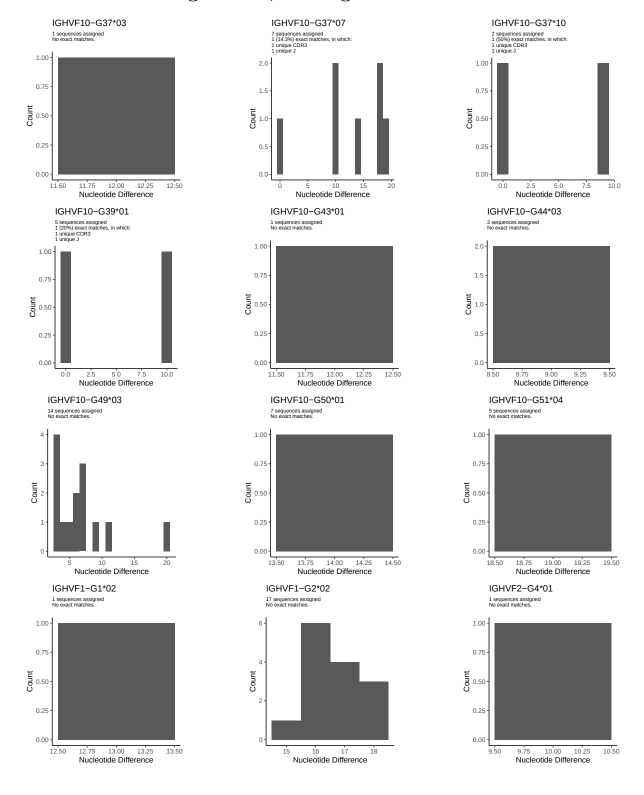
OGRDBstats Report

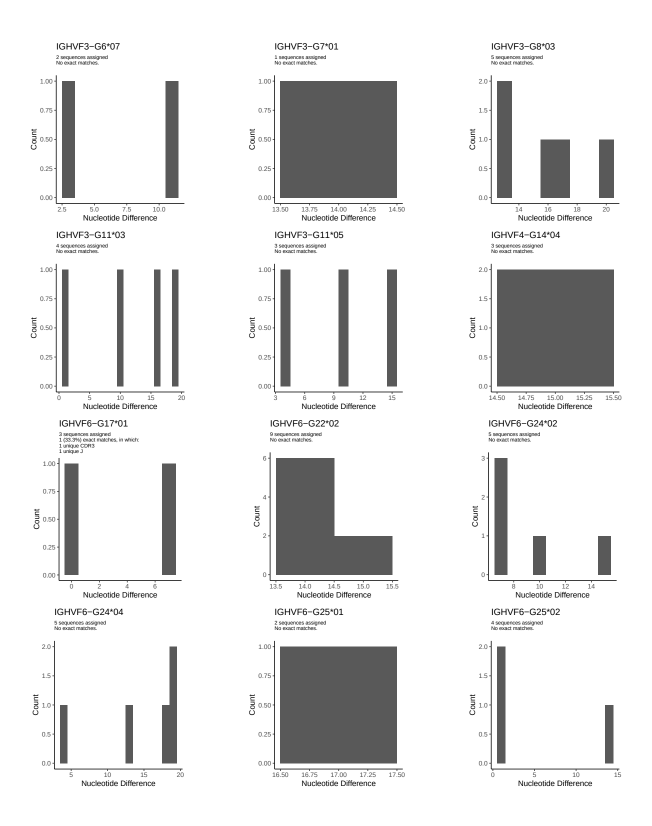
Contents

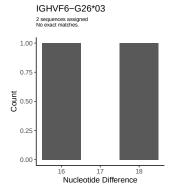
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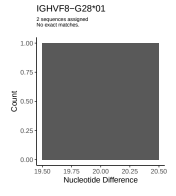
- 1 Novel sequence analysis
- 1.1 CDR3 length distribution, in assignments to novel alleles

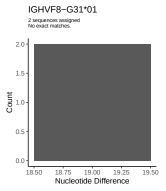
2 Variation from germline, in assignments to each allele

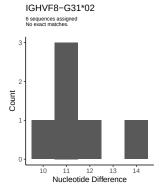




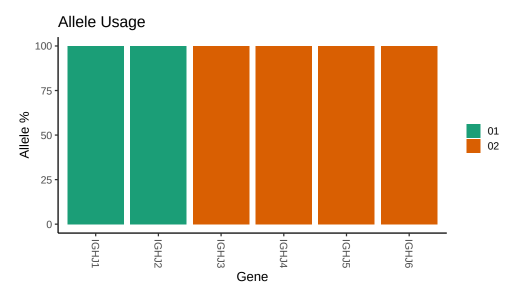








3 Allele usage in potential haplotype anchor genes



4 Haplotype plots

5 Configuration settings

```
## Repertoire file: /work/jenkins/PRJNA248475/M4/M4_lesion/M4_lesion/8a/6c6e7dec5ea4918944a1f
## Germline reference file: /work/jenkins/PRJNA248475/M4/M4_lesion/M4_lesion/M4_lesion/8a/6c6e7dec5ea49
## Novel allele file: /work/jenkins/PRJNA248475/M4/M4_lesion/M4_lesion/M4_lesion/8a/6c6e7dec5ea4918944a
## Species: Homosapiens
## Chain: IGHV
## Segment: V
##
## Novel sequence(s) IGHVF3 G9 04_A319G are not listed in the genotype and will be ignored.
##
## No sequences found for IGHVF3 G9 04_A319G(check SEQUENCE_IMGT in input file)
```