Spectra-trait PLSR example using leaf-level spectra and leaf mass per area (LMA) data from 36 species growing in Rosa rugosa invaded coastal grassland communities in Belgium

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Overview

This is an R Markdown Notebook to illustrate how to retrieve a dataset from the EcoSIS spectral database, choose the "optimal" number of plsr components, and fit a plsr model for leaf-mass area (LMA)

Getting Started

Installation

```
## Loading required package: usethis
## Attaching package: 'remotes'
## The following objects are masked from 'package:devtools':
##
##
       dev_package_deps, install_bioc, install_bitbucket, install_cran,
       install_deps, install_dev, install_git, install_github,
##
##
       install_gitlab, install_local, install_svn, install_url,
       install_version, update_packages
##
  The following object is masked from 'package:usethis':
##
##
##
       git_credentials
##
## Attaching package: 'pls'
## The following object is masked from 'package:stats':
##
##
       loadings
##
## 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
## here() starts at /Users/sserbin/Data/GitHub/PLSR_for_plant_trait_prediction
```

```
##
## Attaching package: 'gridExtra'
## The following object is masked from 'package:dplyr':
##
## combine
```

Setup other functions and options

```
### Setup other functions and options
github_dir <- file.path(here::here(), "R_Scripts")</pre>
source from gh <- TRUE
if (source_from_gh) {
  # Source helper functions from GitHub
  print("*** GitHub hash of functions.R file:")
  devtools::source_url("https://raw.githubusercontent.com/TESTgroup-BNL/PLSR_for_plant_trait_prediction
} else {
  functions <- file.path(github_dir, "functions.R")</pre>
  source(functions)
## [1] "*** GitHub hash of functions.R file:"
## SHA-1 hash of file is 9eddd103e3a468dc4578c28c341b0275f244c04a
# not in
`%notin%` <- Negate(`%in%`)</pre>
# Script options
pls::pls.options(plsralg = "oscorespls")
pls::pls.options("plsralg")
## $plsralg
## [1] "oscorespls"
# Default par options
opar <- par(no.readonly = T)</pre>
# What is the target variable?
inVar <- "LMA_g_m2"</pre>
# What is the source dataset from EcoSIS?
ecosis id <- "9db4c5a2-7eac-4e1e-8859-009233648e89"
# Specify output directory, output_dir
# Options:
# tempdir - use a OS-specified temporary directory
# user defined PATH - e.g. "~/scratch/PLSR"
output_dir <- "tempdir"</pre>
```

Set working directory (scratch space)

[1] "/private/var/folders/xp/h3k9vf3n2jx181ts786_yjrn9c2gjq/T/RtmpFuoMi6"

Grab data from EcoSIS

```
print(paste0("Output directory: ",getwd())) # check wd
URL: https://ecosis.org/package/fresh-leaf-spectra-to-estimate-lma-over-neon-domains-in-
eastern-united-states
## [1] "Output directory: /Users/sserbin/Data/GitHub/PLSR_for_plant_trait_prediction/vignettes"
### Get source dataset from EcoSIS
dat_raw <- get_ecosis_data(ecosis_id = ecosis_id)</pre>
## [1] "**** Downloading Ecosis data ****"
## Downloading data...
## Parsed with column specification:
## cols(
##
     .default = col_double(),
##
     `Latin Species` = col_character(),
##
     ids = col_character(),
     `plot code` = col_character(),
##
##
     `species code` = col_character()
## )
## See spec(...) for full column specifications.
## Download complete!
head(dat_raw)
## # A tibble: 6 x 2,164
     `Cw/EWT (cm3/cm~ `Latin Species` `Leaf area (mm2~ `Leaf calcium c~
##
##
                <dbl> <chr>
                                                   <dbl>
                                                                    <db1>
                                                                   0.0291
## 1
              0.00887 Arrhenatherum ~
                                                    696.
## 2
              0.00824 Bromus sterilis
                                                    447.
                                                                   0.0230
              0.0280 Jacobaea vulga~
## 3
                                                   2418.
                                                                   0.0950
              0.0106 Rubus caesius
## 4
                                                   5719.
                                                                   0.0700
## 5
              0.00851 Arrhenatherum ~
                                                    671.
                                                                   0.0286
              0.0153 Crepis capilla~
                                                   1401.
                                                                   0.0470
## #
    ... with 2,160 more variables: `Leaf magnesium content per leaf area
## #
       (mg/mm2) \ dbl>, \ Leaf mass per area (g/cm2) \ dbl>, \ Leaf nitrogen content
## #
       per leaf area (mg/mm2) \(^2\) <dbl>, \(^1\) Leaf phosphorus content per leaf area
## #
       (mg/mm2) \(^ \cdot \), \(^ \cdot \) potassium content per leaf area (mg/mm2) \(^ \cdot \),
## #
       `Plant height vegetative (cm)` <dbl>, ids <chr>, `plot code` <chr>,
## #
       `species code` <chr>, `350` <dbl>, `351` <dbl>, `352` <dbl>, `353` <dbl>,
## #
       `354` <dbl>, `355` <dbl>, `356` <dbl>, `357` <dbl>, `358` <dbl>,
       `359` <dbl>, `360` <dbl>, `361` <dbl>, `362` <dbl>, `363` <dbl>,
## #
       `364` <dbl>, `365` <dbl>, `366` <dbl>, `367` <dbl>, `368` <dbl>,
## #
## #
       `369` <dbl>, `370` <dbl>, `371` <dbl>, `372` <dbl>, `373` <dbl>,
       `374` <dbl>, `375` <dbl>, `376` <dbl>, `377` <dbl>, `378` <dbl>,
       `379` <dbl>, `380` <dbl>, `381` <dbl>, `382` <dbl>, `383` <dbl>,
## #
       `384` <dbl>, `385` <dbl>, `386` <dbl>, `387` <dbl>, `388` <dbl>,
## #
       `389` <dbl>, `390` <dbl>, `391` <dbl>, `392` <dbl>, `393` <dbl>,
## #
       `394` <dbl>, `395` <dbl>, `396` <dbl>, `397` <dbl>, `398` <dbl>,
       `399` <dbl>, `400` <dbl>, `401` <dbl>, `402` <dbl>, `403` <dbl>,
## #
       `404` <dbl>, `405` <dbl>, `406` <dbl>, `407` <dbl>, `408` <dbl>,
```

```
`409` <dbl>, `410` <dbl>, `411` <dbl>, `412` <dbl>, `413` <dbl>,
## #
       `414` <dbl>, `415` <dbl>, `416` <dbl>, `417` <dbl>, `418` <dbl>,
       `419` <dbl>, `420` <dbl>, `421` <dbl>, `422` <dbl>, `423` <dbl>,
## #
## #
       `424` <dbl>, `425` <dbl>, `426` <dbl>, `427` <dbl>, `428` <dbl>,
       `429` <dbl>, `430` <dbl>, `431` <dbl>, `432` <dbl>, `433` <dbl>,
## #
## #
       `434` <dbl>, `435` <dbl>, `436` <dbl>, `437` <dbl>, `438` <dbl>,
       `439` <dbl>, `440` <dbl>, ...
names(dat raw)[1:40]
    [1] "Cw/EWT (cm3/cm2)"
    [2] "Latin Species"
##
   [3] "Leaf area (mm2)"
##
   [4] "Leaf calcium content per leaf area (mg/mm2)"
    [5] "Leaf magnesium content per leaf area (mg/mm2)"
##
##
    [6]
       "Leaf mass per area (g/cm2)"
##
   [7] "Leaf nitrogen content per leaf area (mg/mm2)"
   [8] "Leaf phosphorus content per leaf area (mg/mm2)"
   [9] "Leaf potassium content per leaf area (mg/mm2)"
## [10] "Plant height vegetative (cm)"
## [11] "ids"
## [12] "plot code"
## [13] "species code"
## [14] "350"
## [15] "351"
## [16] "352"
## [17] "353"
## [18] "354"
## [19] "355"
## [20] "356"
## [21] "357"
## [22] "358"
## [23] "359"
## [24] "360"
## [25] "361"
## [26] "362"
## [27] "363"
## [28] "364"
## [29]
       "365"
## [30] "366"
## [31] "367"
## [32]
       "368"
## [33] "369"
## [34] "370"
## [35] "371"
## [36] "372"
## [37] "373"
## [38] "374"
## [39] "375"
```

[40] "376"

Create full plsr dataset

```
### Create plsr dataset
Start.wave <- 500
End.wave \leftarrow 2400
wv <- seq(Start.wave, End.wave, 1)</pre>
Spectra <- as.matrix(dat_raw[,names(dat_raw) %in% wv])</pre>
colnames(Spectra) <- c(paste0("Wave_",wv))</pre>
sample_info <- dat_raw[,names(dat_raw) %notin% seq(350,2500,1)]</pre>
head(sample_info)
## # A tibble: 6 x 13
     `Cw/EWT (cm3/cm~ `Latin Species` `Leaf area (mm2~ `Leaf calcium c~
##
                <dbl> <chr>
                                                   <dbl>
                                                                     <dbl>
## 1
              0.00887 Arrhenatherum ~
                                                    696.
                                                                    0.0291
## 2
              0.00824 Bromus sterilis
                                                    447.
                                                                    0.0230
## 3
              0.0280 Jacobaea vulga~
                                                   2418.
                                                                    0.0950
## 4
              0.0106 Rubus caesius
                                                   5719.
                                                                    0.0700
## 5
              0.00851 Arrhenatherum ~
                                                    671.
                                                                    0.0286
              0.0153 Crepis capilla~
                                                   1401.
                                                                    0.0470
## # ... with 9 more variables: `Leaf magnesium content per leaf area
       (mg/mm2) <dbl>, Leaf mass per area (g/cm2) <dbl>, Leaf nitrogen content
## #
       per leaf area (mg/mm2)` <dbl>, `Leaf phosphorus content per leaf area
       (mg/mm2)` <dbl>, `Leaf potassium content per leaf area (mg/mm2)` <dbl>,
       `Plant height vegetative (cm)` <dbl>, ids <chr>, `plot code` <chr>,
## #
       `species code` <chr>
sample_info2 <- sample_info %>%
  select(Plant_Species=`Latin Species`,Species_Code=`species code`,Plot=`plot code`,
         LMA_g_cm2=`Leaf mass per area (g/cm2)`)
sample_info2 <- sample_info2 %>%
 mutate(LMA_g_m2=LMA_g_cm2*10000)
head(sample_info2)
## # A tibble: 6 x 5
    Plant Species
                            {\tt Species\_Code\ Plot} \quad {\tt LMA\_g\_cm2} \ {\tt LMA\_g\_m2}
##
     <chr>>
##
                            <chr>
                                         <chr>
                                                    <dbl>
                                                              <dbl>
                                         DC1
                                                  0.00342
## 1 Arrhenatherum elatius Arrela
                                                               34.2
                                         DC1
                                                  0.00282
## 2 Bromus sterilis
                                                               28.2
                            Broste
                                         DC1
                                                  0.00417
## 3 Jacobaea vulgaris
                            Jacvul
                                                               41.7
## 4 Rubus caesius
                                         DC1
                                                  0.00566
                                                               56.6
                            Rubcae
## 5 Arrhenatherum elatius Arrela
                                         DC2
                                                  0.00361
                                                               36.1
## 6 Crepis capillaris
                                         DC2
                                                               28.3
                            Creves
                                                  0.00283
plsr_data <- data.frame(sample_info2,Spectra)</pre>
rm(sample_info, sample_info2, Spectra)
```

Example data cleaning.

Create cal/val datasets

```
method <- "dplyr" #base/dplyr
# base R - a bit slow
# dplyr - much faster
split_data <- create_data_split(approach=method, split_seed=7529075, prop=0.8,
                                group variables="Species Code")
names(split_data)
## [1] "cal_data" "val_data"
cal.plsr.data <- split_data$cal_data
head(cal.plsr.data)[1:8]
##
         Plant_Species Species_Code Plot LMA_g_cm2 LMA_g_m2 Wave_500 Wave_501
## 1 Ammophila arenaria
                              Ammare MC2 0.01679492 167.9492 0.135785 0.13685
## 2 Ammophila arenaria
                              Ammare WC3 0.01844376 184.4376 0.151750 0.15275
## 3 Ammophila arenaria
                              Ammare MC4 0.02030190 203.0190 0.156830 0.15790
                              Ammare ZC2 0.01591894 159.1894 0.144450 0.14525
## 4 Ammophila arenaria
## 5 Ammophila arenaria
                              Ammare ZC1 0.01483469 148.3469 0.147665 0.14910
## 6 Ammophila arenaria
                              Ammare ZC3 0.01802409 180.2409 0.130885 0.13175
    Wave 502
## 1 0.138150
## 2 0.154150
## 3 0.159065
## 4 0.146220
## 5 0.150330
## 6 0.132750
val.plsr.data <- split_data$val_data</pre>
head(val.plsr.data)[1:8]
##
            Plant_Species Species_Code Plot
                                              LMA_g_cm2 LMA_g_m2
                                                                    Wave_500
                                Jacvul WC2 0.003551614 35.51614 0.06736887
## 184 Jacobaea vulgaris
## 185 Potentilla reptans
                                Potrep WC2 0.005586320 55.86320 0.07125000
## 186
            Rubus caesius
                                Rubcae WC2 0.005803902 58.03902 0.05993560
## 187
           Urtica dioica
                                Urtdio WC2 0.005215705 52.15705 0.06508300
                                Ammare WC3 0.018443757 184.43757 0.15175000
## 188 Ammophila arenaria
                                Jacvul WC3 0.004980002 49.80002 0.06805547
## 189
      Jacobaea vulgaris
##
         Wave_501
                    Wave 502
## 184 0.06870667 0.07014220
## 185 0.07235000 0.07368350
## 186 0.06162000 0.06352233
## 187 0.06625000 0.06758350
## 188 0.15275000 0.15415000
## 189 0.06938000 0.07093553
rm(split_data)
# Datasets:
print(paste("Cal observations: ",dim(cal.plsr.data)[1],sep=""))
## [1] "Cal observations: 183"
print(paste("Val observations: ",dim(val.plsr.data)[1],sep=""))
## [1] "Val observations: 73"
```

```
cal_hist_plot <- qplot(cal.plsr.data[,paste0(inVar)],geom="histogram",</pre>
                        main = paste0("Calibration Histogram for ",inVar),
                        xlab = pasteO(inVar),ylab = "Count",fill=I("grey50"),col=I("black"),alpha=I(.7))
val_hist_plot <- qplot(val.plsr.data[,paste0(inVar)],geom="histogram",</pre>
                        main = paste0("Validation Histogram for ",inVar),
                        xlab = paste0(inVar),ylab = "Count",fill=I("grey50"),col=I("black"),alpha=I(.7))
histograms <- grid.arrange(cal_hist_plot, val_hist_plot, ncol=2)
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
## `stat bin()` using `bins = 30`. Pick better value with `binwidth`.
                                                  Validation Histogram for LMA_g_m2
   Calibration Histogram for LMA_g_m2
 20
 15
                     LMA_g_m2
                                                                     LMA_g_m2
ggsave(filename = file.path(outdir,paste0(inVar,"_Cal_Val_Histograms.png")), plot = histograms,
       device="png", width = 30, height = 12, units = "cm", dpi = 300)
# output cal/val data
write.csv(cal.plsr.data,file=file.path(outdir,paste0(inVar,'_Cal_PLSR_Dataset.csv')),row.names=FALSE)
write.csv(val.plsr.data,file=file.path(outdir,paste0(inVar,'_Val_PLSR_Dataset.csv')),row.names=FALSE)
```

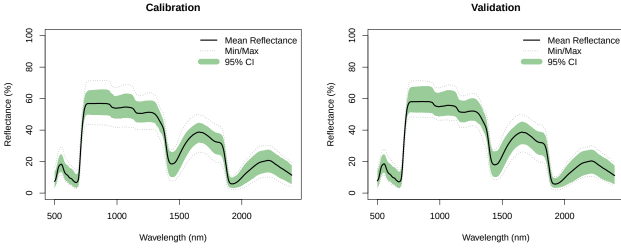
Create calibration and validation PLSR datasets

```
### Format PLSR data for model fitting
cal_spec <- as.matrix(cal.plsr.data[, which(names(cal.plsr.data) %in% paste0("Wave_",wv))])</pre>
cal.plsr.data <- data.frame(cal.plsr.data[, which(names(cal.plsr.data) %notin% paste0("Wave_",wv))],</pre>
                             Spectra=I(cal_spec))
head(cal.plsr.data)[1:5]
          Plant_Species Species_Code Plot LMA_g_cm2 LMA_g_m2
## 1 Ammophila arenaria
                                      MC2 0.01679492 167.9492
                               Ammare
## 2 Ammophila arenaria
                               Ammare
                                       WC3 0.01844376 184.4376
## 3 Ammophila arenaria
                               Ammare MC4 0.02030190 203.0190
## 4 Ammophila arenaria
                               Ammare ZC2 0.01591894 159.1894
## 5 Ammophila arenaria
                               Ammare ZC1 0.01483469 148.3469
                               Ammare ZC3 0.01802409 180.2409
## 6 Ammophila arenaria
val_spec <- as.matrix(val.plsr.data[, which(names(val.plsr.data) %in% paste0("Wave_",wv))])</pre>
val.plsr.data <- data.frame(val.plsr.data[, which(names(val.plsr.data) %notin% paste0("Wave_",wv))],</pre>
                             Spectra=I(val spec))
head(val.plsr.data)[1:5]
```

```
##
            Plant_Species Species_Code Plot
                                             LMA_g_cm2 LMA_g_m2
## 184
       Jacobaea vulgaris
                               Jacvul
                                       WC2 0.003551614
                                                        35.51614
## 185 Potentilla reptans
                               Potrep
                                       WC2 0.005586320
                                                        55.86320
## 186
            Rubus caesius
                               Rubcae
                                       WC2 0.005803902
                                                        58.03902
## 187
            Urtica dioica
                               Urtdio WC2 0.005215705 52.15705
## 188 Ammophila arenaria
                               Ammare
                                       WC3 0.018443757 184.43757
  189
        Jacobaea vulgaris
                                Jacvul
                                       WC3 0.004980002 49.80002
```

plot cal and val spectra

```
par(mfrow=c(1,2)) # B, L, T, R
f.plot.spec(Z=cal.plsr.data$Spectra,wv=seq(Start.wave,End.wave,1),plot_label="Calibration")
f.plot.spec(Z=val.plsr.data$Spectra,wv=seq(Start.wave,End.wave,1),plot_label="Validation")
```



```
## quartz_off_screen
## 3
dev.off();
## pdf
## 2
par(mfrow=c(1,1))
```

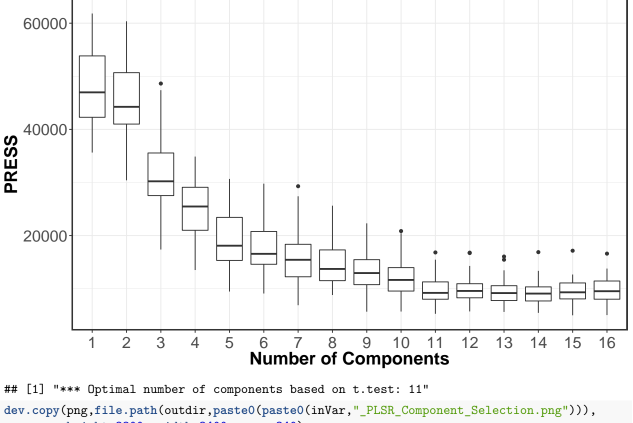
Use permutation to determine the optimal number of components

```
### Use permutation to determine the optimal number of components
if(grepl("Windows", sessionInfo()$running)){
  pls.options(parallel = NULL)
} else {
  pls.options(parallel = parallel::detectCores()-1)
}
```

```
method <- "firstMin" #pls, firstPlateau, firstMin
random_seed <- 7529075
seg <- 80
maxComps <- 16
iterations <- 50
prop <- 0.70
if (method=="pls") {
 nComps <- find_optimal_components(dataset=cal.plsr.data, method=method, maxComps=maxComps,
                                    seg=seg, random_seed=random_seed)
  print(paste0("*** Optimal number of components: ", nComps))
} else {
  nComps <- find_optimal_components(dataset=cal.plsr.data, method=method, maxComps=maxComps,
                                    iterations=iterations, seg=seg, prop=prop,
                                    random_seed=random_seed)
}
## [1] "*** Running permutation test. Please hang tight, this can take awhile ***"
## [1] "Options: 16 50 80 0.7"
## Running interation 1
## Running interation 2
## Running interation 3
## Running interation 4
## Running interation 5
## Running interation 6
## Running interation 7
## Running interation 8
## Running interation 9
## Running interation 10
## Running interation 11
## Running interation 12
## Running interation 13
## Running interation 14
## Running interation 15
## Running interation 16
## Running interation 17
## Running interation 18
## Running interation 19
## Running interation 20
## Running interation 21
## Running interation 22
## Running interation 23
## Running interation 24
```

```
## Running interation 25
```

- ## Running interation 26
- ## Running interation 27
- ## Running interation 28
- ## Running interation 29
- ## Running interation 30
- ## Running interation 31
- ## Running interation 32
- ## Running interation 33
- **O**
- ## Running interation 34
- ## Running interation 35
- ## Running interation 36
- ## Running interation 37
- ## Running interation 38
- ## Running interation 39
- ## Running interation 40
- ## Running interation 41
- ## Running interation 42
- ## Running interation 43
- ## Running interation 44
- ## Running interation 45
- ## Running interation 46
- ## Running interation 47
- ## Running interation 48
- ## Running interation 49
- ## Running interation 50
- ## No id variables; using all as measure variables



Fit final model

```
### Fit final model - using leave-one-out cross validation
plsr.out <- plsr(as.formula(paste(inVar,"~","Spectra")),scale=FALSE,ncomp=nComps,validation="LOO",</pre>
                  trace=FALSE,data=cal.plsr.data)
fit <- plsr.out$fitted.values[,1,nComps]</pre>
pls.options(parallel = NULL)
# External validation fit stats
par(mfrow=c(1,2)) # B, L, T, R
RMSEP(plsr.out, newdata = val.plsr.data)
## (Intercept)
                     1 comps
                                   2 comps
                                                3 comps
                                                              4 comps
                                                                            5 comps
##
         37.79
                       32.71
                                     30.36
                                                   23.51
                                                                21.58
                                                                              18.46
##
       6 comps
                     7 comps
                                   8 comps
                                                             10 comps
                                                                           11 comps
                                                9 comps
##
         15.89
                       15.44
                                     15.52
                                                   15.19
                                                                15.14
                                                                              13.68
```

```
plot(RMSEP(plsr.out,estimate=c("test"),newdata = val.plsr.data), main="MODEL RMSEP",
     xlab="Number of Components", ylab="Model Validation RMSEP", lty=1, col="black", cex=1.5, lwd=2)
box(1wd=2.2)
R2(plsr.out, newdata = val.plsr.data)
## (Intercept)
                      1 comps
                                     2 comps
                                                    3 comps
                                                                  4 comps
                                                                                 5 comps
      -0.06195
##
                      0.20461
                                     0.31467
                                                    0.58911
                                                                  0.65365
                                                                                 0.74649
##
       6 comps
                      7 comps
                                     8 comps
                                                    9 comps
                                                                 10 comps
                                                                                11 comps
       0.81222
                      0.82276
                                     0.82084
                                                    0.82841
##
                                                                  0.82945
                                                                                 0.86090
plot(R2(plsr.out,estimate=c("test"),newdata = val.plsr.data), main="MODEL R2",
     xlab="Number of Components", ylab="Model Validation R2", lty=1, col="black", cex=1.5, lwd=2)
box(1wd=2.2)
                     MODEL RMSEP
                                                                            MODEL R2
                                                        0.8
    35
Model Validation RMSEP
                                                    Model Validation R2
    30
                                                        0.4
    25
                                                        0.2
    20
                                                        0.0
    15
               2
                             6
                                    8
                                                                   2
                                                                                        8
        0
                                           10
                                                            0
                                                                                               10
                    Number of Components
                                                                        Number of Components
dev.copy(png,file.path(outdir,paste0(paste0(inVar,"_Validation_RMSEP_R2_by_Component.png"))),
          height=2800, width=4800, res=340)
## quartz_off_screen
                     3
dev.off();
## pdf
##
par(opar)
```

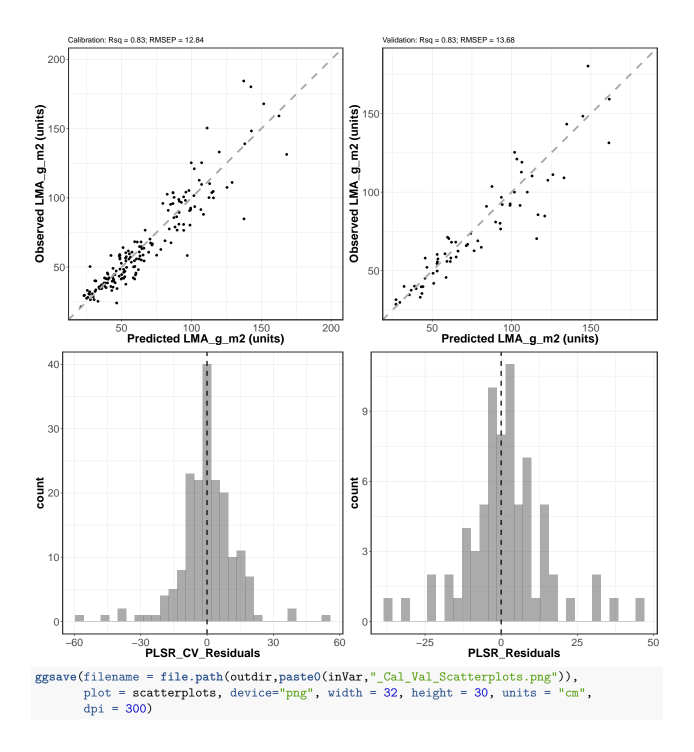
PLSR fit observed vs. predicted plot data

##

Plant_Species Species_Code Plot LMA_g_cm2 LMA_g_m2 PLSR_Predicted

```
## 1 Ammophila arenaria
                              Ammare MC2 0.01679492 167.9492
                                                                      154.1892
                              Ammare WC3 0.01844376 184.4376
                                                                      147.0878
## 2 Ammophila arenaria
## 3 Ammophila arenaria
                              Ammare MC4 0.02030190 203.0190
                                                                      153.8674
## 4 Ammophila arenaria
                              Ammare ZC2 0.01591894 159.1894
                                                                      161.6047
## 5 Ammophila arenaria
                              Ammare ZC1 0.01483469 148.3469
                                                                      144.9268
                              Ammare ZC3 0.01802409 180.2409
                                                                      148.2100
## 6 Ammophila arenaria
     PLSR CV Predicted PLSR CV Residuals
## 1
              151.7161
                              -16.233027
## 2
              137.3863
                              -47.051273
## 3
              144.2584
                              -58.760574
              162.6250
                                 3.435614
## 5
                               -5.436767
              142.9101
## 6
              142.5160
                              -37.724928
cal.R2 <- round(pls::R2(plsr.out)[[1]][nComps],2)</pre>
cal.RMSEP <- round(sqrt(mean(cal.plsr.output$PLSR_CV_Residuals^2)),2)</pre>
val.plsr.output <- data.frame(val.plsr.data[, which(names(val.plsr.data) %notin% "Spectra")],
                              PLSR_Predicted=as.vector(predict(plsr.out,
                                                                newdata = val.plsr.data,
                                                                 ncomp=nComps, type="response")[,,1]))
val.plsr.output <- val.plsr.output %>%
  mutate(PLSR_Residuals = PLSR_Predicted-get(inVar))
head(val.plsr.output)
##
          Plant_Species Species_Code Plot
                                             LMA_g_cm2 LMA_g_m2 PLSR_Predicted
                              Jacvul WC2 0.003551614 35.51614
                                                                        43.51586
## 1 Jacobaea vulgaris
## 2 Potentilla reptans
                              Potrep WC2 0.005586320 55.86320
                                                                        61.41726
## 3
          Rubus caesius
                                                                        45.55789
                              Rubcae WC2 0.005803902 58.03902
## 4
          Urtica dioica
                              Urtdio WC2 0.005215705 52.15705
                                                                        46.65139
## 5 Ammophila arenaria
                              Ammare WC3 0.018443757 184.43757
                                                                       147.08781
      Jacobaea vulgaris
                              Jacvul WC3 0.004980002 49.80002
                                                                        53.09532
##
     PLSR_Residuals
## 1
           7.999719
## 2
           5.554059
## 3
         -12.481126
## 4
          -5.505664
## 5
         -37.349758
## 6
           3.295298
val.R2 <- round(pls::R2(plsr.out,newdata=val.plsr.data)[[1]][nComps],2)</pre>
val.RMSEP <- round(sqrt(mean(val.plsr.output$PLSR_Residuals^2)),2)</pre>
rng_quant <- quantile(cal.plsr.output[,inVar], probs = c(0.001, 0.999))</pre>
cal_scatter_plot <- ggplot(cal.plsr.output, aes(x=PLSR_CV_Predicted, y=get(inVar))) +</pre>
  theme_bw() + geom_point() + geom_abline(intercept = 0, slope = 1, color="dark grey",
                                           linetype="dashed", size=1.5) + xlim(rng quant[1],
                                                                                rng_quant[2]) +
  ylim(rng_quant[1], rng_quant[2]) +
  labs(x=paste0("Predicted ", paste(inVar), " (units)"),
       y=paste0("Observed ", paste(inVar), " (units)"),
       title=paste0("Calibration: ", paste0("Rsq = ", cal.R2), "; ", paste0("RMSEP = ",
                                                                              cal.RMSEP))) +
  theme(axis.text=element_text(size=18), legend.position="none",
        axis.title=element_text(size=20, face="bold"),
```

```
axis.text.x = element_text(angle = 0, vjust = 0.5),
        panel.border = element rect(linetype = "solid", fill = NA, size=1.5))
cal_resid_histogram <- ggplot(cal.plsr.output, aes(x=PLSR_CV_Residuals)) +</pre>
  geom_histogram(alpha=.5, position="identity") +
  geom_vline(xintercept = 0, color="black",
             linetype="dashed", size=1) + theme_bw() +
  theme(axis.text=element text(size=18), legend.position="none",
        axis.title=element text(size=20, face="bold"),
        axis.text.x = element_text(angle = 0, vjust = 0.5),
        panel.border = element_rect(linetype = "solid", fill = NA, size=1.5))
rng_quant <- quantile(val.plsr.output[,inVar], probs = c(0.001, 0.999))</pre>
val_scatter_plot <- ggplot(val.plsr.output, aes(x=PLSR_Predicted, y=get(inVar))) +</pre>
  theme_bw() + geom_point() + geom_abline(intercept = 0, slope = 1, color="dark grey",
                                          linetype="dashed", size=1.5) + xlim(rng_quant[1],
                                                                               rng_quant[2]) +
  ylim(rng_quant[1], rng_quant[2]) +
  labs(x=paste0("Predicted ", paste(inVar), " (units)"),
       y=paste0("Observed ", paste(inVar), " (units)"),
       title=paste0("Validation: ", paste0("Rsq = ", val.R2), "; ", paste0("RMSEP = ",
                                                                            val.RMSEP))) +
  theme(axis.text=element text(size=18), legend.position="none",
        axis.title=element_text(size=20, face="bold"),
        axis.text.x = element text(angle = 0, vjust = 0.5),
        panel.border = element_rect(linetype = "solid", fill = NA, size=1.5))
val_resid_histogram <- ggplot(val.plsr.output, aes(x=PLSR_Residuals)) +</pre>
  geom_histogram(alpha=.5, position="identity") +
  geom_vline(xintercept = 0, color="black",
            linetype="dashed", size=1) + theme_bw() +
  theme(axis.text=element_text(size=18), legend.position="none",
        axis.title=element_text(size=20, face="bold"),
        axis.text.x = element_text(angle = 0, vjust = 0.5),
        panel.border = element_rect(linetype = "solid", fill = NA, size=1.5))
# plot cal/val side-by-side
scatterplots <- grid.arrange(cal_scatter_plot, val_scatter_plot, cal_resid_histogram,</pre>
                             val_resid_histogram, nrow=2, ncol=2)
## Warning: Removed 6 rows containing missing values (geom_point).
## Warning: Removed 6 rows containing missing values (geom_point).
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
```



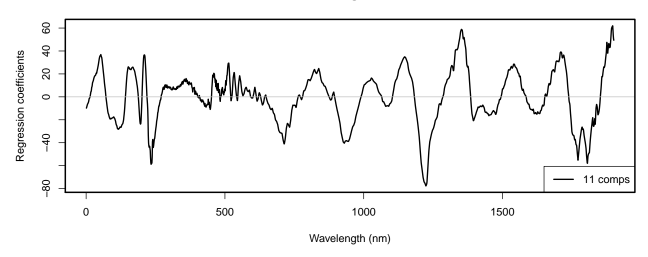
Generate Coefficient and VIP plots

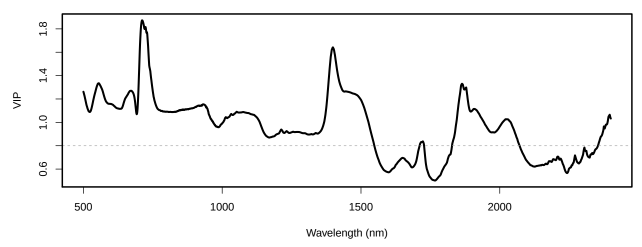
```
vips <- VIP(plsr.out)[nComps,]

par(mfrow=c(2,1))
plot(plsr.out, plottype = "coef",xlab="Wavelength (nm)",
        ylab="Regression coefficients",legendpos = "bottomright",
        ncomp=nComps,lwd=2)
box(lwd=2.2)</pre>
```

```
plot(seq(Start.wave,End.wave,1),vips,xlab="Wavelength (nm)",ylab="VIP",cex=0.01)
lines(seq(Start.wave,End.wave,1),vips,lwd=3)
abline(h=0.8,lty=2,col="dark grey")
box(lwd=2.2)
```

LMA_g_m2





```
## quartz_off_screen
## 3
dev.off();
```

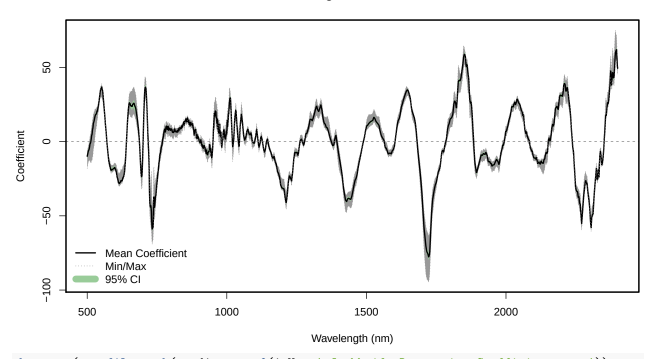
pdf ## 2

Jackknife validation

```
if(grepl("Windows", sessionInfo()$running)){
  pls.options(parallel =NULL)
} else {
  pls.options(parallel = parallel::detectCores()-1)
jk.plsr.out <- pls::plsr(as.formula(paste(inVar, "~", "Spectra")), scale=FALSE,
                         center=TRUE, ncomp=nComps, validation="LOO", trace=FALSE,
                         jackknife=TRUE,
                         data=cal.plsr.data)
pls.options(parallel = NULL)
Jackknife_coef <- f.coef.valid(plsr.out = jk.plsr.out, data_plsr = cal.plsr.data,</pre>
                               ncomp = nComps)
Jackknife intercept <- Jackknife coef[1,,,]</pre>
Jackknife_coef <- Jackknife_coef[2:dim(Jackknife_coef)[1],,,]</pre>
interval <-c(0.025, 0.975)
Jackknife_Pred <- val.plsr.data$Spectra %*% Jackknife_coef +</pre>
  matrix(rep(Jackknife_intercept, length(val.plsr.data[,inVar])), byrow=TRUE,
         ncol=length(Jackknife_intercept))
Interval Conf <- apply(X = Jackknife Pred, MARGIN = 1, FUN = quantile,</pre>
                       probs=c(interval[1], interval[2]))
Interval_Pred <- apply(X = Jackknife_Pred, MARGIN = 1, FUN = quantile,</pre>
                       probs=c(interval[1], interval[2]))
sd_mean <- apply(X = Jackknife_Pred, MARGIN = 1, FUN =sd)</pre>
sd_res <- sd(val.plsr.output$PLSR_Residuals)</pre>
sd_tot <- sqrt(sd_mean^2+sd_res^2)</pre>
val.plsr.output$LCI <- Interval_Pred[1,]</pre>
val.plsr.output$UCI <- Interval_Pred[2,]</pre>
val.plsr.output$LPI <- val.plsr.output$PLSR_Predicted-1.96*sd_tot
val.plsr.output$UPI <- val.plsr.output$PLSR_Predicted+1.96*sd_tot
head(val.plsr.output)
##
          Plant Species Species Code Plot
                                             LMA_g_cm2 LMA_g_m2 PLSR_Predicted
## 1 Jacobaea vulgaris
                              Jacvul WC2 0.003551614 35.51614
                                                                        43.51586
## 2 Potentilla reptans
                              Potrep WC2 0.005586320 55.86320
                                                                        61.41726
## 3
          Rubus caesius
                              Rubcae WC2 0.005803902 58.03902
                                                                        45.55789
## 4
          Urtica dioica
                              Urtdio WC2 0.005215705 52.15705
                                                                        46.65139
## 5 Ammophila arenaria
                              Ammare WC3 0.018443757 184.43757
                                                                       147.08781
## 6 Jacobaea vulgaris
                              Jacvul WC3 0.004980002 49.80002
                                                                        53.09532
     PLSR_Residuals
                          LCI
                                    UCI
                                               LPI
                                                         UPI
##
## 1
           7.999719 42.58086 44.15724 16.70642 70.32530
## 2
           5.554059 60.10507 62.52674 34.59536 88.23916
         -12.481126 44.66849 48.22967 18.70489 72.41090
## 3
## 4
         -5.505664 45.70375 47.84938 19.82512 73.47765
## 5
         -37.349758 145.09309 148.61694 120.18052 173.99510
## 6
           3.295298 52.40880 53.97806 26.28498 79.90565
```

Jackknife coefficient plot

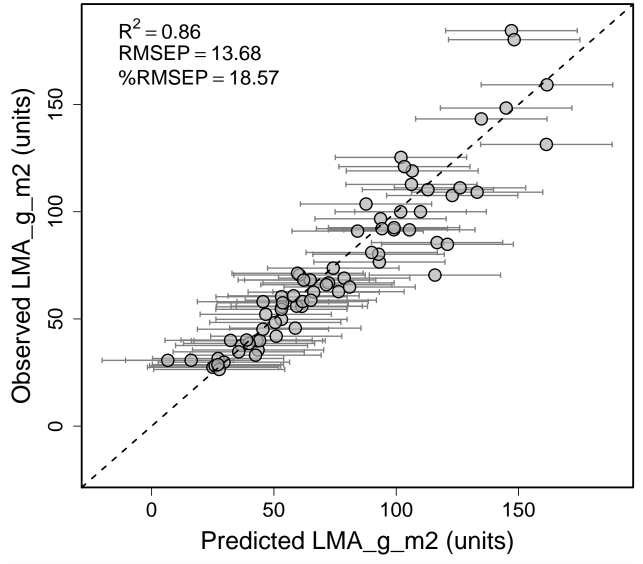
Jackknife regression coefficients



```
## quartz_off_screen
## 3
dev.off();
## pdf
## 2
```

Jackknife validation plot

```
lwd=1.6, xlim=c(rng_vals[1], rng_vals[2]), ylim=c(rng_vals[1], rng_vals[2]),
    err="x", pch=21, col="black", pt.bg=alpha("grey70",0.7), scol="grey50",
    cex=2, xlab=paste0("Predicted ", paste(inVar), " (units)"),
    ylab=paste0("Observed ", paste(inVar), " (units)"),
    cex.axis=1.5,cex.lab=1.8)
abline(0,1,lty=2,lw=2)
legend("topleft", legend=expr, bty="n", cex=1.5)
box(lwd=2.2)
```



```
## quartz_off_screen
## 3
dev.off();
```

pdf ## 2

Output jackknife results

```
out.jk.coefs <- data.frame(Iteration=seq(1,length(Jackknife_intercept),1),</pre>
                          Intercept=Jackknife_intercept,t(Jackknife_coef))
head(out.jk.coefs)[1:6]
##
        Iteration Intercept
                            Wave_500 Wave_501
                                                  Wave_502 Wave_503
               1 18.33909 -7.580446 -6.724083 -5.886226 -4.984744
## Seg 1
                2 21.22164 -8.574931 -7.084795 -6.255716 -5.384000
## Seg 2
## Seg 3
                3 19.63843 -18.104491 -17.260522 -16.154983 -14.960119
## Seg 4
               4 15.90905 -10.715594 -9.874766 -8.926979 -8.007834
## Seg 5
                5 17.51805 -8.952143 -8.305344 -7.136167 -6.221407
                6 12.18563 -7.702160 -7.128890 -6.532276 -5.840220
## Seg 6
write.csv(out.jk.coefs,file=file.path(outdir,
                                    pasteO(inVar,'_Jackkife_PLSR_Coefficients.csv')),
         row.names=FALSE)
```

Create core PLSR outputs

```
print(paste("Output directory: ", outdir))
## [1] "Output directory: /var/folders/xp/h3k9vf3n2jx181ts786_yjrn9c2gjq/T//RtmpFuoMi6"
# Observed versus predicted
write.csv(cal.plsr.output,file=file.path(outdir,
                                          pasteO(inVar,'_Observed_PLSR_CV_Pred_',
                                                 nComps,'comp.csv')),
          row.names=FALSE)
# Validation data
write.csv(val.plsr.output,file=file.path(outdir,
                                          paste0(inVar, '_Validation_PLSR_Pred_',
                                                 nComps,'comp.csv')),
          row.names=FALSE)
# Model coefficients
coefs <- coef(plsr.out,ncomp=nComps,intercept=TRUE)</pre>
write.csv(coefs,file=file.path(outdir,
                               pasteO(inVar,'_PLSR_Coefficients_',
                                       nComps,'comp.csv')),
          row.names=TRUE)
# PLSR VIP
write.csv(vips,file=file.path(outdir,
                              paste0(inVar, '_PLSR_VIPs_',
                                      nComps,'comp.csv')))
```

Confirm files were written to temp space

```
print("**** PLSR output files: ")
```

```
## [1] "**** PLSR output files: "
list.files(outdir)[grep(pattern = inVar, list.files(outdir))]
    [1] "LMA_g_m2_Cal_PLSR_Dataset.csv"
##
   [2] "LMA_g_m2_Cal_Val_Histograms.png"
##
    [3] "LMA_g_m2_Cal_Val_Scatterplots.png"
##
##
    [4] "LMA_g_m2_Cal_Val_Spectra.png"
   [5] "LMA_g_m2_Coefficient_VIP_plot.png"
##
    [6] "LMA_g_m2_Jackkife_PLSR_Coefficients.csv"
##
    [7] "LMA_g_m2_Jackknife_Regression_Coefficients.png"
##
   [8] "LMA_g_m2_Observed_PLSR_CV_Pred_11comp.csv"
##
   [9] "LMA_g_m2_PLSR_Coefficients_11comp.csv"
## [10] "LMA_g_m2_PLSR_Component_Selection.png"
## [11] "LMA_g_m2_PLSR_Validation_Scatterplot.png"
## [12] "LMA_g_m2_PLSR_VIPs_11comp.csv"
## [13] "LMA_g_m2_Val_PLSR_Dataset.csv"
## [14] "LMA_g_m2_Validation_PLSR_Pred_11comp.csv"
```

[15] "LMA_g_m2_Validation_RMSEP_R2_by_Component.png"