

Alakazam: Using sequencing quality scores

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The `alakazam` package includes a set of functions to inspect the sequencing quality.

Example data

Load example data:

```
library(alakazam)
library(dplyr)
library(aирr)

db <- read_rearrangement(system.file("extdata", "example_quality.tsv", package="alakazam"))
fastq_file <- system.file("extdata", "example_quality.fastq", package="alakazam")
```

Load quality scores

This method allows to add the quality scores to the repertoire `data.frame` as strings.

```
original_cols <- colnames(db)
db <- readFastqDb(db, fastq_file, style="both", quality_sequence=TRUE)
new_cols <- setdiff(colnames(db), original_cols)
db[,new_cols] %>% head()

## # A tibble: 1 x 4
##   quality_num           quality quality_alignment_num quality_alignment
##   <chr>                 <chr>      <chr>                <chr>
## 1 90,90,90,90,90,90,90,90~ {{~ 90,90,90,90,90,90~ {{~ {{~ {{~ {{~
```

The function `readFastq` takes as main inputs a repertoire `data.frame` (`db`) and a path to the corresponding `.fastq` file (`fastq_file`). The sequencing quality scores will be merged into the `data.frame` by `sequence_id`. The newly added columns are: `quality_num`, `quality`, `quality_alignment_num`, `quality_alignment`. The other fields, contain the ASCII quality scores in the form of a vector, where values are comma separated, and - or . positions have value " " (blank).

After loading the quality scores with `readFastqDb`, `getPositionQuality` can be used to generate a `data.frame` of sequencing quality values per position.

```
quality <- getPositionQuality(db, sequence_id="sequence_id",
                                sequence="sequence_alignment",
                                quality_num="quality_alignment_num")

## Warning in FUN(X[[i]], ...): NAs introduced by coercion

head(quality)

##   position quality_alignment_num      sequence_id nt
## 1         1                     90 CGCTTTTCGGATTGGAA C
## 2         2                     90 CGCTTTTCGGATTGGAA A
## 3         3                     90 CGCTTTTCGGATTGGAA G
## 4         4                     90 CGCTTTTCGGATTGGAA C
## 5         5                     90 CGCTTTTCGGATTGGAA T
## 6         6                     90 CGCTTTTCGGATTGGAA G

min_pos <- min(quality$position)
max_pos <- max(quality$position)

ggplot(quality, aes(x=position,
                     y=quality_alignment_num,
                     color=nt)) +
  geom_point() +
  coord_cartesian(xlim=c(110,120)) +
  xlab("IMGT position") +
  ylab("Sequencing quality") +
  scale_fill_gradient(low = "light blue", high = "dark red") +
  scale_x_continuous(breaks=c(min_pos:max_pos)) +
  alakazam::baseTheme()

## Warning: Removed 27 rows containing missing values or values outside the scale range
## (`geom_point()`).
```

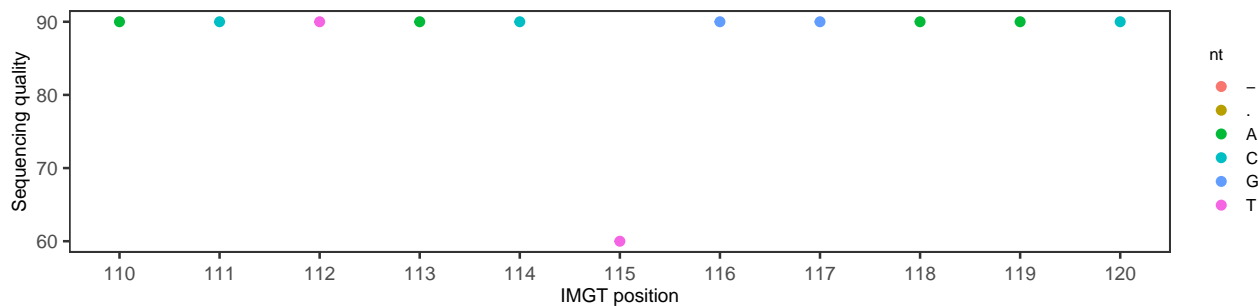


Figure 1: Sequence quality per IMGT position for one sequence.

You can add use the quality `data.frame` to complement analysis performed with other tools from the Immcantation framework. For example, you could inspect the sequencing quality of novel polymorphisms identified with `tigger`, or the sequencing quality in mutated/unmutated regions.

Mask low quality positions

Use `maskPositionsByQuality` to mask low quality positions. Positions with a sequencing quality < `min_quality` will be replaced with an 'N'. A message will show the number of sequences in db that had at least one position masked.

```
db <- maskPositionsByQuality(db, min_quality=70,  
                               sequence="sequence_alignment",  
                               quality="quality_alignment_num")  
  
## Number of masked sequences: 1
```