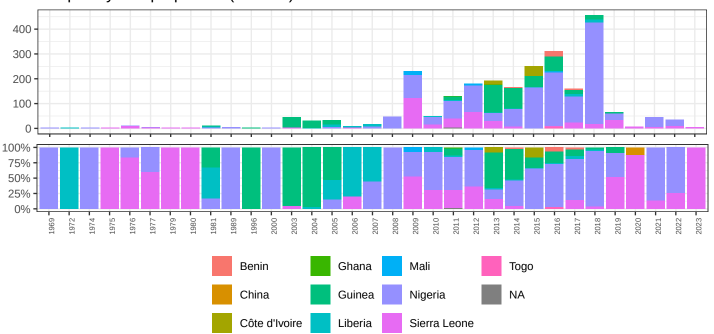
Background

Lassa has two segments "L" and "S" from Chapter 16 of "Molecular Virology of Human Pathogenic Viruses" by Wang-Shick Ryu

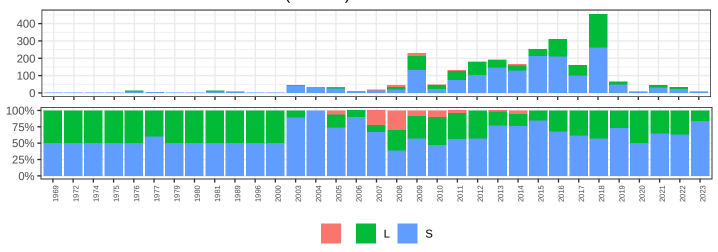


Exploratory Graphics

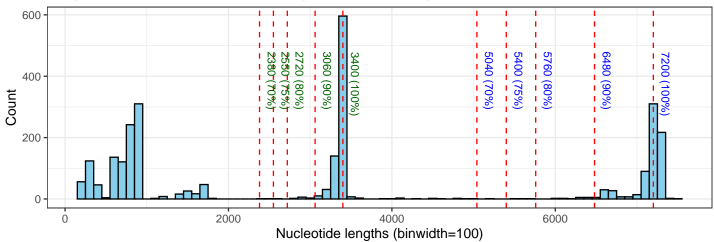
Frequency and proportion (n=2527)



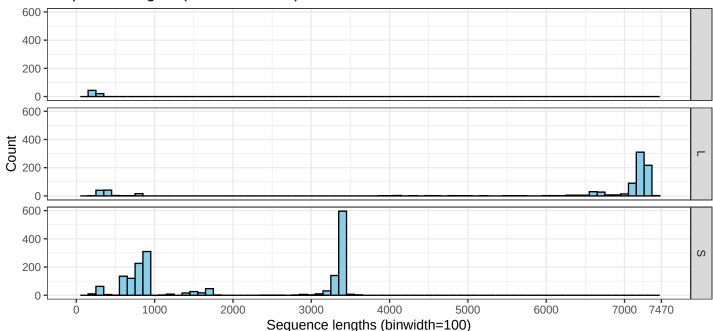
Lassa entries with collection date (n=2527)



Diagnostic plot to estimate min-length filter for phylogenetic analysis (all data = 2700)



Sequence lengths (all data = 2700)



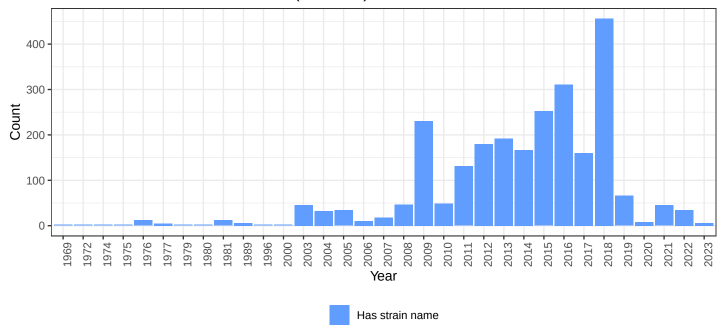
- Total lassa records = 2700.
- Total L lassa records = 846
- Total S lassa records = 1789
- Non L or S lassa records = 65

Lassa strain name

Is setting the strain name helpful for lassa? It looks like a majority of the strain names are the GenBank accession anyways.

```
nrow(data)
## [1] 2700
strainset=data %>%
  subset(strain != accession) %>%
  nrow(.)
cdata %>%
  ggplot(., aes(x=col_year, fill=strain != accession)) +
 geom_bar() +
  theme_bw() +
 theme(
    axis.text.x = element_text(angle=90, vjust=1, hjust=1),
   legend.position = "bottom",
   legend.title = element_blank()
 labs(
    title=paste("Lassa entries with collection date (n = ",nrow(cdata),")", sep=""),
    x="Year", y="Count"
  scale_fill_manual(values = c("TRUE" = "#619CFF", "FALSE" = "#F8766D"),
                    labels = c("TRUE" = "Has strain name", "FALSE" = "Uses accession"))
```

Lassa entries with collection date (n = 2527)



- Total lassa records = 2700. Notice how this includes samples that lack a collection date.
- Total lassa records where strain does not equal accession = 2694
- Percentage with strain names 99.78%
- Pulling in "strain=(.*)" filled in more samples (as compared to 49%)

Strain name duplicates

Are there more than 2 strain name duplicates? More than the S and L segments?

```
library(gt)
count_names <- data %>%
 group_by(strain) %>%
 summarize(n=n(), average_length=mean(length)) %>%
 arrange(desc(n))
more_names <- count_names %>%
 filter(n>2)
# Print top 10
more_names %>%
 head(10) %>%
 gt() %>%
  tab_header(title = "Top 10 Strains Name Duplicates by Count") %>%
 fmt_number(columns = c(n, average_length), decimals = 0) %>%
  cols_label(
   strain = "Strain Name",
   n = "Count Duplicates",
   average_length = "Average Length"
```

Top 10 Strains Name Duplicates by Count

Strain Name	Count Duplicates	Average Length
Josiah	15	3,878
AV	4	5,319
$LASV_3523$	4	2,508
$LASV_3604$	4	2,525
LASV_3609	4	2,500
$LASV_3625$	4	2,524
$LASV_3629$	4	2,510
$LASV_{-}3630$	4	2,524
$LASV_3706$	4	2,516
LASV_3711	4	2,506

- Percentage with strain names 99.78%
- Number of strain names that have 1 sequence record: 576
- Number of strain names that have 2 sequence records: 944
- Number of strain names that have more than 2 sequence records: 70

Table 2: Top 25 most frequent sequence submitters with their region and countries

		countries
361		Sierra Leone, Liberia, Nigeria
254	Africa	Nigeria
214	Africa	Nigeria
193	Africa	Nigeria, Sierra Leone, Liberia, Guinea
179	Africa	Guinea
170	Africa	Nigeria
156	Africa	Nigeria
143	Africa	Nigeria, Guinea
132	Africa	Guinea
106	Africa	Sierra Leone
68	Africa	Sierra Leone, Guinea
63	Africa	Nigeria
57	NA	NA
56	Africa	Liberia, Nigeria, Sierra Leone, Guinea
52	Africa	Côte d'Ivoire
40	Africa	Nigeria
40	Africa	Sierra Leone
36	Africa	Sierra Leone
35	Africa	Nigeria
33	Africa	Guinea
32	Africa	Liberia, Guinea
31	Africa	Benin
22	Africa	Mali
20	Africa	Nigeria, Benin
18	Africa	Guinea
	214 193 179 170 156 143 132 106 68 63 57 56 52 40 40 36 35 33 32 31 22	361 Africa 254 Africa 214 Africa 193 Africa 179 Africa 170 Africa 156 Africa 143 Africa 106 Africa 68 Africa 57 NA 56 Africa 52 Africa 40 Africa 36 Africa 35 Africa 32 Africa 31 Africa 22 Africa 20 Africa