

# Package ‘NestLink’

June 24, 2025

**Type** Package

**Title** NestLink an R data package to guide through Engineered Peptide Barcodes for In-Depth Analyzes of Binding Protein Ensembles

**Version** 1.25.0

**Depends** R (>= 3.6), AnnotationHub (>= 2.15), ExperimentHub (>= 1.0), Biostrings (>= 2.51), gplots (>= 3.0), protViz (>= 0.4), ShortRead (>= 1.41)

**Imports** grDevices, graphics, stats, utils

**Description** Provides next-generation sequencing (NGS) and mass spectrometry (MS) sample data, code snippets and replication material used for developing NestLink. The NestLink approach is a protein binder selection and identification technology able to biophysically characterize thousands of library members at once without handling individual clones at any stage of the process. Data were acquired on NGS and MS platforms at the Functional Genomics Center Zurich.

**License** GPL

**VignetteBuilder** knitr

**Suggests** BiocStyle (>= 2.2), DT, ggplot2, knitr, rmarkdown, testthat, specL, lattice, scales

**NeedsCompilation** no

**biocViews** ExperimentHub, ExperimentData, SequencingData, MassSpectrometryData, ReproducibleResearch

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|              |                                                                          |
|--------------|--------------------------------------------------------------------------|
| .ssrc.mascot | <i>computes the correlation of predicted and measured retention time</i> |
|--------------|--------------------------------------------------------------------------|

---

## Description

this helper function computes a linear model between redicted and measured retention time of the as imput set given identified peptides.

TODO(cp): consider moving this method to the protViz package.

## Usage

```
.ssrc.mascot(x, scores = c(10, 20, 40, 50), ...)
```

## Arguments

|        |                                                          |
|--------|----------------------------------------------------------|
| x      | as,data.frame <b>mascot</b> generated data.frame object. |
| scores | default is c(10, 20, 40, 50).                            |
| ...    | passed to the plot function.                             |

**Value**

a plot and summary

**Author(s)**

Christian Panse <cp@fgcz.ethz.ch>, 2017,2019

**Examples**

```
library(ExperimentHub)
eh <- ExperimentHub();
load(query(eh, c("NestLink", "F255744.RData"))[[1]])
.ssrc.mascot(F255744, scores = 15)
```

---

|                    |                                                               |
|--------------------|---------------------------------------------------------------|
| compose_GPGx8cTerm | <i>Compose a peptide with a defined AA sequence frequency</i> |
|--------------------|---------------------------------------------------------------|

---

**Description**

composes, out of an as input given amino acid distribution, a randomly sampled amino acid sequence. [compose\\_GPGx8cTerm](#), [compose\\_GSx7cTerm](#), and [compose\\_GPx10R](#) belong to three groups composing different flycode (peptide) construction. The construction is given in the function name. For example, GPGx8cTerm, composes a flycode having as prefix GPG followed by eight (x8) amino acids followed by a cTerm sequence. The different construction will have different detectability properties as mass range and hydrophobicity values.

**Usage**

```
compose_GPGx8cTerm(pool = c(rep("A", 12), rep("S", 0), rep("T", 12),
  rep("N", 12), rep("Q", 12), rep("D", 8), rep("E", 0), rep("V", 12),
  rep("L", 0), rep("F", 0), rep("Y", 8), rep("W", 0), rep("G", 12),
  rep("P", 12)), cTerm = c("VFR", "VSR", "VFGIR", "VSGER"))
```

**Arguments**

|       |                 |
|-------|-----------------|
| pool  | AA distributen. |
| cTerm | c-Terms         |

**Value**

a AA sequence

**Author(s)**

Christian Panse <cp@fgcz.ethz.ch> 2015

**Examples**

```
set.seed(1)
compose_GPGx8cTerm()
(FlyCodes <- replicate(10, compose_GPGx8cTerm()))
plot(parentIonMass(FlyCodes) ~ssrc(FlyCodes))
```

compose\_GPx10R

*Compose a peptide with a defined AA sequence***Description**

composes, out of an as input given amino acid distribution, a randomly sampled amino acid sequence. `compose_GPGx8cTerm`, `compose_GSx7cTerm`, and `compose_GPx10R` belong to three groups composing different flycode (peptide) construction. The construction is given in the function name. For example, `GPGx8cTerm`, composes a flycode having as prefix GPG followed by eight (x8) amino acids followed by a cTerm sequence. The different construction will have different detectability properties as mass range and hydrophobicity values.

**Usage**

```
compose_GPx10R(aa_pool1, aa_pool2)
```

**Arguments**

```
aa_pool1      AA distributen.
aa_pool2      AA distributen.
```

**Value**

```
a AA sequence
```

**Author(s)**

```
Christian Panse <cp@fgcz.ethz.ch> 2015
```

**Examples**

```
set.seed(1)
aa_pool_1_2_9_10 <- c(rep('A', 8), rep('S', 7), rep('T', 7), rep('N', 6),
  rep('Q', 6), rep('D', 8), rep('E', 8), rep('V', 9), rep('L', 6), rep('F', 5),
  rep('Y', 9), rep('W', 6), rep('G', 15), rep('P', 0))

aa_pool_3_8 <- c(rep('A', 5), rep('S', 4), rep('T', 5), rep('N', 2),
  rep('Q', 2), rep('D', 8), rep('E', 8), rep('V', 7), rep('L', 5), rep('F', 4),
  rep('Y', 6), rep('W', 4), rep('G', 12), rep('P', 28))

compose_GPx10R(aa_pool_1_2_9_10, aa_pool_3_8)
(FlyCodes <- replicate(10, compose_GPx10R(aa_pool_1_2_9_10, aa_pool_3_8)))
plot(parentIonMass(FlyCodes) ~ssrc(FlyCodes))
```

---

|                   |                                                        |
|-------------------|--------------------------------------------------------|
| compose_GSx7cTerm | <i>Compose a FlyCode GSx7cTerm Amino Acid Sequence</i> |
|-------------------|--------------------------------------------------------|

---

## Description

composes, out of an as input given amino acid distribution, a randomly sampled amino acid sequence. `compose_GPGx8cTerm`, `compose_GSx7cTerm`, and `compose_GPx10R` belong to three groups composing different flycode (peptide) construction. The construction is given in the function name. For example, `GPGx8cTerm`, composes a flycode having as prefix GPG followed by eight (x8) amino acids followed by a cTerm sequence. The different construction will have different detectability properties as mass range and hydrophobicity values.

## Usage

```
compose_GSx7cTerm(pool = c(rep("A", 18), rep("S", 6), rep("T", 12),
  rep("N", 1), rep("Q", 1), rep("D", 11), rep("E", 11), rep("V", 12),
  rep("L", 2), rep("F", 1), rep("Y", 4), rep("W", 1), rep("G", 8), rep("P",
  12)), cTerm = c("WR", "WLTVR", "WQEGGR", "WQSR", "WLR"))
```

## Arguments

|       |                                |
|-------|--------------------------------|
| pool  | a vector of amino acids.       |
| cTerm | a vector of a sequence suffix. |

## Value

a amino acid sequence, e.g., GSAPTTVFGWLTVR.

## Author(s)

Christian Panse <cp@fgcz.ethz.ch> 2015

## Examples

```
sample.size <- 100
#
## Compose a GSXXXXXX(WR|WLTVR|WQGGGR|WQSR|WLR) peptide
set.seed(2)
FC.GSx7cTerm <- replicate(sample.size, compose_GSx7cTerm())
## Some Sanity Checks
table(FC.GSx7cTerm)
stopifnot(length(FC.GSx7cTerm) == 100)
FC.PATTERN <- "^GS[ASTNQDEFVLYWGP]{7}(WR|WLTVR|WQEGGR|WLR|WQSR)$"
stopifnot(
  length(FC.GSx7cTerm[grep(FC.PATTERN, FC.GSx7cTerm)])
  == sample.size)
```

F255744

*F255744 Mascot Search results***Description**

F255744 Mascot Search results

**Author(s)**

Pascal Egloff &lt;p.egloff@imm.uzh.ch&gt;

**See Also**[F255744](#)**Examples**

```
library(ExperimentHub)
eh <- ExperimentHub(); load(query(eh, c("NestLink", "F255744.RData"))[[1]])
class(F255744)
hist(F255744$RTINSECONDS)
hist(F255744$RTINSECONDS[F255744$pep_score > 20])
```

getExperimentHubFilename

*getExperimentHubFilename***Description**

getExperimentHubFilename

**Usage**

getExperimentHubFilename(filename)

**Arguments**

filename            of the aws s3 blob.

**Value**

the file name of the local ExperimentHub.

**Examples**

```
f1 <- system.file("extdata", "metadata.csv", package="NestLink")
metadata <- read.csv(f1, stringsAsFactors=FALSE)
  metadata$Title

lapply(metadata$RDataPath, getExperimentHubFilename)
```

---

|       |                            |
|-------|----------------------------|
| getFC | <i>Read FlyCodes (FCs)</i> |
|-------|----------------------------|

---

## Description

A wrapper function for reading the flycodes using ExperimentHub. The files are used for demonstrating the detectability of the AA sequences. The wrapper functions are extended by columns [ssrc](#) prediction and the [parentIonMass](#). The column ESP\_Prediction was generated by using the service from <https://genepattern.broadinstitute.org>.

## Usage

```
getFC(pattern = "^GS[ASTNQDEFVLYWGP]{7}(WR|WLTVR|WQEGGR|WLR|WQSR)$",  
      filename = NULL)
```

## Arguments

|          |                                                                                                                                                                     |
|----------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| pattern  | a regular expression FlyCode pattern                                                                                                                                |
| filename | a two column tab separated file containing a peptide sequence and an ESP value. default is NULL which reads the data provided by the package through ExperimentHub. |

## Value

a data.frame object of Flycodes

## Author(s)

Christian Panse <cp@fgcz.ethz.ch> 2015, 2018

## Source

- [https://fgcz-gstore.uzh.ch/projects/p1644/analysis\\_20170609\\_o3040/p1644o3482-4\\_S4.extendedFrag\\_uniqNB2FC.txt](https://fgcz-gstore.uzh.ch/projects/p1644/analysis_20170609_o3040/p1644o3482-4_S4.extendedFrag_uniqNB2FC.txt)
- [https://fgcz-gstore.uzh.ch/projects/p1644/analysis\\_20170609\\_o3040/p1644o3482-5\\_S5.extendedFrag\\_uniqNB2FC.txt](https://fgcz-gstore.uzh.ch/projects/p1644/analysis_20170609_o3040/p1644o3482-5_S5.extendedFrag_uniqNB2FC.txt)

## Examples

```
FC <- getFC()  
dim(FC)
```

---

`getNB`*Read NanoBodies (NBs)*

---

## Description

A wrapper function for reading the flycodes using ExperimentHub. The files are used for demonstrating the detectability of the AA sequences. The wrapper functions are extended by columns `ssrc` prediction and the `parentIonMass`. The column `ESP_Prediction` was generated by using the service from <https://genepattern.broadinstitute.org>.

## Usage

```
getNB(filename = NULL)
```

## Arguments

|                       |                                                                                                                                                                                  |
|-----------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>filename</code> | a two column tab separated file containing a peptide sequence and an ESP value. default is <code>NULL</code> which reads the data provided by the package through ExperimentHub. |
|-----------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

## Value

a `data.frame` object of NBs

## Author(s)

Christian Panse <cp@fgcz.ethz.ch> 2015, 2018, 2019

## Source

- [https://fgcz-gstore.uzh.ch/projects/p1644/analysis\\_20170609\\_o3040/p1644o3482-4\\_S4.extendedFragments\\_uniqNB2FC.txt](https://fgcz-gstore.uzh.ch/projects/p1644/analysis_20170609_o3040/p1644o3482-4_S4.extendedFragments_uniqNB2FC.txt)
- [https://fgcz-gstore.uzh.ch/projects/p1644/analysis\\_20170609\\_o3040/p1644o3482-5\\_S5.extendedFragments\\_uniqNB2FC.txt](https://fgcz-gstore.uzh.ch/projects/p1644/analysis_20170609_o3040/p1644o3482-5_S5.extendedFragments_uniqNB2FC.txt)

## Examples

```
NB <- getNB()
dim(NB)
```

---

nanobodyFlycodeLinking.as.fasta  
*Write FASTA*

---

**Description**

Write FASTA

**Usage**

```
nanobodyFlycodeLinking.as.fasta(x, file = NULL, ...)
```

**Arguments**

|      |                                                                                 |
|------|---------------------------------------------------------------------------------|
| x    | a nanobodyFlycodeLinking S3 object computed by <a href="#">runNGSAnalysis</a> . |
| file | a filename                                                                      |
| ...  | just passed                                                                     |

**Value**

sprintf stream

**Author(s)**

Lennart Opitz, Christian Panse 2018

**Examples**

```
library(ExperimentHub)
eh <- ExperimentHub()
f <- query(eh, c("NestLink", "nanobodyFlycodeLinkage.RData"))[[1]]
load(f)
summary(nanobodyFlycodeLinkage.sample)
nanobodyFlycodeLinking.as.fasta(nanobodyFlycodeLinkage.sample)
```

---

nanobodyFlycodeLinking.summary  
*Object Summaries of S3 class nanobodyFlycodeLinking*

---

**Description**

Object Summaries of S3 class nanobodyFlycodeLinking

**Usage**

```
nanobodyFlycodeLinking.summary(object)
```

**Arguments**

|        |                                                                             |
|--------|-----------------------------------------------------------------------------|
| object | a nanobodyFlycodeLinking class computed by <a href="#">runNGSAnalysis</a> . |
|--------|-----------------------------------------------------------------------------|

**Value**

a data.frame object

**Examples**

```
library(ExperimentHub)
eh <- ExperimentHub()
f <- query(eh, c("NestLink", "nanobodyFlycodeLinkage.RData"))[[1]]
load(f)
summary(nanobodyFlycodeLinkage.sample)
```

---

|                |                                  |
|----------------|----------------------------------|
| NB.unambiguous | <i>Determine unambiguous NBs</i> |
|----------------|----------------------------------|

---

**Description**

Determine unambiguous NBs

**Usage**

```
NB.unambiguous(x = getNB())
```

**Arguments**

x                      a data.frame containing a column peptide

**Value**

a data.frame a data.frame of unambiguously assignable peptides (those, which occur only on one nanobody)

**Examples**

```
NB <- getNB()
dim(NB.unambiguous(NB))
```

---

|           |                             |
|-----------|-----------------------------|
| NB.unique | <i>make NB table unique</i> |
|-----------|-----------------------------|

---

**Description**

make NB table unique

**Usage**

```
NB.unique(x = getNB())
```

**Arguments**

x                      a data.frame

**Value**

a data.frame

**Examples**

```
NB <- getNB()
dim(NB.unique(NB))
```

---

|          |                         |
|----------|-------------------------|
| PGexport | <i>PGexport results</i> |
|----------|-------------------------|

---

**Description**

PGexport results

**Author(s)**

Pascal Egloff <p.egloff@imm.uzh.ch>

**Source**

<https://fgcz-bfabric.uzh.ch>

- Workunit : 158716 - QEXACTIVEHF\_1 20170919\_16\_62465\_n15idx1-3\_6titratecoli.raw 20170919\_05\_62465\_n15idx1-3\_6titratecoli.raw
- Workunit : 158717 - QEXACTIVEHF\_1 20170919\_14\_62466\_n15idx1-3\_7titratesmeg.raw 20170919\_09\_62466\_n15idx1-3\_7titratesmeg.raw

**Examples**

```
# filename <- system.file(
#   "extdata/PGexport2_normalizedAgainstSBstandards_Peptides.csv",
#   package = "NestLink")
library(ExperimentHub)
eh <- ExperimentHub()
filename <- query(eh,
  c("NestLink", "PGexport2_normalizedAgainstSBstandards_Peptides.csv"))[[1]]
P <- read.csv(filename, header = TRUE, sep=';')
P <- P[P$Modifications == '', ]
P <- P[,c('Accession', 'Sequence',
  "X20170919_05_62465_n15idx1.3_6titratecoli",
  "X20170919_16_62465_n15idx1.3_6titratecoli",
  "X20170919_09_62466_n15idx1.3_7titratesmeg",
  "X20170919_14_62466_n15idx1.3_7titratesmeg")]
names(P)<-c('Accession','Sequence','coli1','coli2','smeg1','smeg2')
P<- P[grepl("^P[0-9][A-Z][0-9]", P$Accession), ]

P$FCset_ng <- NA
P$FCset_ng[P$Accession %in% c('P1A4', 'P1B4', 'P1C4',
  'P1D4', 'P1E4', 'P1F4')] <- 92
P$FCset_ng[P$Accession %in% c('P1A5', 'P1B5', 'P1C5',
  'P1D5', 'P1G4', 'P1H4')] <- 295
P$FCset_ng[P$Accession %in% c('P1A6', 'P1B6', 'P1E5',
```

```

'P1F5', 'P1G5', 'P1H5'))] <- 943
P$FCset_ng[P$Accession %in% c('P1C6', 'P1D6', 'P1E6',
'P1F6', 'P1G6', 'P1H6'))] <- 3017

P$coli1 <- (log(P$coli1,2) - mean(log(P$coli1,2))) / sd(log(P$coli1,2))
P$coli2 <- (log(P$coli2,2) - mean(log(P$coli2,2))) / sd(log(P$coli2,2))
P$smeg1 <- (log(P$smeg1,2) - mean(log(P$smeg1,2))) / sd(log(P$smeg1,2))
P$smeg2 <- (log(P$smeg2,2) - mean(log(P$smeg2,2))) / sd(log(P$smeg2,2))

O <- P
b <- boxplot(df<-cbind(P$coli1 - P$coli2, P$coli1 - P$smeg1,
P$coli1 - P$smeg2,P$coli2 - P$smeg1, P$coli2 - P$smeg2,
P$smeg1 - P$smeg2),
  ylab='normalized log2ratios', ylim = c(-1,1), axes=FALSE,
  main=paste("ConcGr = all"))
axis(1, 1:6, c('coli[12]', 'coli1-smeg1', 'coli1-smeg2', 'coli2-smeg1',
'coli2- smeg2', 'smeg[12]'))
abline(h=0, col='red')
box()
axis(2)
axis(3, 1:6, b$n)
outliers.idx <- sapply(1:length(b$group), function(i){
  q <- df[, b$group[i]] == b$out[i];
  text(b$group[i], b$out[i], P[q, 2], pos=4, cex=0.4);
  text(b$group[i], b$out[i], P[q, 1], pos=2, cex=0.4);
  which(q)}
)

```

---

plot\_in\_silico\_LCMS\_map

*plot a LC-MS map of a given set of amino acid sequences*


---

## Description

plot a LC-MS map of a given set of amino acid sequences

## Usage

```
plot_in_silico_LCMS_map(peptides, ...)
```

## Arguments

|          |                               |
|----------|-------------------------------|
| peptides | a vector of pepitdes.         |
| ...      | pass through the plot method. |

## Details

TODO(cp): consider using hexbin using ggplot2 ggplot facet\_wrap aes geom\_point

## Value

gplots::hist2d a gplot 2d histogram

**Author(s)**

Christian Panse

**Examples**

```
set.seed(1)
par(mfrow=c(2,1));
FlyCodes <- replicate(10000, compose_GPGx8cTerm())
rv <- plot_in_silico_LCMS_map(FlyCodes)
```

---

runNGSAnalysis

*NGS linkage workflow*

---

**Description**

performs the NGS filtering workflow to get high quality FlyCode and Nanobody sequences linkage.

**Usage**

```
runNGSAnalysis(file, param)
```

**Arguments**

|       |                                                                 |
|-------|-----------------------------------------------------------------|
| file  | sequence file path                                              |
| param | list of input parameters, explained in details paragraph below. |

**Details**

The elements of the parameter list object is described as follows:

- NB\_Linker1 nucleotide sequence of the linker left to the nanobody.
- NB\_Linker2 nucleotide sequence of the linker right to the nanobody.
- ProteaseSite nucleotide sequence left to the flycode.
- FC\_Linker nucleotide sequence right to the flycode.
- knownNB known nanobody sequences in the experiment.
- nReads number of Reads from the start of fastq file to process.
- minRelBestHitFreq minimal fraction of the dominant nanobody for a specific flycode.
- minConsensusScore minimal fraction per sequence position in nanobody consensus sequence calculation.
- maxMismatch number of accepted mismatches for all pattern search steps.
- minNanobodyLength minimal nanobody length in [nt].
- minFlycodeLength minimal flycode length in [nt].
- FCminFreq minimal number of subreads for a specific flycode to keep it in the analysis.

missing elements are replace by the example provided values.

**Value**

uniqNB2FC dataframe

**Author(s)**

Lennart Opitz <lopitz@fgcz.ethz.ch>, 2019

**Examples**

```
library(ExperimentHub)
eh <- ExperimentHub()
expFile <- query(eh, c("NestLink", "NL42_100K.fastq.gz"))[[1]]
knownNB_File <- query(eh, c("NestLink", "knownNB.txt"))[[1]]
knownNB_data <- read.table(knownNB_File, sep='\t', header = TRUE,
  row.names = 1, stringsAsFactors = FALSE)
knownNB <- Biostrings::translate(DNAStringSet(knownNB_data$Sequence))
names(knownNB) <- rownames(knownNB_data)
knownNB <- sapply(knownNB, toString)
param <- list()
param[['NB_Linker1']] <- "GGCCggcggGGCC"
param[['NB_Linker2']] <- "GCAGGAGGA"
param[['ProteaseSite']] <- "TTAGTCCCAAGA"
param[['FC_Linker']] <- "GGCCaaggaggcCGG"
param[['knownNB']] <- knownNB
param[['nReads']] <- 10000
param[['minRelBestHitFreq']] <- 0.8
param[['minConsensusScore']] <- 0.9
param[['maxMismatch']] <- 1
param[['minNanobodyLength']] <- 348
param[['minFlycodeLength']] <- 33
param[['FCminFreq']] <- 1
runNGSAnalysis(file = expFile[1], param)
```

---

twoPatternReadFilter    *Filter input sequences for two patterns*

---

**Description**

Filter input sequences for two patterns

**Usage**

```
twoPatternReadFilter(reads, leftPattern, rightPattern, maxMismatch,
  prevPatternPos = NULL)
```

**Arguments**

|                |                                                 |
|----------------|-------------------------------------------------|
| reads          | input sequences                                 |
| leftPattern    | left pattern motive.                            |
| rightPattern   | right pattern motive.                           |
| maxMismatch    | maximal number of miss matches.                 |
| prevPatternPos | prev pattern posiotion; default is set to NULL. |

**Value**

list object

**Examples**

```
reads <- DNASTringSet(c('ACTGGGTTT','ACCCTGGGTTT'))
leftPattern <- 'CT'
rightPattern <- 'TTT'
maxMismatch <- 0
twoPatternReadFilter(reads, leftPattern, rightPattern, maxMismatch)
```

WU160118

*WU160118 Mascot Search results***Description**

WU160118 Mascot Search results

**Author(s)**

Christian Panse

**References**

<https://fgcz-bfabric.uzh.ch/bfabric/userlab/show-workunit.html?id=160118>

**See Also**

please read the vignette summaryFASTA.Rmd.

**Examples**

```
library(ExperimentHub)
eh <- ExperimentHub();
load(query(eh, c("NestLink", "WU160118.RData"))[[1]])
class(WU160118)
PATTERN <- "^GS[ASTNQDEFVLYWGP]{7}(WR|WLTVR|WQEGGR|WLR|WQSR)$"
idx <- grepl(PATTERN, WU160118$pep_seq)
WU <- WU160118[idx & WU160118$pep_score > 25,]

library(lattice)
histogram(~RTINSECONDS| datfilename, data = WU, type='count')
```

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