bcRep news

Julia Bischof Julia.Bischof@uksh.de 2016-02-25

Contents

Changes in version 1.3 (2016-02-25)
BUG FIXES
NEW FEATURES AND FUNCTIONS
Changes in version 1.3 (2016-02-15)
BUG FIXES
NEW FEATURES AND FUNCTIONS
Changes in version 1.2.2 (2015-10-28)
BUG FIXES
NEW FEATURES AND FUNCTIONS
Version 1.0 (2015-10-09)

Changes in version 1.3 (2016-02-25)

BUG FIXES

- * readIMGT(): eliminated errors
- * clones(): eliminated error message when no clone was found
- * geneUsage(): eliminated error for JH subgroup usage

NEW FEATURES AND FUNCTIONS

- * sequences.mutation.AA() added (proportions of amino acid mutations)
- * plotSequencesMutationAA() added
- * sequences.mutation.base() added (proportions of bases around mutations)
- * plotSequencesMutationBase() added

Changes in version 1.3 (2016-02-15)

BUG FIXES

- * clones.shared(): reduced computation time and memory; copy number of CDR3 sequences changed
- * sequences.mutation(): added sequence ID's

NEW FEATURES AND FUNCTIONS

- * added NEWS
- * sequences.distance() added (distance/dissimilarity measurements on sequence data)
- * geneUsage.distance() added (distance/dissimilarity measurements on gene usage data)
- * dist.PCoA() and plotDistPCoA() added (Principal coordinate analysis on distance data and visualization)

Changes in version 1.2.2 (2015-10-28)

BUG FIXES

- * plotClonesCopyNumber(): added parameter to plot with and without outliers (clone sizes > 75\% quantile)
- * geneUsage(): Combination of gene usage and functionality didn't work for alleles before
- * sequences.geneComb(): added parallel processing option

NEW FEATURES AND FUNCTIONS

- * added Vignette
- * combineIMGT(): combines IMGT output files from different folders
- * clones.giniIndex(): calculates Gini Index of clones
- * clones.filterSize(): filters clones for their size (number, percentage)
- * clones.filterFuntionality(): filters clones for their funtionality
- * clones.filterJunctionFrame(): filters clones for their junction frame usage
- * compare.aaDistribution(), plotCompareAADistribution(): compare amino acid distribution of different samples
- * compare.trueDiversity(), plotCompareTrueDiversity(): compare richness and diversity of different samples
- * compare.geneUsage(), plotCompareGeneUsage(): compare gene usage of different samples (subgroup, gene, allele)

Version 1.0 (2015-10-09)