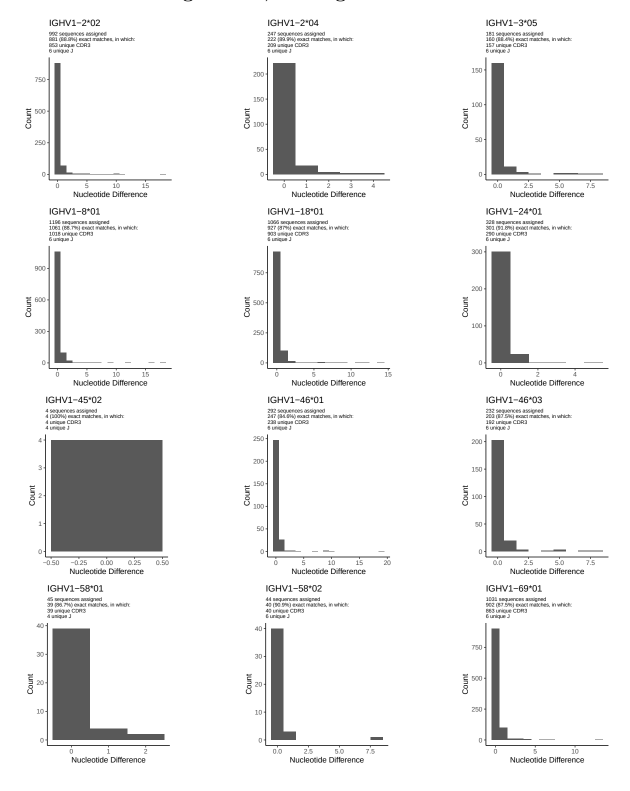
# OGRDBstats Report

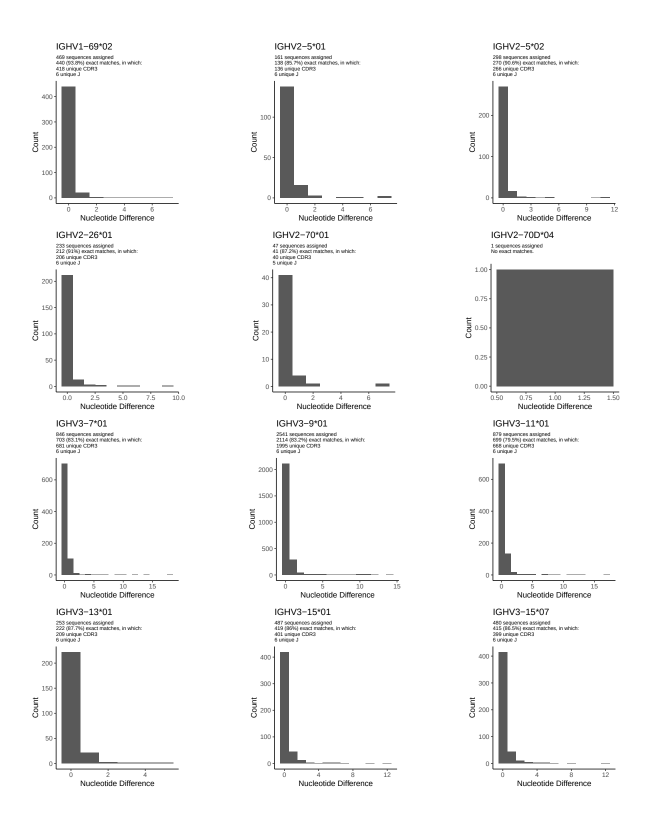
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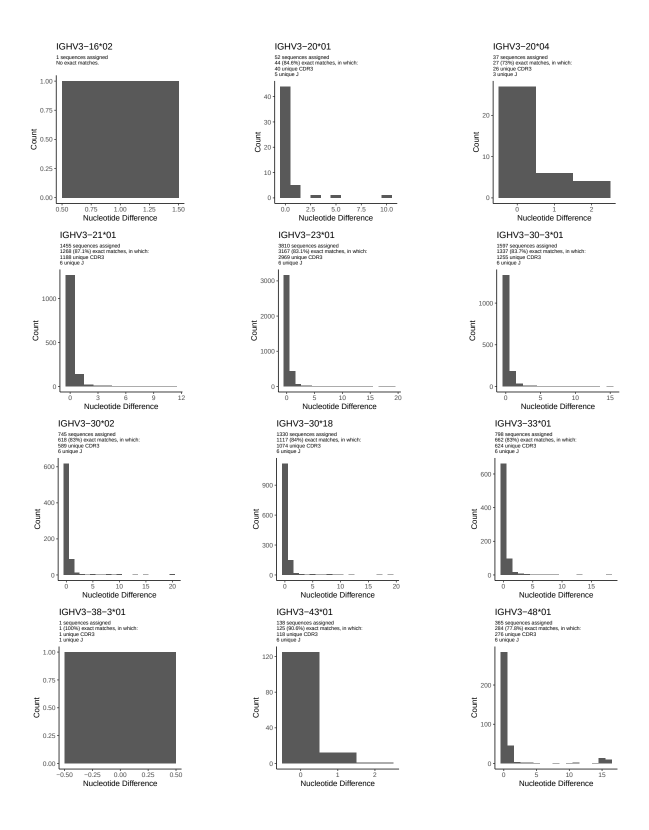
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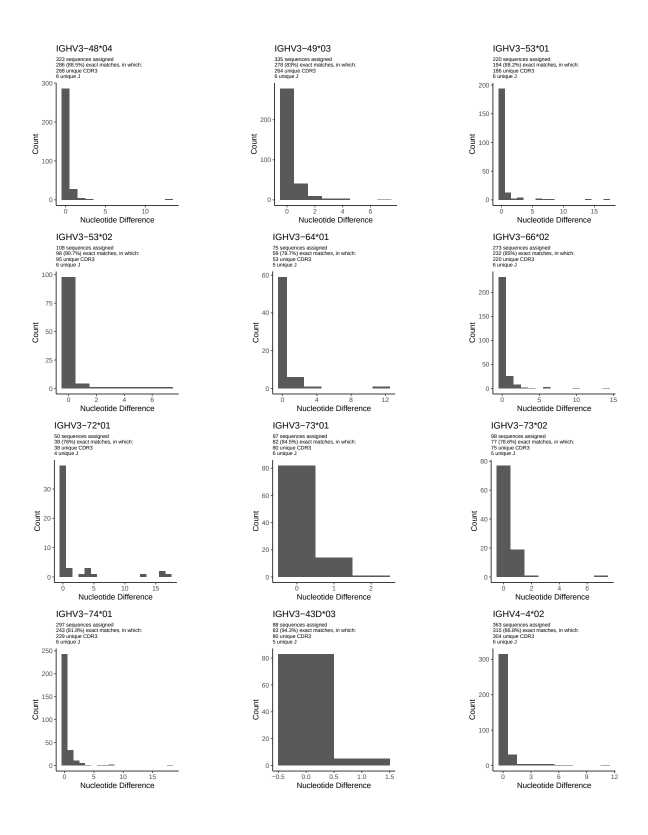
- 1 Novel sequence analysis
- 1.1 CDR3 length distribution, in assignments to novel alleles

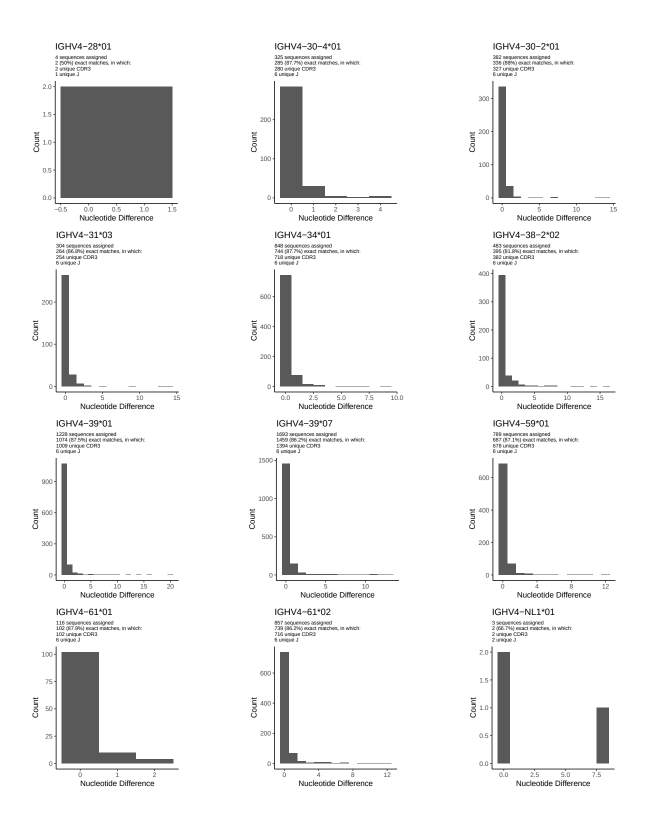
#### 2 Variation from germline, in assignments to each allele

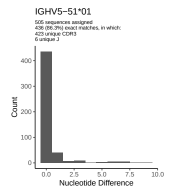


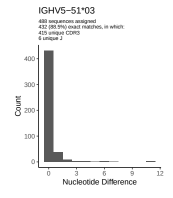


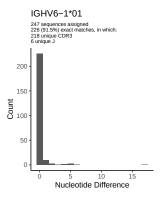


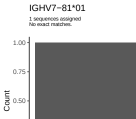












0.75 1.00 1.25 Nucleotide Difference

0.25

0.50

## 3 Allele usage in potential haplotype anchor genes



4 Haplotype plots

#### 5 Configuration settings

```
## Repertoire file: /misc/work/jenkins/PRJEB26509/S25/S25/25/e8/4c27e55e66cf6cc60899fbdf92121f/25_Fir
##

## Germline reference file: /misc/work/jenkins/PRJEB26509/S25/S25/25/d0/9a41a2fad29ae627c4a85bd51b86
##

## Novel allele file:
##

## Species: Homosapiens
##

## Chain: IGHV
##

## Segment: V
##

## Warning no inferred sequences found.
```