Distribution, RS ratio and patterns of novel SHM

Mikhail Shugay, Anna Obraztsova 17/04/2017

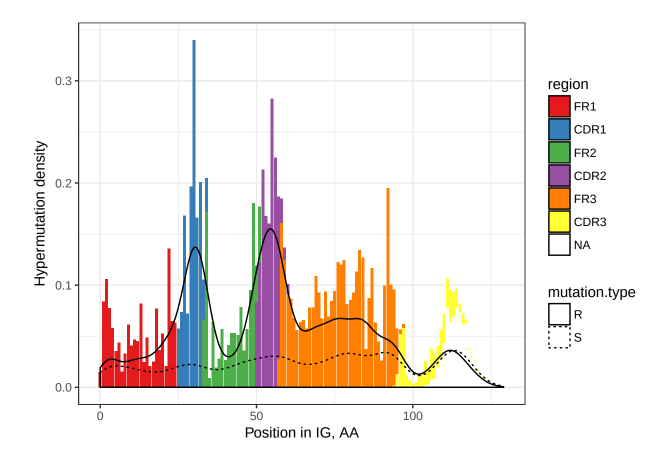
Analysis of substitution type and frequency

Load preprocessed data

Check if we observe well-documented increase in replacement:synonimic hypermutation ratio in CDR regions:

```
ggplot(df) +
  geom_bar(aes(x=pos.aa, weight=weight, fill=region)) +
  geom_density(aes(x=pos.aa, weight=weight, linetype = mutation.type)) +
  ylab("Hypermutation density") +
  xlab("Position in IG, AA") +
  scale_fill_brewer(palette = "Set1") +
  theme_bw()
```

```
## Warning in density.default(x, weights = w, bw = bw, adjust = adjust, kernel
## = kernel, : sum(weights) != 1 -- will not get true density
## Warning in density.default(x, weights = w, bw = bw, adjust = adjust, kernel
## = kernel, : sum(weights) != 1 -- will not get true density
```

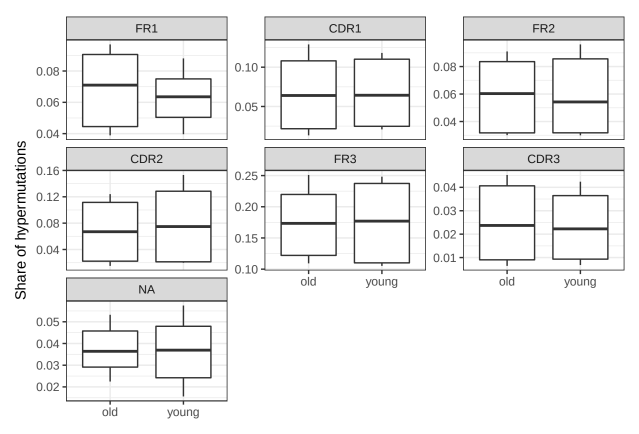


Comparative analysis

Summarize by region and type, compute frequencies and R:S ratio

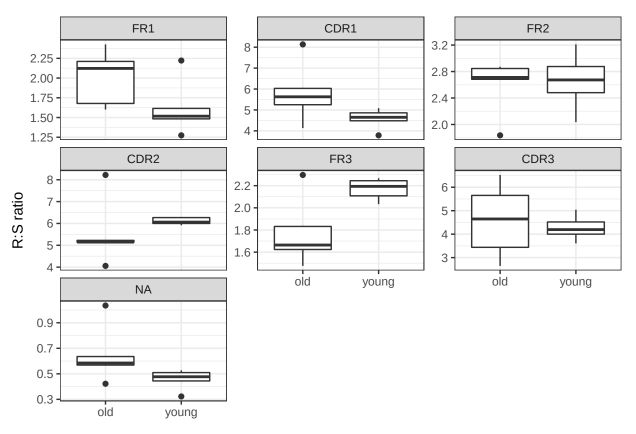
Fraction of errors in each region

```
ggplot(df.1, aes(x=proj, y = freq)) + geom_boxplot() +
facet_wrap(~region, scales = "free_y") +
xlab("") + ylab("Share of hypermutations") +
theme_bw()
```

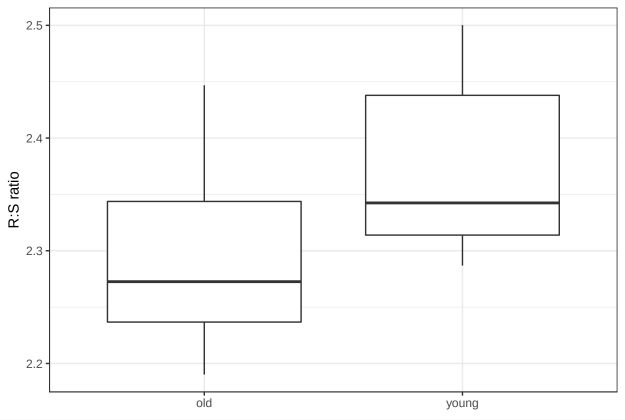


R:S ratio varies greatly by region

```
ggplot(subset(df.1, mutation.type == "R"), aes(x=proj, y = ratio)) +
  geom_boxplot() +
  facet_wrap(~region, scales = "free_y") +
  xlab("") + ylab("R:S ratio") +
  theme_bw()
```



Overall R:S ratio is higher in young



```
t.test(ratio ~ proj, subset(df.2, mutation.type == "R"))
```

Role of age factor can be deduced using generalized linear model for replacement hypermutation probability (binomial family). Probability of replacement hypermutations is increased by $7 \pm 1\%$ in young compared to old $(P < 10^{-6})$

```
df$R <- ifelse(df$mutation.type == "R", 1, 0)
fit <- glm(R ~ region + proj - 1, df, family = binomial())
summary(fit)</pre>
```

```
##
## Call:
## glm(formula = R ~ region + proj - 1, family = binomial(), data = df)
##
## Deviance Residuals:
## Min 1Q Median 3Q Max
```

```
## -1.9512 -1.4509
                     0.6020
                              0.8137
                                       0.9266
##
## Coefficients:
##
             Estimate Std. Error z value Pr(>|z|)
## regionFR1
              0.62334
                         0.02568 24.270
                                           <2e-16 ***
## regionCDR1 1.61618
                         0.03258 49.599
                                           <2e-16 ***
                         0.02932 30.226
              0.88610
## regionFR2
                                           <2e-16 ***
## regionCDR2 1.70720
                         0.03304 51.663
                                           <2e-16 ***
## regionFR3
              0.93541
                         0.01907 49.063
                                           <2e-16 ***
## regionCDR3 1.17395
                         0.04603 25.503
                                           <2e-16 ***
## projyoung
              0.03508
                         0.02043
                                  1.717
                                           0.086 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 72365
                            on 52200 degrees of freedom
## Residual deviance: 57555 on 52193 degrees of freedom
     (3048 observations deleted due to missingness)
## AIC: 57569
##
## Number of Fisher Scoring iterations: 4
```

Substitution patterns

R:S ratio across different substitution patterns at nucleotide level

```
df$mutation.signature <- paste(df$from.nt, df$to.nt, sep = ">")

df.3 <- ddply(df, .(mutation.signature, mutation.type), summarize, count = sum(total.clonotypes))

df.3 <- ddply(df.3, .(mutation.signature), transform, ratio = count / (sum(count) - count))

df.3 <- subset(df.3, mutation.type == "R")

ggplot(df.3, aes(x = mutation.signature, y = ratio, fill = mutation.signature)) +

geom_bar(stat="identity", color="black") +

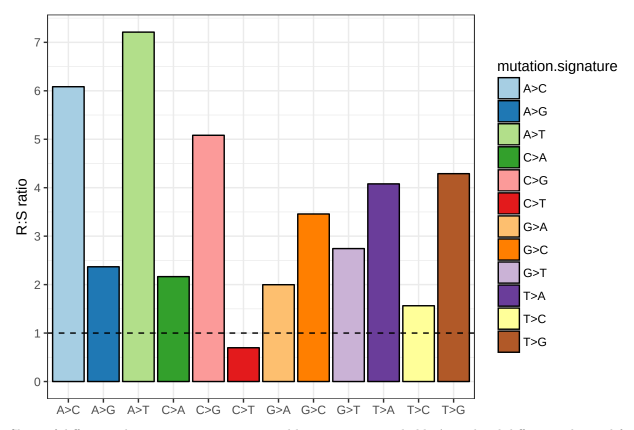
geom_hline(yintercept = 1, linetype="dashed") +

scale_fill_brewer(palette = "Paired") +

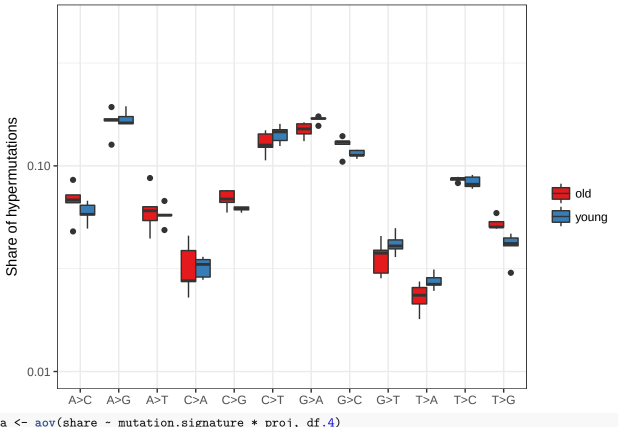
scale_y_continuous("R:S ratio", breaks=0:8) +

xlab("") +

theme_bw()</pre>
```



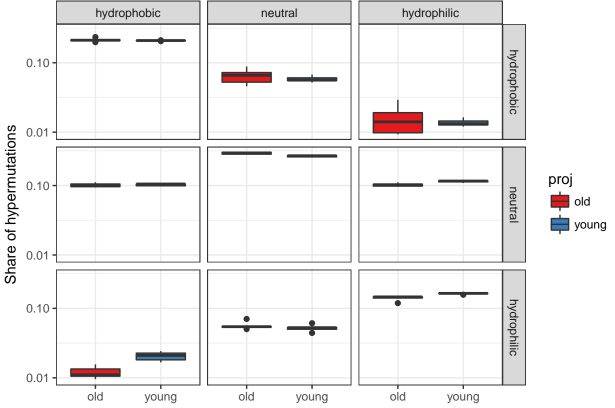
Share of different substitution patterns compared between young and old. Age-related difference observed for certain substitution patterns aross hypermutations.

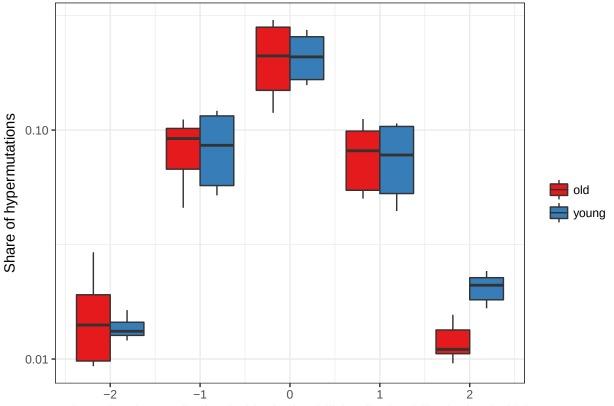


```
a <- aov(share ~ mutation.signature * proj, df.4)
summary(a)</pre>
```

```
## mutation.signature 11 0.2743 0.024934 234.697 <2e-16 ***
## proj 1 0.0000 0.000000 0.000 1.0000
## mutation.signature:proj 11 0.0026 0.000236 2.222 0.0191 *
## Residuals 96 0.0102 0.000106
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Hydropathy change patterns observed at amino acid level. More hydrophilic -> hydrophobic amino acid hypermutations in young compared to old.





Hydropathy change (hydrophobic>hydrophilic) .. (hydrophilic>hydrophobic)

```
a <- aov(share ~ hydrop.x : hydrop.y + hydrop.x : hydrop.y : proj, df.5)
summary(a)</pre>
```