A basic error model for UMI-tagged data

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Control experiment

Some auxiliary functions

```
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
library(ggplot2)
library(RColorBrewer)
library(stringr)
library(nloptr)
library(scales)
select <- dplyr::select
mtypes <- data.frame(mutation.fromto = c("A>C","A>G","A>T","C>A","C>G","C>T","G>A",
                                                "G>C", "G>T", "T>A", "T>C", "T>G"),
                        \label{eq:mutation.type} = c("A>C,T>G","A>G,T>C","A>T,T>A","C>A,G>T",
                                                    "C>G,G>C","C>T,G>A","C>T,G>A","C>G,G>C",
                                                    "C>A,G>T","A>T,T>A","A>G,T>C","A>C,T>G"))
read_variant_table <- function(file_name, pos_filter = function(x) T,</pre>
                                count_filter = function(x) T) {
  .df <- read.table(file_name, header=T, sep="\t", stringsAsFactors = F)</pre>
  .df <- subset(.df, !grepl("^[DI]", mutation) & coverage > 100 & freq < 0.45)
  .df$mutation.fromto <- unlist(lapply(str_split(.df$mutation, ":"), function(x) x[2]))
  .df$mutation.pos <- as.integer(unlist(lapply(str_split(.df$mutation, ":"),
                                                 function(x) str_sub(x[1], 2, nchar(x[1]))))
  .df <- merge(.df, mtypes)</pre>
  .df %>%
    filter(pos_filter(mutation.pos) & count_filter(count)) %>%
    select(mutation.pos, mutation.type, count, coverage)
}
```

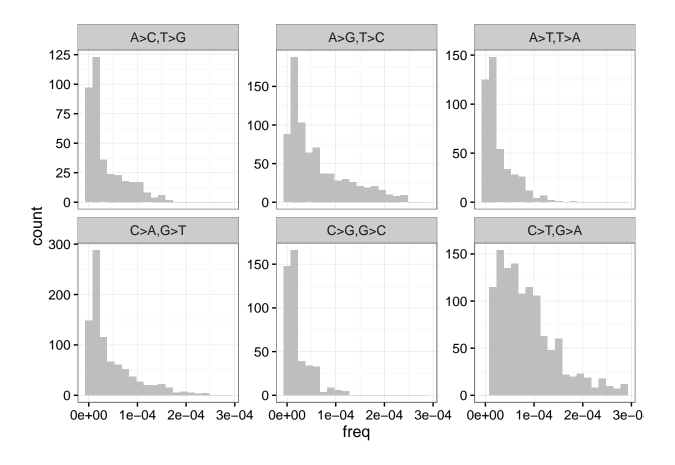
Load data from 2 independent experiments with library preparation performed using 10 different PCR assays: a known synthetic template sequence that was UMI-tagged and amplified using a set of different polymerases. At this stage a single event is a combination of substituted nucleotide, its substitution, position in template, sample (polymerase) and project (replica). Samples are obtained by grouping all events by substitution type which is one of 6 from and to nucleotide combinations that can be observed when not knowing the exact strand at which a given error has happened. Note that the latter is done because in most practical applications this information is hard to obtain.

```
df <- data.frame()</pre>
samples <- c("encyclo", "kappa-hf-taq", "phusion", "sd-hs", "snp-detect",</pre>
                    "taq-hs", "tersus", "tersus-snp-buff", "truseq", "velox")
for (proj in c("73", "82")) {
  for (sample in samples) {
    file_name <- paste("data/25_8_polerr", proj, ".", sample,
                        ".variant.caller.txt", sep = "")
    # ignore positions with no errors
    # this includes primers/barcodes
    .df <- read_variant_table(file_name,</pre>
                                count filter=function(x) x>0)
    .df$proj <- proj
    .df$sample <- sample
    df <- rbind(df, .df)</pre>
  }
}
```

Plot error frequencies grouped by substitution type

```
df.1 <- df %>%
    select(mutation.type, count, coverage) %>%
    mutate(freq = count / coverage) %>%
    # winsorize data
    group_by(mutation.type) %>%
    mutate(q5 = quantile(freq, 0.05), q95 = quantile(freq, 0.95)) %>%
    filter(freq >= q5 & freq <= q95) %>%
    select(mutation.type, freq)

ggplot(df.1, aes(x=freq)) + geom_histogram(fill="grey", bins=20) +
    facet_wrap(~mutation.type, scales="free_y") +
    theme_bw()
```



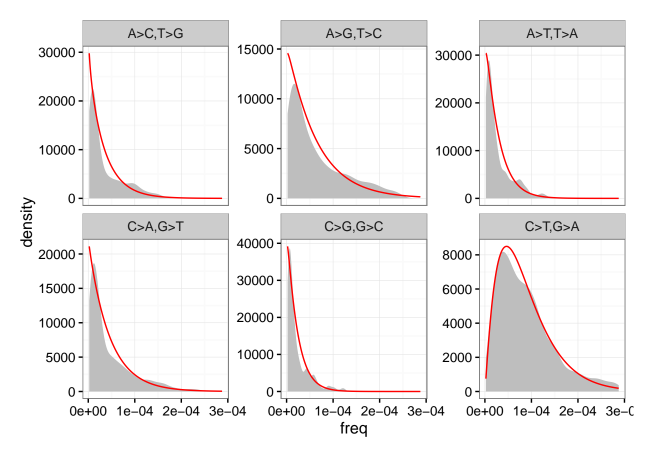
Estimating error rate distribution parameters for various substitution types

Fit frequency distributions with a Beta model.

```
betanll <- function(pars, data){</pre>
  alpha <- pars[[1]]</pre>
  beta <- pars[[2]]</pre>
  return (-sum(dbeta(data, shape1 = alpha, shape2 = beta, log = T)))
}
param_guess = function(data) {
  m = mean(data)
  v = sd(data) ^ 2
  c(m, (1 - m)) * (m * (1 - m) / v - 1)
fit_beta <- function(data) {</pre>
  list(nloptr(x0 = param_guess(data), eval_f = betanll,
              1b = c(0,0), data = data,
              opts = list(algorithm = "NLOPT_LN_SBPLX",
                           maxeval = 1e5))$solution)
}
fit.params <- df.1 %>%
  group_by(mutation.type) %>%
  summarize(fit = fit_beta(freq))
```

```
fit.params$alpha <- unlist(lapply(fit.params$fit, function(x) x[1]))
fit.params$beta <- unlist(lapply(fit.params$fit, function(x) x[2]))
fit.params$fit <- NULL</pre>
```

Checking goodness of fit. Note that poor fit in the vicinity of 0 can be attributed to sampling effect under finite coverage.



The probability of error p_T for each substitution type group T is fitted with a Beta distribution

```
p_T \sim Beta(\alpha_T, \beta_T)
```

with parameters

```
print(fit.params)
```

```
## # A tibble: 6 × 3
##
    mutation.type
                       alpha
                                 beta
##
            <fctr>
                       <dbl>
                                 <dbl>
## 1
           A>C,T>G 0.9278882 26823.24
## 2
           A>G,T>C 1.0295689 16394.12
## 3
           A>T,T>A 1.0838020 38974.43
## 4
           C>A,G>T 0.9957058 21862.23
## 5
           C>G,G>C 1.0584582 48029.97
## 6
           C>T,G>A 2.1314829 24344.82
```

Thus the count of an error of a certain type n_T at the coverage N is distributed as

```
n_T \sim BetaBinom(N, \alpha_T, \beta_T)
```

Benchmark using an independent control sample

Load control data from UMI-tagged amplicon sequencing of control donor DNA

Observed (black) and fitted (red) distribution of error ocurrences by error count

```
library(TailRank)
```

Loading required package: oompaBase

```
alpha <- fit.params$alpha
names(alpha) <- fit.params$mutation.type
beta <- fit.params$beta
names(beta) <- names(alpha)

compute_p <- function(count, coverage, mutation.type) {
   mapply(function(k, n, a, b) dbb(k, n, a, b),
        count, coverage, alpha[mutation.type], beta[mutation.type])
}</pre>
```

```
df.control.1 <- df.control %>%
  group_by(mutation.type) %>%
  mutate(total = n()) %>%
  group_by(count, mutation.type) %>%
  mutate(coverage.med = as.numeric(median(coverage))) %>%
  group_by(count, mutation.type, coverage.med, total) %>%
  summarise(freq = n()) %>%
  mutate(freq.fit = total * dbb(count, coverage.med,
                                alpha[mutation.type], beta[mutation.type]))
fig2 <- ggplot(df.control.1, aes(x = count)) +
  geom_line(aes(y=freq), linetype="dashed") +
  geom_point(aes(y=freq)) +
  geom_line(aes(y=round(freq.fit)), color="red") +
  scale_x_continuous("Erroneous variant count", limits=c(0,6), breaks = 0:6) +
  ylab("Occurrences") +
  facet_wrap(~mutation.type) +
  theme_bw()
save(file = "../figures/fig2.Rda", fig2)
fig2
```

Warning: Removed 5 rows containing missing values (geom_point).

