SOFTWARE TOOL ARTICLE

**REVISED** TreeSummarizedExperiment: a S4 class for data with hierarchical structure [version 2; peer review: 3 approved]

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**Abstract**

Data organized into hierarchical structures (e.g., phylogenies or cell types) arises in several biological fields. It is therefore of interest to have data containers that store the hierarchical structure together with the biological profile data, and provide functions to easily access or manipulate data at different resolutions. Here, we present TreeSummarizedExperiment, a R/S4 class that extends the commonly used SingleCellExperiment class by incorporating tree representations of rows and/or columns (represented by objects of the phylo class). It follows the convention of the SummarizedExperiment class, while providing links between the assays and the nodes of a tree to allow data manipulation at arbitrary levels of the tree. The package is designed to be extensible, allowing new functions on the tree (phylo) to be contributed. As the work is based on the SingleCellExperiment class and the phylo class, both of which are popular classes used in many R packages, it is expected to be able to interact seamlessly with many other tools.

**Keywords**

SummarizedExperiment, tree, microbiome, hierarchical structure

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**Open Peer Review**

**Reviewer Status** ✔ ✔ ✔

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Introduction

Biological data arranged into a hierarchy occurs in several fields. A notable example is in microbial survey studies, where the microbiome is profiled with amplicon sequencing or whole genome shotgun sequencing, and microbial taxa are organized as a tree according to their similarities in the genomic sequence or the evolutionary history. Also, a tree might be used in single cell cytometry or RNA-seq data, with nodes representing cell sub-populations at different granularities. Currently, phyloseq and SingleCellExperiment are popular classes used in the analysis of microbial data and single cell data, respectively. The former supports the information pertaining to the hierarchical structure that is available as the phylo class (e.g., phylogenetic tree), and the latter is derived from the SummarizedExperiment class (defined in the SummarizedExperiment package), which is widely used as a standardized container across many Bioconductor packages. Since the data structures in these fields share similarities, we were motivated to develop an S4 class, TreeSummarizedExperiment, that not only leverages the facilities from the SummarizedExperiment class, but also bridges the functionality from the phylo class, which is available from the ape package and has been imported in more than 200 R packages.

We define TreeSummarizedExperiment by extending the SingleCellExperiment class, so that it is a member of the SummarizedExperiment family, and thus benefits from the comprehensive Bioconductor ecosystem (e.g., iSEE, SEtools, and ggbiom). At the same time, all slots of the phyloseq class have their corresponding slots in the TreeSummarizedExperiment class, which enables convenient conversion between these classes. Furthermore, we allow the link between profile data and nodes of the tree, including leaves and internal nodes, which is useful for algorithms in the downstream analysis that need to access internal nodes of the tree (e.g., treeclimbR).

Overall, the class TreeSummarizedExperiment is provided as a standalone R package, analogous to SummarizedExperiment and SingleCellExperiment. Thus, it is convenient for R package developers to import it and build downstream data analyses or visualizations on it. Also, it is flexible to combine with R packages that are linked to the SummarizedExperiment family or the phylo tree class, which enables R package users to explore data with the support of other tools.

Methods

Implementation

The structure of TreeSummarizedExperiment. The structure of the TreeSummarizedExperiment class is shown in Figure 1.

Compared to the SingleCellExperiment objects, TreeSummarizedExperiment has five additional slots:

- rowTree: the hierarchical structure on the rows of the assays.
- rowLinks: the link information between rows of the assays and the rowTree.
- colTree: the hierarchical structure on the columns of the assays.
- colLinks: the link information between columns of the assays and the colTree.
- referenceSeq (optional): the sequence information for rows of the assays.

The rowTree and/or colTree can be left empty (NULL) if no trees are available; in this case, the rowLinks and/or colLinks are also set to NULL. The referenceSeq is an optional slot to store the sequence data of...
features either as DNAStringSet or DNAStringSetList. All other TreeSummarizedExperiment slots are inherited from SingleCellExperiment.

The rowTree and colTree slots require the tree to be an object of the phylo class. If a tree is available in an alternative format, it can often be converted to a phylo object using dedicated R packages (e.g., treeio).

Functions in the TreeSummarizedExperiment package fall in two main categories: operations on the TreeSummarizedExperiment object or operations on the tree (phylo) objects. The former includes constructors and accessors, and the latter serves as “components” to be assembled as accessors or functions that manipulate the TreeSummarizedExperiment object. Given that more than 200 R packages make use of the phylo class, there are many resources (e.g., ape) for users to manipulate the small “pieces” in addition to those provided in TreeSummarizedExperiment.

The toy datasets as the example data

We generate a toy dataset that has observations of 6 entities collected from 4 samples as an example to show how to construct a TreeSummarizedExperiment object.

```r
library(TreeSummarizedExperiment)

# assays data (typically, representing observed data from an experiment)
assay_data <- rbind(rep(0, 4), matrix(1:20, nrow = 5))
colnames(assay_data) <- paste0("sample", 1:4)
rownames(assay_data) <- paste0("entity", seq_len(6))
assay_data
```

```
# sample1 sample2 sample3 sample4
# entity1  0     0     0     0
# entity2  1     6     11    16
# entity3  2     7     12    17
# entity4  3     8     13    18
```
The information of entities and samples are given in the `row_data` and `col_data`, respectively.

```r
# row data (feature annotations)
row_data <- data.frame(
  Kingdom = "A",
  Phylum = rep(c("B1", "B2"), c(2, 4)),
  Class = rep(c("C1", "C2", "C3"), each = 2),
  OTU = paste0("D", 1:6),
  row.names = rownames(assay_data),
  stringsAsFactors = FALSE)

# column data (sample annotations)
col_data <- data.frame(
  gg = c(1, 2, 3, 3),
  group = rep(LETTERS[1:2], each = 2),
  row.names = colnames(assay_data),
  stringsAsFactors = FALSE)
```

The hierarchical structure of the 6 entities and 4 samples are denoted as `row_tree` and `col_tree`, respectively. The two trees are phylo objects randomly created with rtree from the package ape. Note that the row tree has 5 rather than 6 leaves; this is used later to show that multiple rows in the assays are allowed to map to a single node in the tree.

```r
library(ape)

# The first toy tree
set.seed(12)
row_tree <- rtree(5)

# The second toy tree
set.seed(12)
col_tree <- rtree(4)

# change node labels
col_tree$tip.label <- colnames(assay_data)
col_tree$node.label <- c("All", "GroupA", "GroupB")
```

We visualize the tree using the package `ggtree` (v. 2.2.4)\(^1\). Here, the internal nodes of the `row_tree` have no labels as shown in Figure 2.
library(ggtree)
library(ggplot2)

# Visualize the row tree
ggtree(row_tree, size = 2, branch.length = "none") + 
    geom_text2(aes(label = node), color = "darkblue", 
                hjust = -0.5, vjust = 0.7, size = 4) + 
    geom_text2(aes(label = label), color = "darkorange", 
                hjust = -0.1, vjust = -0.7, size = 4)

The `col_tree` has labels for internal nodes as shown in Figure 3.

# Visualize the column tree
ggtree(col_tree, size = 2, branch.length = "none") + 
    geom_text2(aes(label = node), color = "darkblue", 
                hjust = -0.5, vjust = 0.7, size = 4) + 
    geom_text2(aes(label = label), color = "darkorange", 
                hjust = -0.1, vjust = -0.7, size = 4) + 
    ylim(c(0.8, 4.5)) + 
    xlim(c(0, 2.2))

Figure 2. The structure of the row tree. The node labels and the node numbers are in orange and blue text, respectively.

Figure 3. The structure of the column tree. The node labels and the node numbers are in orange and blue text, respectively.
The construction of `TreeSummarizedExperiment`

The `TreeSummarizedExperiment` class is used to store the toy data generated in the previous section: `assay_data`, `row_data`, `col_data`, `col_tree` and `row_tree`. To correctly store data, the link information between the rows (or columns) of `assay_data` and the nodes of the `row_tree` (or `col_tree`) can be provided via a character vector `rowNodeLab` (or `colNodeLab`), with length equal to the number of rows (or columns) of the assays; otherwise the row (or column) names are used. Tree data takes precedence to determine entities included during the creation of the `TreeSummarizedExperiment` object; columns and rows with labels that are not present among the node labels of the tree are removed with warnings. The link data between the assays tables and the tree data is automatically generated during the construction.

The row and column trees can be included simultaneously during the construction of a `TreeSummarizedExperiment` object. Here, the column names of `assay_data` can be found in the node labels of the column tree, which enables the link to be created between the column dimension of `assay_data` and the column tree `col_tree`. If the row names of `assay_data` are not in the node labels of `row_tree`, we would need to provide their corresponding node labels (`row_lab`) to `rowNodeLab` in the construction of the object. It is possible to map multiple rows or columns to a node, for example, the same leaf label is used for the first two rows in `row_lab`.

```r
# all column names could be found in the node labels of the column tree
all(colnames(assay_data) %in% c(col_tree$tip.label, col_tree$node.label))
```

```r
## [1] TRUE
```

```r
# provide the node labels in rowNodeLab
tip_lab <- row_tree$tip.label
row_lab <- tip_lab[c(1, 1:5)]
row Lab
```

```r
## [1] "t3" "t3" "t2" "t1" "t5" "t4"
```

```r
both_tse <- TreeSummarizedExperiment(assays = list(Count = assay_data),
                                      rowData = row_data,
                                      colData = col_data,
                                      rowTree = row_tree,
                                      rowNodeLab = row_lab,
                                      colTree = col_tree)
```

```r
both_tse
```

```r
## class: TreeSummarizedExperiment
## dim: 6 4
## metadata(0):
## assays(1): Count
## rownames(6): entity1 entity2 ... entity5 entity6
## rowData names(4): Kingdom Phylum Class OTU
## colnames(4): sample1 sample2 sample3 sample4
## colData names(2): gg group
## reducedDimNames(0):
## altExpNames(0):
## rowLinks: a LinkDataFrame (6 rows)
## rowTree: 1 phylotree(s) (5 leaves)
## colLinks: a LinkDataFrame (4 rows)
## colTree: 1 phylotree(s) (4 leaves)
```

When printed on screen, `TreeSummarizedExperiment` objects display information as the parent `SingleCell-Experiment` class followed by four additional lines for `rowLinks`, `rowTree`, `colLinks` and `colTree`. 
The accessor functions

Slots inherited from the `SummarizedExperiment` class can be accessed in the standard way (e.g., via accessors `assays()`, `rowData()`, `colData()` and `metadata()`). These functions are both getters and setters. To clarify, getters and setters are functions for users to retrieve and to overwrite data from the corresponding slots, respectively. Here, accessors for `TreeSummarizedExperiment` are both getters and setters unless specifically mentioned.

For new slots, we provide `rowTree` (and `colTree`) to access the row (column) trees, and `rowLinks` (and `colLinks`) as getters to retrieve the link information between `assays` and the row (column) tree. If the tree is not available, the corresponding link data is `NULL`.

```r
# access trees
rowTree(both_tse)
```

```r
## Phylogenetic tree with 5 tips and 4 internal nodes.
## Tip labels:
##  t3, t2, t1, t5, t4
## Rooted; includes branch lengths.

## access the link data
(r_link <- rowLinks(both_tse))
```

```r
## LinkDataFrame with 6 rows and 5 columns
##   nodeLab nodeLab_alias nodeNum isLeaf whichTree
##        <character>   <character> <integer> <logical> <character>
## entity1       t3       alias_1         1      TRUE   phylo
## entity2       t3       alias_1         1      TRUE   phylo
## entity3       t2       alias_2         2      TRUE   phylo
## entity4       t1       alias_3         3      TRUE   phylo
## entity5       t5       alias_4         4      TRUE   phylo
## entity6       t4       alias_5         5      TRUE   phylo
```

```r
(c_link <- colLinks(both_tse))
```

```r
## LinkDataFrame with 4 rows and 5 columns
##   nodeLab nodeLab_alias nodeNum isLeaf whichTree
##        <character>   <character> <integer> <logical> <character>
## sample1     sample1       alias_1         1      TRUE   phylo
## sample2     sample2       alias_2         2      TRUE   phylo
## sample3     sample3       alias_3         3      TRUE   phylo
## sample4     sample4       alias_4         4      TRUE   phylo
```

The link data objects are of the `LinkDataFrame` class, which extends the `DataFrame` class from `S4Vectors` with the restriction that it has five columns:

- `nodeLab`: the labels of nodes on the tree
• nodeLab_alias: the alias labels of nodes on the tree
• nodeNum: the numbers of nodes on the tree
• isLeaf: whether the node is a leaf node
• whichTree: which tree the row/col is linked to

The data in colLinks() is updated automatically with the change of colTree().

```r
# remove the column tree
colTree(both_tse) <- NULL
# the colLinks() is updated accordingly
colLinks(both_tse)

## NULL

# colTree works as a setter
colTree(both_tse) <- col_tree
colLinks(both_tse)

## LinkDataFrame with 4 rows and 5 columns
##   nodeLab   nodeNum nodeLab_alias    isLeaf   whichTree
## 1  sample1         1       alias_1      TRUE       phylo
## 2  sample2         2       alias_2      TRUE       phylo
## 3  sample3         3       alias_3      TRUE       phylo
## 4  sample4         4       alias_4      TRUE       phylo
```

The subsetting function

A `TreeSummarizedExperiment` object can be subset in two different ways: to subset by rows or columns, and `subsetByNode` to retrieve row and/or columns that correspond to nodes of a tree. As the numeric ID of a node changes with the cut of a phylo tree, to keep track of the original data, we do not prune the tree structure in the subsetting. Below, we can see that rowLinks and rowData are updated to have the same number of rows as assays.

```r
sub_tse <- both_tse[1:2, 1]
sub_tse
```

```r
## class: TreeSummarizedExperiment
## dim: 2 1
## metadata(0):
## assays(1): Count
## rownames(2): entity1 entity2
## rowData names(4): Kingdom Phylum Class OTU
## colnames(1): sample1
## colData names(2): gg group
## reducedDimNames(0):
## altExpNames(0):
## rowLinks: a LinkDataFrame (2 rows)
## rowTree: 1 phylo tree(s) (5 leaves)
## colLinks: a LinkDataFrame (1 rows)
## colTree: 1 phylo tree(s) (4 leaves)
```

# the row data
rowData(sub_tse)
To subset by nodes, we specify the node by its node label or node number. Here, *entity1* and *entity2* are both mapped to the same node *t3*, so both of them are retained.

```r
data_tse <- subsetByNode(x = both_tse, rowNode = "t3")
rowLinks(data_tse)
```

Subsetting simultaneously in both dimensions is also allowed.

```r
data_tse <- subsetByNode(x = both_tse, rowNode = "t3", colNode = c("sample1", "sample2"))
assays(data_tse)[[1]]
```

### Changing the tree

The current tree can be replaced by a new one using `changeTree`. Here, we don’t use `rowTree()` to do the replacement because the new tree has node labels that cannot match with row names of the `TreeSummarizedExperiment` object. If the hierarchical information is available as a `data.frame` with each column representing a taxonomic level (e.g., *row_data*), we provide `toTree` to convert it into a `phylo` object that is further visualized in Figure 4.
Figure 4. The structure of the taxonomic tree that is generated from the taxonomic table.

```r
# convert it to a phylo tree
taxa_tree <- toTree(data = taxa)

# Viz the new tree

ggtree(taxa_tree) +
  geom_text2(aes(label = node), color = "darkblue",
             hjust = -0.5, vjust = 0.7, size = 4) +
  geom_text2(aes(label = label), color = "darkorange",
             hjust = -0.1, vjust = -0.7, size = 4) +
  geom_point2()
```

If the nodes of the two trees have a different set of labels, a vector mapping the nodes of the new tree must be provided in `rowNodeLab`.

```r
taxa_tse <- changeTree(x = both_tse, rowTree = taxa_tree,
                       rowNodeLab = taxa["OTU"])
taxa_tse
```

```text
## class: TreeSummarizedExperiment
## dim: 6 4
## metadata(0):
## assays(1): Count
## rownames(6): entity1 entity2 ... entity5 entity6
## rowData names(4): Kingdom Phylum Class OTU
```
## colnames(4): sample1 sample2 sample3 sample4
## colData names(2): gg group
## reducedDimNames(0):
## altExpNames(0):
## rowLinks: a LinkDataFrame (6 rows)
## rowTree: 1 phylo tree(s) (6 leaves)
## colLinks: a LinkDataFrame (4 rows)
## colTree: 1 phylo tree(s) (4 leaves)

### rowLinks(taxa_tse)

### LinkDataFrame with 6 rows and 5 columns
###             nodeLab nodeLab_alias nodeNum isLeaf whichTree
###          <character>   <character> <integer> <logical> <character>
### entity1          D1       alias_1         1      TRUE       phylo
### entity2          D2       alias_2         2      TRUE       phylo
### entity3          D3       alias_3         3      TRUE       phylo
### entity4          D4       alias_4         4      TRUE       phylo
### entity5          D5       alias_5         5      TRUE       phylo
### entity6          D6       alias_6         6      TRUE       phylo

### Aggregation

Since it may be of interest to report or analyze observed data at multiple resolutions based on the provided tree(s), the `TreeSummarizedExperiment` package offers functionality to flexibly aggregate data to arbitrary levels of a tree.

### The column dimension.

Here, we demonstrate the aggregation functionality along the column dimension. The desired aggregation level is given in the `colLevel` argument, which can be specified using node labels (orange text in Figure 3) or node numbers (blue text in Figure 3). Furthermore, the summarization method used to aggregate multiple values can be specified via the argument `colFun`.

```r
# use node labels to specify colLevel
agg_col <- aggTSE(x = taxa_tse,
                   colLevel = c("GroupA", "GroupB"),
                   colFUN = sum)
# or use node numbers to specify colLevel
agg_col <- aggTSE(x = taxa_tse, colLevel = c(6, 7), colFUN = sum)
```

```r
assays(agg_col)[[1]]
```

```
##         alias_6 alias_7
## entity1       0      0
## entity2       7     27
## entity3      11      31
## entity4      15      35
## entity5      19      39
## entity6      23     43
```

The `rowData` does not change, but the `colData` is updated to reflect the metadata information that remains valid for the individual nodes after aggregation. For example, the column `group` has the A value for `GroupA` because the descendant nodes of `GroupA` all have the value A; whereas the column `gg` has the NA value for `GroupA` because the descendant nodes of `GroupA` have different values, (1 and 2).

```r
# before aggregation
colData(taxa_tse)
```
## DataFrame with 4 rows and 2 columns
##                gg       group
##         <numeric> <character>
## sample1         1           A
## sample2         2           A
## sample3         3           B
## sample4         3           B

# after aggregation
colData(agg_col)
## DataFrame with 2 rows and 2 columns
##                gg       group
##         <numeric> <character>
## alias_6        NA           A
## alias_7         3           B

The colLinks is also updated to link the new rows of assays tables to the corresponding nodes of the column tree (Figure 3).

# the link data is updated
colLinks(agg_col)
## LinkDataFrame with 2 rows and 5 columns
##             nodeLab nodeLab_alias   nodeNum    isLeaf   whichTree
##         <character>   <character> <integer> <logical> <character>
## alias_6      GroupA       alias_6         6     FALSE       phylo
## alias_7      GroupB       alias_7         7     FALSE       phylo

The row dimension. Similarly, we can aggregate rows to phyla by providing the names of the internal nodes that represent the phylum level (see taxa_one below).

# the phylum level
taxa <- c(taxa_tree$tip.label, taxa_tree$node.label)
(taxa_one <- taxa[startsWith(taxa, "Phylum:"))]

# aggregation
agg_taxa <- aggTSE(x = taxa_tse, rowLevel = taxa_one, rowFUN = sum)
assays(agg_taxa)[[1]]
##                  sample1 sample2 sample3 sample4
## alias_8          1       6      11      16
## alias_10        14      34      54      74

Users are nonetheless free to choose nodes from different taxonomic ranks for each final aggregated row. Note that it is not necessary to use all original rows during the aggregation process. Similarly, it is entirely possible for a row to contribute to multiple aggregated rows.

# A mixed level
taxa_mix <- c("Class:C3", "Phylum:B1")
agg_any <- aggTSE(x = taxa_tse, rowLevel = taxa_mix, rowFUN = sum)
rowData(agg_any)
## DataFrame with 2 rows and 4 columns
##                Kingdom Phylum     Class       OTU
##         <character> <character> <character> <logical>
## alias_12         A         B2        C3        NA
## alias_8          A         B1         C1        NA
Both dimensions. The aggregation on both dimensions could be performed in one step, in which case users can specify the order of aggregation; either rows first (rowFirst = TRUE) or columns first (rowFirst = FALSE). The aggregate functions for the row and the column dimension can be provided via rowFun and colFun, respectively. Additionally, parallel computation is enable by providing BPPARAM with a BiocParallelParam object.

```r
tag both <- aggTSE(x = both_tse,
                   rowLevel = 7:9, rowFun = sum,
                   colLevel = 6:7, colFun = mean,
                   rowFirst = FALSE)
```

As expected, we obtain a table with 3 rows representing the aggregated row nodes 7, 8, and 9 (rowLevel = 7:9) and 2 columns representing the aggregated column nodes 6 and 7 (colLevel = 6:7).

```r
assays(agg_both)[[1]]
```

```
#   alias_6 alias_7
# alias_7   8.0  28.0
# alias_8  19.5  49.5
# alias_9  12.0  32.0
```

Relations operating on the phylo object

Next, we highlight some functions to manipulate and/or to extract information from the phylo object. Further operations can be found in other packages, such as `ape`, `tidytree`. These functions are useful for users who wish to develop more functions for the `TreeSummarizedExperiment` class.

To show these functions, we use the tree shown in Figure 5.

```r
data("tinyTree")
ggtree(tinyTree, branch.length = "none") +
  geom_text2(aes(label = label), hjust = -0.1, size = 3) +
  geom_text2(aes(label = node), vjust = -0.8,
             hjust = -0.2, color = "blue", size = 3)
```

**Figure 5.** An example tree with node labels and numbers in black and blue texts, respectively.
Conversion of the node label and the node number  The translation between the node labels and node numbers can be achieved by the function `convertNode`.

```r
convertNode(tree = tinyTree, node = c(12, 1, 4))
## [1] "Node_12" "t2"      "t9"
```

```r
convertNode(tree = tinyTree, node = c("t4", "Node_18"))
##       t4 Node_18
##        5      18
```

Find the descendants  To get descendants that are at the leaf level, we could set the argument `only.leaf = TRUE` for the function `findDescendant`.

```r
# only the leaf nodes
findDescendant(tree = tinyTree, node = 17, only.leaf = TRUE)
## $Node_17
## [1] 4 5 6
```

When `only.leaf = FALSE`, all descendants are returned.

```r
# all descendant nodes
findDescendant(tree = tinyTree, node = 17, only.leaf = FALSE)
## $Node_17
## [1] 4 5 6 18
```

More functions.  We list some functions that might also be useful in Table 1. More functions are available in the package, and we encourage users to develop and contribute their own functions to the package.

<table>
<thead>
<tr>
<th>Functions</th>
<th>Goal</th>
</tr>
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<tbody>
<tr>
<td>printNode</td>
<td>print out the information of nodes</td>
</tr>
<tr>
<td>countNode</td>
<td>count the number of nodes</td>
</tr>
<tr>
<td>distNode</td>
<td>give the distance between a pair of nodes</td>
</tr>
<tr>
<td>matTree</td>
<td>list paths of a tree</td>
</tr>
<tr>
<td>findAncestor</td>
<td>find ancestor nodes</td>
</tr>
<tr>
<td>findChild</td>
<td>find child nodes</td>
</tr>
<tr>
<td>findSibling</td>
<td>find sibling nodes</td>
</tr>
<tr>
<td>shareNode</td>
<td>find the first node shared in the paths of nodes to the root</td>
</tr>
<tr>
<td>unionLeaf</td>
<td>find the union of descendant leaves</td>
</tr>
<tr>
<td>trackNode</td>
<td>track nodes by adding alias labels to a phylo object</td>
</tr>
<tr>
<td>isLeaf</td>
<td>test whether a node is a leaf node</td>
</tr>
<tr>
<td>showNode</td>
<td>print out nodes of a tree</td>
</tr>
<tr>
<td>addLabel</td>
<td>label nodes of a tree</td>
</tr>
<tr>
<td>joinNode</td>
<td>represent descendant nodes with their ancestor nodes</td>
</tr>
</tbody>
</table>
Custom functions for the *TreeSummarizedExperiment* class

Here, we show examples of how to write custom functions for *TreeSummarizedExperiment* objects. To extract data corresponding to specific leaves, we created a function `subsetByLeaf` by combining functions working on the `phylo` class (e.g., `convertNode`, `keep.tip`, `trackNode`) with the accessor function `subsetByNode`. Here, `convertNode` and `trackNode` are available in *TreeSummarizedExperiment*, and `keep.tip` is from the *ape* package. Since the numeric identifier of a node is changed after pruning a tree with `keep.tip`, `trackNode` is provided to track the node and further update links between the rectangular assay matrices and the new tree.

```r
# tse: a TreeSummarizedExperiment object
# rowLeaf: specific leaves
subsetByLeaf <- function(tse, rowLeaf) {
  # if rowLeaf is provided as node labels, convert them to node numbers
  if (is.character(rowLeaf)) {
    rowLeaf <- convertNode(tree = rowTree(tse), node = rowLeaf)
  }
  # subset data by leaves
  sse <- subsetByNode(tse, rowNode = rowLeaf)

  # update the row tree
  ## -------------- new tree: drop leaves ----------
  oldTree <- rowTree(sse)
  newTree <- ape::keep.tip(phy = oldTree, tip = rowLeaf)

  ## -------------- update the row tree ----------
  # track the tree
  track <- trackNode(oldTree)
  track <- ape::keep.tip(phy = track, tip = rowLeaf)

  # update the row tree:
  # 1. get the old alias label and update it to the new node label
  # 2. provide the new node label as rowNodeLab to update the row tree
  oldAlias <- rowLinks(sse)$nodeLab_alias
  newNode <- convertNode(tree = track, node = oldAlias)
  newLab <- convertNode(tree = newTree, node = newNode)

  changeTree(x = sse, rowTree = newTree, rowNodeLab = newLab)
}
```

The row tree is updated; after subsetting, it has only two leaves, t2 and t3.

```r
(both_sse <- subsetByLeaf(tse = both_tse, rowLeaf = c("t2", "t3")))
```

```r
## class: TreeSummarizedExperiment
## dim: 3 4
## metadata(0):
## assays(1): Count
## rownames(3): entity1 entity2 entity3
## rowData names(4): Kingdom Phylum Class OTU
## colnames(4): sample1 sample2 sample3 sample4
## colData names(2): gg group
## reducedDimNames(0):
## altExpNames(0):
## rowLinks: a LinkDataFrame (3 rows)
## rowTree: 1 phylo tree(s) (2 leaves)
## colLinks: a LinkDataFrame (4 rows)
## colTree: 1 phylo tree(s) (4 leaves)
```
## LinkDataFrame with 3 rows and 5 columns
##             nodeLab nodeLab_alias nodeNum isLeaf whichTree
##         <character>   <character> <integer> <logical> <character>
## entity1          t3       alias_1         1      TRUE       phylo
## entity2          t3       alias_1         1      TRUE       phylo
## entity3          t2       alias_2         2      TRUE       phylo

### Operation
The TreeSummarizedExperiment package can be installed by following the standard installation procedures of Bioconductor packages.

```r
# install BiocManager
if (!requireNamespace("BiocManager", quietly = TRUE))
  install.packages("BiocManager")

# install TreeSummarizedExperiment package
BiocManager::install("TreeSummarizedExperiment")
```

Minimum system requirements is R version 3.6 (or later) on a Mac, Windows or Linux system. The version of TreeSummarizedExperiment should be later than 1.99.10, which is available in Bioconductor 3.13.

### Use cases
**HMP 16S rRNA-seq data**
To demonstrate the functionality of TreeSummarizedExperiment, we use it to store and manipulate a microbial dataset. We further show exploratory graphics using the available functions designed for SummarizedExperiment objects in other packages (e.g., scater), or customized functions from popular visualization packages (e.g., ggplot2).  

```r
# Packages providing dataset
library(HMP16SData)

# Package to do parallel computation
library(BiocParallel)

# Packages to manipulate data extracted from TreeSummarizedExperiment
library(tidyr)
library(dplyr)

# Packages providing visualization
library(ggplot2)
library(scales)
library(ggtree)
library(scater)
library(cowplot)
```

The Human Microbiome Project (HMP) 16S rRNA sequencing data, v35, is downloaded using the R package HMP16SData, which contains survey data of samples collected at five major body sites in the variable regions 3–5. v35 is available as a SummarizedExperiment object via the ExperimentHub.

```r
(v35 <- V35())
```
The storage of HMP 16S rRNA-seq data

We store the phylogenetic tree as the `rowTree`. Links between nodes of the tree and rows of `assays` are automatically generated in the construction of the `TreeSummarizedExperiment` object, and are stored as `rowLinks`. Rows of the `assays` matrices that do not have a match to nodes of the tree are removed with warnings.

```r
(tse_phy <- TreeSummarizedExperiment(
  assays = assays(v35), 
  rowData = rowData(v35), 
  colData = colData(v35), 
  rowTree = metadata(v35)$phylogeneticTree, 
  metadata = metadata(v35)["experimentData"]
))

## Warning: 47 row(s) couldn’t be matched to the tree and are/is removed.
```

## Exploratory graphics

Here, we show `TreeSummarizedExperiment` working seamlessly with `SEtools` (v.1.2.0) to prepare data for the exploratory graphics. Since all operational taxonomic units (OTUs) in the sample belong to Bacteria in the `SUPERKINGDOM` level, we can calculate the sequencing depths by aggregating counts to the `SUPERKINGDOM` level. The resultant `TreeSummarizedExperiment` object `agg_total` is further converted into a data frame `df_total` with selected columns (`HMP_BODY_SITE` and `RUN_CENTER`) from the column data.
library(SEtools)

agg_total <- aggSE(x = tse_phy, by = "SUPERKINGDOM",
                   assayFun = sum)

# The assays data and selected columns of the row/col data are merged into a
# data frame
df_total <- meltSE(agg_total, genes = rownames(agg_total),
                   colDat.columns = c("HMP_BODY_SITE", "RUN_CENTER"))

head(df_total)

##    feature    sample          HMP_BODY_SITE RUN_CENTER Count
## 1  Bacteria 700013549  Gastrointestinal Tract        BCM  5295
## 2  Bacteria 700014386  Gastrointestinal Tract     BCM,BI 10811
## 3  Bacteria 700014403                      Oral     BCM,BI 12312
## 4  Bacteria 700014409                      Oral     BCM,BI 20355
## 5  Bacteria 700014412                      Oral     BCM,BI 14021
## 6  Bacteria 700014415                      Oral     BCM,BI 17157

To make harmonized figures with ggplot2 (v. 3.3.2)\textsuperscript{13}, we customized a theme to be applied to several
plots in this section.

# Customized the plot theme
prettify <- theme_bw(base_size = 10) + theme(
  panel.spacing = unit(0, "lines"),
  axis.text = element_text(color = "black"),
  axis.text.x = element_text(angle = 45, hjust = 1),
  axis.title = element_text(size = 8),
  legend.key.size = unit(3, "mm"),
  legend.spacing.x = unit(1, "mm"),
  plot.title = element_text(hjust = 0.5),
  legend.text = element_text(size = 8),
  legend.position = "bottom",
  strip.background = element_rect(colour = "black", fill = "gray90"),
  strip.text.x = element_text(color = "black", size = 10),
  strip.text.y = element_text(color = "black", size = 10))

From Figure 6, we note that more samples were collected from the oral site than other body sites.

# Figure: (the number of samples) VS (centers)
ggplot(df_total) +
  geom_bar(aes(RUN_CENTER, fill = HMP_BODY_SITE),
           position = position_dodge()) +
  labs(title = "The number of samples across centers", y = "") +
  scale_fill_brewer(palette = "Set1") +
  prettify

Figure 7 shows that the sequencing depth of each sample across different coordination centers are quite similar.
Within the coordination center, samples collected from Skin are more variable in the sequencing depth
than those from other body sites.

# Figure: (the sequencing depths) VS (centers)
ggplot(df_total) +
  geom_boxplot(aes(x = RUN_CENTER, y = Count, fill = HMP_BODY_SITE),
               position = position_dodge()) +
  labs(title = "The sequencing depths of samples") +
  scale_y_continuous(trans = log10_trans()) +
  scale_fill_brewer(palette = "Set1") +
  labs(y = "Total counts") +
  prettify
### Dimensionality Reduction

We visualize samples in reduced dimensions to see whether those from the same body site are similar to each other. Three dimensionality reduction techniques are available in the package `scater` (v. 1.16.2), including principal component analysis (PCA)\(^\text{15}\), t-distributed Stochastic Neighbor Embedding (t-SNE)\(^\text{16}\), and uniform manifold approximation and projection (UMAP)\(^\text{17}\). Since `TreesummarizedExperiment` extends the `SingleCellExperiment` class, functions from `scater`\(^\text{13}\) can be used directly. Here, we first apply PCA and t-SNE on data at the

---

**Figure 6.** The number of samples from different research centers. Samples collected at different body sites are in different colors.

**Figure 7.** The sequencing depth of samples from different research centers. Samples collected at different body sites are in different colors.
OTU level, and select the one better clustering the samples to apply on data aggregated at coarser taxonomic levels to see whether the resolution affects the separation of samples.

**PCA and t-SNE at the OTU level** The PCA is performed on the log-transformed counts that are stored in the `assays` matrix with the name `logcounts`. In practice, data normalization is usually applied prior to the downstream analysis, to address bias or noise introduced during the sampling or sequencing process (e.g., uneven sampling depth). Here, the library size is highly variable (Figure 7) and non-zero OTUs vary across body sites. It is difficult to say what is the optimal normalization strategy, and the use of an inappropriate normalization method might introduce new biases. The discussion of normalization is outside the scope of this work. To keep it simple, we will visualize data without further normalization.

In Figure 8, we see that the Oral samples are distinct from those of other body sites. Samples from Skin, Urogenital Tract, Airways and Gastrointestinal Tract are not separated very well in the first two principal components.

```r
# log-transformed data
assays(tse_phy)$logcounts <- log(assays(tse_phy)$Count + 1)

# run PCA at the OTU level
tse_phy <- runPCA(tse_phy, name="PCA_OTU", exprs_values = "logcounts")

# plot samples in the reduced dimensions
plotReducedDim(tse_phy, dimred = "PCA_OTU", 
colour_by = "HMP_BODY_SITE") +
  labs(title = "PCA at the OTU level") +
  guides(fill = guide_legend(override.aes = list(size=2.5, alpha = 1)))
```

![PCA at the OTU level](image)

**Figure 8.** Principal component analysis (PCA) plot of samples using data at the OTU level. The first two principal components (PCs) are plotted. Each point represents a sample. Samples are coloured according to the body sites.
The separation is well improved with the use of t-SNE in Figure 9. Samples from Oral, Gastrointestinal Tract, and Urogenital Tract form distinct clusters. Skin samples and airways samples still overlap.

```r
# run t-SNE at the OTU level
tse_phy <- runTSNE(tse_phy, name="TSNE_OTU", exprs_values = "logcounts")

# plot samples in the reduced dimensions
tsne_otu <- plotReducedDim(tse_phy, dimred = "TSNE_OTU",
                            colour_by = "HMP_BODY_SITE") +
  labs(title = "t-SNE at the OTU level") +
  theme(plot.title = element_text(hjust = 0.5)) +
  scale_fill_brewer(palette = "Set1") +
  labs(fill = "Body sites") +
  guides(fill = guide_legend(override.aes = list(size=2.5, alpha = 1))) +
  prettify

# plot oral samples
is_oral <- colData(tse_phy)$HMP_BODY_SITE %in% "Oral"
colData(tse_phy)$from_plaque <- grepl(pattern = "Plaque",
                                       colData(tse_phy)$HMP_BODY_SUBSITE)

# Oral samples
plotReducedDim(tse_phy[, is_oral], dimred = "TSNE_OTU",
               colour_by = "from_plaque") +
  guides(fill = guide_legend(override.aes = list(size=2.5, alpha = 1))) +
  prettify
```

Notably, there are two well-separated clusters labelled as oral samples. The smaller cluster includes samples from the Supragingival Plaque and Subgingival Plaque sites, while the larger cluster includes samples from other oral sub-sites (Figure 10).

```r
is_oral <- colData(tse_phy)$HMP_BODY_SITE %in% "Oral"
colData(tse_phy)$from_plaque <- grepl(pattern = "Plaque",
                                       colData(tse_phy)$HMP_BODY_SUBSITE)

# Oral samples
plotReducedDim(tse_phy[, is_oral], dimred = "TSNE_OTU",
               colour_by = "from_plaque") +
  guides(fill = guide_legend(override.aes = list(size=2.5, alpha = 1))) +
  prettify
```

Figure 9. t-SNE plot of samples using data at the OTU level. The first two t-SNE components are plotted. Each point represents a sample. Samples are coloured according to the body site.
Figure 10. t-distributed Stochastic Neighbor Embedding (t-SNE) plot of samples from the oral site using
data at the OTU level. The two t-SNE components computed are plotted. Each point is a sample. Samples from the ‘supragingival or subgingival Plaque’ are in orange, and those from other oral sub-sites are in blue.

t-SNE on broader taxonomic levels

To organize data at different taxonomic levels, we first replace the phylogenetic tree with the taxonomic tree that is generated from the taxonomic table. Due to the existence of polyphyletic groups, a tree structure cannot be generated. For example, the Alteromonadaceae family is from different orders: Alteromonadales and Oceanospirillales.

```r
# taxonomic tree
tax_order <- c("SUPERKINGDOM", "PHYLUM", "CLASS", "ORDER", "FAMILY", "GENUS", "CONSENSUS_LINEAGE")
tax_0 <- data.frame(rowData(tse_phy)[, tax_order])
tax_loop <- detectLoop(tax_tab = tax_0)

# show loops that are not caused by NA
head(tax_loop[!is.na(tax_loop$child), ])
```

<table>
<thead>
<tr>
<th>#</th>
<th>parent</th>
<th>child</th>
<th>parent_column</th>
<th>child_column</th>
</tr>
</thead>
<tbody>
<tr>
<td>35</td>
<td>Alteromonadales</td>
<td>Alteromonadaceae</td>
<td>ORDER</td>
<td>FAMILY</td>
</tr>
<tr>
<td>36</td>
<td>Oceanospirillales</td>
<td>Alteromonadaceae</td>
<td>ORDER</td>
<td>FAMILY</td>
</tr>
<tr>
<td>37</td>
<td>Rhizobiales</td>
<td>Rhodobacteraceae</td>
<td>ORDER</td>
<td>FAMILY</td>
</tr>
<tr>
<td>38</td>
<td>Rhodobacterales</td>
<td>Rhodobacteraceae</td>
<td>ORDER</td>
<td>FAMILY</td>
</tr>
<tr>
<td>39</td>
<td>Chromatiales</td>
<td>Sinobacteraceae</td>
<td>ORDER</td>
<td>FAMILY</td>
</tr>
<tr>
<td>40</td>
<td>Xanthomonadaceas</td>
<td>Sinobacteraceae</td>
<td>ORDER</td>
<td>FAMILY</td>
</tr>
</tbody>
</table>

To resolve the loops, we add a suffix to the polyphyletic genus with resolveLoop. For example, Ruminococcus belonging to the Lachnospiraceae and the Ruminococccaeae families become Ruminococcus_1 and Ruminococcus_2, respectively. A phylo tree is created afterwards using toTree.
tax_1 <- resolveLoop(tax_tab = tax_0)
tax_tree <- toTree(data = tax_1)

# change the tree
tse_tax <- changeTree(x = tse_phy, rowTree = tax_tree,
                      rowNodeLab = rowData(tse_phy)$CONSENSUS_LINEAGE)

The separation of samples from different body sites appears to be worse when the data on broader resolution is used (Figure 11).

# aggregation data to all internal nodes
# Parallel computation is performed with BPPARAM
tse_agg <- aggTSE(x = tse_tax,
                  rowLevel = tax_tree$node.label,
                  whichassay = "Count",
                  BPPARAM = MulticoreParam(),
                  message = FALSE)

# log-transform count
assays(tse_agg)$logcounts <- log(assays(tse_agg)[[1]] + 1)

Specifically, we loop over each taxonomic rank and generate a t-SNE representation using data aggregated at that taxonomic rank level.

Figure 11. t-SNE plot of samples using data at different taxonomic levels. The two t-SNE components computed are plotted. Each point is a sample. Samples are colored according to the body sites.
tax_rank <- c("GENUS", "FAMILY", "ORDER", "CLASS", "PHYLUM")
names(tax_rank) <- tax_rank
fig_list <- lapply(tax_rank, FUN = function(x) {
  # nodes represent the specific taxonomic level
  xx <- startsWith(rowLinks(tse_agg)$nodeLab, x)

  # run t-SNE on the specific level
  xx_tse <- runTSNE(tse_agg, name = paste0("TSNE_", x),
                   exprs_values = "logcounts",
                   subset_row = rownames(tse_agg)[xx])

  # plot samples in the reduced dimensions
  plotReducedDim(xx_tse, dimred = paste0("TSNE_", x),
                  colour_by = "HMP_BODY_SITE",
                  point_size = 0.5) +
                  labs(title = x) +
                  prettify +
                  theme(legend.position = "none") +
                  scale_fill_brewer(palette = "Set1") +
                  guides(fill = guide_legend(override.aes = list(size=2.5)))
})

legend <- get_legend(
  # create some space to the left of the legend
  tsne_otu +
  theme(legend.box.margin = margin(0, 0, 0, 35),
        legend.position = "right")
)
plot_grid(plotlist = fig_list,
          legend, nrow = 2)

CyTOF data
Here, a mass cytometry (CyTOF) dataset\cite{19} is used to show the application of \textit{TreeSummarizedExperiment} on single cell data. The data was available initially as a \textit{SummarizedExperiment} object, and became a \textit{TreeSummarizedExperiment} object after the incorporation of a tree on cells. Data was then aggregated along nodes of the tree to provide data at different resolutions. The data visualization was finally performed as heatmaps along with the tree using the R package \textit{ggtree}.

# packages for visualization
library(ggplot2)
library(ggtree)
library(ggnewscale)
library(RColorBrewer)

# packages for data download and preprocess
library(HDCytoData)
library(diffcyt)
library(ape)

# packages for data manipulation
library(dplyr)
library(tidyverse)
library(tibble)

The mass cytometry (CyTOF) dataset\cite{19} is downloaded from the R package \textit{HDCytoData}\cite{20}. The data has 16 samples (eight pairs) of peripheral blood cell collected from eight healthy individuals. Each pair consists of
one unstimulated sample, and one sample stimulated with B cell receptor/Fc receptor cross-linker (BCR-XL). The data contains expressions of 24 protein markers: 10 surface lineage markers (type) and 14 intracellular signaling functional markers (state), from 172791 cells.

```r
# download data
d_se <- Bodenmiller_BCR_XL_SE()

# Extract data of protein markers
# surface lineage markers: type
# intracellular signaling functional markers: state
is_ab <- colData(d_se)$marker_class %in% c("type", "state")
d_se <- d_se[, is_ab]

## class: SummarizedExperiment
## dim: 172791 24
## metadata(2): experiment_info n_cells
## assays(1): exprs
## rownames: NULL
## rowData names(4): group_id patient_id sample_id population_id
## colnames(24): CD3 CD45 ... HLA-DR CD7
## colData names(3): channel_name marker_name marker_class
```

We preprocess the data and cluster cells using the workflow from the `diffcyt` package[21,22]. According to the median expressions of lineage markers per cluster, a tree `cytof_hclust` is then constructed by applying the hierarchical clustering on the cell cluster level, using only the “type” gene.

```r
# Transform data
d_se <- transformData(d_se)

# Include a random seed to generate a reproducible clustering
# Use cluster IDs as row names
rownames(d_se) <- rowData(d_se)$cluster_id

# Generate a tree with cell clusters as leaves
md <- assay(d_medians)[, metadata(d_medians)$id_type_markers]$id_type_markers
cytof_hclust <- hclust(dist(md, method = "manhattan"), method = "mcquitty")
```

The data `d_se` is converted from `SummarizedExperiment` to `TreeSummarizedExperiment` to provide a `rowTree` slot for the storage the tree information.

```r
# The tree format: convert from hclust to phylo; label internal nodes
cytof_tree <- as.phylod(cytof_hclust)
cytof_tree <- addLabel(tree = cytof_tree, on = "internal")

# Construct a TreeSummarizedExperiment object
lse <- as(d_se, "TreeSummarizedExperiment")
rowTree(lse) <- cytof_tree
```

In `lse`, multiple rows (cells) are mapped to a leaf (a cell cluster) of the tree.
# Data corresponding to the cluster_1
subsetByNode(lse, rowNode = "cluster_1")

## class: TreeSummarizedExperiment
## dim: 2461 24
## metadata(3): experiment_info n_cells MST
## assays(1): exprs
## rownames(2461): cluster_1 cluster_1 ... cluster_1 cluster_1
## rowData names(5): group_id patient_id sample_id population_id
##   cluster_id
## colnames(24): CD3 CD45 ... HLA-DR CD7
## colData names(3): channel_name marker_name marker_class
## reducedDimNames(0):
## altExpNames(0):
## rowLinks: a LinkDataFrame (2461 rows)
## rowTree: 1 phylo tree(s) (49 leaves)
## colLinks: NULL
## colTree: NULL

**Data aggregation**
We split the data into two `TreeSummarizedExperiment` objects: `lse_type` for lineage markers and `lse_state` for functional markers, to perform aggregation in different ways. For `lse_type`, the marker median expression is calculated over all samples to compare expression patterns of lineage markers across all cell clusters. For `lse_state`, the marker median expression is computed on individual samples, to enable comparison between stimulated and unstimulated samples across clusters.

```
# Split TSE: lineage markers and functional markers
lse_type <- lse[, colData(lse)$marker_class == "type"]
lse_state <- lse[, colData(lse)$marker_class == "state"]

# All nodes of the tree
nodes <- showNode(tree = cytof_tree, only.leaf = FALSE)
length(nodes)
```

```
# Calculate marker median expressions for clusters
tse_type <- aggTSE(x = lse_type, rowLevel = nodes, rowFun = median)
# use node labels instead of node alias as row names
rownames(tse_type) <- rowLinks(tse_type)$nodeLab
# row --> node; column --> marker
dim(tse_type)
```

```
# Calculate marker median expressions for clusters separately on each sample
tse_state <- aggTSE(x = lse_state, rowLevel = nodes, rowFun = median,
                   rowBlock = "sample_id")
# use node labels instead of node alias as row names
rownames(tse_state) <- rowLinks(tse_state)$nodeLab
# row --> node per sample; column --> marker
dim(tse_state)
```

```
# [1] 1552 14
```
After aggregation, `tse_type` and `tse_state` have 97 and 1552 rows, respectively. The former has each row representing a cell cluster that is mapped to a node of the tree; the latter has each row representing a cell cluster in a sample.

In the downloaded data, cells are annotated with cell types (`population_id` in `rowData()`). As clustering is not perfect, cells within a cluster are not expected to have exactly the same cell type. Therefore, we would like to annotate a cell cluster with the cell type that the majority of cells (> 60%) belong to, or `mixed` if none of cell types has more than > 60% cells. Note, internal nodes of the tree `cytof_tree` are considered as cell clusters at broader resolution than leaf nodes. To annotate an internal node, we need to first find all cells that are mapped to its descendant leaves, and then take the cell type shared by its majority of cells.

```r
# Find descendant leaves of all nodes; Leaves return themselves
desd_leaf <- findDescendant(tree = cytof_tree, node = nodes, only.leaf = TRUE, self.include = TRUE)

# For example, Node_90 has two descendant leaves: 32 & 33 (node number)
desd_leaf[[90]]

## Node_90 Node_90
##      32      33
```

## Decide cell type for each node according to majorities of cells belong to it

```r
treshold <- 0.6
ct <- sapply(desd_leaf, FUN = function(x) {
  # Data of cells belong to the descendant leaves (x) of a node
  xse <- lse[rowLinks(lse)$nodeNum %in% x, ]

  # Percentages of cell types
  tx <- table(rowData(xse)$population_id)
  pr <- tx/sum(tx)

  # The cell type shared by the majority of cells
  ind <- which(pr > threshold)
  if (!length(ind)) {return("mixed")}
  rownames(tx)[ind]
})
head(ct)

##     cluster_1    cluster_10    cluster_11    cluster_12    cluster_13
## "NK cells" "CD8 T-cells" "NK cells" "CD8 T-cells" "monocytes"
##     cluster_14
## "B-cells IgM-"
```

```r
rowData(tse_type)$population_id <- ct[rownames(tse_type)]
rowData(tse_state)$population_id <- ct[rownames(tse_state)]
```

**Visualization**

The `cytof_tree` tree is considered as a hierarchical structure organizing cell clusters at different granularities. So, an internal node is a cell cluster that incorporates several cell clusters represented by its descendant leaves. Here, we are interested in exploring the expression profile of markers at different resolutions.

We customize a function `treeHM` (below) to draw a tree along with three heatmaps as **Figure 12**. The function is created mainly based on `ggtree` and `gheatmap` from the R package `ggtree`. The use of different color palettes for heatmaps is enable by `scale_fill_*` (from `ggplot2`) and `new_scale_fill()` (from `ggnewscale`).
In Figure 12, the expression of pS6 appears different mainly in three cell clusters (cluster_6, cluster_7 and cluster_14) between stimulated and unstimulated samples. These three clusters all belong to B cells. Also, monocytes seem to have slightly higher pS6 in stimulated samples than in unstimulated samples. cluster_18 is labeled as CD8 T-cells, but it is more similar to monocytes in the expression pattern of lineage markers.
Figure 12. The median expression of markers across 49 cell clusters. Leaves of the tree are labeled with their corresponding cell clusters. Cell types (population_id) of leaves are shown in the first heatmap. Median expressions of ten lineage markers on each leaf are shown in the middle heatmap. Cell clusters in the same branch show similar expression patterns of lineage markers. The right heatmap is about the median expression of a functional marker pS6 in clusters (rows) per sample (column).

We manipulate cytof_tree by merging its three branches as three internal nodes to creates a new tree agg_tree (see Figure 13). ShareNode is used to find the first shared node on paths from specific nodes to the root.

```r
# find branch nodes of the three branches
B_node <- shareNode(tree = cytof_tree, node = c("cluster_6", "cluster_7"))
CD4_node <- shareNode(tree = cytof_tree, node = c("cluster_36", "cluster_31"))
mct_node <- shareNode(tree = cytof_tree, node = c("cluster_34", "cluster_28"))

# Nodes are labeled in red (see Figure)
agg_node <- c(B_node, CD4_node, mct_node)
agg_label <- names(agg_node)

# Set the specific nodes as leaves
agg_tree <- asLeaf(tree = cytof_tree, node = agg_label)

# Generate a figure to compare both trees
both_tree <- list("cytof_tree" = cytof_tree, "agg_tree" = agg_tree)
class(both_tree) <- "multiPhylo"
ggtree(both_tree, ladderize = FALSE) +
  facet_wrap(~.id, scales = "free") +
  geom_tiplab(size = 1.8, align = TRUE) +
  geom_point2(aes(subset = label %in% agg_label),
              color = "red", size = 2) +
  xlim(c(0, 10))
```
We replace the `rowTree()` with the new tree `agg_tree`, and update Figure 12 to get 14. Also, other functional markers, e.g., pNFkB, can be visualized instead of pS6, which we do not show here.

```r
rowTree(tse_type) <- rowTree(tse_state) <- agg_tree
treeHM(tse_type, tse_state, "pS6")
```

**Figure 13. Comparison of two trees: cytof_tree and agg_tree.** Three branches that are connected to the three red nodes in `cytof_tree` are merged, and are presented as dashed lines in `agg_tree`.

**Figure 14. The median expression of markers across 26 cell clusters.** This figure is similar to Figure 12 except that B cells (Node_51), monocytes (Node_53) and CD4 T-cells (Node_55) are now visualized at a broader resolution.
Overall, with `TreeSummarizedExperiment`, single-cell users can over-cluster cells into many cell subpopulations, customize visualization functions to explore data at the high resolution, and finally merge clusters with similar profiles to a suitable resolution to perform downstream analysis.

**Summary**

`TreeSummarizedExperiment` is an S4 class in the family of `SummarizedExperiment` classes, which enables it to work seamlessly with many other packages in Bioconductor. It integrates the `SummarizedExperiment` and the `phylo` class, facilitating data access or manipulation at different resolutions of the hierarchical structure. By providing additional functions for the `phylo` class, we support users to customize functions for the `TreeSummarizedExperiment` class in their workflows.

**Data availability**

**Underlying data**

Human Microbiome Project data (v35) and mass cytometry (CyTOF) dataset were used for the presented use cases. They can be downloaded using the R package `HMP16SData` and `HDCytoData`, respectively.

**Software availability**

The TreeSummarizedExperiment package is available at:


Source code of the development version of the package is available at:

https://github.com/fionarhuang/TreeSummarizedExperiment

Archived source code as at time of publication: http://doi.org/10.5281/zenodo.4046096

**Author information**

RH developed the software with contributions from FGME. All authors participated in the conceptualization of the software. RH, CS and MDR drafted the manuscript with review and editing from KCRA, SCH, GY and FGME. All authors read and approved the final manuscript.

**Acknowledgments**

We thank Héctor Corrada Bravo, LeviWaldron, Hervé Pagès, Martin Morgan, Federico Marini, Jayaram Kancherla, Domenick Braccia, Vince Carey, Kasper D Hansen, Davide Risso, Daniel van Twisk, Marcel Ramos and other members of the Bioconductor community for their helpful suggestions.

**References**


Shila Ghazanfar
Cancer Research UK Cambridge Institute, Li Ka Shing Centre, University of Cambridge, Cambridge, UK

The authors have done an excellent job in addressing each of my questions and implementing feature suggestions where appropriate.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: statistics, high throughput genomics data analysis, single cell genomics analysis, spatial gene expression analysis

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Leo Lahti
Department of Computing, Faculty of Technology, University of Turku, Turku, Finland

Approved.

Competing Interests: Since I reviewed this article in October 2020, I have started collaboration
with the authors of this manuscript. This had not influenced my original review, and I think the feedback in that original review has been adequately addressed now.

**Reviewer Expertise:** Bioinformatics, open research software, R/Bioconductor, microbiome research, statistical machine learning

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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**Version 1**

Reviewer Report 23 November 2020

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Matthew Ritchie

The Walter and Eliza Hall Institute of Medical Research, Parkville, Vic, Australia

Huang et al. describe the TreeSummarizedExperiment package, which provides well-designed S4 infrastructure that couples the phylo and SingleCellExperiment classes to create a container for high-throughput data that can be organised in a tree-like structure.

The article is structured like a vignette, providing an overview of the class (Figure 1) and stepping the reader through the process of setting up a TreeSummarizedExperiment object and accessing and assigning data to its various slots, firstly for a toy data set and then for data from the Human Microbiome Project.

The article is very clearly written, and the authors demonstrate the ability to use TreeSummarizedExperiment objects in conjunction with other established software for dealing with trees (e.g. ggtree and tidyTree) or dimensionality reduction of high-throughput data (e.g. scater). One topic that I was interested to read more about was its potential use in a single cell RNA-seq analysis. Perhaps use cases for such applications can be added as future work. The TreeSummarizedExperiment package has been available from Bioconductor since May 2019 and it has been downloaded > 2.4K times, which indicates it is being taken up by the community.

Minor issues:
- Affiliation 5: missing 'O' in 'Oxford'.
- 'Functions operating on the phylo object.' section, sentence 2, missing word: 'such as ape [and] tidytree.'
Is the rationale for developing the new software tool clearly explained? Yes

Is the description of the software tool technically sound? Yes

Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others? Yes

Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool? Yes

Are the conclusions about the tool and its performance adequately supported by the findings presented in the article? Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Transcriptomics (bulk and single cell)

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 02 Mar 2021

**Ruizhu Huang**, Department of Molecular Life Sciences, University of Zurich, Zurich, Switzerland

1. **Affiliation 5: missing 'O' in 'Oxford'.**

   Thank you. The typo is fixed.

2. **'Functions operating on the phylo object.' section, sentence 2, missing word: 'such as ape [and] tidytree.'**

   The missing word is added now.

**Competing Interests:** No competing interests were disclosed.

Reviewer Report 12 November 2020

https://doi.org/10.5256/f1000research.29440.r73184
Shila Ghazanfar

Cancer Research UK Cambridge Institute, Li Ka Shing Centre, University of Cambridge, Cambridge, UK

Huang and colleagues have written a software article presenting `TreeSummarizedExperiment` [currently version 1.6.0], a Bioconductor package aimed at providing an S4 class for omics data with hierarchical tree structure. The `TreeSummarizedExperiment` class builds on the popular `SingleCellExperiment` object class, with additional slots in which hierarchical structure, in the form of phylo objects, can be added for features (rows) and observations (columns). In addition to the object class, the package contains several functions for manipulating these objects, ranging from getting/setting/resetting the tree slots, aggregating across rows and/or columns, and various analytical tasks operating on the phylo objects.

The article is well written with clear motivation and description of the package, and addresses an important problem of performing analysis of high dimensional hierarchically structured data using object-oriented programming. I have a few further comments and questions that may improve the breadth of use of `TreeSummarizedExperiment` by the research community.

- Is there a way to simply include an argument for `aggValue()` that would swap the order to columns first and rows second, rather than requiring the user to perform two distinct operations?

- The new slots, rowTree, colTree, rowLinks and colLinks are 'getter' accessors but not currently 'setter' functions. I can imagine a popular use-case among users with an already constructed object of class `SummarizedExperiment` or `SingleCellExperiment` would be to simply use as, "TreeSummarizedExperiment") and then attempt to add the additional slots, for example as the output of hclust(). I would suggest prioritising converting these functions to both 'getter' and 'setter', or perhaps adding a constructor usage for `TreeSummarizedExperiment` for objects that are already `SummarizedExperiment` or `SingleCellExperiment`, if possible.

- I'm interested in how `TreeSummarizedExperiment` would work in the case where the hierarchical structure is not a typical single tree, but comprising of multiple distinct tree structures. An example of such is single cell (or single clone) lineage data where there exists a tree structure within each experimental condition, but not between cells from different conditions. Would the `colTree` slot correspond to a list of trees in this case?

- How would one go about combining different `TreeSummarizedExperiment` objects? Do the typical `cbind()` and `rbind()` operations have meaning here? In which cases are they not to be used?

- I would be interested in getting to a clustered heatmap as an example of visualisation for the `TreeSummarizedExperiment`, either implemented using `ggplot2/ggtree`, or other packages like `ComplexHeatmap`?
How would the tree structure information storage scale in terms of the number of rows and columns, or in the hierarchical structures?

Minor/cosmetic
- typo in affiliation 5.
- legend cut off in Figures 6, 7, and 8.

Is the rationale for developing the new software tool clearly explained?
Yes

Is the description of the software tool technically sound?
Yes

Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?
Yes

Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?
Yes

Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** statistics, high throughput genomics data analysis, single cell genomics analysis, spatial gene expression analysis

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

---

Author Response 02 Mar 2021

**Ruizhu Huang,** Department of Molecular Life Sciences, University of Zurich, Zurich, Switzerland

Thank you for your comments!
1. **swap the order to columns first and rows second, rather than requiring the user to perform two distinct operations?**

    aggValue() is now deprecated and replaced by a new function, aggTSE(), that allows users to swap the order of aggregation and define different functions for the row and
2. **The new slots, rowTree, colTree, rowLinks and colLinks are 'getter' accessors but not currently 'setter' functions.** I can imagine a popular use-case among users with an already constructed object of class SummarizedExperiment or SingleCellExperiment would be to simply use as(TreeSummarizedExperiment) and then attempt to add the additional slots, for example as the output of hclust(). I would suggest prioritising converting these functions to both 'getter' and 'setter', or perhaps adding a constructor usage for TreeSummarizedExperiment for objects that are already SummarizedExperiment or SingleCellExperiment, if possible.

rowTree and colTree are now both setters and getters. When the row/column tree is replaced, the rowLinks/colLinks is updated automatically. To avoid breaking links between assays and trees, we don't recommend users to modify the rowLinks/colLinks data. Therefore, rowLinks/colLinks are still kept as getters.

3. **I'm interested in how TreeSummarizedExperiment would work in the case where the hierarchical structure is not a typical single tree, but comprising of multiple distinct tree structures.** An example of such is single cell (or single clone) lineage data where there exists a tree structure within each experimental condition, but not between cells from different conditions. Would the colTree slot correspond to a list of trees in this case?

Yes, it's possible to have a list of trees in the rowTree/colTree. In the rowLinks/colLinks, we have added a new column (whichTree) to give information about which row/column tree a row/column is mapped to. We have also added a new vignette describing how to combine multiple TSEs. (https://www.bioconductor.org/packages/devel/bioc/vignettes/TreeSummarizedExperiment/inst/doc/The_combination_of_multiple_TSEs.html)

4. **How would one go about combining different TreeSummarizedExperiment objects?** Do the typical cbind() and rbind() operations have meaning here? In which cases are they not to be used?

rbind() and cbind() are now implemented for TreeSummarizedExperiment objects. To rbind() multiple TSEs successfully, it's required that the TSEs agree in the column dimension to have the same colTree() and colLinks(). Similarly, cbind() would require TSEs to have the same rowTree() and rowLinks(). More detailed information is available in the new vignette about combining multiple TSEs. (https://www.bioconductor.org/packages/devel/bioc/vignettes/TreeSummarizedExperiment/inst/doc/The_combination_of_multiple_TSEs.html)

5. **I would be interested in getting to a clustered heatmap as an example of visualisation for the TreeSummarizedExperiment, either implemented using ggplot2/ggtree, or other packages like ComplexHeatmap?**
We have added a new use case of TSE on CyTOF data, and customized a visualization function based on `ggtree`, `ggplot2` and `ggnewscale` to plot a clustered heatmap.

6. **How would the tree structure information storage scale in terms of the number of rows and columns, or in the hierarchical structures.**

   We store the tree structure as a phylo object. The size of a phylo object is quite small even for a tree with $10^6$ leaves (about 90 Mb). To set up the link between rows/columns to a tree, it takes only a few seconds even for $10^6$ rows to a tree with $10^6$ leaves.

7. **typo in affiliation 5. legend cut off in Figures 6, 7, and 8.**

   The typo and the legend cut off are fixed.

*Competing Interests:* No competing interests were disclosed.
software development and its use by others; the documentation and examples are sufficient for getting started with and interpreting outputs of the new class for anyone who has the technical skills that are needed to utilize this work.

**Major**

1. Efficiency of the new method could be discussed further; does this scale up to population level cohorts that have thousands of samples are increasing hierarchical resolutions?

2. How easy it would be to incorporate further supporting information on the rows and columns, for instance on DNA/RNA sequence information?

3. The class is very generic; is the idea that this package can be used as such in (hierarchical) single-cell experiments, microbiome research, and potentially other fields that have little overlap currently? Or is this package meant to be a fundamental structure that can be further extended in more specific application domains? Some more discussion on these aspects could help to contextualize the new class.

**Minor**

1. "ssay_data" -> "assay_data"

**Is the rationale for developing the new software tool clearly explained?**

Yes

**Is the description of the software tool technically sound?**

Yes

**Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?**

Yes

**Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?**

Yes

**Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?**

Yes

**Competing Interests:** I recently discussed possible further extensions with the authors of this work. The discussion was based on my own initiative as I am working on related topics, and at that time I did not know that they had (already) submitted this manuscript for review. I do not know the authors, and we have no ongoing collaboration.

**Reviewer Expertise:** Bioinformatics, open research software, R/Bioconductor, microbiome research, statistical machine learning

I confirm that I have read this submission and believe that I have an appropriate level of
expertise to confirm that it is of an acceptable scientific standard.

Author Response 02 Mar 2021

Ruizhu Huang, Department of Molecular Life Sciences, University of Zurich, Zurich, Switzerland

Thank you for reviewing our work.

1. Efficiency of the new method could be discussed further; does this scale up to population level cohorts that have thousands of samples are increasing hierarchical resolutions?

TreeSummarizedExperiment (TSE) inherits the slots of the SummarizedExperiment (SE) & SingleCellExperiment (SCE) classes, and adds new slots like rowTree, colTree, rowLinks, colLinks, referenceSeq. For operations involving the inherited slots, TSE works similarly as SE and SCE. For the new slots, the data manipulation depends on the functions that users have applied on the tree object (of class phylo). These functions might be from TSE or outside TSE. For functions from TSE, either those working on the phylo tree (e.g., findDescendant, convertNode, matTree, addLabel) or those working on TSE (e.g., rowTree, colTree, rowLinks, colLinks, changeTree), takes only seconds even for a tree with up to 100,000 nodes.

2. How easy it would be to incorporate further supporting information on the rows and columns, for instance on DNA/RNA sequence information?

TSE now has a slot referenceSeq to store the sequence information of features (rows).

3. The class is very generic; is the idea that this package can be used as such in (hierarchical) single-cell experiments, microbiome research, and potentially other fields that have little overlap currently? Or is this package meant to be a fundamental structure that can be further extended in more specific application domains? Some more discussion on these aspects could help to contextualize the new class.

Currently, there is not much overlap in the community across fields, e.g, single-cell experiments, microbiome research. But, we do see that they share similarities in data structures, and can potentially share synergies in data visualization or analysis. We provide TSE as a standalone R package like SummarizedExperiment and SingleCellExperiment, and propose it as a convenient starting point to create R packages for downstream analysis or visualization of data with tree structures. We are open to update our work or receive pull requests if new features (or slots) required in a specific field are feasible to be integrated to TreeSummarizedExperiment. For example, a new optional slot referenceSeq(), which was requested mainly for microbiome data to store RNA/DNA sequencing information, has been developed by Félix G.M. Ernst, and the PR has been accepted in TreeSummarizedExperiment.
4. "ssay_data" -> "assay data"

The typo is fixed.

**Competing Interests:** No competing interests were disclosed.

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