

# Package ‘TrIdent’

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**Type** Package

**BiocType** Software

**Title** TrIdent - Transduction Identification

**Version** 0.99.3

**Description** The `TrIdent` R package automates the analysis of transductomics data by detecting, classifying, and characterizing read coverage patterns associated with potential transduction events. Transductomics is a DNA sequencing-based method for the detection and characterization of transduction events in pure cultures and complex communities. Transductomics relies on mapping sequencing reads from a viral-like particle (VLP)-fraction of a sample to contigs assembled from the metagenome (whole-community) of the same sample. Reads from bacterial DNA carried by VLPs will map back to the bacterial contigs of origin creating read coverage patterns indicative of ongoing transduction.

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TrIdent-package      *TrIdent - Transduction Identification*

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**Description**

Automatic detection, classification and characterization of transduction events in transductomics datasets using read coverage pattern-matching.

Please see [Transductomics: sequencing-based detection and analysis of transduced DNA in pure cultures and microbial communities] (<https://microbiomejournal.biomedcentral.com/articles/10.1186/s40168-020-00935-5>) for more information on the transductomics method, data and analysis workflow.

**Details**

The three main functions in TrIdent are:

1. [TrIdentClassifier](#) performs the pattern-matching, classification and characterization of read coverage patterns on contigs.
2. [plotTrIdentResults](#) plots the results from `TrIdentClassifier()`
3. [specializedTransductionID](#) searches contigs classified as Prophage-like by `TrIdentClassifier()` for potential specialized transduction

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**See Also**

Useful links:

- <https://github.com/jlmaier12/TrIdent>
- <https://jlmaier12.github.io/TrIdent/>
- Report bugs at <https://github.com/jlmaier12/TrIdent/issues>

---

allPatternMatches      *Collects pattern-match information for all classifications*

---

**Description**

Collects pattern information associated with all contigs classified as Prophage-like, Sloping and HighCovNoPattern.

**Usage**

```
allPatternMatches(bestMatchList, classifSummTable)
```

**Arguments**

bestMatchList    Classifications made with patternMatcher function.

classifSummTable

Classification summary table with whole-community:VLP-fraction read coverage ratios calculated.

**Value**

List

---

allProphageLikeClassifs  
*Collects Prophage-like classification pattern-match information*

---

**Description**

Collects pattern information associated with all contigs classified as Prophage-like.

**Usage**

```
allProphageLikeClassifs(bestMatchList)
```

**Arguments**

bestMatchList    Classifications made with patternMatcher function.

**Value**

List

---

allSlopingClassifs	<i>Collects Sloping classification pattern-match information</i>
--------------------	--

---

**Description**

Collects pattern information associated with all contigs classified as Sloping in the patternMatcher function.

**Usage**

```
allSlopingClassifs(bestMatchList)
```

**Arguments**

bestMatchList    Classifications made with patternMatcher function.

**Value**

List

---

blockBuilder	<i>Builds prophage-like block patterns</i>
--------------	--

---

**Description**

Build and translate a block pattern going off the left side, right side and full length of the contig.

**Usage**

```
blockBuilder(viralSubset, windowSize, minBlockSize, maxBlockSize)
```

**Arguments**

viralSubset    A subset of the read coverage pileup that pertains only to the contig currently being assessed

windowSize    The window size used to re-average read coverage pileups

minBlockSize    The minimum size of the prophage-like block pattern. Default is 10000 bp.

maxBlockSize    The maximum size of the prophage-like block pattern. Default is NA.

**Value**

List containing three objects

---

blockTranslator	<i>Full block pattern-translator</i>
-----------------	--------------------------------------

---

### Description

Translates full block-pattern across a contig. Translate the pattern 1000 bp at a time. Stop translating when the pattern is 5000 bp from the end of the contig.

### Usage

```
blockTranslator(viralSubset, bestMatchInfo, windowSize, pattern)
```

### Arguments

viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
bestMatchInfo	The information associated with the current best pattern-match.
windowSize	The window size used to re-average read coverage pileups
pattern	A vector containing the values associated with the block pattern

### Value

List

---

changeSlope	<i>Change slope of sloping pattern</i>
-------------	--

---

### Description

Change the value of the slope used for the sloping pattern-match

### Usage

```
changeSlope(
  leftOrRight,
  slopeBottom,
  halfToMaxReadCov,
  cov,
  viralSubset,
  windowSize
)
```

**Arguments**

leftOrRight	Generate pattern for negative slope (left to right, i.e. 'Left') or positive slope (right to left, i.e. 'Right')
slopeBottom	The value for the bottom of the sloping value
halfToMaxReadCov	Half of the max VLP-fraction read coverage divided by 10
cov	The value for the top of the slope
viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
windowSize	The window size used to re-average read coverage pileup

**Value**

List

---

changeSlopeWStart	<i>Change slope of sloping pattern with initial start</i>
-------------------	---

---

**Description**

Change the value of the slope used for the sloping with start pattern-match

**Usage**

```
changeSlopeWStart(
  leftOrRight,
  slopeBottom,
  slopeBottomChange,
  cov,
  viralSubset,
  windowSize
)
```

**Arguments**

leftOrRight	Generate pattern for negative slope (left to right, i.e. 'Left') or positive slope (right to left, i.e. 'Right')
slopeBottom	The value for the bottom of the sloping value
slopeBottomChange	The value used to increase the bottom of the slope
cov	The value for the top of the slope
viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
windowSize	The window size used to re-average read coverage pileup

**Value**

List

---

changeWindowSize	<i>Change the read coverage rolling mean window size</i>
------------------	--

---

**Description**

Re-averages window sizes of read coverage averages. Start with 100bp windows always. Cannot make window size less than 100bp.

**Usage**

```
changeWindowSize(cleanPileup, windowSize)
```

**Arguments**

cleanPileup	A read coverage dataset that has been cleaned and reformatted.
windowSize	The number of base pairs to average coverage values over. Options are 100, 500, 1000, or 2000 only!

**Value**

Dataframe

---

contigClassSumm	<i>Summarizes pattern-match information</i>
-----------------	---

---

**Description**

Summarizes the classifications made in the patternMatcher() function into a dataframe.

**Usage**

```
contigClassSumm(bestMatchList)
```

**Arguments**

bestMatchList	Classifications made with patternMatcher function.
---------------	--

**Value**

dataframe



---

fullSlope	<i>Sloping pattern builder</i>
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---

**Description**

Build a sloping pattern that consists of a sloping line spanning the contig being assessed. The line slopes from left to right. The slope of the line is changed, but the pattern is not translated across the contig.

**Usage**

```
fullSlope(viralSubset, windowSize, minSlope)
```

**Arguments**

viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
windowSize	The window size used to re-average read coverage pileup
minSlope	The minimum slope value to test for sloping patterns

**Value**

List containing two objects

---

leftRightBlockTranslator	<i>Translate left and right block patterns across contig</i>
--------------------------	--

---

**Description**

Translates left and right block patterns across contigs 1000 bp at a time

**Usage**

```
leftRightBlockTranslator(  
    viralSubset,  
    pattern,  
    leftOrRight,  
    windowSize,  
    minReadCov,  
    cov,  
    bestMatchInfo,  
    minBlockSize  
)
```

**Arguments**

viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
pattern	The pattern vector being translated
leftOrRight	Is the left or right block pattern being translated
windowSize	The window size used to re-average read coverage pileups
minReadCov	The baseline value used for the region outside of the block pattern (either 0 or the minimum VLP-fraction read coverage for the contig)
cov	The height value currently being used for the block pattern
bestMatchInfo	The information associated with the current best pattern-match.
minBlockSize	The minimum size of the Prophage-like block pattern. Default is 10,000 bp.

**Value**

List

---

makeBlockPattern	<i>Make block patterns for pattern-matching</i>
------------------	---

---

**Description**

Make full, left and right block patterns for Prophage-like classifications

**Usage**

```
makeBlockPattern(
  viralSubset,
  windowSize,
  fullLeftRight,
  blockLength,
  nonBlock,
  minReadCov,
  cov
)
```

**Arguments**

viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
windowSize	The window size used to re-average read coverage pileups
fullLeftRight	The block pattern variation being built
blockLength	Maximum block pattern length
nonBlock	Maximum non-block pattern length
minReadCov	Either 0 or the minimum VLP-fraction read coverage value
cov	The height value of the block pattern

**Value**

List containing two objects

---

makeFullSlopes	<i>Make full slope patterns</i>
----------------	---------------------------------

---

**Description**

Makes slope patterns sloping either left to right (Left) or right to left (right) across the contig being assessed.

**Usage**

```
makeFullSlopes(leftOrRight, viralSubset, newMax, minReadCov, windowSize)
```

**Arguments**

leftOrRight	Generate pattern for negative slope (left to right, i.e. 'Left') or positive slope (right to left, i.e. 'Right')
viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
newMax	A value for the top of the sloping pattern that is slightly higher than the maximum coverage value on the viralSubset
minReadCov	Minimum read coverage value of the viralSubset
windowSize	The window size used to re-average read coverage pileups

**Value**

List

---

makeSlopesWStarts	<i>Make slope patterns with starts</i>
-------------------	--

---

**Description**

Makes slope patterns sloping either left to right (Left) or right to left (right) across the contig being assessed. Slope patterns contain an initiation point.

**Usage**

```
makeSlopesWStarts(leftOrRight, viralSubset, newMax, minReadCov, windowSize)
```

**Arguments**

leftOrRight	Generate pattern for negative slope (left to right, i.e. 'Left') or positive slope (right to left, i.e. 'Right')
viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
newMax	A value for the top of the sloping pattern that is slightly higher than the maximum coverage value on the viralSubset
minReadCov	Minimum read coverage value of the viralSubset
windowSize	The window size used to re-average read coverage pileup

**Value**

List

---

NARemover

*NA remover*

---

**Description**

Removes NAs from dataframe.

**Usage**

```
NARemover(x)
```

**Arguments**

x                    dataset with potential NAs

**Value**

Dataframe

**See Also**

<https://stackoverflow.com/questions/18142117/how-to-replace-nan-value-with-zero-in-a-huge-data-frame/18143097#18143097>

---

noPattern	<i>No pattern pattern-match</i>
-----------	---------------------------------

---

**Description**

A horizontal line at the mean or median coverage should be an optimal pattern-match if the contig read coverage displays no sloping or block patterns

**Usage**

```
noPattern(viralSubset)
```

**Arguments**

viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
-------------	--

**Value**

List

---

patternBuilder	<i>Pattern-builder</i>
----------------	------------------------

---

**Description**

Builds the pattern (vector) associated with the best pattern-match' for each contig classified as Prophage-like, Sloping, or HighCovNoPattern.

**Usage**

```
patternBuilder(viralSubset, classifList, classification, i)
```

**Arguments**

viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
classifList	A list containing pattern match information associated with all classified contigs.
classification	The contig's classification assigned by the TrIdentClassifier function
i	The list index associated with each contig's pattern-match information

**Value**

Vector

---

patternMatcher      *Main pattern-matching function*

---

### Description

Creates the viralSubset, representative of one contig, that is used as input for each individual pattern-matching function. After the information associated with the best match for each pattern is obtained, the pattern with the smallest match score is used to classify the contig being assessed. Prior to the pattern-matching, contigs smaller than the minContigLength and contigs without 5,000 bp of 10x read coverage are removed.

### Usage

```
patternMatcher(
  VLPpiledup,
  WCPiledup,
  windowSize,
  minBlockSize,
  maxBlockSize,
  minContigLength,
  minSlope,
  verbose
)
```

### Arguments

VLPpiledup	A table containing contig names, coverages averaged over 100 bp windows, and contig positions associated with mapping VLP-fraction reads to whole-community contigs
WCPiledup	A table containing contig names, coverages averaged over 100 bp windows, and contig positions associated with mapping whole-community reads to whole-community contigs
windowSize	The window size used to re-average read coverage datasets
minBlockSize	The minimum size of the prophage-like block pattern. Default is 10,000 bp.
maxBlockSize	The maximum size of the prophage-like block pattern. Default is NA
minContigLength	The minimum contig size (in bp) to perform pattern-matching on. Must be at least 20,000 bp. Default is 30,000 bp.
minSlope	The minimum slope value to test for sloping patterns
verbose	TRUE or FALSE. Print progress messages to console. Default is TRUE.

### Value

List containing three objects.

---

patternMatchSize      *Pattern-match size calculator*

---

**Description**

Calculate the size (bp) of the matching region for Prophage-like and Sloping patterns

**Usage**

```
patternMatchSize(classifSumm, classifList, windowSize, verbose)
```

**Arguments**

classifSumm	Classification summary table
classifList	A list containing pattern match information associated with all contig classifications
windowSize	The window size used to re-average read coverage pileups
verbose	TRUE or FALSE. Print progress messages to console. Default is TRUE.

**Value**

dataframe

---

pileupFormatter      *Correctly formats pileup files.*

---

**Description**

Places columns in correct order and renames columns. Cleans the contig labels to remove excess information after whitespace.

**Usage**

```
pileupFormatter(pileup)
```

**Arguments**

pileup	A table containing contig names, read coverages averaged over 100 bp windows, and contig positions
--------	--

**Value**

dataframe

---

plotTrIdentResults      *Plot read coverage graphs of contigs classified as Prophage-like, Sloping, or HighCovNoPattern*

---

### Description

Plot the read coverages of a contig and its associated pattern-match for Prophage-like, Sloping and HighCovNoPattern classifications. Returns a list of ggplot objects.

### Usage

```
plotTrIdentResults(
  VLPpileup,
  WCPileup,
  TrIdentResults,
  matchScoreFilter,
  saveFilesTo
)
```

### Arguments

VLPpileup	VLP-fraction pileup file generated by mapping sequencing reads from a sample's ultra-purified VLP-fraction mapped to the sample's whole-community metagenome assembly. The pileup file MUST have the following format: * V1: Contig accession * V2: Mapped read coverage values averaged over 100 bp windows * V3: Starting position (bp) of each 100 bp window. Restarts from 0 at the start of each new contig. * V4: Starting position (bp) of each 100 bp window. Does NOT restart at the start of each new contig.
WCPileup	A whole-community pileup file generated by mapping sequencing reads from a sample's whole-community mapped to the sample's whole-community metagenome assembly. The pileup file MUST have the following format: * V1: Contig accession * V2: Mapped read coverage values averaged over 100 bp windows * V3: Starting position (bp) of each 100 bp window. Restarts from 0 at the start of each new contig. * V4: Starting position (bp) of each 100 bp window. Does NOT restart at the start of each new contig.
TrIdentResults	Output from 'TrIdentClassifier()'.
matchScoreFilter	Optional, Filter plots using the normalized pattern match-scores. A suggested filtering threshold is provided by 'TrIdentClassifier()' if 'suggFiltThresh=TRUE'.
saveFilesTo	Optional, Provide a path to the directory you wish to save output to. A folder will be made within the provided directory to store results.

### Value

Large list containing ggplot objects



**Examples**

```
data("VLPFractionSamplePileup")
data("WholeCommunitySamplePileup")
data("TrIdentSampleOutput")

patternMatches <- plotTrIdentResults(
  VLPpiledup = VLPFractionSamplePileup,
  WCPiledup = WholeCommunitySamplePileup,
  TrIdentResults = TrIdentSampleOutput
)
```

---

prophageLikeBorders     *Prophage-like border finder*

---

**Description**

Find borders of Prophage-like patterns with more specificity than pattern-matching using 100 bp window pileups and sliding standard deviation technique.

**Usage**

```
prophageLikeBorders(viralSubset, classificationPatterns, i, windowSize)
```

**Arguments**

viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
classificationPatterns	The pattern match information associated with each contig classified as Prophage-like, Sloping, or HighCovNoPattern
i	The index for the contig currently being assessed
windowSize	The window size used to re-average read coverage pileups

**Value**

List

---

prophageLikeElevation *Determine Prophage-like read coverage elevation in whole-community*

---

### Description

Determines whether a detected Prophage-like genetic element has read coverage in the whole-community that is either elevated or depressed compared to the average read coverage of the non-prophage region.

### Usage

```
prophageLikeElevation(
  classifSummTable,
  prophageLikeClassifList,
  VLPpileup,
  WCPileup,
  windowSize,
  verbose
)
```

### Arguments

classifSummTable	Classification summary table
prophageLikeClassifList	A list containing pattern match information associated with all contigs classified as Prophage-like.
VLPpileup	A table containing contig names, coverages averaged over 100 bp windows, and contig positions associated with mapping VLP-fraction reads to whole-community contigs
WCPileup	A table containing contig names, coverages averaged over 100 bp windows, and contig positions associated with mapping whole-community reads to whole-community contigs
windowSize	The window size used to re-average read coverage pileups
verbose	TRUE or FALSE. Print progress messages to console. Default is TRUE.

### Value

dataframe

---

prophageLikeZoom	<i>Prophage-like pattern zoom</i>
------------------	-----------------------------------

---

**Description**

'Zoom-in' on (aka subset) desired region surrounding block pattern.

**Usage**

```
prophageLikeZoom(viralSubset, classificationPatterns, i, zoom, windowSize)
```

**Arguments**

viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
classificationPatterns	The pattern match information associated with each contig classified as Prophage-like, sloping, or HighCovNoPattern
i	The index for the contig currently being assessed
zoom	The number of rows outside the start and stop positions of the block pattern to zoom-in on
windowSize	The window size used to re-average read coverage pileups

**Value**

Dataframe

---

resultsHisto	<i>Create histogram of normalized pattern-match scores</i>
--------------	--

---

**Description**

Plots a histogram of normalized match scores for all Prophage-like, Sloping and HighCovNoPattern classifications and colors the plot based on the classifications. A suggested filtering threshold is provided for filtering results based on the quality of the pattern-match.

**Usage**

```
resultsHisto(summaryList, suggFiltThresh)
```

**Arguments**

summaryList	Classification summary table filtered to only include contigs with Prophage-like, Sloping and HighCovNoPattern classifications
suggFiltThresh	TRUE or FALSE, Suggest a filtering threshold on the output pattern-match score histogram. Default is FALSE.

**Value**

ggplot object

---

slopeSumm	<i>Summarize slopes for sloping classifications</i>
-----------	---

---

**Description**

Add slope information for sloping classifications to summary table

**Usage**

```
slopeSumm(classifSumm, slopingClassifList, windowSize)
```

**Arguments**

classifSumm	Classification summary table
slopingClassifList	A list containing pattern match information associated with all contigs classified as sloping.
windowSize	The window size used to re-average read coverage pileups

**Value**

dataframe

---

slopeTranslator	<i>Sloping pattern translator</i>
-----------------	-----------------------------------

---

**Description**

Translates a sloping pattern containing the initial jump-up in read coverage across a contig. Translate the pattern 1000 bp at a time. Stop translating when the pattern left on the contig reaches 20,000 bp.

**Usage**

```
slopeTranslator(
  viralSubset,
  bestMatchInfo,
  windowSize,
  slopeChange,
  leftOrRight
)
```

**Arguments**

viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
bestMatchInfo	The pattern-match information associated with the current best pattern match.
windowSize	The window size used to re-average read coverage pileups
slopeChange	A list containing pattern vector, slope value, and value of slope bottom
leftOrRight	The direction of the sloping pattern. Either "Left" for left to right (neg) slopes or "Right" for right to left (pos) slopes.

**Value**

List

---

slopeWithStart	<i>Sloping pattern with an initial jump-up in read coverage</i>
----------------	---

---

**Description**

Build, translate, and change slope of sloping pattern with slope start

**Usage**

```
slopeWithStart(viralSubset, windowSize, minSlope)
```

**Arguments**

viralSubset	A subset of the read coverage pileup that pertains only to the contig currently being assessed
windowSize	The window size used to re-average read coverage pileups
minSlope	The minimum slope value to test for sloping patterns

**Value**

List containing two objects

---

specializedTransductionID

*Identify potential specialized transduction events on contigs classified as Prophage-like*

---

### Description

Search contigs classified as Prophage-like for dense read coverage outside of the pattern-match borders that may indicate specialized transduction. Returns a list with the first object containing a summary table and the second object containing a list of plots of with associated specialized transduction search results. If the plot is green, it has been identified as having potential specialized transduction.

### Usage

```
specializedTransductionID(
  VLPpfileup,
  TrIdentResults,
  specificContig,
  noReadCov = 500,
  specTransLength = 2000,
  matchScoreFilter,
  logScale = FALSE,
  verbose = TRUE,
  SaveFilesTo
)
```

### Arguments

VLPpfileup	VLP-fraction pileup file generated by mapping sequencing reads from a sample's ultra-purified VLP-fraction mapped to the sample's whole-community metagenome assembly. The pileup file MUST have the following format: * V1: Contig accession * V2: Mapped read coverage values averaged over 100 bp windows * V3: Starting position (bp) of each 100 bp window. Restarts from 0 at the start of each new contig. * V4: Starting position (bp) of each 100 bp window. Does NOT restart at the start of each new contig.
TrIdentResults	Output from 'TrIdentClassifier()'
specificContig	Optional, Search a specific contig classified as Prophage-like ("NODE_1").
noReadCov	Number of basepairs of zero read coverage encountered before specialized transduction searching stops. Default is 500. Must be at least 100.
specTransLength	Number of basepairs of non-zero read coverage needed for specialized transduction to be considered. Default is 2000. Must be at least 100.
matchScoreFilter	Optional, Filter plots using the normalized pattern match-scores. A suggested filtering threshold is provided by 'TrIdentClassifier()' if 'suggFiltThresh=TRUE'.

logScale	TRUE or FALSE, display VLP-fraction read coverage in log10 scale. Default is FALSE.
verbose	TRUE or FALSE. Print progress messages to console. Default is TRUE.
SaveFilesTo	Provide a path to the directory you wish to save output to. 'specializedTransductionID()' will make a folder within the provided directory to store results.

**Value**

Large list containing two objects

**Examples**

```
data("VLPfractionSamplePileup")
data("TrIdentSampleOutput")

specTransduction <- specializedTransductionID(
  VLPpileup = VLPfractionSamplePileup,
  TrIdentResults = TrIdentSampleOutput
)

specTransductionNODE62 <- specializedTransductionID(
  VLPpileup = VLPfractionSamplePileup,
  TrIdentResults = TrIdentSampleOutput,
  specificContig = "NODE_62"
)
```

---

specTransductionPlot *Specialized transduction plot*

---

**Description**

Plot search results of 'specializedTransductionID()'

**Usage**

```
specTransductionPlot(
  viralSubsetZoom,
  startPosBp,
  endPosBp,
  SpecTransLeft,
  specTransRight,
  contigName,
  classifPatternMatches,
  i,
  specTransSumm,
  logScale,
  classifSumm
)
```

**Arguments**

viralSubsetZoom	contig subset surrounding Prophage-like pattern-match
startPosBp	Left border position
endPosBp	Right border position
SpecTransLeft	End position of spec transduction on left border
specTransRight	End position of spec transduction on right border
contigName	The reference name of the contig currently being assessed (i.e "NODE_1")
classifPatternMatches	The pattern match information associated with each contig classified as prophage-like, sloping, or HighCovNoPattern
i	The index for the contig currently being assessed
specTransSumm	Results for spec transduction search
logScale	If TRUE, coverage is plotted in log10. If FALSE, raw coverage values are plotted. Default is FALSE.
classifSumm	The summary information associated with each contig classified as Prophage-like, Sloping, or HighCovNoPattern

**Value**

ggplot object

---

specTransductionSearch

*Specialized transduction search and plot*

---

**Description**

Search contigs classified as prophage-like for potential specialized transduction and return the plot visualizing the search results.

**Usage**

```
specTransductionSearch(
  contigName,
  VLPpiledup,
  classifPatternMatches,
  classifSumm,
  windowSize,
  i,
  noReadCov,
  specTransLength,
  logScale
)
```



**Arguments**

contigName	The reference name of the contig currently being assessed (i.e "NODE_1")
VLPpiledup	A table containing contig names, coverages averaged over 100 bp windows, and contig positions associated with mapping VLP-fraction reads to whole-community contigs
classifPatternMatches	The pattern match information associated with each contig classified as prophage-like, sloping, or HighCovNoPattern
classifSumm	The summary information associated with each contig classified as Prophage-like, Sloping, or HighCovNoPattern
windowSize	The window size used to re-average read coverage pileups
i	The index for the contig currently being assessed
noReadCov	How many bp of no read coverage are encountered before searching stops? Default is 500.
specTransLength	How many bp of read coverage to look for outside of prophage borders? Default is 2000.
logScale	If TRUE, coverage is plotted in log10. If FALSE, raw coverage values are plotted. Default is FALSE.

**Value**

List containing two objects

---

TrIdentClassifier	<i>Classify contigs as Prophage-like, Sloping, HighCovNoPattern, and NoPattern</i>
-------------------	--

---

**Description**

Performs all the pattern-matching and summarizes the results into a list. The first item in the list is a table consisting of the summary information of all the contigs that passed through pattern-matching (i.e were not filtered out). The second item in the list is a table consisting of the summary information of all contigs that were classified via pattern-matching. The third item in the list contains the pattern-match information associated with each contig in the previous table. The fourth object in the list is a table containing the contigs that were filtered out prior to pattern-matching. The fifth item is the windowSize used for the search.

**Usage**

```
TrIdentClassifier(
  VLPpiledup,
  WCPiledup,
  windowSize = 1000,
  minBlockSize = 10000,
```

```

    maxBlockSize = Inf,
    minContigLength = 30000,
    minSlope = 0.001,
    suggFiltThresh = FALSE,
    verbose = TRUE,
    SaveFilesTo
)

```

## Arguments

VLPpileup	VLP-fraction pileup file generated by mapping sequencing reads from a sample's ultra-purified VLP-fraction mapped to the sample's whole-community metagenome assembly. The pileup file MUST have the following format: * V1: Contig accession * V2: Mapped read coverage values averaged over 100 bp windows * V3: Starting position (bp) of each 100 bp window. Restarts from 0 at the start of each new contig. * V4: Starting position (bp) of each 100 bp window. Does NOT restart at the start of each new contig.
WCPileup	A whole-community pileup file generated by mapping sequencing reads from a sample's whole-community mapped to the sample's whole-community metagenome assembly. The pileup file MUST have the following format: * V1: Contig accession * V2: Mapped read coverage values averaged over 100 bp windows * V3: Starting position (bp) of each 100 bp window. Restarts from 0 at the start of each new contig. * V4: Starting position (bp) of each 100 bp window. Does NOT restart at the start of each new contig.
windowSize	The number of basepairs to average read coverage values over. Options are 100, 200, 500, 1000 ONLY. Default is 1000.
minBlockSize	The minimum size (in bp) of the Prophage-like block pattern. Default is 10000. Must be at least 1000.
maxBlockSize	The maximum size (in bp) of the Prophage-like block pattern. Default is NA (no maximum).
minContigLength	The minimum contig size (in bp) to perform pattern-matching on. Must be at least 25000. Default is 30000.
minSlope	The minimum slope value to test for sloping patterns. Default is 0.001 (i.e minimum change of 10x read coverage over 100,000 bp).
suggFiltThresh	TRUE or FALSE, Suggest a filtering threshold for TrIdent classifications based on the normalized pattern-match scores. Default is FALSE.
verbose	TRUE or FALSE. Print progress messages to console. Default is TRUE.
SaveFilesTo	Optional, Provide a path to the directory you wish to save output to. A folder will be made within the provided directory to store results.

## Value

Large list containing 5 objects

## Examples

```
data("VLPFractionSamplePileup")
data("WholeCommunitySamplePileup")

TrIdent_results <- TrIdentClassifier(
  VLPpileup = VLPFractionSamplePileup,
  WCPileup = WholeCommunitySamplePileup
)
```

---

TrIdentSampleOutput    *TrIdentSampleOutput*

---

## Description

The TrIdentClassifier output from the VLPFractionSamplePileup and WholeCommunitySamplePileup files run with default parameters Report...

## Usage

```
data('TrIdentSampleOutput')
```

## Format

## 'TrIdentSampleOutput' A list with 6 objects:

**SummaryTable** A dataframe containing classifications for all contigs that were processed with pattern-matching

**CleanedSummaryTable** SummaryTable dataframe filtered to remove contigs that recieved a 'None' classification

**PatternMatchInfo** A list of lists containing pattern-match information for each classified contig

**FilteredOutContigTable** A dataframe containing names of contigs that were filtered out prior to pattern-matching

**windowSize** windowSize used in TrIdentClassifier function (1000)

**ResultHistogram** a histogram displaying the overall abundance and quality of pattern-matches in addition to the composition of classifications. The displayed pattern-match scores are normalized by dividing each score by its associated contig length. The scores are normalized to visualize the overall quality of pattern-matching for the entire dataset.

## Details

A list object produced by the TrIdentClassifier function run on the VLPFractionSamplePileup and WholeCommunitySamplePileup files run with default parameters

---

VLPFractionSamplePileup

*VLP-Fraction of Sample Dataset*

---

### Description

A subset of contigs from the raw VLP-fraction read coverage pileup file generated from BMap's pileup.sh. Report...

### Usage

```
data('VLPFractionSamplePileup')
```

### Format

```
## 'VLPFractionSamplePileup' A data frame with 10,805 rows and 4 columns:
```

**V1** Contig accession

**V2** Mapped read coverage averaged over a 100 bp window size

**V3** Starting position (bp) of each 100 bp window. Restarts from 0 at the start of each new contig.

**V4** Starting position (bp) of each 100 bp window. Does NOT restart at the start of each new contig.

### Details

This dataset represents one half of a complete transductions dataset which is comprised of two parts—a whole-community fraction and a viral-like particle (VLP)-fraction. This dataset represents the VLP fraction and was generated by purifying VLPs from a conventional mouse fecal homogenate using CsCl density gradient ultracentrifugation. The VLP-fraction extracted DNA was sequenced with Illumina (paired-end mode, 150 bp reads) The sequencing reads were mapped to the associated whole-community assembly using BMap. The bmap.sh bincov parameter with covbinsize=100 was used to create a pileup file with 100 bp windows. A subset of 10 contigs from the pileup file were selected for this sample dataset. The contigs were chosen because their associated read coverage patterns in the VLP-fraction exemplify TrIdent's pattern-matching functionality across classifications: NODE\_617: Prophage-like, active/abundant, with spec transduction NODE\_135: Prophage-like, off one side of contig, no spec transduction NODE\_352: Sloping, left to right slope NODE\_1088: Sloping, right to left slope NODE\_2060: Sloping, right to left slope with start NODE\_1401: None, no pattern match NODE\_62: Prophage-like, with spec transduction NODE\_368: Prophage-like, not homogeneously integrated/present, no spec transduction NODE\_560: HighCovNoPattern NODE\_1165: None, filtered out To access the sequencing data used to generate this pileup file and for additional details on the assembly and mapping parameters, refer to the reference below: Reference: Kleiner, M., Bushnell, B., Sanderson, K.E. et al. Transductions: sequencing-based detection and analysis of transduced DNA in pure cultures and microbial communities. *Microbiome* 8, 158 (2020). <https://doi.org/10.1186/s40168-020-00935-5>

### Source

<<https://microbiomejournal.biomedcentral.com/articles/10.1186/s40168-020-00935-5>>

---

VLptoWCRatioCalc      *VLP-fraction:whole-community read coverage ratio calculator*

---

### Description

Calculate the VLP-fraction:whole-community read coverage ratio for every contig using the median read coverage values. If the ratio is greater than 2 (i.e VLP-fraction read coverage is, on average, at least double the whole-community read coverage), then the contig is classified as HighCovNoPattern

### Usage

```
VLptoWCRatioCalc(classifSumm, WCpileup, VLPpileup)
```

### Arguments

classifSumm	Classification summary table
WCpileup	A table containing contig names, coverages averaged over 100 bp windows, and contig positions associated with mapping whole-community reads to whole-community contigs
VLPpileup	A table containing contig names, coverages averaged over 100 bp windows, and contig positions associated with mapping VLP-fraction reads to whole-community contigs

### Value

dataframe

---

WholeCommunitySamplePileup  
*Whole-Community Fraction of Sample Dataset*

---

### Description

A subset of contigs from the raw whole-community fraction read coverage pileup file generated during read mapping. Report...

### Usage

```
data('WholeCommunitySamplePileup')
```

**Format**

## 'WholeCommunitySamplePileup' A data frame with 10,805 rows and 4 columns:

**V1** Contig accession

**V2** Mapped read coverage averaged over a 100 bp window size

**V3** Starting position (bp) of each 100 bp window. Restarts from 0 at the start of each new contig.

**V4** Starting position (bp) of each 100 bp window. Does NOT restart at the start of each new contig.

**Details**

This dataset represents one half of a complete transductions dataset which is comprised of two parts—a whole-community fraction and a viral-like particle (VLP)-fraction. This dataset represents the whole-community fraction and was generated from a conventional mouse fecal homogenate. The whole-community extracted DNA was sequenced with Illumina (paired-end mode, 150 bp reads) after which the metagenome was assembled. The sequencing reads were mapped to the assembled contigs using BMAP. The bmap.sh bincov parameter with covbinsize=100 was used to create a pileup file with 100 bp windows. A subset of 10 contigs from the pileup file were selected for this sample dataset. The contigs were chosen because their associated read coverage patterns in the VLP-fraction exemplify TrIdent's pattern-matching and characterization functionality across classifications: NODE\_617: Prophage-like, active/abundant, with spec transduction NODE\_135: Prophage-like, off one side of contig, no spec transduction NODE\_352: Sloping, left to right slope NODE\_1088: Sloping, right to left slope NODE\_2060: Sloping, right to left slope with start NODE\_1401: None, no pattern match NODE\_62: Prophage-like, with spec transduction NODE\_368: Prophage-like, not homogeneously integrated/present, no spec transduction NODE\_560: HighCovNoPattern NODE\_1165: None, filtered out To access the sequencing data used to generate this pileup file and for additional details on the assembly and mapping parameters, refer to the reference below: Reference: Kleiner, M., Bushnell, B., Sanderson, K.E. et al. Transductions: sequencing-based detection and analysis of transduced DNA in pure cultures and microbial communities. *Microbiome* 8, 158 (2020). <https://doi.org/10.1186/s40168-020-00935-5>

**Source**

<<https://microbiomejournal.biomedcentral.com/articles/10.1186/s40168-020-00935-5>>

---

zeroCountSearch

*Counts zero values to the left and right of prophage-like borders*

---

**Description**

Checks to see at which point the number of consecutive zero values to the left and right of the prophage-like pattern match borders equals the noReadCov parameter

**Usage**

zeroCountSearch(startOrEnd, viralSubsetZoom, startOrEndPosRow, noReadCov)

**Arguments**

- `startOrEnd`      searching the start (left side) or end (right side) of the prophage-like pattern-match
- `viralSubsetZoom`      viralSubset dataframe subsetting to 50,000 bp outside the pattern match borders
- `startOrEndPosRow`      The row index of the start or end position of the prophage-like pattern match
- `noReadCov`      How many bp of no read coverage are encountered before specialized transduction searching stops? Default is 500.

**Value**

List

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