Alakazam: Amino acid physicochemical property analysis

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The alakazam package includes a set of function to analyze the physicochemical properties of Ig and TCR amino acid sequences. Of particular interest is the analysis of CDR3 properties, which this vignette will demonstate. However, the same process can be applied to other regions simply by altering the sequence data column used.

Wu YC, et al. High-throughput immunoglobulin repertoire analysis distinguishes between human IgM memory and switched memory B-cell populations. Blood 116, 1070-8 (2010).

Wu YC, et al. The relationship between CD27 negative and positive B cell populations in human peripheral blood. Front Immunol 2, 1-12 (2011).

Example data

A small example Change-O database, ExampleDb, is included in the alakazam package.

```
# Load required packages
library(alakazam)
library(dplyr)

# Subset example data
data(ExampleDb)
db <- ExampleDb[ExampleDb$SAMPLE == "+7d", ]</pre>
```

Calculate the properties of amino acid sequences

Multiple amino acid physicochemical properties can be obtained with the function aminoAcidProperties. The available properties are:

- length: total amino acid count
- gravy: grand average of hydrophobicity

- bulkiness: average bulkiness
- polarity: average polarity
- aliphatic: normalized aliphatic index
- charge: normalized net charge
- acidic: acidic side chain content
- basic: basic side chain residue content
- aromatic: aromatic side chain content

This example demonstrates how to calculate all of the available amino acid properties from DNA sequences found in JUNCTION column of the Change-O file previously loaded.

Translation of the DNA sequences to amino acid sequences is accomplished by specifying the nt=TRUE argument. To reduce the junction sequence to the CDR3 sequence we specify the argument trim=TRUE which will strip the first and last codon (the conserved residues) prior to analysis. The prefix CDR3 is added to the output column names using the label="CDR3" argument.

```
db_props <- aminoAcidProperties(db, seq="JUNCTION", nt=TRUE, trim=TRUE,</pre>
                                 label="CDR3")
# The full set of properties are calculated by default
dplyr::select(db_props[1:3, ], starts_with("CDR3"))
##
        CDR3_AA_LENGTH CDR3_AA_GRAVY CDR3_AA_BULK CDR3_AA_ALIPHATIC
## 1001
                    29
                            0.1724138
                                          14.12345
                                                            0.8034483
## 1002
                    29
                           -0.3482759
                                          14.69034
                                                            0.6724138
## 1003
                    26
                           -0.9884615
                                          13.96154
                                                            0.5653846
##
        CDR3 AA POLARITY CDR3 AA CHARGE CDR3 AA BASIC CDR3 AA ACIDIC
                              0.03902939
## 1001
                8.168966
                                             0.1034483
                                                            0.06896552
## 1002
                8.255172
                              2.21407038
                                             0.2068966
                                                            0.06896552
## 1003
                8.873077
                              1.11045407
                                             0.2307692
                                                            0.15384615
##
        CDR3 AA AROMATIC
## 1001
              0.06896552
## 1002
              0.27586207
## 1003
              0.19230769
# Define a ggplot theme for all plots
tmp_theme <- theme_bw() + theme(legend.position="bottom")</pre>
# Generate plots for a four of the properties
g1 <- ggplot(db_props, aes(x=ISOTYPE, y=CDR3_AA_LENGTH)) + tmp_theme +
    ggtitle("CDR3 length") +
    xlab("Isotype") + ylab("Amino acids") +
    scale_fill_manual(name="Isotype", values=IG_COLORS) +
    geom_boxplot(aes(fill=ISOTYPE))
g2 <- ggplot(db_props, aes(x=ISOTYPE, y=CDR3_AA_GRAVY)) + tmp_theme +
    ggtitle("CDR3 hydrophobicity") +
    xlab("Isotype") + ylab("GRAVY") +
```

```
scale_fill_manual(name="Isotype", values=IG_COLORS) +
    geom_boxplot(aes(fill=ISOTYPE))
g3 <- ggplot(db_props, aes(x=ISOTYPE, y=CDR3_AA_BASIC)) + tmp_theme +
    ggtitle("CDR3 basic residues") +
    xlab("Isotype") + ylab("Basic residues") +
    scale y continuous(labels=scales::percent) +
    scale fill manual(name="Isotype", values=IG COLORS) +
    geom_boxplot(aes(fill=ISOTYPE))
g4 <- ggplot(db_props, aes(x=ISOTYPE, y=CDR3_AA_ACIDIC)) + tmp_theme +
    ggtitle("CDR3 acidic residues") +
    xlab("Isotype") + ylab("Acidic residues") +
    scale_y_continuous(labels=scales::percent) +
    scale_fill_manual(name="Isotype", values=IG_COLORS) +
    geom_boxplot(aes(fill=ISOTYPE))
# Plot in a 2x2 grid
gridPlot(g1, g2, g3, g4, ncol=2)
                  CDR3 length
                                                           CDR3 basic residues
                                                30%
   30
                                             Basic residues
 Amino acids
   25
                                                20%
                                                10%
   10
                                                 0%
                                                                 IgD
                   IgD
                            lgG
                                     IgM
                                                                         IgG
          IgA
                                                        IgΑ
                                                                                  IgM
                                                                   Isotype
                     Isotype
         Isotype 📋 IgA 📙 IgD 🛑 IgG 🛑 IgM
                                                       Isotype 🖨 IgA 🖨 IgD 🖨 IgG 🖨 IgM
              CDR3 hydrophobicity
                                                           CDR3 acidic residues
    1
                                                30%
                                              Acidic residues
    0
 GRAVY
                                                20%
                                                10%
                                                 0%
   -2
                            IgG
                                     ΙgΜ
                                                                 IgD
                                                                                  ΙgΜ
          IgA
                   IgD
                                                        IgA
                                                                         IgG
                     Isotype
                                                                   Isotype
         Isotype 📋 IgA 📙 IgD 🚔 IgG 🚔 IgM
                                                      Isotype 📋 IgA 📙 IgD 🚞 IgG 텛 IgM
```

Obtaining properties individually

A subset of the properties may be calculated using the property argument of aminoAcidProperties. For example, calculations may be restricted to only the grand average of hydrophobicity (gravy) index and normalized net charge (charge) by specifying property=c("gravy", "charge").

Using user defined scales

Each property has a default scale setting, but users may specify alternate scales if they wish. The following example shows how to import and use the Kidera et al, 1985 hydrophobicity scale and the Murrary et al, 2006 pK values from the **seqinr** package instead of the defaults for calculating the GRAVY index and net charge.

```
# Load the relevant data objects from the seginr package
library(seqinr)
data(aaindex)
data(pK)
h <- aaindex[["KIDA850101"]]$I
p <- setNames(pK[["Murray"]], rownames(pK))</pre>
# Rename the hydrophobicity vector to use single-letter codes
names(h) <- translateStrings(names(h), ABBREV_AA)</pre>
db_props <- aminoAcidProperties(db, seq="JUNCTION", property=c("gravy", "charge"),</pre>
                                 nt=TRUE, trim=TRUE, label="CDR3",
                                 hydropathy=h, pK=p)
dplyr::select(db props[1:3, ], starts with("CDR3"))
        CDR3_AA_GRAVY CDR3_AA_CHARGE
##
## 1001
          -0.06551724
                           -0.0661116
## 1002
                            2.0664863
           0.10482759
## 1003
           0.13807692
                            1.0370349
```

Getting vectors of individual properties

The aminoAcidProperties function provides a convenient wrapper for calculating multiple properties at once from a data.frame. If a vector of a specific property is required this may be accomplished using one of the worker functions:

- gravy: grand average of hydrophobicity
- bulk: average bulkiness
- polar: average polarity
- aliphatic: aliphatic index
- charge: net charge
- countPatterns: counts the occurrence of patterns in amino acid sequences

The input to each function must be vector of amino acid sequences.

```
# Translate junction DNA sequences to amino acids and trim first and last codons
cdr3 <- translateDNA(db$JUNCTION[1:3], trim=TRUE)</pre>
# Grand average of hydrophobicity
gravy(cdr3)
## [1] 0.1724138 -0.3482759 -0.9884615
# Average bulkiness
bulk(cdr3)
## [1] 14.12345 14.69034 13.96154
# Average polarity
polar(cdr3)
## [1] 8.168966 8.255172 8.873077
# Normalized aliphatic index
aliphatic(cdr3)
## [1] 0.8034483 0.6724138 0.5653846
# Unnormalized aliphatic index
aliphatic(cdr3, normalize=FALSE)
## [1] 23.3 19.5 14.7
# Normalized net charge
charge(cdr3)
## [1] 0.03902939 2.21407038 1.11045407
# Unnormalized net charge
charge(cdr3, normalize=FALSE)
```

[1] 0.03902939 2.21407038 1.11045407 # Count of acidic amino acids # Takes a named list of regular expressions countPatterns(cdr3, c(ACIDIC="[DE]"), label="CDR3") ## CDR3_ACIDIC ## 1 0.06896552 ## 2 0.06896552 ## 3 0.15384615

Default scales

The following references were used for the default physicochemical scales:

- Aliphatic index:
 - Ikai AJ. Thermostability and aliphatic index of globular proteins. J Biochem 88, 1895-1898 (1980).
- Bulkiness scale:
 - Zimmerman JM, Eliezer N, Simha R. The characterization of amino acid sequences in proteins by statistical methods. J Theor Biol 21, 170-201 (1968).
- Hydrophobicity scale:
 - Kyte J, Doolittle RF. A simple method for displaying the hydropathic character of a protein. J Mol Biol 157, 105-32 (1982).
- pK values:
 - http://emboss.sourceforge.net/apps/cvs/emboss/apps/iep.html
- Polarity scale:
 - Grantham R. Amino acid difference formula to help explain protein evolution. Science 185, 862-864 (1974).