# polyfid

Load required libraries, load and merge MAGERI results.

```
library(plyr); library(ggplot2); library(seqLogo); library(Biostrings); library(reshape2); library(gplo
## Warning: package 'ggplot2' was built under R version 3.2.4
## Loading required package: grid
## Loading required package: BiocGenerics
## Loading required package: parallel
## Attaching package: 'BiocGenerics'
## The following objects are masked from 'package:parallel':
##
##
       clusterApply, clusterApplyLB, clusterCall, clusterEvalQ,
##
       clusterExport, clusterMap, parApply, parCapply, parLapply,
##
       parLapplyLB, parRapply, parSapply, parSapplyLB
## The following objects are masked from 'package:stats':
##
##
       IQR, mad, xtabs
## The following objects are masked from 'package:base':
##
##
       anyDuplicated, append, as.data.frame, as.vector, cbind,
##
       colnames, do.call, duplicated, eval, evalq, Filter, Find, get,
##
       grep, grepl, intersect, is.unsorted, lapply, lengths, Map,
##
       mapply, match, mget, order, paste, pmax, pmax.int, pmin,
       pmin.int, Position, rank, rbind, Reduce, rownames, sapply,
##
##
       setdiff, sort, table, tapply, union, unique, unlist, unsplit
## Loading required package: S4Vectors
## Loading required package: stats4
##
## Attaching package: 'S4Vectors'
## The following object is masked from 'package:plyr':
##
##
       rename
## Loading required package: IRanges
```

```
## The following object is masked from 'package:plyr':
##
##
       desc
## Loading required package: XVector
##
## Attaching package: 'XVector'
## The following object is masked from 'package:plyr':
##
##
       compact
## Warning: package 'gplots' was built under R version 3.2.4
##
## Attaching package: 'gplots'
## The following object is masked from 'package: IRanges':
##
##
       space
## The following object is masked from 'package:stats':
##
       lowess
df.meta <- read.table("metadata.txt", header=T, sep = "\t")</pre>
df <- data.frame()</pre>
for (project in levels(df.meta$project)) {
  for (sample in levels(df.meta$sample)) {
    fname <- paste(project, sample, "variant.caller.txt", sep =".")</pre>
    if (file.exists(fname)) {
      df.1 <- read.table(fname, header=T, sep="\t")</pre>
      df.1$project <- project</pre>
      df.1$sample <- sample</pre>
      df <- rbind(df, df.1)
    }
  }
}
df$project <- as.factor(df$project)</pre>
df$sample <- as.factor(df$sample)</pre>
df <- merge(df, df.meta, all.x=T, all.y=F)</pre>
template <- "TGGGATCCATTATCGGCGGCGAATTTACCACCATTGAAAACCAGCCGTGGTTTGCGGCGATTTATCGTCGTCATCGTGGCGGCAGCGTGA
```

##

## Attaching package: 'IRanges'

#### Error rates

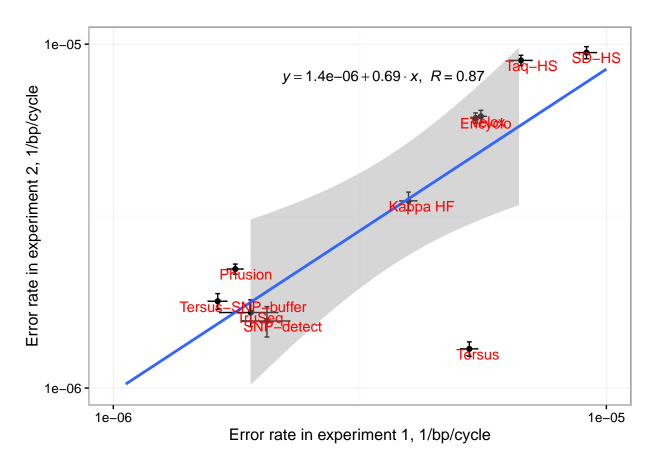
Overall error rates

```
df.er <- ddply(df, .(project, name), summarize,</pre>
               mismatches = sum(count.major), umi.count = round(mean(coverage)),
               err.rate = mismatches / umi.count / nchar(template) / mean(cycles),
               delta = sqrt(mismatches / umi.count * (1 - mismatches / umi.count) / umi.count) / nchar(
               err.lb = err.rate - 1.96 * delta, err.ub = err.rate + 1.96 * delta)
print(df.er)
       project
                             name mismatches umi.count
                                                            err.rate
## 1
      polerr73
                          Encyclo
                                        7759
                                                558336 5.428379e-06
## 2
     polerr73
                         Kappa HF
                                                240548 3.968802e-06
                                        2444
## 3
                          Phusion
      polerr73
                                        2902
                                                512908 1.768105e-06
## 4
      polerr73
                            SD-HS
                                        1696
                                                 72692 9.113795e-06
## 5
                                                 57755 2.049336e-06
      polerr73
                      SNP-detect
                                         303
## 6
                                                 78646 6.710251e-06
     polerr73
                          Taq-HS
                                        1351
## 7
      polerr73
                                        2430
                                                180029 5.272588e-06
                          Tersus
      polerr73 Tersus-SNP-buffer
                                        2035
                                                488168 1.628378e-06
## 9
                                                 43380 1.899997e-06
      polerr73
                          TruSeq
                                         211
## 10 polerr73
                           Velox
                                        6089
                                                427259 5.566918e-06
## 11 polerr82
                         Encyclo
                                        3127
                                                200300 6.098274e-06
## 12 polerr82
                        Kappa HF
                                        1040
                                                116021 3.501521e-06
## 13 polerr82
                          Phusion
                                        3349
                                                471735 2.218539e-06
## 14 polerr82
                            SD-HS
                                        2294
                                                 94810 9.451469e-06
## 15 polerr82
                      SNP-detect
                                         372
                                                 92765 1.566458e-06
                                                133952 8.970097e-06
## 16 polerr82
                          Taq-HS
                                        3076
## 17 polerr82
                           Tersus
                                        1655
                                                497406 1.299712e-06
## 18 polerr82 Tersus-SNP-buffer
                                        1391
                                                303834 1.788343e-06
                                         483
                                                113803 1.657881e-06
## 19 polerr82
                          TruSeq
## 20 polerr82
                            Velox
                                        2234
                                                141485 6.167836e-06
##
             delta
                          err.1b
                                       err.ub
## 1 6.119677e-08 5.308433e-06 5.548325e-06
      7.987141e-08 3.812255e-06 4.125350e-06
     3.272858e-08 1.703957e-06 1.832253e-06
     2.187056e-07 8.685132e-06 9.542458e-06
     1.174220e-07 1.819188e-06 2.279483e-06
## 6
     1.809874e-07 6.355515e-06 7.064986e-06
     1.062355e-07 5.064366e-06 5.480810e-06
     3.602183e-08 1.557775e-06 1.698980e-06
      1.304827e-07 1.644251e-06 2.155743e-06
## 10 7.083125e-08 5.428088e-06 5.705747e-06
## 11 1.081998e-07 5.886203e-06 6.310346e-06
## 12 1.080898e-07 3.289665e-06 3.713377e-06
## 13 3.819992e-08 2.143667e-06 2.293411e-06
## 14 1.949324e-07 9.069401e-06 9.833536e-06
## 15 8.105407e-08 1.407592e-06 1.725324e-06
## 16 1.598672e-07 8.656758e-06 9.283437e-06
## 17 3.189512e-08 1.237197e-06 1.362226e-06
```

## 18 4.783996e-08 1.694577e-06 1.882109e-06 ## 19 7.527596e-08 1.510340e-06 1.805422e-06 ## 20 1.294599e-07 5.914094e-06 6.421577e-06 Error rate consistency

```
df.er.cast <- dcast(df.er, name ~ project,</pre>
                    value.var = "err.rate")
df.er.cast2 <- dcast(df.er, name ~ project,</pre>
                    value.var = "delta")
df.er.cast <- merge(df.er.cast, df.er.cast2, by = "name")</pre>
m <- lm(polerr73.x ~ polerr82.x, df.er.cast);</pre>
eq <- substitute(italic(y) == a + b \%.% italic(x)*","~~italic(R)~"="~r,
     list(a = format(coef(m)[1], digits = 2),
          b = format(coef(m)[2], digits = 2),
          r = format(sqrt(summary(m)$r.squared), digits = 2)))
lbl<-as.character(as.expression(eq))</pre>
ggplot(df.er.cast, aes(x=polerr73.x, y=polerr82.x)) + geom_point() +
 geom errorbarh(aes(xmax=polerr73.x+1.96*polerr73.y,xmin=polerr73.x-1.96*polerr73.y)) +
  geom_errorbar(aes(ymax=polerr82.x+1.96*polerr82.y,ymin=polerr82.x-1.96*polerr82.y)) +
  geom_smooth(method = "lm", fullrange = T) +
  geom_text(aes(label=name), vjust=1, hjust = .3, color="red") +
  annotate("text", x = 2e-6, y = 8e-6, label = lbl, hjust=-0.1, parse = TRUE)+
  scale_x_log10(name="Error rate in experiment 1, 1/bp/cycle", limits=c(1e-6,1e-5)) +
  scale_y_log10(name="Error rate in experiment 2, 1/bp/cycle", limits=c(1e-6,1e-5)) +
 theme bw()
```

## Warning: Removed 2 rows containing missing values (geom\_smooth).



Combine error rates from a pair of experiments.

```
df.coverage.summary <- ddply(df, .(name, project), summarize, coverage = mean(coverage))
df.coverage.summary <- ddply(df.coverage.summary, .(name), summarize, coverage = sum(coverage))
df <- ddply(df, .(name, mutation), summarize, count.major = sum(count.major))
df <- merge(df, df.coverage.summary, by = "name")</pre>
```

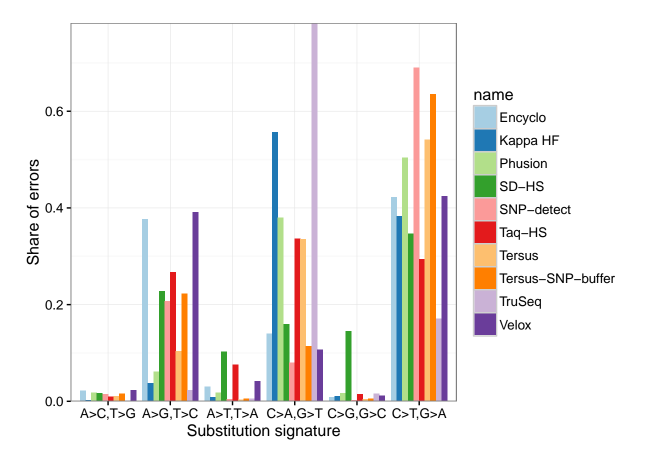
# Error substitution patterns

Parse mutation signatures (needed for further analysis)

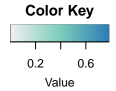
```
df$mut.split <- sapply(df$mutation, function(x) strsplit(as.character(x),"[S:>]"))
df$mutation.pos <- as.integer(sapply(df$mut.split, function(x) x[2]))
df$mutation.from <- sapply(df$mut.split, function(x) x[3])
df$mutation.to <- sapply(df$mut.split, function(x) x[4])
df$mut.split <- NULL</pre>
```

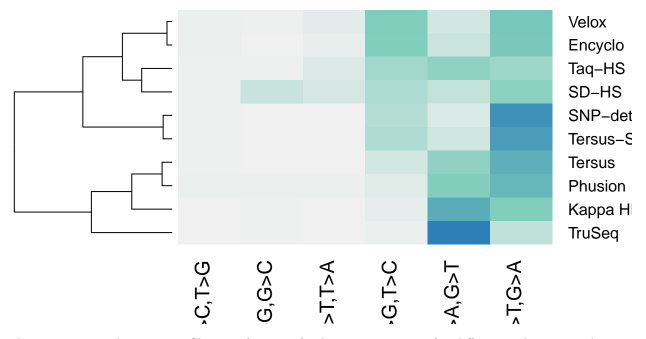
Substitution signature preferences

```
df$mutation.signature <- paste(df$mutation.from, df$mutation.to, sep =">")
sign.rep <- data.frame(mutation.signature = c("A>C","A>G","A>T","C>A","C>G","C>T","G>A","G>C","G>T","T>
```



```
## 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"
```





This goes to supplementary. Clear preference of substitution patters for different polymerases, but no position-related trend.

```
ggplot(df, aes(x = mutation.pos, weight = count.major / coverage, fill = mutation.from)) +
  geom_histogram(bins = nchar(template)) + scale_fill_brewer("Substituted base", palette = "Set1") +
  xlab("Position on template") + ylab("Error rate") +
  facet_wrap(~name) + theme_bw()#, scales = "free_y") + theme_bw()
```



```
a <- aov(count.major / coverage ~ mutation.from * name + mutation.pos, df)
summary(a)
                       Df
                            Sum Sq
                                     Mean Sq F value
                                                     Pr(>F)
                       3 5.840e-07 1.946e-07 44.781 < 2e-16 ***
## mutation.from
## name
                       9 1.064e-06 1.182e-07 27.207 < 2e-16 ***
## mutation.pos
                       1 1.000e-08 1.036e-08
                                              2.384
                                                        0.123
## mutation.from:name
                      27 5.010e-07 1.857e-08
                                              4.273 1.72e-12 ***
## Residuals
                     2121 9.217e-06 4.350e-09
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

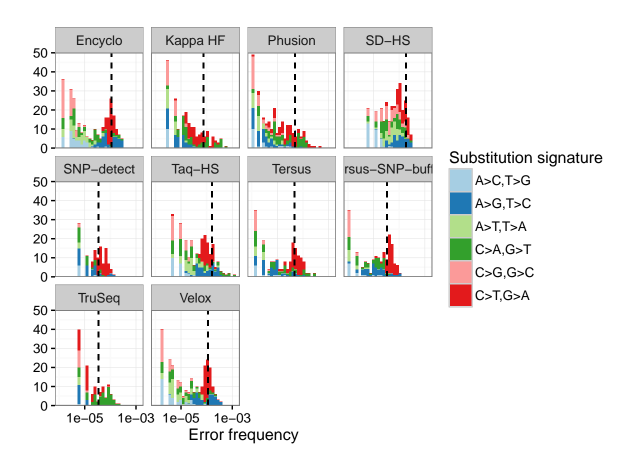
## Error hotspot context pattern

```
df.mean.err <- ddply(df, .(name), summarize, mean.err.rate = sum(count.major) / mean(coverage) / nchar()

df <- merge(df, df.mean.err, all.x=T, all.y=F)

ggplot(df) +
    geom_histogram(aes(x = count.major / coverage, fill = mutation.signature.rep)) +
    geom_linerange(aes(x = mean.err.rate, ymin = 0, ymax=50), linetype = "dashed", color="black") +
    scale_fill_brewer("Substitution signature", palette = "Paired") +
    scale_x_log10("Error frequency") +
    scale_y_continuous("", expand=c(0,0)) + facet_wrap(~name) + theme_bw()</pre>
```

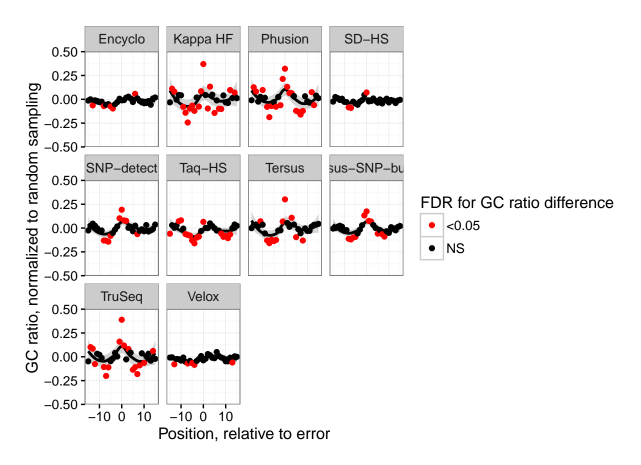
## `stat\_bin()` using `bins = 30`. Pick better value with `binwidth`.



#facet\_wrap(~name, scales = "free\_y") + theme\_bw()

## GC context profile

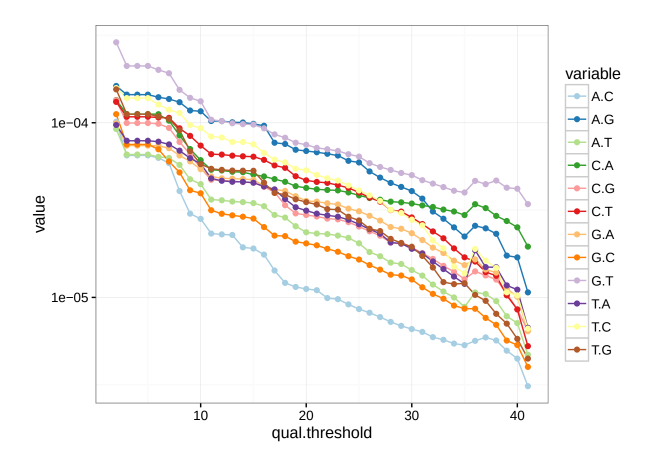
```
write.table(df.context, file = "context.txt", quote = F, sep = "\t", row.names = F)
system("groovy ProcessContext.groovy")
df.context.profile <- read.table("context.proc.txt", header=T, sep="\t")</pre>
df.context.normalized <- merge(subset(df.context.profile, name != "random"),</pre>
                                subset(df.context.profile, name == "random"),
                                by = "pos")
df.context.normalized <- ddply(df.context.normalized, .(pos, name.x), transform,</pre>
                                pval = prop.test(x = value.y, n = sum.y, p = value.x / sum.x,
                                                 alternative = "two.sided", correct = F)[[3]])
df.context.normalized$pval <- p.adjust(df.context.normalized$pval)</pre>
ggplot(df.context.normalized, aes(x = pos - window.size,
                                   y = value.x / sum.x - value.y / sum.y)) +
  geom_smooth(colour="black") + geom_point(aes(color = factor(ifelse(pval < 0.05, "<0.05", "NS")))) +</pre>
  facet_wrap(~name.x) + scale_colour_manual("FDR for GC ratio difference", values=c("red", "black")) +
  scale_y_continuous("GC ratio, normalized to random sampling", limits = c(-0.5, 0.5), expand=c(0,0)) +
  theme_bw()
```



```
df.q <- read.table("mmqc.txt", header=T,sep="\t")
library(reshape2)</pre>
```

```
df.q <- melt(df.q, id.vars = "qual.threshold")

ggplot(df.q, aes(x=qual.threshold, color=variable, y=value)) +
  geom_line() + geom_point() + scale_y_log10() +
  scale_color_brewer(palette = "Paired") + theme_bw()</pre>
```



```
df.q$subset <- ifelse(df.q$qual.threshold >= 35, "high-quality", "low-quality")
df.q$signature <- ifelse(df.q$variable %in% c("G.T", "C.A"), "TrueSeq", "other")

#ggplot(df.q, aes(x=subset, group=interaction(subset, signature), fill=signature, y = value)) +
# geom_boxplot()</pre>
```