fqacalc

A Floristic Quality Assessment Calculator for R

This package provides functions for calculating Floristic Quality Assessment (FQA) metrics using regional FQA databases that have been reviewed and certified as ecological planning models by the U.S. Army Corps of Engineers (USACE). These databases are stored in a sister R package, fqadata. Both packages were developed for the USACE by the U.S. Army Engineer Research and Development Center's Environmental Laboratory.

To complete this tutorial interactively, follow along in R studio.

Installation

You can install the development version of fqacalc from GitHub with:

```
# install.packages("devtools")
devtools::install_github("ifoxfoot/fqacalc")

#attach packages required for this tutorial
library(fqacalc) #for FQA calculations
library(stringr) #for string manipulation
library(dplyr) #for data manipulation

#>

#> Attaching package: 'dplyr'

#> The following objects are masked from 'package:stats':

#>

#> filter, lag

#> The following objects are masked from 'package:base':

#>

#> intersect, setdiff, setequal, union
```

Package Data

fqacalc contains all regional FQA databases that have been either fully approved or approved with reservations for use by the U.S. Army Corps of Engineers. By referencing these databases, the package can assign a Coefficient of Conservatism (or C Value) to each plant species that the user inputs. A list of regional FQA databases can be viewed using the db_names() function, and specific FQA databases can be accessed using the view_db() function. Below is an example of how to view one of the regional databases.

```
#view a list of all available databases
head(db names())
#>
                                                       recommendation
                                                                               notes
                                        fqa_db
#> 1
           atlantic_coastal_pine_barrens_2018 Yes, with reservations
                                                                                <NA>
                          chicago\_region\_2017
#> 2
                                                                   YES
                                                                                <NA>
#> 3
                                 colorado 2020
                                                                                <NA>
#> 4
           dakotas_excluding_black_hills_2017 Previously Certified
                                                                                <NA>
                                 delaware_2013 Yes, with reservations
                                                                                <NA>
#> 6 eastern_great_lakes_hudson_lowlands_2018
                                                                   YES Wetlands Only
#>
#> 1
                                         Faber-Langendoen, D., Cameron, D., Gilman, A. V., Metzler, K. J
```

```
#> 2
#> 3
#> 4 The Northern Great Plains Floristic Quality Assessment Panel. 2001. Coefficients of conservatism f
                                                                   Chamberlain, S. J., & Ingram, H.
#> 6
                                      Faber-Langendoen, D., Cameron, D., Gilman, A. V., Metzler, K. J
#store the Colorado database as an object
colorado <- view db("colorado 2020")</pre>
#view it
head(colorado)
#> # A tibble: 6 x 13
    name
                name ~1 acronym accep~2 family nativ~3 c w wetla~4 physi~5
#>
    <chr>
                <chr> <chr> <chr> <chr> <chr> <chr> <chr> <dbl> <dbl> <dbl> <chr>
#> 1 ABIES BIFO~ accept~ ABBI3 Abies ~ Pinac~ native
                                                       5 1 FACU
#> 2 ABIES LASI~ synonym <NA> Abies ~ Pinac~ native
                                                         5
                                                               1 FACU
                                                                         tree
#> 3 ABIES CONC~ accept~ ABCO Abies ~ Pinac~ native
                                                         5 NA <NA>
#> 4 ABRONIA EL~ accept~ ABEL Abroni~ Nycta~ native
                                                                         forb
                                                         4 NA <NA>
#> 5 ABRONIA FR~ accept~ ABFR2 Abroni~ Nycta~ native
                                                        6 NA <NA>
                                                                         forb
                                                     9
#> 6 ABRONIA GL~ accept~ ABAR Abroni~ Nycta~ native
                                                            NA <NA>
                                                                         forb
#> # ... with 3 more variables: duration <chr>, common_name <chr>, fqa_db <chr>,
#> # and abbreviated variable names 1: name_origin, 2: accepted_scientific_name,
#> # 3: nativity, 4: wetland_indicator, 5: physiognomy
```

Smith, P

fqacalc also comes with a site assessment from Crooked Island, Michigan, downloaded from the Universal FQA Calculator. The data set is called crooked_island and is used in this tutorial to demonstrate how the package works. When calculating metrics for crooked_island, use the 'michigan_2014' regional database.

```
#view the data
head(crooked island)
#> acronym common_name
                                            name
                            Abies balsamea
#> 1 ABIBAL
             balsam fir
#> 2 AMMBRE marram grass Ammophila breviligulata
#> 3 ANTELE white camas
                               Anticlea elegans
#> 4 ARCUVA
             bearberry Arctostaphylos uva-ursi
#> 5 ARTCAM
                wormwood
                           Artemisia campestris
#> 6 CALEPI
               reedgrass Calamagrostis epigeios
#print the dimensions (35 rows and 3 columns)
dim(crooked_island)
#> [1] 35 3
#view the documentation for the data set (bottom right pane of R studio)
?crooked_island
#load the data set into local environment
crooked_island <- crooked_island</pre>
```

Reading Site Assessment Data into R

Site assessments can be read into R for analysis using base R or the readxl package (for .xls or .xlsx files).

If the site assessment is a csv file, it can be read in using read.csv(). For example, code to read in data might look like my_data <- read.csv("path/to/my/data.csv"). If the site assessment is in an Excel file, it can be read in with the same code, but replace read.csv() with read_excel().

In order to calculate FQA metrics using fqacalc, the site assessment data must be in the following format:

- 1. The data must have either a column named name containing scientific names of plant species, or a column named acronym containing acronyms of plant species. Different regional FQA databases use different naming conventions and have different ways of creating acronyms (and some don't have acronyms!) so be sure to look at the relevant regional database to check that the site assessment is using the same conventions. Names/acronyms do not have to be in the same case, but otherwise must exactly match their counterpart in the regional FQA database in order to be recognized by fqacalc functions.
- 2. If the user is calculating cover-weighted metrics, the data must have another column containing cover values and it must be called cover. If the cover values are in percent cover, they must be between 0-100. If they are in a cover class, such as the Braun-Blanquet classification system, they must be correct for that class or else they won't be recognized. See the section on cover-weighted functions to learn more about cover classes.
- 3. If the user is calculating cover-weighted metrics for a transect containing multiple plots, the data should also have a column containing the plot ID. The plot ID column can have any name, and it can contain numbers or characters, as long as the IDs are exactly the same within plots but distinct between plots.

In this case, each observation is one row, containing the species name or acronym, the cover value, and the plot ID. It might look something like this:

plot_id	name	cover
1	Plant A	20
1	Plant B	50
2	Plant C	35
2	Plant D	45

Functions that Match Plant Species from Site Assessments to Regional FQA Databases

fqacalc contains two functions that help the user understand how the data they input matches up to the regional database: accepted_entries() and unassigned_plants(). accepted_entries() is a function that shows which plant species in the input data frame are successfully matched to species in the regional database, and unassigned_plants() shows which species are matched but don't have a C value stored in the regional database.

What happens when a plant species is not in the regional FQA database?

accepted_enteries shows which species are recognized, but it also provides warnings when a species is not recognized. To demonstrate this we can add a mistake to the crooked island data set.

```
#introduce a typo
mistake_island <- crooked_island %>%
   mutate(name = str_replace(name, "Abies balsamea", "Abies blahblah"))

#store accepted entries
accepted_entries <- accepted_entries(#this is the data</pre>
```

Now, when we use accepted_entries() to see which species were matched to the regional data set, we can see that we received a message about the species 'ABIES BLAHBLAH' being discarded and we can also see that the accepted entries data set we created only has 34 entries instead of the expected 35 entries.

What happens when plant species don't have C values?

In some cases, a plant species from the site assessment can be matched to the regional database, but the species is not associated with any C Value. Plant species that are matched but have no C Value will be excluded from FQA metric calculation but they can *optionally* be included in other metrics like species richness, relative cover, relative frequency, relative importance, and mean wetness, as well as any summarizing functions containing these metrics. This option is denoted with the allow_no_c argument.

unassigned_plants() is a function that shows the user which plant species have not been assigned a C Value.

```
#To see unassigned_plants in action we're going to Montana!
#first create a df of plants to input
no_c_plants<- data.frame(name = c("ABRONIA FRAGRANS",</pre>
                                   "ACER GLABRUM",
                                   "ACER GRANDIDENTATUM",
                                   "ACER PLATANOIDES"))
#then create a df of unassigned plants
unassigned plants(no c plants, key = "name", db = "montana 2017")
#> montana_2017 does not have wetness coefficients, wetland metrics cannot be calculated.
#>
                                       name origin acronym accepted scientific name
#> 1
        ABRONIA\ FRAGRANS\ accepted\_scientific\_name
                                                       <NA>
                                                                    Abronia fragrans
#> 2 ACER GRANDIDENTATUM accepted scientific name
                                                       <NA>
                                                                 Acer grandidentatum
                       nativity c w wetland_indicator physiognomy duration
#>
            family
#> 1 Nyctaginaceae
                                                    <NA>
                                                                 <NA>
                         native NA NA
                                                                          <NA>
#> 2
         Aceraceae undetermined NA NA
                                                    <NA>
                                                                 <NA>
                                                                          <NA>
                     common name
                                        fqa_db
#> 1 Fragrant White Sand-Verbena montana_2017
#> 2
                  Bigtooth Maple montana_2017
```

The function returns two species that are in the 'montana_2017' databases but aren't assigned a C Value.

How will duplicates be treated?

If the site assessment data contains duplicate species, they will be excluded from certain FQA metrics. For example, species richness counts the number of unique species, so duplicates are not allowed. Generally,

duplicates are excluded for all unweighted (inventory) metrics but can optionally be included in coverweighted metrics and are always included in relative metrics.

Duplicate behavior in cover-weighted functions is controlled by the allow_duplicates argument and the plot_id argument. If allow_duplicates = FALSE, no duplicate species will be allowed at all, no matter how plot_id is set. If allow_duplicates = TRUE and the plot_id argument is set, duplicate species will be allowed if they are in different plots.

If there are duplicates, and the user is attempting to perform a cover-weighted calculation where duplicates are not allowed, the duplicated species will be condensed into one entry with an aggregate cover value. A message will notify the user if this occurs. See this example.

Will synonyms be recognized?

Some regional FQA databases include accepted scientific names as well as commonly used synonyms. As long as these synonyms are in the regional database, they will be recognized by fqacalc functions. There are a few important rules regarding synonyms.

- 1. If both the synonym and the accepted name are used in a site assessment, the synonym will be converted to the accepted name and both observations will only count as *one* species.
- 2. If the site assessment data contains a name that is listed as a synonym to one species and an accepted name to a different species, it will default to the species with the matching accepted name.
- 3. If the site assessment contains a species that is listed as a synonym to multiple species in the regional FQA database, this entry will *not* be included! To include the species, enter the accepted scientific name instead of the synonym.

In all of these cases, fqacalc functions will print messages to warn the user about synonym issues. See this example:

Unweighted (Inventory) FQI Metrics

fqacalc contains a variety of functions that calculate Total Species Richness, Native Species Richness, Mean C, Native Mean C, Total FQI, Native FQI, and Adjusted FQI. All of these functions eliminate duplicate species and species that cannot be found in the regional database. All but Total Species Richness and Native Species Richness automatically eliminate species that are not associated with a C Value.

Function Arguments In general, all of these metric functions have the same arguments.

- x: A data frame containing a list of plant species. This data frame *must* have one of the following columns: name or acronym.
- key: A character string representing the column that will be used to join the input x with the regional FQA database. If a value is not specified the default is name. name and acronym are the only acceptable values for key.
- db: A character string representing the regional FQA database to use. See db_names() for a list of potential values.
- native: native Boolean (TRUE or FALSE). If TRUE, calculate metrics using only native species.

Additionally, species_richness() and all_metrics() have an argument called allow_no_c. If allow_no_c = TRUE than species that are in the regional FQA database but don't have C Values will be included. If allow_no_c is FALSE, then these species will be omitted. This argument is also found in mean_w() and all of the relative functions.

```
#total mean c
mean_c(crooked_island, key = "acronym", db = "michigan_2014", native = FALSE)
#> [1] 5.371429

#native mean C
mean_c(crooked_island, key = "acronym", db = "michigan_2014", native = TRUE)
#> [1] 6.714286

#total FQI
FQI(crooked_island, key = "acronym", db = "michigan_2014", native = FALSE)
#> [1] 31.7778

#native FQI
FQI(crooked_island, key = "acronym", db = "michigan_2014", native = TRUE)
#> [1] 35.52866

#adjusted FQI (always includes both native and introduced species)
adjusted_FQI(crooked_island, key = "acronym", db = "michigan_2014")
#> [1] 60.0544
```

Functions And finally, all metrics() prints all of the metrics in a data frame format.

```
#a summary of all metrics (always includes both native and introduced)
#can optionally include species with no C value
#--if TRUE, this species will count in species richness and mean wetness metrics
all_metrics(crooked_island, key = "acronym", db = "michigan_2014", allow_no_c = TRUE)
#>
                             metrics
                                         values
#> 1
              Total Species Richness 35.0000000
#> 2
             Native Species Richness 28.0000000
#> 3
        Introduced Species Richness 7.0000000
#> 4
        % of Species with no C Value 0.0000000
#> 5
        % of Species with 0 C Value 20.0000000
#> 6
       % of Species with 1-3 C Value 8.5714286
      % of Species with 4-6 C Value 34.2857143
     % of Species with 7-10 C Value 37.1428571
#> 8
                              Mean C 5.3714286
#> 9
#> 10
                       Native Mean C 6.7142857
#> 11
                           Total FQI 31.7778000
#> 12
                          Native FQI 35.5286605
#> 13
                        Adjusted FQI 60.0543971
#> 14
                        Mean Wetness 0.7142857
#> 15
                 Native Mean Wetness 0.8571429
#> 16
                       % Hydrophytes 37.1428571
```

All of the functions are documented with help pages.

```
#if in R studio, running this line of code should bring up a help page in bottom right pane ?all_metrics
```

Cover-Weighted Functions

Cover-Weighted Functions calculate the same metrics but they are weighted by species abundance. Therefore, the input data frame must also have a column named cover containing cover values. Cover values can be continuous (i.e. percent cover) or classed (e.g. using the Braun-Blanquet method).

The following tables describe how cover classes are converted to percent cover. Internally, cover-weighted functions convert cover classes to the percent cover midpoint. For this reason, using percent cover is recommended over using cover classes.

Braun-Blanquet Classes	% Cover Range	Midpoint
+	<1%	0.1
1	< 5%	2.5
2	5-25%	15
3	25 50%	37.5
4	50 - 75%	62.5
4	75 - 100%	87.5

Carolina Veg Survey Classes	% Cover Range	Midpoint
1	< 0.1	0.1
2	0 - 1%	0.5
3	1-2%	1.5
4	2-5%	3.5

Carolina Veg Survey Classes	% Cover Range	Midpoint
5	5-10%	7.5
6	10-25%	17.5
7	25-50%	37.5
8	50 - 75%	62.5
9	75 - 95%	85
10	95 - 100%	97.5
10	95-100%	97.5

Daubenmire Classes	% Cover Range	Midpoint
1	0-5%	2.5
2	5-25%	15
3	25 50%	37.5
4	50 - 75%	62.5
5	75- $95%$	85
6	95 - 100%	97.5

USFS Ecodata Classes	% Cover Range	Midpoint
1	<1%	0.5
3	1.1 - 5%	3
10	5.1 - 15%	10
20	15.1 - 25%	20
30	25.1-35%	30
40	35.1 - 45%	40
50	45.1 - 55%	50
60	55.1-65%	60
70	65.1 - 75%	70
80	75.1 - 85%	80
90	85.1 - 95%	90
98	95.1-100%	98

Cover-Weighted functions come in two flavors: Transect-level and plot-level. Transect-level metrics are those that calculate a metric for an entire transect, which typically includes multiple plots. transect_summary and plot_summary are both always calculated at the transect-level. Plot-level metrics calculate a metric for a single plot. cover_mean_c and cover_FQI can be transect-level or plot-level. It is up to the user to decide if they are calculating a transect-level or a plot-level metric.

To calculate <code>cover_mean_c</code> and <code>cover_FQI</code> at the transect-level, set <code>allow_duplicate = TRUE</code>, because different plots along the transect may contain the same species. It is also recommended to include a plot ID column and set the <code>plot_id</code> argument to be equal to that column name. This will allow duplicate species between plots but not allow duplication within plots.

To calculate cover_mean_c and cover_FQI at the plot-level, set allow_duplicate = FALSE. There is no need to set the plot_id argument because duplicate species will not be allowed in any circumstance.

If duplicated species are found where they are not supposed to be, the duplicated entries will only be counted once and their cover values will be added together. The user will also recieve a message stating duplicates have been removed.

Function Arguments Cover-Weighted Functions have a few additional arguments:

- cover_class: A character string representing the cover method used. Acceptable cover methods are: "percent_cover", "carolina_veg_survey", "braun-blanquet", "doubinmire", and "usfs_ecodata". "percent_cover" is the default and is recommended.
- allow_duplicates: Boolean (TRUE or FALSE). If TRUE, allow duplicate entries of the same species. If FALSE, do not allow species duplication. See cover-weighted function description. Setting allow_duplicates to TRUE is best for calculating metrics for multiple plots which potentially contain the same species. Setting allow_duplicates to FALSE is best for calculating metrics for a single plot, where each species is entered once along with its total cover value.
- plot_id: A character string representing the column in x that contains plot identification values. plot_id is a required argument in plot_summary, where it acts as a grouping variable. plot_id is optional for cover-weighted functions and frequency functions. If plot_id is set in a cover-weighted function or a frequency function, it only prevents duplicates from occurring in the same plot. It does not act as a grouping variable.

```
#first make a hypothetical plot with cover values
plot <- data.frame(acronym = c("ABEESC", "ABIBAL", "AMMBRE", "ANTELE"),</pre>
                   name = c("Abelmoschus esculentus",
                            "Abies balsamea", "Ammophila breviligulata",
                            "Anticlea elegans; zigadenus glaucus"),
                   cover = c(50, 4, 20, 30))
#now make up a transect
transect <- data.frame(acronym = c("ABEESC", "ABIBAL", "AMMBRE",</pre>
                                    "AMMBRE", "ANTELE", "ABEESC", "ABIBAL", "AMMBRE"),
                       cover = c(50, 4, 20, 30, 30, 40, 7, 60),
                       plot_id = c(1, 1, 1, 1, 2, 2, 2, 2))
#plot cover mean c (no duplicates allowed)
cover_mean_c(plot, key = "acronym", db = "michigan_2014",
             native = FALSE, cover_class = "percent_cover",
             allow_duplicates = FALSE)
#> [1] 4.923077
#transect cover mean c (duplicates allowed along a transect, unless in the same plot)
cover_mean_c(transect, key = "acronym", db = "michigan_2014",
             native = FALSE, cover_class = "percent_cover",
             allow_duplicates = TRUE, plot_id = "plot_id")
#> Duplicate entries detected in the same plot. Duplicates in the same plot will be counted once. Cover
#> [1] 6.394834
*cover-weighted FQI (again, you can choose to allow duplicates or not depending on if species are in a
cover_FQI(transect, key = "acronym", db = "michigan_2014", native = FALSE,
          cover_class = "percent_cover",
          allow_duplicates = TRUE)
#> [1] 11.66145
#transect summary function (always allows duplicates)
transect_summary(transect, key = "acronym", db = "michigan_2014")
#>
                             metrics
                                         values
#> 1
              Total Species Richness 4.0000000
```

```
Native Species Richness 3.0000000
#> 3
        Introduced Species Richness
                                     1.0000000
       % of Species with no C Value 0.0000000
        % of Species with 0 C Value 25.0000000
      % of Species with 1-3 C Value 25.0000000
#> 7
      % of Species with 4-6 C Value 0.0000000
#> 8
     % of Species with 7-10 C Value 50.0000000
#> 9
                             Mean C 5.7500000
#> 10
                      Native Mean C 7.666667
              Cover-Weighted Mean C 5.8307255
#> 11
#> 12
       Cover-Weighted Native Mean C 9.4665127
#> 13
                          Total FQI 11.5000000
#> 14
                         Native FQI 13.2790562
#> 15
                  Cover-Weighted FQI 11.6614509
          Cover-Weighted Native FQI 16.3964810
#> 16
#> 17
                       Adjusted FQI 66.3952810
#> 18
                       Mean Wetness 1.7500000
#> 19
                 Native Mean Wetness 0.666667
#> 20
                       % Hydrophytes 12.5000000
```

Functions There is also a plot summary function that summarizes plots along a transect. Data is input as a single data frame containing species per plot. This data frame must also have a column representing the plot that the species was observed in.

Because it is sometimes useful to calculate the total amount of bare ground or un-vegetated water in a plot, the user can also choose to include bare ground or water. To get this feature to work, the user must set another argument:

• allow_non_veg: Boolean (TRUE or FALSE). If TRUE, allow input to contain un-vegetated ground and un-vegetated water.

If allow_non_veg is true, the user can include "UNVEGETATED GROUND" or "UNVEGETATED WATER" along with plant species. They can also use acronyms "GROUND" or "WATER".

```
#print transect to view structure of data
transect_unveg <- data.frame(acronym = c("GROUND", "ABEESC", "ABIBAL", "AMMBRE",</pre>
                                          "ANTELE", "WATER", "GROUND", "ABEESC",
                                          "ABIBAL", "AMMBRE"),
                             cover = c(60, 50, 4, 20, 30, 20, 20, 40, 7, 60),
                             quad_id = c(1, 1, 1, 1, 1, 2, 2, 2, 2, 2))
#plot summary of a transect (duplicates are allowed, unless they are in the same plot)
plot_summary(x = transect_unveg, key = "acronym", db = "michigan_2014",
             cover_class = "percent_cover",
             plot_id = "quad_id")
#> Species c("GROUND", "WATER", "GROUND") does not have a wetness coefficient. It will be omitted from
     quad_id species_richness native_species_richness mean_wetness
                                                                      mean_c
#> 1
          1
                                                    3
                                                          1.750000 5.750000
                            4
#> 2
           2
                            3
                                                    2
                                                           3.333333 4.3333333
                                      FQI native_FQI cover_FQI native_cover_FQI
     native_mean_c cover_mean_c
                       4.923077 11.500000 13.279056 9.846154
#> 1
          7.666667
                                                                        16.42241
#> 2
          6.500000
                       5.803738 7.505553
                                            9.192388 10.052370
                                                                        13.10786
   adjusted_FQI percent_ground_cover percent_water_cover
```

```
#> 1 66.39528 60 NA
#> 2 53.07228 20 20
```

Relative Functions

Relative functions calculate relative frequency, relative coverage, and relative importance for each species, phylognomic group, or family. fqacalc also contains a species summary function that produces a summary of each species' relative metrics in a data frame. Relative functions always allow duplicate species observations. If a plot ID column is indicated using the plot_id argument, duplicates will not be allowed if they occur in the same plot. Relative functions also always allow "ground" and "water" to be included.

Relative functions have one additional argument which tells the functions what to calculate the relative value of:

• col: A character string equal to 'species', 'family', or 'physiog'.

Relative functions do not distinguish between native and introduced.

```
#To calculate the relative value of a tree
#relative frequency
relative_frequency(transect, key = "acronym", db = "michigan_2014",
             col = "physiog")
     physiognomy relative_frequency
#> 1
                               37.5
            forb
           grass
                               37.5
#> 2
#> 3
                               25.0
            tree
#can also include bare ground and water in the data
\textit{\#here transect\_unveg is data containing ground and water defined previously}
relative_frequency(transect_unveg, key = "acronym", db = "michigan_2014",
              col = "physiog")
#>
            physiognomy relative frequency
#> 1 Unvegetated Ground
#> 2 Unvegetated Water
                                         10
#> 3
                                         30
                   forb
                  grass
                                         20
#> 4
#> 5
                                         20
                   tree
#relative cover
relative_cover(transect, key = "acronym", db = "michigan_2014",
               col = "family", cover_class = "percent_cover")
#>
            family relative_cover
#> 1
         Malvaceae
                        37.344398
#> 2 Melanthiaceae
                        12.448133
#> 3
         Pinaceae
                        4.564315
#> 4
          Poaceae
                        45.643154
#relative importance
relative importance(transect, key = "acronym", db = "michigan 2014",
                    col = "species", cover_class = "percent_cover")
#>
                        name relative importance
#> 1 ABELMOSCHUS ESCULENTUS
                                         31.17220
```

```
ABIES BALSAMEA
                                     14.78216
#> 3 AMMOPHILA BREVILIGULATA
                                     41.57158
          ANTICLEA ELEGANS
#> 4
                                     12.47407
#species summary (including ground and water)
species_summary(transect_unveg, key = "acronym", db = "michigan_2014",
              cover_class = "percent_cover")
    acronym
                             name nativity c w frequency coverage
#> 1 ABEESC ABELMOSCHUS ESCULENTUS introduced 0 5
                                                        2
#> 2 ABIBAL
                                    native 3 0
                                                        2
             ABIES BALSAMEA
                                                                11
#> 3 AMMBRE AMMOPHILA BREVILIGULATA
                                     native 10 5
                                                        2
                                                                80
#> 4 ANTELE ANTICLEA ELEGANS
                                     native 10 -3
                                                        1
                                                                30
UNVEGETATED GROUND
                                       <NA> O NA
                                                        2
                                                                80
                                       <NA> O NA
                                                         1
                                                                20
   relative_frequency relative_cover relative_importance
#> 1
                  20
                          28.938907
                                             24.469453
#> 2
                   20
                           3.536977
                                             11.768489
#> 3
                   20
                          25.723473
                                             22.861736
#> 4
                   10
                           9.646302
                                              9.823151
#> 5
                   20
                          25.723473
                                             22.861736
#> 6
                   10
                           6.430868
                                             8.215434
#physiognomy summary (including ground and water)
physiog_summary(transect_unveg, key = "acronym", db = "michigan_2014",
              cover_class = "percent_cover")
#>
           physiognomy frequency coverage relative_frequency relative_cover
#> 1
                 forb
                            3
                                  120
                                                      30
                                                              38.585209
#> 2
                             2
                                    80
                                                      20
                                                              25.723473
                grass
#> 3
                 tree
                             2
                                    11
                                                      20
                                                              3.536977
                             2
                                    80
                                                      20
                                                              25.723473
#> 4 Unvegetated Ground
#> 5 Unvegetated Water
                            1
                                    20
                                                      10
                                                              6.430868
   relative_importance
#> 1
             34.292605
#> 2
             22.861736
#> 3
             11.768489
#> 4
             22.861736
              8.215434
#> 5
```

Wetness metric

fqacalc has one wetness metric function called mean_w, which calculates the mean wetness coefficient. The wetness coefficient is based off of the wetland indicator status. Negative wetness coefficients indicate a stronger affinity for wetlands, while positive wetness coefficients indicate an affinity for uplands.

mean_w can optionally include species without a C value, as long as they do have a wetness coefficient.

```
#mean wetness
mean_w(crooked_island, key = "acronym", db = "michigan_2014", allow_no_c = FALSE)
#> [1] 0.7142857
```

The End