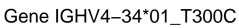
OGRDBstats Report

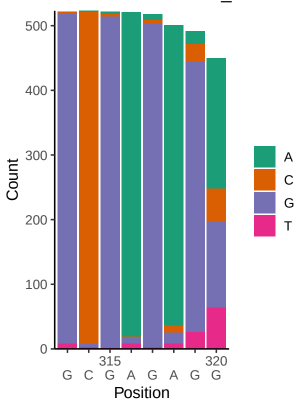
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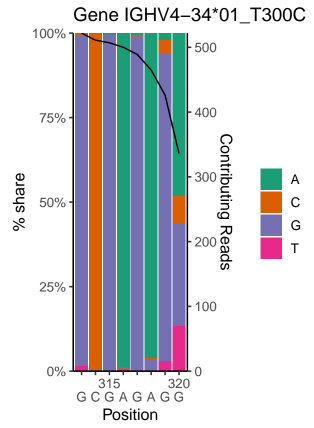
1 Novel sequence analysis

1.1 End-nucleotide composition

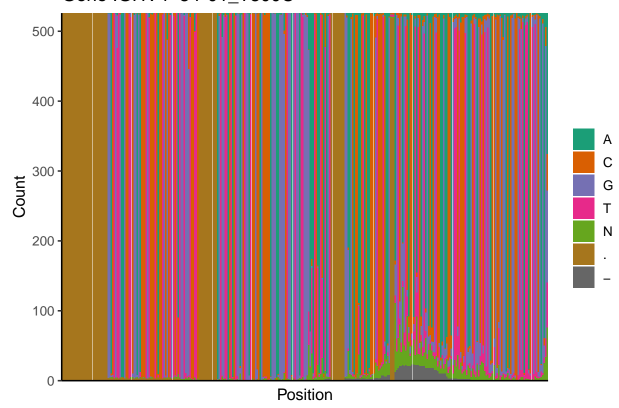




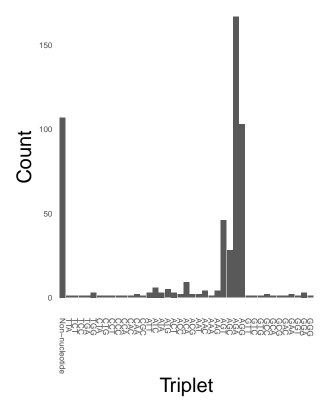
1.2 Per-nucleotide consensus where previous nucleotides match the consensus



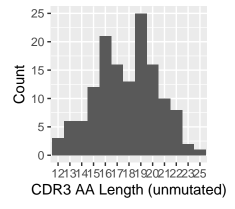
1.3 Whole-sequence composition of each assigned read Gene $IGHV4-34*01_T300C$



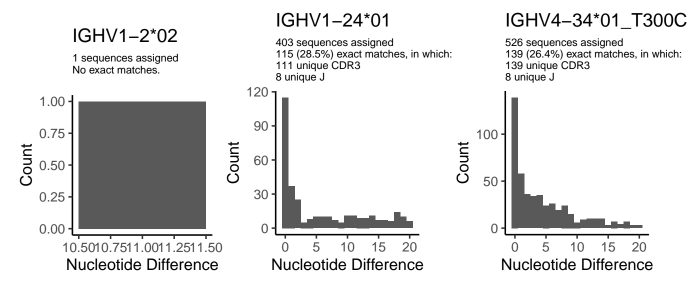
1.4 Final three nucleotides: frequency of each observed triplet IGHV4-34*01_T300C- Final 3 nucleotides as a triplet



1.5 CDR3 length distribution, in assignments to novel alleles $IGHV4-34*01_T300C$



2 Variation from germline, in assignments to each allele



3	Allele usage in potential haplotype anchor genes

4 Configuration settings