

# Package ‘defrostR’

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**Type** Package

**Title** A tool for keeping up with amphibian taxonomy

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**Description** This package provides a workflow for combining amphibian taxonomic data sets from different sources that use different taxonomic nomenclature, by suggesting the most current nomenclature used by the Amphibian Species of the World online reference for a given input query of species names. It can also be used to extract specific systematic information (e.g. species downstream of a given taxonomic level).

**Depends** R (>= 3.0.1), XML

**Suggests** knitr, rmarkdown

**VignetteBuilder** knitr

**License** GPL (>= 2.0)

**Encoding** UTF-8

**LazyData** true

**RoxygenNote** 6.0.1

**URL** <http://github.com/hcliedtke/defrostR>

**BugReports** <http://github.com/hcliedtke/defrostR/issues>

**NeedsCompilation** no

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amphweb

*AmphibiaWeb Taxonomy*


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### Description

Data table of taxonomy downloaded from <http://amphibiaweb.org/>. The dataset contains a list of species and species names used by AmphibiaWeb as well as higher level taxonomy and more (IUCN red list status etc.). A version is stored internally for the purpose of running examples, but users should be aware that this may be outdated depending both on how recently the defrostR package as well as the AmphibiaWeb website has been updated.

### Usage

```
amphweb
```

### Format

R dataset

### Examples

```
head(amphweb)
amphweb<-defrostR::amphweb
```

---

aswStats

*Get summary statistics of current taxonomy*


---

### Description

This function uses the ASW taxonomy table created with the function `getTaxonomy()` and summarizes counts of units per taxonomic level

### Usage

```
aswStats(asw_taxonomy = defrostR::asw_taxonomy, verbose = F, Order = NA,
         Superfamily = NA, Family = NA, Subfamily = NA, Genus = NA)
```

### Arguments

asw_taxonomy	output of the function <code>frostScraper()</code> default will take defrostR's stored version, which might be outdated
verbose	logical arguments can be turned on to summarize at given taxonomic levels or to give numbers of each unit per taxonomic level. default is switched off (FALSE)
Order	string to specify summary statistics of only a specific order
Superfamily	string to specify summary statistics of only a specific superfamily
Family	string to specify summary statistics of only a specific Family
Subfamily	string to specify summary statistics of only a specific subfamily
Genus	string to specify summary statistics of only a specific genus

**Details**

One logical arguments can be turned on to summarize at given taxonomic levels or to give numbers of each unit per taxonomic level

**Value**

returns either a data frame or list of summary statistics

**Examples**

```
aswStats()  
aswStats(verbose=TRUE, Family="Rhacophoridae")
```

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asw_synonyms	<i>Amphibian Species of the World synonyms</i>
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**Description**

Data table generated using the getSynonyms() function. The dataset contains a full list of synonyms listed on the Amphibian Species of the World (ASW) website and the current accepted name of the species. A version is stored internally for the purpose of running examples, but users should be aware that this may be outdated depending both on how recently the defrostR package as well as the ASW website has been updated.

**Usage**

```
asw_synonyms
```

**Format**

R dataset

**Details**

dataset last updated on 12th July 2017

**Examples**

```
head(asw_synonyms)  
str(asw_synonyms)  
asw_synonyms<-defrostR::asw_synonyms
```

---

asw\_taxonomy

*Amphibian Species of the World Taxonomy*


---

### Description

Data table generated using the `getTaxonomy()` function. The dataset contains a full list of species listed on the Amphibian Species of the World (ASW) website along with all higher level taxonomy and the URLs for each species. A version is stored internally for the purpose of running examples, but users should be aware that this may be outdated depending both on how recently the `defrostR` package as well as the ASW website has been updated.

### Usage

```
asw_taxonomy
```

### Format

R dataset

### Details

dataset last updated on 12th July 2017

### Examples

```
head(asw_taxonomy)
str(asw_taxonomy)
asw_taxonomy<-defrostR::asw_taxonomy
```

---

defrost

*Submit query to update nomenclature according to the ASW taxonomy*


---

### Description

This function takes a query (a list of taxon names to be assessed) and the amphibian species of the world synonym table (preferably generated with the function `getSynonyms()`, or loaded from the dataset stored internally) as input and returns an updated list of names.

### Usage

```
defrost(query, asw = defrostR::asw_synonyms, interactive = F,
        return.no.matches = F)
```

**Arguments**

query	vector of taxon names to be processed (can also be tip labels of a phylogeny for example)
asw	amphibian species of the world synonym reference table on which to base new names on. Default is to use defrostR's internally stored data set that may not be the most up-to-date.
interactive	logical argument (default=FALSE) of whether to allow an "on the fly" decision to be made on what name to take if synonym matches multiple names. TRUE will return all possible names for a given query
return.no.matches	logical argument of whether to leave taxa not found in the reference table blank or whether to fill in the names provided by the query. default is FALSE

**Details**

Two logical arguments can be turned on to a) allow an "on the fly" decision to be made on what name to take if synonym matches multiple names (if not, it will return all possible names) and to b) return the original name if no match is found

**Value**

this function returns a data frame with the following information/columns: original/input names, "stripped" names with no formatting, status of what action has been taken, updated names as recommended by the reference table

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getSynonyms	<i>Compile dataframe of all synonyms listed on Amphibia Species of the World</i>
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**Description**

This function looks up any listed synonyms on the ASW website (<http://research.amnh.org/vz/herpetology/amphibia/index.html>). It takes a minimum of one argument, the asw\_taxonomy table generated with the function getTaxonomy(). If a full search is performed, this can take quite long (looking through ~7000 websites for upwards of 20 000 synonyms), but additional arguments can be included to restrict searches to a specified taxonomic group.

**Usage**

```
getSynonyms(asw_taxonomy = defrostR::asw_taxonomy, Order = NA,
             Superfamily = NA, Family = NA, Subfamily = NA, Genus = NA,
             Species = NA)
```

**Arguments**

asw_taxonomy	the ASW taxonomy table obtained with getTaxonomy(). If no table is provided, the default is to use defrostR's internally stored version. WARNING! this version may be outdated
Order	limit search to a user-specified amphibian order
Superfamily	limit search to a user-specified amphibian superfamily

Family	limit search to a user-specified amphibian family
Subfamily	limit search to a user-specified amphibian subfamily
Genus	limit search to a user-specified amphibian genus
Species	limit search to a user-specified amphibian species

### Details

Users may experience issues with umlauts that are not supported by their system language. On Mac OSX, this can be changed by running the following line of code in R:

```
system("defaults write org.R-project.R force.LANG en_US.UTF-8")
```

and then restarting the session. Read more here: [https://cran.r-project.org/bin/macosx/RMacOSX-FAQ.html#Internationalization-of-the-R\\_002eapp](https://cran.r-project.org/bin/macosx/RMacOSX-FAQ.html#Internationalization-of-the-R_002eapp)

### Value

returns a dataframe listing all species and their listed synonyms

### Examples

```
#to get a full list of synonyms for all species, run: asw_synonyms<-getSynonyms()
breviceptid_synonyms<-getSynonyms(Family="Brevicipitidae")
```

---

getTaxonomy	<i>Construct taxonomy used by Amphibian Species of the World</i>
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---

### Description

This function takes no arguments. It cycles through the ASW website (<http://research.amnh.org/vz/herpetology/amphibia>) in an attempt to scrap all taxonomic information from Order to Species. As there are 5 taxonomic levels within each order (Superfamily/Family/Subfamily/Genus/Species), one can expect the function to require 1 initial loop followed by 4 more, but it is designed to keep going until all lineages have reached the species level.

### Usage

```
getTaxonomy()
```

### Value

returns a dataframe listing all species and their taxonomic backbone as well as all URLs per species.

### Examples

```
#run asw_taxonomy_table<-getTaxonomy() to scrape the ASW website for the most current taxonomy
```

---

synonymReport	<i>Get summary report of "defrosted" query</i>
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**Description**

This function produces a report of an undertaken defrost() query

**Usage**

```
synonymReport(defrosted, verbose = F)
```

**Arguments**

defrosted	output of the function defrost()
verbose	logical arguments can be turned on to list species which result in conflict. Default=FALSE

**Details**

One logical arguments can be turned on to summarize in terms of numbers of cases, or to provide species names of conflicting issues.

**Value**

returns either a data frame or list of summary statistics

**Examples**

```
amphweb<-defrostR::amphweb
head(amphweb$species)
amphweb.bufo<-amphweb[amphweb$family=="Bufo",]
bufo.defrosted<-defrost(query=amphweb.bufo$species)
synonymReport(bufo.defrosted)
synonymReport(bufo.defrosted,)
```

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