

# Package ‘planttfhunter’

March 25, 2025

**Title** Identification and classification of plant transcription factors

**Version** 1.6.0

**Date** 2022-03-10

**Description** planttfhunter is used to identify plant transcription factors (TFs) from protein sequence data and classify them into families and subfamilies using the classification scheme implemented in PlantTFDB. TFs are identified using pre-built hidden Markov model profiles for DNA-binding domains. Then, auxiliary and forbidden domains are used with DNA-binding domains to classify TFs into families and subfamilies (when applicable). Currently, TFs can be classified in 58 different TF families/subfamilies.

**License** GPL-3

**URL** <https://github.com/almeidasilvaf/planttfhunter>

**BugReports** <https://support.bioconductor.org/t/planttfhunter>

**biocViews** Software, Transcription, FunctionalPrediction, GenomeAnnotation, FunctionalGenomics, HiddenMarkovModel, Sequencing, Classification

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**SystemRequirements** HMMER <<http://hmmer.org/>>

**Imports** Biostrings, SummarizedExperiment, utils, methods

**Suggests** BiocStyle, covr, sessioninfo, knitr, rmarkdown, testthat (>= 3.0.0)

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**Depends** R (>= 4.2.0)

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planttfhunter-package *planttfhunter: Identification and classification of plant transcription factors*

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## Description

planttfhunter is used to identify plant transcription factors (TFs) from protein sequence data and classify them into families and subfamilies using the classification scheme implemented in Plant-TFDB. TFs are identified using pre-built hidden Markov model profiles for DNA-binding domains. Then, auxiliary and forbidden domains are used with DNA-binding domains to classify TFs into families and subfamilies (when applicable). Currently, TFs can be classified in 58 different TF families/subfamilies.

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## See Also

Useful links:

- <https://github.com/almeidasilvaf/planttfhunter>
- Report bugs at <https://support.bioconductor.org/t/planttfhunter>

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annotate_pfam	<i>Annotate proteins sequences with PFAM domains</i>
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### Description

PFAM domains are assigned to each sequence using HMMER.

### Usage

```
annotate_pfam(seq = NULL, evaluate = 1e-05)
```

### Arguments

seq	An AAStringSet object as returned by <code>Biostrings::readAAStringSet()</code> . The sequences in this object must represent only the translated sequences of primary (or longest) transcripts.
evaluate	Numeric indicating the E-value threshold for <code>hmmsearch</code> to be used for domains without pre-defined domain cutoffs. Only valid if parameter <code>mode = 'local'</code> . Default: 1e-05.

### Value

A 2-column data frame with the variables **Gene** and **Domain**, which contain gene IDs and domain IDs, respectively.

### Examples

```
data(gsu)
seq <- gsu[1:5]
if(hmmer_is_installed()) {
  annotate_pfam(seq)
}
```

---

classification_scheme	<i>Data frame of TF family classification scheme</i>
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### Description

The classification scheme is the same as the one used by PlantTFDB.

### Usage

```
data(classification_scheme)
```

**Format**

A data frame with the following variables:

**Family** TF family name.

**Subfamily** TF subfamily name.

**DBD** DNA-binding domain

**Auxiliary** Auxiliary domain

**Forbidden** Forbidden domain

**References**

Jin, J., Tian, F., Yang, D. C., Meng, Y. Q., Kong, L., Luo, J., & Gao, G. (2016). PlantTFDB 4.0: toward a central hub for transcription factors and regulatory interactions in plants. *Nucleic acids research*, gkw982.

**Examples**

```
data(classification_scheme)
```

---

classify_tfs	<i>Identify TFs and classify them in families</i>
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**Description**

Identify TFs and classify them in families

**Usage**

```
classify_tfs(domain_annotation = NULL)
```

**Arguments**

domain\_annotation

A 2-column data frame with the gene ID in the first column and the domain ID in the second column.

**Value**

A 2-column data frame with the variables **Gene** and **Family** representing gene ID and TF family, respectively.

**Examples**

```
data(gsu_annotation)
domain_annotation <- gsu_annotation
families <- classify_tfs(domain_annotation)
```

---

get_tf_counts	<i>Get TF frequencies for each species as a SummarizedExperiment object</i>
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---

## Description

This function identifies and classifies TFs, and returns TF counts for each family as a SummarizedExperiment object

## Usage

```
get_tf_counts(proteomes, species_metadata = NULL)
```

## Arguments

**proteomes** List of **AAStringSet** objects

**species\_metadata** (Optional) A data frame containing species names in row names (names must match element names in the **proteomes** list), and species metadata (e.g., taxonomic information, ecological information) in columns. If **NULL**, the colData of the SummarizedExperiment object will be empty.

## Value

A SummarizedExperiment object containing transcription factor frequencies per family in each species, as well as species metadata (if **species\_metadata** is not **NULL**).

## Examples

```
data(gsu)

set.seed(123)
# Pick random subsets of 100 genes to simulate other species
proteomes <- list(
  Gsu1 = gsu[sample(names(gsu), 50, replace = FALSE)],
  Gsu2 = gsu[sample(names(gsu), 50, replace = FALSE)],
  Gsu3 = gsu[sample(names(gsu), 50, replace = FALSE)],
  Gsu4 = gsu[sample(names(gsu), 50, replace = FALSE)]
)

# Create species metadata
species_metadata <- data.frame(
  row.names = names(proteomes),
  Division = "Rhodophyta",
  Origin = c("US", "Belgium", "China", "Brazil")
)

# Get SummarizedExperiment object
if(hmmer_is_installed()) {
  se <- get_tf_counts(proteomes, species_metadata)
}
```

---

 gsu

*Protein sequences of the algae species Galdieria sulphuraria*


---

**Description**

Data obtained from PLAZA Diatoms. Only genes containing domains used for TF family classification were kept for package size issues.

**Usage**

```
data(gsu)
```

**Format**

An AAStringSet object as returned by `Biostrings::readAAStringSet()`.

**References**

Osuna-Cruz, C. M., Bilcke, G., Vancaester, E., De Decker, S., Bones, A. M., Winge, P., ... & Vandepoele, K. (2020). The *Seminavis robusta* genome provides insights into the evolutionary adaptations of benthic diatoms. *Nature communications*, 11(1), 1-13.

**Examples**

```
data(gsu)
```

---

 gsu\_annotation

*Domain annotation for the algae species Galdieria sulphuraria The data set was created using the function annotate\_pfam() in local mode.*


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**Description**

Domain annotation for the algae species *Galdieria sulphuraria*

The data set was created using the function `annotate_pfam()` in local mode.

**Usage**

```
data(gsu_annotation)
```

**Format**

A 2-column data frame with the following variables:

**Gene** Gene ID

**Annotation** Domain ID or domain name when ID is not available in PFAM

**Examples**

```
data(gsu_annotation)
```

---

gsu_families	<i>TFs families of the algae species Galdieria sulphuraria The data set was created using the funcion classify_tfs().</i>
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---

**Description**

TFs families of the algae species Galdieria sulphuraria  
The data set was created using the funcion `classify_tfs()`.

**Usage**

```
data(gsu_families)
```

**Format**

A 2-column data frame with the following variables:

**Gene** Gene ID

**Family** TF family

**Examples**

```
data(gsu_families)
```

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hmmr_is_installed	<i>Check if HMMER is installed</i>
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**Description**

Check if HMMER is installed

**Usage**

```
hmmr_is_installed()
```

**Value**

Logical indicating whether HMMER is installed or not.

**Examples**

```
hmmr_is_installed()
```

---

`tf_counts`*TF counts per family in 4 simulated species*

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**Description**

Simulated species were created by sampling 100 genes from the example data set **gsu** with after `set.seed(123)`.

**Usage**

```
data(tf_counts)
```

**Format**

A `SummarizedExperiment` with TF frequencies per family in each species in **assay** and species metadata in **colData**.

**Examples**

```
data(tf_counts)
```



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