

Package ‘Ibex’

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Title Methods for BCR single-cell embedding

Version 1.3.0

Description Implementation of the Ibex algorithm for single-cell embedding based on BCR sequences. The package includes a standalone function to encode BCR sequence information by amino acid properties or sequence order using tensorflow-based autoencoder. In addition, the package interacts with SingleCellExperiment or Seurat data objects.

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BugReports <https://github.com/BorchLab/Ibex/issues>

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Ibex-package	<i>Ibex: Methods for BCR single-cell embedding</i>
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Description

Ibex implements methods for embedding B-cell receptor (BCR) sequences from single-cell assays into a continuous latent space. It supports amino-acid property-based and sequence-order encodings via a TensorFlow autoencoder, and interoperates with common single-cell containers such as **SingleCellExperiment** and **SeuratObject**.

Details

Key features

- Encode BCR sequence information using biochemical properties or raw sequence order (TensorFlow autoencoder).
- Interoperate with **SingleCellExperiment** and **SeuratObject** for downstream analysis and visualization.
- Utilities for loading pretrained models and managing dependencies in an isolated **basilisk** environment.

Getting started

```
browseVignettes("Ibex")
```

Models and caching Pretrained encoders can be retrieved with `aa.model.loader()`, which validates against internal metadata and caches downloaded artifacts; see the function help for cache location and behavior.

Python/TensorFlow note Ibex uses **basilisk** to provision an isolated Python environment at runtime; no manual setup is usually required.

Author(s)

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See Also

<https://github.com/BorchLab/Ibex>

<https://github.com/BorchLab/Ibex/issues>

combineExpandedBCR *combineBCR for CDR1/2/3 sequences*

Description

This function enhances BCR processing by incorporating additional sequence information from CDR1 and CDR2 regions before applying the BCR combination logic. The function depends on `scRepertoire::combineBCR()`.

Usage

```
combineExpandedBCR(
  input.data,
  samples = NULL,
  ID = NULL,
  call.related.clones = TRUE,
  threshold = 0.85,
  removeNA = FALSE,
  removeMulti = FALSE,
  filterMulti = TRUE,
  filterNonproductive = TRUE
)
```

Arguments

<code>input.data</code>	List of filtered contig annotations.
<code>samples</code>	Character vector. Labels of samples (required).
<code>ID</code>	Character vector. Additional sample labeling (optional).
<code>call.related.clones</code>	Logical. Whether to call related clones based on nucleotide sequence and V gene. Default is TRUE.
<code>threshold</code>	Numeric. Normalized edit distance for clone clustering. Default is 0.85.
<code>removeNA</code>	Logical. Whether to remove any chain without values. Default is FALSE.
<code>removeMulti</code>	Logical. Whether to remove barcodes with more than two chains. Default is FALSE.
<code>filterMulti</code>	Logical. Whether to select the highest-expressing light and heavy chains. Default is TRUE.
<code>filterNonproductive</code>	Logical. Whether to remove nonproductive chains. Default is TRUE.

Value

A list of consolidated BCR clones with expanded CDR sequences.

See Also

`scRepertoire::combineBCR()`

Examples

```
#' # Get Data
ibex_vdj <- get(data("ibex_vdj"))

combined.BCR <- combineExpandedBCR(list(ibex_vdj),
                                   samples = "Sample1",
                                   filterNonproductive = TRUE)
```

CoNGAfy

Reduce a Single-Cell Object to Representative Cells

Description

This function generates a single-cell object with a reduced representation of RNA expression by clone. The approach is inspired by the method introduced in **CoNGA**. Users can generate either a mean representation of features by clone or identify a representative cell using count-based minimal Euclidean distance. Please read and cite the original work by the authors of CoNGA.

Usage

```
CoNGAfy(
  input.data,
  method = "dist",
  features = NULL,
  assay = "RNA",
  meta.carry = c("CTaa", "CTgene")
)
```

Arguments

<code>input.data</code>	A single-cell dataset in Seurat or SingleCellExperiment format.
<code>method</code>	Character. Specifies the method to reduce the dataset: <ul style="list-style-type: none"> • "mean" - Computes the mean expression of selected features across cells in each clonotype. • "dist" - Uses PCA reduction to identify the cell with the minimal Euclidean distance within each clonotype group.
<code>features</code>	Character vector. Selected genes for the reduction. If NULL (default), all genes are used.
<code>assay</code>	Character. The name of the assay or assays to include in the output. Defaults to the active assay.
<code>meta.carry</code>	Character vector. Metadata variables to carry over from the input single-cell object to the output.

Value

A reduced single-cell object where each clonotype is represented by a single cell.

Examples

```
#' # Get Data
ibex_example <- get(data("ibex_example"))

ibex.clones <- CoNGAfy(ibex_example,
                      method = "dist")

ibex.clones <- CoNGAfy(ibex_example,
                      method = "mean")
```

filter.cells	<i>Filter Single-Cell Data Based on CDR3 Sequences</i>
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Description

This function subsets a Seurat or SingleCellExperiment object, removing cells where the CTaa column is missing or contains unwanted patterns.

Usage

```
filter.cells(sc.obj, chain)
```

Arguments

sc.obj	A Seurat or SingleCellExperiment object.
chain	Character. Specifies the chain type ("Heavy" or "Light").

Value

A filtered Seurat or SingleCellExperiment object.

ibex_example	<i>A SingleCellExperiment object with 200 randomly-sampled B cells with BCR sequences from the 10x Genomics 2k_BEAM-Ab_Mouse_HEL_5pv2 dataset.</i>
--------------	--

Description

This object includes normalized gene expression values, metadata annotations, and B cell clonotype information derived from 10x V(D)J sequencing. It is intended as a small example dataset for testing and demonstration purposes.

Format

A SingleCellExperiment object with 32,285 genes (rows) and 200 cells (columns).

assays List of matrices containing expression values: counts (raw counts) and logcounts (log-transformed).

rowData Empty in this example (no gene-level annotations).

colData A DataFrame with 14 columns of cell metadata, including: - orig.ident: Original sample identity. - nCount_RNA: Total number of counts per cell. - nFeature_RNA: Number of detected genes per cell. - cloneSize: Size of each clone. - ident: Cluster assignment.

reducedDims Contains dimensionality reductions: PCA, pca, and apca.

altExp One alternative experiment named BEAM containing additional expression data.

 Ibex_matrix

Ibex Matrix Interface

Description

This function runs the Ibex algorithm to generate latent vectors from input data. The output can be returned as a matrix, with options to choose between deep learning autoencoders or geometric transformations based on the BLOSUM62 matrix.

Usage

```
Ibex_matrix(
  input.data,
  chain = c("Heavy", "Light"),
  method = c("encoder", "geometric"),
  encoder.model = c("CNN", "VAE", "CNN.EXP", "VAE.EXP"),
  encoder.input = c("atchleyFactors", "crucianiProperties", "kideraFactors", "MSWHIM",
    "tScales", "OHE"),
  geometric.theta = pi/3,
  species = "Human",
  verbose = TRUE
)
```

Arguments

input.data	Input data, which can be: <ul style="list-style-type: none"> • A Single Cell Object in Seurat or SingleCellExperiment format • The output of <code>scRepertoire::combineBCR()</code> or <code>combineExpandedBCR()</code> • [Experimental] A character vector of amino acid sequences. The chain parameter specifies whether these are heavy or light chain sequences. For expanded models (CNN.EXP/VAE.EXP), sequences should be formatted as CDR1-CDR2-CDR3 separated by hyphens. If the vector is named, the names will be used as row names in the output.
chain	Character. Specifies which chain to analyze: <ul style="list-style-type: none"> • "Heavy" for the heavy chain • "Light" for the light chain

method	Character. The algorithm to use for generating latent vectors: <ul style="list-style-type: none"> "encoder" - Uses deep learning autoencoders "geometric" - Uses geometric transformations based on the BLOSUM62 matrix
encoder.model	Character. The type of autoencoder model to use: <ul style="list-style-type: none"> "CNN" - CDR3 Convolutional Neural Network-based autoencoder "VAE" - CDR3 Variational Autoencoder "CNN.EXP" - CDR1/2/3 CNN "VAE.EXP" - CDR1/2/3 VAE
encoder.input	Character. Specifies the input features for the encoder model. Options include: <ul style="list-style-type: none"> Amino Acid Properties: "atchleyFactors", "crucianiProperties", "kideraFactors", "MSWHIM", "tScales", "zScales" "OHE" for One Hot Encoding
geometric.theta	Numeric. Angle (in radians) for the geometric transformation. Only used when method = "geometric".
species	Character. Default is "Human" or "Mouse".
verbose	Logical. Whether to print progress messages. Default is TRUE.

Value

A matrix of latent vectors generated by the specified method.

See Also

[immApex::propertyEncoder\(\)](#), [immApex::geometricEncoder\(\)](#)

Examples

```
# Get Data
ibex_example <- get(data("ibex_example"))

# Using the encoder method with a variational autoencoder
ibex_values <- Ibex_matrix(ibex_example,
                          chain = "Heavy",
                          method = "encoder",
                          encoder.model = "VAE",
                          encoder.input = "atchleyFactors")

# Using the geometric method with a specified angle
ibex_values <- Ibex_matrix(ibex_example,
                          chain = "Heavy",
                          method = "geometric",
                          geometric.theta = pi)

# Using a character vector of amino acid sequences
sequences <- c("CARDYW", "CARDSSGYW", "CARDTGYW")
ibex_values <- Ibex_matrix(sequences,
                          chain = "Heavy",
                          method = "geometric")
```

ibex_vdj	<i>Full filtered_annotated_contig.csv from the 10x 2k_BEAM-Ab_Mouse_HEL_5pv2</i>
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Description

This dataset contains single-cell V(D)J sequencing annotations from the 10x Genomics BEAM-Ab Mouse dataset. It includes V(D)J gene calls, CDR regions, productivity information, and clonotype assignments for each contig.

Format

A data frame with 6 rows and 35 columns:

barcode Character. Unique cell barcode.

is_cell Logical. Whether the barcode is identified as a cell.

contig_id Character. Unique identifier for each contig.

high_confidence Logical. Whether the contig is high confidence.

length Integer. Length of the contig.

chain Character. Chain type (e.g., IGH, IGK).

v_gene Character. V gene annotation.

d_gene Character. D gene annotation.

j_gene Character. J gene annotation.

c_gene Character. C gene annotation.

full_length Logical. Whether the contig is full-length.

productive Logical. Whether the contig is productive.

fwr1 Character. Amino acid sequence for Framework Region 1.

fwr1_nt Character. Nucleotide sequence for FWR1.

cdr1 Character. Amino acid sequence for CDR1.

cdr1_nt Character. Nucleotide sequence for CDR1.

fwr2 Character. Amino acid sequence for FWR2.

fwr2_nt Character. Nucleotide sequence for FWR2.

cdr2 Character. Amino acid sequence for CDR2.

cdr2_nt Character. Nucleotide sequence for CDR2.

fwr3 Character. Amino acid sequence for FWR3.

fwr3_nt Character. Nucleotide sequence for FWR3.

cdr3 Character. Amino acid sequence for CDR3.

cdr3_nt Character. Nucleotide sequence for CDR3.

fwr4 Character. Amino acid sequence for FWR4.

fwr4_nt Character. Nucleotide sequence for FWR4.

reads Integer. Number of reads supporting the contig.

umis Integer. Number of UMIs supporting the contig.

raw_clonotype_id Character. Clonotype ID from 10x output.

raw_consensus_id Character. Consensus ID from 10x output.

exact_subclonotype_id Integer. Exact subclonotype grouping.

runIbex

*Ibex Single-Cell Calculation***Description**

This function applies the Ibex algorithm to single-cell data, integrating seamlessly with Seurat or SingleCellExperiment pipelines. The algorithm generates latent dimensions using deep learning or geometric transformations, storing the results in the dimensional reduction slot. runIbex will automatically subset the single-cell object based on amino acid sequences present for the given chain selection.

Usage

```
runIbex(
  sc.data,
  chain = "Heavy",
  method = "encoder",
  encoder.model = "VAE",
  encoder.input = "atchleyFactors",
  geometric.theta = pi,
  reduction.name = "Ibex",
  species = "Human",
  verbose = TRUE
)
```

Arguments

sc.data	A single-cell dataset, which can be: <ul style="list-style-type: none"> • A Seurat object • A SingleCellExperiment object
chain	Character. Specifies the chain to analyze: <ul style="list-style-type: none"> • "Heavy" for the heavy chain • "Light" for the light chain
method	Character. Algorithm to use for generating latent dimensions: <ul style="list-style-type: none"> • "encoder" - Uses deep learning autoencoders • "geometric" - Uses geometric transformations based on the BLOSUM62 matrix
encoder.model	Character. The type of autoencoder model to use: <ul style="list-style-type: none"> • "CNN" - CDR3 Convolutional Neural Network-based autoencoder • "VAE" - CDR3 Variational Autoencoder • "CNN.EXP" - CDR1/2/3 CNN • "VAE.EXP" - CDR1/2/3 VAE
encoder.input	Character. Input features for the encoder model: <ul style="list-style-type: none"> • Amino Acid Properties: "atchleyFactors", "crucianiProperties", "kideraFactors", "MSWHIM", "tScales" • "OHE" - One Hot Encoding

<code>geometric.theta</code>	Numeric. Angle (in radians) for geometric transformation. Used only when <code>method = "geometric"</code> .
<code>reduction.name</code>	Character. The name to assign to the dimensional reduction. This is useful for running Ibex with multiple parameter settings and saving results under different names.
<code>species</code>	Character. Default is "Human" or "Mouse".
<code>verbose</code>	Logical. Whether to print progress messages. Default is TRUE.

Value

An updated Seurat or SingleCellExperiment object with Ibex dimensions added to the dimensional reduction slot.

Examples

```
# Get Data
ibex_example <- get(data("ibex_example"))

# Using the encoder method with a variational autoencoder
ibex_example <- runIbex(ibex_example,
  chain = "Heavy",
  method = "encoder",
  encoder.model = "VAE",
  encoder.input = "atchleyFactors")

# Using the geometric method with a specified angle
ibex_example <- runIbex(ibex_example,
  chain = "Heavy",
  method = "geometric",
  geometric.theta = pi)
```

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