Package 'awst'

January 27, 2025

Title Asymmetric Within-Sample Transformation

Version 1.14.0

Description We propose an Asymmetric Within-Sample Transformation (AWST) to regularize RNA-seq read counts and reduce the effect of noise on the classification of samples. AWST comprises two main steps: standardization and smoothing. These steps transform gene expression data to reduce the noise of the lowly expressed features, which suffer from background effects and low signal-to-noise ratio, and the influence of the highly expressed features, which may be the result of amplification bias and other experimental artifacts.

```
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```

Encoding UTF-8

RoxygenNote 7.1.1

URL https://github.com/drisso/awst

BugReports https://github.com/drisso/awst/issues

Imports stats, methods, SummarizedExperiment

Suggests airway, ggplot2, testthat, EDASeq, knitr, BiocStyle, RefManageR, sessioninfo, rmarkdown

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Author Davide Risso [aut, cre, cph] (https://orcid.org/0000-0001-8508-5012), Stefano Pagnotta [aut, cph] (https://orcid.org/0000-0002-8298-9777)

Maintainer Davide Risso <risso.davide@gmail.com>

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awst

Asymmetric Within-Sample Transformation

Description

This function implements the asymmetric within-sample transformation described in Risso and Pagnotta (2019). The function includes two steps: a standardization step and a asymmetric winsorization step. See details.

Usage

```
## S4 method for signature 'matrix'
awst(x, poscount = FALSE, full_quantile = FALSE, sigma0 = 0.075, lambda = 13)
## S4 method for signature 'SummarizedExperiment'
awst(
    x,
    poscount = FALSE,
    full_quantile = FALSE,
    sigma0 = 0.075,
    lambda = 13,
    expr_values = "counts",
    name = "awst"
)
```

Arguments

| X | a matrix of (possibly normalized) RNA-seq read counts or a 'SummarizedExperiment'. |
|---------------|---------------------------------------------------------------------------------------------------------------------------------------------------|
| poscount | a logical value indicating whether positive counts only should be used for the standardization step. |
| full_quantile | a logical value indicating whether the data have been normalized with the full-quantile normalization. In this case, computations can be sped up. |
| sigma0 | a multiplicative constant to be applied to the smoothing function. |
| lambda | a parameter that controls the growth rate of the smoothing function. |
| expr_values | integer scalar or string indicating the assay that contains the matrix to use as input. |
| name | string specifying the name of the assay to be used to store the results of the transformation. |

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Details

The standardization step is based on a log-normal distribution of the high-intensity genes. Optionally, only positive counts can be used in this step (this option is especially useful for single-cell data). The winsorization step is controlled by two parameters, sigma0 and lambda, which control the growth rate of the winsorization function.

Value

if 'x' is a matrix, it returns a matrix of transformed values, with genes in rows and samples in column. If 'x' is a 'SummarizedExperiment', it returns a 'SummarizedExperiment' with the transformed value in the 'name' slot.

Methods (by class)

- matrix: the input is a matrix of (possibly normalized) counts
- SummarizedExperiment: the input is a SummarizedExperiment with (possibly normalized) counts in one of its assays.

References

Risso and Pagnotta (2019). Within-sample standardization and asymmetric winsorization lead to accurate classification of RNA-seq expression profiles. Manuscript in preparation.

Examples

```
x <- matrix(data = rpois(100, lambda=5), ncol=10, nrow=10)
awst(x)</pre>
```

gene_filter

Gene filtering based on heterogeneity

Description

This function filters out genes that show a low heterogeneity, as measured by Shannon's entropy.

Usage

```
## S4 method for signature 'matrix'
gene_filter(
    X,
    from = min(x, na.rm = TRUE),
    to = max(x, na.rm = TRUE),
    nBins = 20,
    heterogeneity_threshold = 0.1
)

## S4 method for signature 'SummarizedExperiment'
gene_filter(
    X,
    from = min(assay(x, awst_values), na.rm = TRUE),
```

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```
to = max(assay(x, awst_values), na.rm = TRUE),
nBins = 20,
heterogeneity_threshold = 0.1,
awst_values = "awst"
)
```

Arguments

values to use as input.

Details

Shannon's entropy is computed on the categorized data after AWST transformation. Those genes that show a lower entropy than the predefined threshold are deemed to carry too low information to be useful for the classification of the samples, and are hence removed.

Value

if 'x' is a matrix, it returns a filtered matrix. If 'x' is a 'SummarizedExperiment', it returns a filtered 'SummarizedExperiment'

Methods (by class)

- matrix: the input is a matrix of awst-transformed values.
- SummarizedExperiment: the input is a SummarizedExperiment with awst-transformed values in one of its assays.

References

Risso and Pagnotta (2019). Within-sample standardization and asymmetric winsorization lead to accurate classification of RNA-seq expression profiles. Manuscript in preparation.

Examples

```
set.seed(222)
x <- matrix(rpois(75, lambda=5), ncol=5, nrow=15)
a <- awst(x)
gene_filter(a)</pre>
```

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