basic

```
library(data.table)
library(dplyr)
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:data.table':
##
##
       between, first, last
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
       intersect, setdiff, setequal, union
library(stringr)
library(ggplot2)
library(NMF)
## Loading required package: pkgmaker
## Loading required package: registry
##
## Attaching package: 'pkgmaker'
## The following object is masked from 'package:base':
##
##
       isNamespaceLoaded
## Loading required package: rngtools
## Loading required package: cluster
## NMF - BioConductor layer [NO: missing Biobase] | Shared memory capabilities [NO: bigmemory] | Cores
     To enable the Bioconductor layer, try: install.extras('
##
## NMF
## ') [with Bioconductor repository enabled]
    To enable shared memory capabilities, try: install.extras('
## NMF
## ')
library(reshape2)
##
## Attaching package: 'reshape2'
## The following objects are masked from 'package:data.table':
##
##
       dcast, melt
library(parallel)
library(RColorBrewer)
library(scales)
```

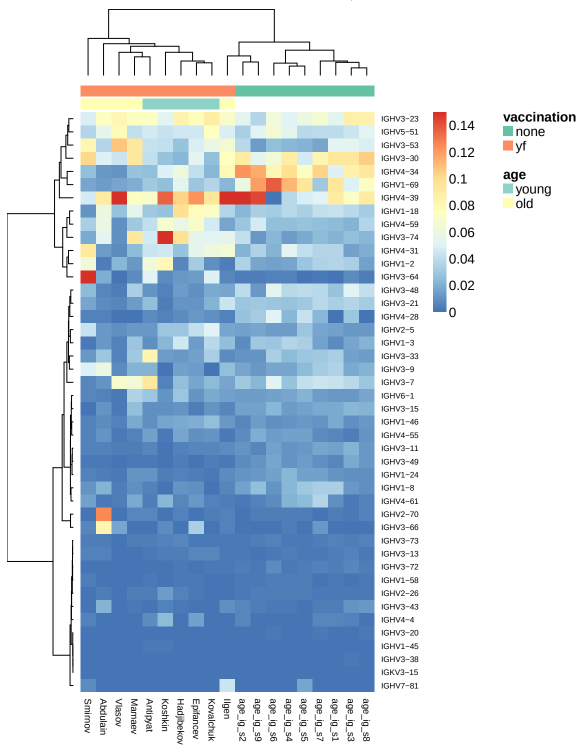
```
select = dplyr::select
summarise = dplyr::summarise

load("mixcr_processed_downsampled.Rda")
#load("mixcr_processed.Rda")
#dt.clones <- dt.clones %>% filter(replica == 1)
#dt.clones <- dt.clones %>% filter(replica == 2 | group == "control")
#dt.clones <- rbind(filter(dt.clones, vaccination != 'yf'),
# mutate(filter(dt.clones, vaccination == 'yf'), freq = freq/2))</pre>
```

V usage

```
dt.vusage = dt.clones %>%
  group by (sample name, vaccination, age, v) %>%
  summarise(freq = sum(freq)) %>% filter(freq > 0.001)
mat.vusage = dt.vusage %>%
  dcast(sample_name + vaccination + age ~ v, fill = 0)
## Using freq as value column: use value.var to override.
dt.annot = mat.vusage[,2:3]
dt.annot$age = factor(dt.annot$age, levels = c('young', 'old'))
rownames(mat.vusage) = mat.vusage$sample_name
mat.vusage = mat.vusage[,4:ncol(mat.vusage)]
ann_colors = c()
i = 2
for (annR in colnames(dt.annot)) {
 n = length(unique(dt.annot[[annR]]))
  tmp = list(x = c(brewer.pal(n, name = paste0("Set",i))[1:n]))
 names(tmp) = annR
 ann_colors = c(ann_colors, tmp)
  i = i + 1
}
## Warning in brewer.pal(n, name = paste0("Set", i)): minimal value for n is 3, returning requested pal
pdf("figures/p1.pdf", width = 6, height = 8)
aheatmap(pmin(as.matrix(t(mat.vusage)), 0.15),
         hclustfun = "ward",
         annCol = dt.annot, annColors = ann colors,
         scale = "none")
## The "ward" method has been renamed to "ward.D"; note new "ward.D2"
## The "ward" method has been renamed to "ward.D"; note new "ward.D2"
dev.off()
## pdf
##
```

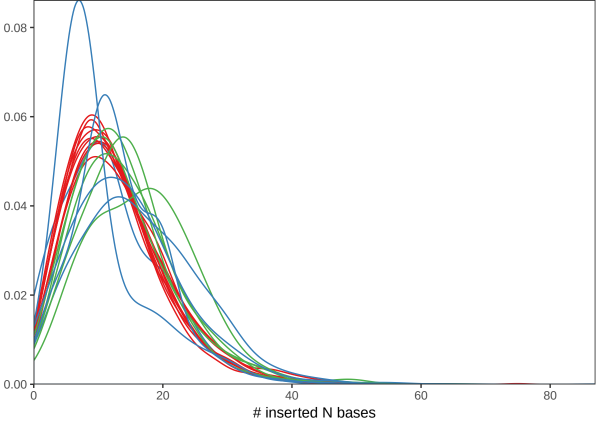
The "ward" method has been renamed to "ward.D"; note new "ward.D2"
The "ward" method has been renamed to "ward.D"; note new "ward.D2"



VDJ junction structure

```
dt.ins = dt.clones %>% filter(!is.na(dEnd)) %>%
  mutate(nIns = dStart - vEnd + jStart - dEnd) %>%
  group_by(sample_name) %>%
  mutate(freq = freq/sum(freq))

p2a=ggplot(dt.ins, aes(x = nIns, group = sample_name, weight = freq, color = group)) +
  geom_density(adjust = 2) +
  #scale_x_continuous(limits=c(-0.001,50)) +
  scale_color_brewer(guide = F, palette = "Set1") +
  scale_x_continuous("# inserted N bases", expand = c(0,0)) + scale_y_continuous("", expand = c(0,0)) +
  theme_bw() +
  theme(panel.grid.major = element_blank(), panel.grid.minor = element_blank())
p2a
```



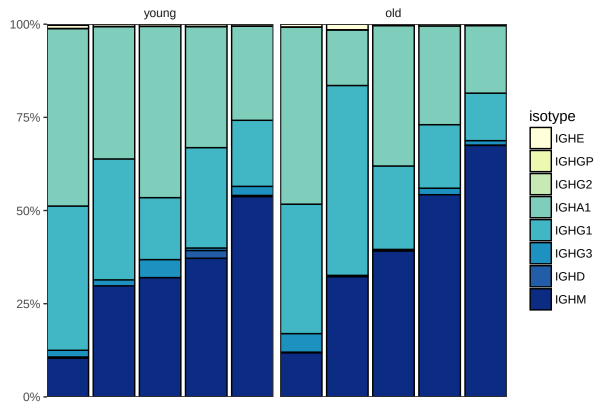
```
dt.ins.s = dt.ins %>%
  group_by(group, sample_name) %>%
  summarise(nInsS = sum(nIns * freq))

p2b=ggplot(dt.ins.s, aes(x = group, group = group, y = nInsS, fill = group)) +
  geom_boxplot() +
  #scale_x_continuous(limits=c(-0.001,50)) +
  scale_fill_brewer("Group", palette = "Set1") +
  xlab("") + ylab("") +
  theme_bw() +
```

```
theme(aspect=1,
        panel.grid.major = element_blank(), panel.grid.minor = element_blank(),
        axis.text.x = element_blank(), axis.ticks.x = element_blank())
p2b
16
                                                                 Group
14
                                                                    control
                                                                    yf-old
                                                                    yf-young
12
a=aov(nInsS~group,dt.ins.s)
summary(a)
##
               Df Sum Sq Mean Sq F value Pr(>F)
                           5.014
                                    2.365 0.126
## group
                2 10.03
## Residuals
               16 33.93
                            2.121
TukeyHSD(a, "group")
##
     Tukey multiple comparisons of means
##
       95% family-wise confidence level
##
## Fit: aov(formula = nInsS ~ group, data = dt.ins.s)
##
## $group
##
                         diff
                                      lwr
                                               upr
                                                       p adj
## yf-old-control
                     0.5826824 \ -1.5131551 \ 2.678520 \ 0.7569211 
## yf-young-control 1.7655678 -0.3302696 3.861405 0.1065288
## yf-young-yf-old 1.1828855 -1.1935708 3.559342 0.4237665
```

Isotype usage

```
dt.iso = dt.clones %>%
  filter(vaccination != "none", isotype != "", !is.na(isotype)) %>%
  group_by(sample_name, age, isotype) %>%
  dplyr::summarise(freq = sum(freq)) %>%
  group by(sample name) %>%
  mutate(freq = freq / sum(freq))
dt.iso$sample_name = factor(dt.iso$sample_name,
                            with(dt.iso %>% filter(isotype == "IGHM"), sample_name[order(freq)]))
dt.iso$isotype = factor(dt.iso$isotype, levels = rev(c("IGHM", "IGHD", "IGHG3", "IGHG1", "IGHA1", "IGHG
dt.iso$age = factor(dt.iso$age, levels = c('young', 'old'))
levels = dt.iso <- filter(dt.iso, !is.na(isotype))</pre>
p3=ggplot(dt.iso, aes(x=sample_name, fill = isotype, y = freq)) +
  geom_bar(position = "stack", stat = "identity", color = "black") +
  facet_wrap(~age, scales = "free_x") +
  scale_x_discrete("",expand = c(0,0)) +
  scale_y_continuous("", expand = c(0,0),labels = percent) +
  scale_fill_brewer(palette = "YlGnBu") +
  theme bw() +
  theme(panel.grid.major = element_blank(), panel.grid.minor = element_blank(),
  axis.text.x = element_blank(), axis.ticks.x = element_blank(),
  strip.background = element_blank())
рЗ
```

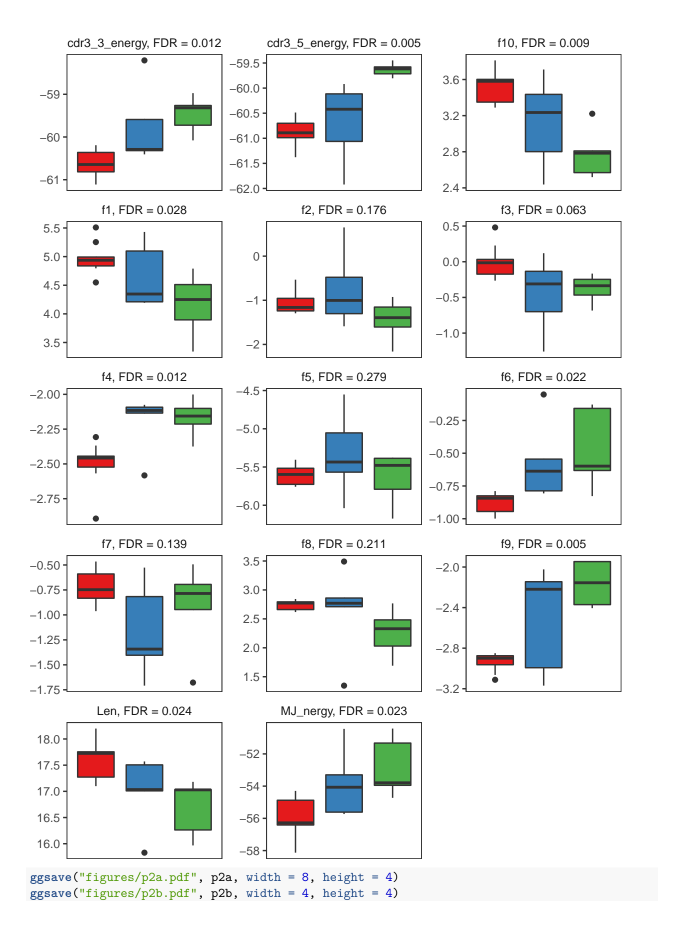


```
dt.p = data.table()
for (i in unique(dt.iso$isotype)) {
  a = aov(freq~age, dt.iso %>% filter(isotype == i))
 p = summary(a)[[1]][["Pr(>F)"]][1]
 dt.p = rbind(dt.p,
 data.table(isotype = i, p.adj = p))
dt.p$p.adj = p.adjust(dt.p$p.adj, method="BH")
print(dt.p)
##
      isotype
                 p.adj
## 1:
       IGHA1 0.7341534
## 2:
        IGHE 0.7655145
## 3:
       IGHG1 0.8968585
## 4: IGHG3 0.7655145
## 5: IGHGP 0.7341534
## 6:
       IGHM 0.7655145
## 7:
        IGHD 0.7341534
## 8:
       IGHG2 0.7341534
Amino acid feature analysis
dt.clones.cdr3prop = dt.clones %>%
  group_by(sample_name, group, cdr3aa) %>%
  dplyr::summarise(freq = sum(freq))
dt.cdr3.flat = unique(dt.clones.cdr3prop$cdr3aa) %>%
  strsplit("") %>%
  mclapply(function(x) data.table(aa = x, cdr3aa = paste0(x, collapse = "")), mc.cores = 1) %>%
 rbindlist
dt.cdr3.flat.ann = dt.cdr3.flat %>%
  merge(fread("kidera.txt") %>% mutate(Len = 1) %>% melt, allow.cartesian = T) %>%
  group_by(cdr3aa, variable) %>%
  summarise(value = sum(value))
## Using aa as id variables
dt.clones.cdr3prop2 = dt.clones.cdr3prop %>%
  merge(dt.cdr3.flat.ann, by = "cdr3aa", allow.cartesian = T)
dt.clones.cdr3prop.s.1 = dt.clones.cdr3prop2 %>%
  group_by(sample_name, group, variable) %>%
  dplyr::summarise(value = sum(value #/ ifelse(variable == "Len", 1, nchar(cdr3aa))
                        * freq))
CDR3 center amino acids properties
dt.clones.cdr33prop = dt.clones %>%
  mutate(cdr3aa = str_sub(cdr3aa, round(nchar(cdr3aa)/2)-1, round(nchar(cdr3aa)/2)+1)) %>%
```

```
dt.clones.cdr33prop = dt.clones %>%
  mutate(cdr3aa = str_sub(cdr3aa, round(nchar(cdr3aa)/2)-1, round(nchar(cdr3aa)/2)+1)) %>%
  group_by(sample_name, group, cdr3aa) %>%
  dplyr::summarise(freq = sum(freq))
```

```
dt.clones.cdr35prop = dt.clones %>%
  mutate(cdr3aa = str_sub(cdr3aa, round(nchar(cdr3aa)/2)-2, round(nchar(cdr3aa)/2)+2)) %%
  filter(nchar(cdr3aa) == 5) %>%
  group_by(sample_name, group, cdr3aa) %>%
  dplyr::summarise(freq = sum(freq))
dt.clones.cdr3prop.2 = rbind(dt.clones.cdr33prop, dt.clones.cdr35prop)
dt.cdr3.flat.2 = unique(dt.clones.cdr3prop.2$cdr3aa) %>%
  strsplit("") %>%
  mclapply(function(x) data.table(aa = x, cdr3aa = paste0(x, collapse = "")), mc.cores = 60) %>%
 rbindlist
dt.energy = fread("prop_mj96t3.txt") %>% mutate(energy = select(., -aa) %>% rowSums())
dt.cdr3.flat.ann.2 = dt.cdr3.flat.2 %>%
  merge(dt.energy, allow.cartesian = T) %>%
  group by(cdr3aa) %>%
  summarise(energy = sum(energy))
dt.clones.cdr3prop.s.2 = dt.clones.cdr3prop.2 %>%
  merge(dt.cdr3.flat.ann.2, by = "cdr3aa", allow.cartesian = T) %>%
  mutate(len = nchar(cdr3aa)) %>%
  group_by(sample_name, group, len) %>%
  summarise(cdr3_energy = sum(energy #/ ifelse(variable == "Len", 1, nchar(cdr3aa))
                        * freq)) %>%
  mutate(value = cdr3_energy/len, variable = paste0("cdr3_",
                                                    ifelse(len == 3, '3', '5'),
                                                     "_energy")) %>%
  select(-len, -cdr3_energy)
dt.clones.cdr3prop.s <- rbind(dt.clones.cdr3prop.s.1, as_tibble(dt.clones.cdr3prop.s.2))</pre>
## Warning in bind_rows_(x, .id): binding factor and character vector,
## coercing into character vector
## Warning in bind_rows_(x, .id): binding character and factor vector,
## coercing into character vector
dt.p.pair = data.table()
pairs = list(c("yf-old", "yf-young"), c("yf-old", "control"), c("yf-young", "control"))
for (v in unique(dt.clones.cdr3prop.s$variable)) {
  for (pair in pairs){
    p1 = dt.clones.cdr3prop.s %>% filter(variable == v, group == pair[1])
    p2 = dt.clones.cdr3prop.s %>% filter(variable == v, group == pair[2])
    p = wilcox.test(p1$value, p2$value)$p.value
    dt.p.pair = rbind(dt.p.pair,
                 data.table(variable = v, group1 = pair[1], group2 = pair[2], p.adj = p))
  }
}
dt.p.pair$p.adj = p.adjust(dt.p.pair$p.adj, method="BH")
print(filter(dt.p.pair, p.adj < 0.05))</pre>
##
           variable group1
                               group2
                                           p.adj
                 f1 yf-young control 0.01398601
## 1
## 2
                 f3 yf-young control 0.03873050
```

```
## 3
                 f4 yf-young control 0.01864802
## 4
                     yf-old control 0.01398601
                 f6
## 5
                 f6 yf-young control 0.02937063
## 6
                 f8 yf-young control 0.03873050
                 f9 yf-young control 0.01048951
## 7
## 8
                f10 yf-young control 0.01048951
## 9
           MJ_nergy yf-young control 0.01864802
## 10
                Len yf-young control 0.01864802
## 11 cdr3_3_energy yf-young control 0.01048951
## 12 cdr3_5_energy
                    yf-old yf-young 0.03030303
## 13 cdr3_5_energy yf-young control 0.01048951
dt.p = data.table()
for (v in unique(dt.clones.cdr3prop.s$variable)) {
  a = aov(value~group,dt.clones.cdr3prop.s %>% filter(variable == v))
  p = summary(a)[[1]][["Pr(>F)"]][1]
  dt.p = rbind(dt.p,
               data.table(variable = v, p.adj = p))
}
dt.p$p.adj = p.adjust(dt.p$p.adj, method="BH")
dt.clones.cdr3prop.s1 <- dt.clones.cdr3prop.s %>%
         merge(dt.p) %>%
         mutate(variable2 = paste0(variable, ", FDR = ", round(p.adj, 3)))
\#dt.clones.cdr3prop.s1\$variable2 = factor(dt.clones.cdr3prop.s1\$variable2, levels = c("f1, FDR = 0.121")
#
                                                                                       "f4, FDR = 0.079"
#
                                                                                       "f7, FDR = 0.079"
                                                                                       "f10, FDR = 0.028
#
p4=ggplot(dt.clones.cdr3prop.s1,
       aes(x = group, fill = group, y = value)) +
  geom_boxplot() +
  scale_fill_brewer(guide = F, palette = "Set1") +
  facet_wrap(~variable2, scales = "free_y", ncol=3) + xlab("") + ylab("") +
  theme_bw() +
  theme(panel.grid.major = element_blank(), panel.grid.minor = element_blank(),
        axis.text.x = element_blank(), axis.ticks.x = element_blank(),
        strip.background = element_blank())
p4
```



```
ggsave("figures/p3.pdf", p3, width = 8, height = 4)
ggsave("figures/p4.pdf", p4, width = 4*2, height = 3*3)
```