Spectra-trait PLSR example using leaf-level spectra and leaf mass per area (LMA) data from more than 40 species grassland species comprising both herbs and graminoids.

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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). In this example, the plants were cultivated in an outdoor setting in the botanical garden of the KIT using 40x40 cm pots with an standardized substrate. The data was measured on a weekly basis (the timestamp is included in the dataset).

### **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</pre>
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 d8af92cac6fa9c1337442724485b786577bc6259
# not in
`%notin%` <- Negate(`%in%`)
# 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 <- "SLA_g_cm"</pre>
# What is the source dataset from EcoSIS?
ecosis id <- "3cf6b27e-d80e-4bc7-b214-c95506e46daa"
```

Set working directory (scratch space)

## [1] "Output directory: /private/var/folders/xp/h3k9vf3n2jx181ts786\_yjrn9c2gjq/T/RtmpzH4y20"

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(),
     `growth form` = col_character(),
##
##
     species = col_character(),
##
     timestamp = col_character()
## )
## See spec(...) for full column specifications.
## Download complete!
head(dat_raw)
## # A tibble: 6 x 2,114
     `Anthocyanin co~ `Anthocyanin co~ `Carotenoid con~ `Carotenoid con~
##
                                                                     <dbl>
##
                <dbl>
                                  <dbl>
                                                   <dbl>
## 1
              0.00106
                                 0.997
                                                 0.00799
                                                                      7.49
## 2
              0.00357
                                  1.22
                                                 0.0221
                                                                      7.53
## 3
              0.00252
                                  1.14
                                                 0.0188
                                                                      8.55
## 4
                                                                     11.5
              0.00310
                                  2.26
                                                 0.0158
## 5
              0.00412
                                  1.73
                                                 0.0216
                                                                      9.08
## 6
                                                                      8.66
              0.00397
                                  1.02
                                                 0.0336
     ... with 2,110 more variables: `Chlorophyll concentration (mg/g)` <dbl>,
       `Chlorophyll content ( g/cm )` <dbl>, `LDMC (g/g)` <dbl>, `LFA (mg/cm
       )` <dbl>, `LWC (mg/cm )` <dbl>, `SLA (g/cm )` <dbl>, `growth form` <chr>,
## #
       species <chr>, timestamp <chr>, `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>, `441` <dbl>, `442` <dbl>,
## #
## #
       '443' <dbl>, '444' <dbl>, '445' <dbl>, '446' <dbl>, '447' <dbl>,
       `448` <dbl>, `449` <dbl>, `450` <dbl>, `451` <dbl>, `452` <dbl>,
       `453` <dbl>, `454` <dbl>, `455` <dbl>, `456` <dbl>, `457` <dbl>,
## #
       `458` <dbl>, `459` <dbl>, `460` <dbl>, `461` <dbl>, `462` <dbl>,
## #
## #
       '463' <dbl>, '464' <dbl>, '465' <dbl>, '466' <dbl>, '467' <dbl>,
## #
       `468` <dbl>, `469` <dbl>, `470` <dbl>, `471` <dbl>, `472` <dbl>,
       `473` <dbl>, `474` <dbl>, `475` <dbl>, `476` <dbl>, `477` <dbl>,
## #
       `478` <dbl>, `479` <dbl>, `480` <dbl>, `481` <dbl>, `482` <dbl>,
## #
```

```
`483` <dbl>, `484` <dbl>, `485` <dbl>, `486` <dbl>, `487` <dbl>,
     `488` <dbl>, `489` <dbl>, `490` <dbl>, ...
names(dat_raw)[1:40]
   [1] "Anthocyanin concentration (mg/g)" "Anthocyanin content ( g/cm )"
##
   [3] "Carotenoid concentration (mg/g)"
                                            "Carotenoid content (g/cm)"
##
   [5] "Chlorophyll concentration (mg/g)" "Chlorophyll content ( g/cm )"
  [7] "LDMC (g/g)"
                                            "LFA (mg/cm )"
##
  [9] "LWC (mg/cm )"
                                            "SLA (g/cm )"
## [11] "growth form"
                                            "species"
## [13] "timestamp"
                                            "400"
## [15] "401"
                                            "402"
## [17] "403"
                                            "404"
## [19] "405"
                                            "406"
## [21] "407"
                                            "408"
## [23] "409"
                                            "410"
## [25] "411"
                                            "412"
## [27] "413"
                                            "414"
## [29] "415"
                                            "416"
## [31] "417"
                                            "418"
## [33] "419"
                                            "420"
## [35] "421"
                                            "422"
## [37] "423"
                                            "424"
## [39] "425"
                                            "426"
```

## Create full plsr dataset

```
### Create plsr dataset
## cleanup any missing
if (any(is.na(dat_raw))) {
   dat_raw <- na.omit(dat_raw)
}
# remove suspect high values
dat_raw <- dat_raw %>%
   filter(`SLA (g/cm )`<=500)

Start.wave <- 500
End.wave <- 2400
wv <- seq(Start.wave,End.wave,1)
Spectra <- as.matrix(dat_raw[,names(dat_raw) %in% wv])
colnames(Spectra) <- c(paste0("Wave_",wv))
sample_info <- dat_raw[,names(dat_raw) %notin% seq(350,2500,1)]
head(sample_info)</pre>
```

```
## # A tibble: 6 x 13
##
     Anthocyanin co~ `Anthocyanin co~ `Carotenoid con~ `Carotenoid con~
                <dbl>
                                 <dbl>
                                                   <dbl>
                                                                    <dbl>
## 1
              0.00106
                                 0.997
                                                 0.00799
                                                                     7.49
## 2
              0.00357
                                 1.22
                                                 0.0221
                                                                     7.53
## 3
              0.00252
                                 1.14
                                                 0.0188
                                                                     8.55
## 4
              0.00310
                                 2.26
                                                 0.0158
                                                                    11.5
## 5
              0.00412
                                 1.73
                                                 0.0216
                                                                     9.08
## 6
              0.00397
                                 1.02
                                                0.0336
                                                                     8.66
```

```
## # ... with 9 more variables: `Chlorophyll concentration (mg/g)` <dbl>,
      `Chlorophyll content ( g/cm )` <dbl>, `LDMC (g/g)` <dbl>, `LFA (mg/cm
      ) ' <dbl>, 'LWC (mg/cm ) ' <dbl>, 'SLA (g/cm ) ' <dbl>, 'growth form' <chr>,
## #
      species <chr>, timestamp <chr>
sample_info2 <- sample_info %>%
  select(Plant_Species=species,Growth_Form=`growth form`,timestamp,SLA_g_cm=`SLA (g/cm )`)
head(sample_info2)
## # A tibble: 6 x 4
    Plant Species
                            Growth Form timestamp
                                                        SLA_g_cm
##
     <chr>>
                            <chr>
                                        <chr>
                                                           <dbl>
## 1 Calamagrostis epigejos graminoid 5/25/2016 12:20
                                                            107.
## 2 Anthoxanthum odoratum graminoid
                                       5/27/2016 8:40
                                                            293.
## 3 Alopecurus pratensis graminoid
                                        5/27/2016 9:23
                                                            220.
## 4 Festuca ovina
                            graminoid 5/27/2016 9:23
                                                            137.
                            graminoid 5/27/2016 9:42
## 5 Agrostis capillaris
                                                            237.
## 6 Aegopodium podagraria forb
                                        5/25/2016 12:20
                                                            388.
plsr_data <- data.frame(sample_info2,Spectra)</pre>
rm(sample_info,sample_info2,Spectra)
```

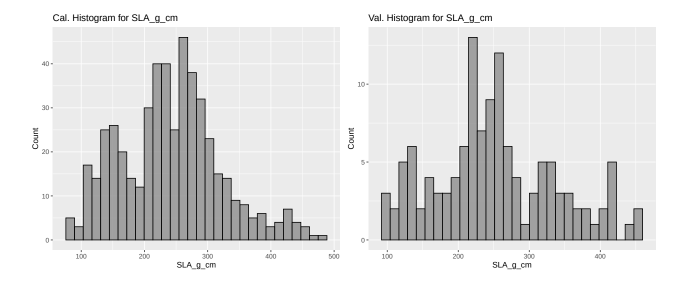
### Create cal/val datasets

```
### Create cal/val datasets
## Make a stratified random sampling in the strata USDA_Species_Code and Domain
method <- "base" #base/dplyr
# base R - a bit slow
# dplyr - much faster
split_data <- create_data_split(approach=method, split_seed=2356812, prop=0.8,</pre>
                                group_variables="Plant_Species")
                           Cal: 80%
## Calamagrostis epigejos
## Anthoxanthum odoratum
                          Cal: 80%
## Alopecurus pratensis Cal: 80%
## Festuca ovina Cal: 78.9473684210526%
## Agrostis capillaris
                        Cal: 82.3529411764706%
## Aegopodium podagraria
                          Cal: 80%
## Arrhenatherum elatius
                          Cal: 82.3529411764706%
## Arctium lappa
                  Cal: 83.3333333333333333
                 Cal: 78.9473684210526%
## Urtica dioica
## Cirsium arvense
                    Cal: 80%
## Geranium pratense
                       Cal: 81.25%
## Geum urbanum
                 Cal: 80%
## Digitalis purpurea
                       Cal: 81.25%
## Stellaria media Cal: 77.777777777778%
```

```
## Trisetum flavescens
                     Cal: 80%
## Trifolium pratense
                      Cal: 80.9523809523809%
## Geranium robertianum Cal: 78.5714285714286%
## Plantago major
                  Cal: 85.7142857142857%
## Nardus stricta
                  Cal: 78.9473684210526%
## Lamium purpureum
                    Cal: 77.777777777778%
## Clinopodium vulgare
                       Cal: 78.5714285714286%
## Poa annua
             Cal: 75%
## Campanula rotundifolia
                         Cal: 78.5714285714286%
## Taraxacum spec.
                   Cal: 80%
## Digitaria sanguinalis
                        Cal: 85.7142857142857%
## Holcus lanatus
                  Cal: 82.3529411764706%
## Lapsana communis
                    Cal: 75%
## Apera spica-venti
                     Cal: 80%
## Alopecurus geniculatus
## Bromus hordeaceus
                     Cal: 80%
## Phalaris arundinaceae
                        Cal: 81.25%
## Thlaspi arvense Not enough observations
## Origanum vulgare
                    Cal: 77.777777777778%
## Pulicaria dysenterica
                         Cal: 79.1666666666667%
## Deschampsia cespitosa
                         Cal: 80%
## Cirsium acaule
                  Cal: 80%
## Brachypodium sylvaticum Cal: 80%
## Centaurium erythraea Cal: 77.77777777778%
## Luzula multiflora
                     Cal: 78.5714285714286%
                       Cal: 78.5714285714286%
## Filipendula ulmaria
## Anthyllis vulneraria
                        Cal: 75%
## Medicago lupulina Cal: 75%
## Scirpus sylvaticus Cal: 77.77777777778%
names(split_data)
## [1] "cal_data" "val_data"
cal.plsr.data <- split_data$cal_data</pre>
```

head(cal.plsr.data)[1:8]

```
##
              Plant Species Growth Form
                                              timestamp SLA_g_cm
                                                                   Wave 500
## 1 Calamagrostis epigejos
                             graminoid 5/25/2016 12:20 106.6500 0.09180559
## 2 Anthoxanthum odoratum
                             graminoid 5/27/2016 8:40 293.3565 0.09022668
## 3
      Alopecurus pratensis graminoid 5/27/2016 9:23 220.2703 0.07998340
## 4
              Festuca ovina graminoid 5/27/2016 9:23 137.1220 0.05205080
## 5
                             graminoid 5/27/2016 9:42 237.4237 0.06695127
       Agrostis capillaris
                                   forb 5/25/2016 12:20 388.2384 0.04091566
## 6 Aegopodium podagraria
##
       Wave 501
                 Wave 502
                             Wave 503
## 1 0.09293251 0.09417092 0.09552863
## 2 0.09125158 0.09237300 0.09359694
## 3 0.08109460 0.08231389 0.08365015
## 4 0.05256869 0.05314560 0.05378355
## 5 0.06766205 0.06845248 0.06932220
## 6 0.04169865 0.04257613 0.04355737
val.plsr.data <- split_data$val_data</pre>
head(val.plsr.data)[1:8]
              Plant_Species Growth_Form
                                              timestamp SLA_g_cm
                                                                   Wave_500
## 9
              Urtica dioica
                                   forb 5/25/2016 12:37 284.6788 0.04716736
           Stellaria media
                                   forb 5/25/2016 13:21 418.4284 0.05694278
## 14
## 21 Alopecurus pratensis graminoid 6/1/2016 11:32 218.2117 0.08135086
## 41 Alopecurus pratensis
                             graminoid
                                         6/8/2016 8:37 216.7568 0.10062342
                                          6/8/2016 9:05 231.5292 0.08099724
## 43
       Agrostis capillaris
                              graminoid
## 44 Aegopodium podagraria
                                          6/7/2016 9:05 311.4018 0.03778815
                                   forb
##
       Wave_501
                 Wave_502
                              Wave 503
## 9 0.04781633 0.04854276 0.04935320
## 14 0.05811729 0.05940497 0.06080936
## 21 0.08249180 0.08373915 0.08509719
## 41 0.10190706 0.10330054 0.10480538
## 43 0.08178586 0.08265099 0.08360108
## 44 0.03845043 0.03919155 0.04001581
rm(split_data)
print(paste("Cal observations: ",dim(cal.plsr.data)[1],sep=""))
## [1] "Cal observations: 490"
print(paste("Val observations: ",dim(val.plsr.data)[1],sep=""))
## [1] "Val observations: 124"
cal_hist_plot <- qplot(cal.plsr.data[,paste0(inVar)],geom="histogram",</pre>
                       main = paste0("Cal. Histogram for ",inVar),
                       xlab = paste0(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("Val. Histogram for ",inVar),
                       xlab = paste0(inVar),ylab = "Count",fill=I("grey50"),
                       col=I("black"),alpha=I(.7))
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`.
```

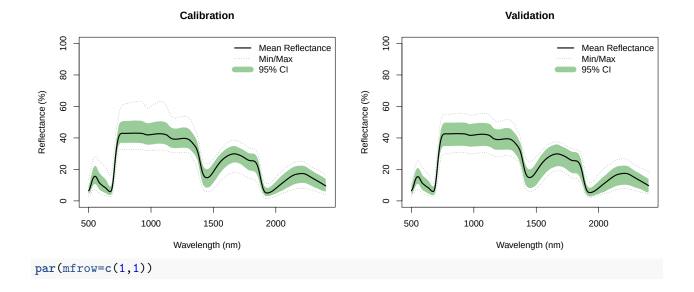


#### Create calibration and validation PLSR datasets

```
### 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))],
                             Spectra=I(cal_spec))
head(cal.plsr.data)[1:5]
##
              Plant_Species Growth_Form
                                               timestamp SLA_g_cm CalVal
## 1 Calamagrostis epigejos
                               graminoid 5/25/2016 12:20 106.6500
                                                                      Cal
      Anthoxanthum odoratum
                                                                      Cal
                               graminoid 5/27/2016 8:40 293.3565
## 3
       Alopecurus pratensis
                               graminoid 5/27/2016 9:23 220.2703
                                                                      Cal
## 4
              Festuca ovina
                               graminoid 5/27/2016 9:23 137.1220
                                                                      Cal
## 5
        Agrostis capillaris
                               graminoid 5/27/2016 9:42 237.4237
                                                                      Cal
## 6
     Aegopodium podagraria
                                    forb 5/25/2016 12:20 388.2384
                                                                      Cal
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 Growth_Form
                                               timestamp SLA_g_cm CalVal
## 9
              Urtica dioica
                                    forb 5/25/2016 12:37 284.6788
                                                                      Val
                                                                      Val
## 14
            Stellaria media
                                    forb 5/25/2016 13:21 418.4284
## 21
       Alopecurus pratensis
                               graminoid
                                          6/1/2016 11:32 218.2117
                                                                      Val
       Alopecurus pratensis
                                           6/8/2016 8:37 216.7568
                                                                      Val
## 41
                               graminoid
                               graminoid
## 43
        Agrostis capillaris
                                           6/8/2016 9:05 231.5292
                                                                      Val
## 44 Aegopodium podagraria
                                           6/7/2016 9:05 311.4018
                                                                      Val
                                    forb
```

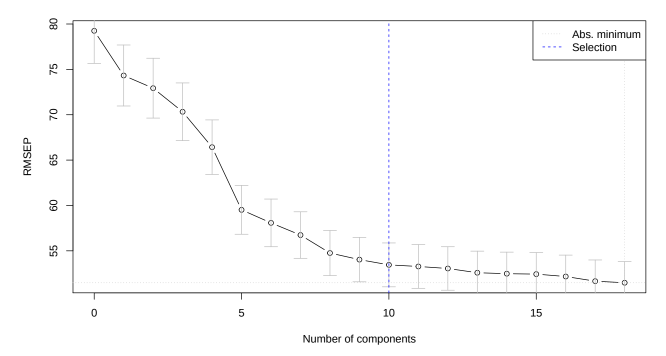
#### plot cal and val spectra

```
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")
```



### Use Jackknife permutation to determine optimal number of components

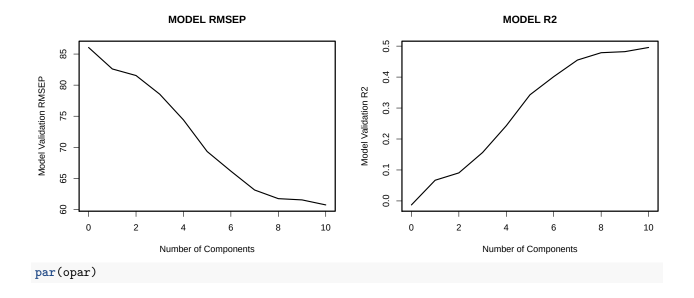
```
### Use permutation to determine the optimal number of components
if(grepl("Windows", sessionInfo()$running)){
  pls.options(parallel = NULL)
} else {
  pls.options(parallel = parallel::detectCores()-1)
method <- "pls" #pls, custom, lowestPRESS</pre>
random seed <- 2356812
seg <- 100
maxComps <- 18
iterations <- 30
if (method=="pls") {
  # pls package approach - faster but estimates more components....
  nComps <- find optimal components(method=method, maxComps=maxComps, seg=seg,
                                    random seed=random seed)
  print(paste0("*** Optimal number of components: ", nComps))
} else {
  # custom method - slow but generally finds the smallest number of components
  nComps <- find_optimal_components(method=method, maxComps=maxComps, iterations=iterations,
                                     seg=seg, prop=0.70,
                                     random_seed=random_seed)
}
```



## [1] "\*\*\* Optimal number of components: 10"

#### Fit final model

```
segs <- 100
plsr.out <- plsr(as.formula(paste(inVar,"~","Spectra")),scale=FALSE,ncomp=nComps,validation="CV",
                 segments=segs, segment.type="interleaved",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
                                                                           5 comps
                                                3 comps
                                                             4 comps
         86.06
                       82.60
                                    81.55
                                                  78.54
                                                                             69.32
##
                                                                74.40
##
       6 comps
                    7 comps
                                  8 comps
                                                9 comps
                                                             10 comps
         66.16
                       63.13
                                    61.74
                                                  61.53
                                                                60.73
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
##
      -0.01288
                    0.06681
                                  0.09056
                                                0.15636
                                                             0.24295
                                                                           0.34288
       6 comps
                    7 comps
##
                                  8 comps
                                                9 comps
                                                             10 comps
       0.40138
                    0.45499
                                  0.47875
                                                0.48216
                                                             0.49563
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)
```

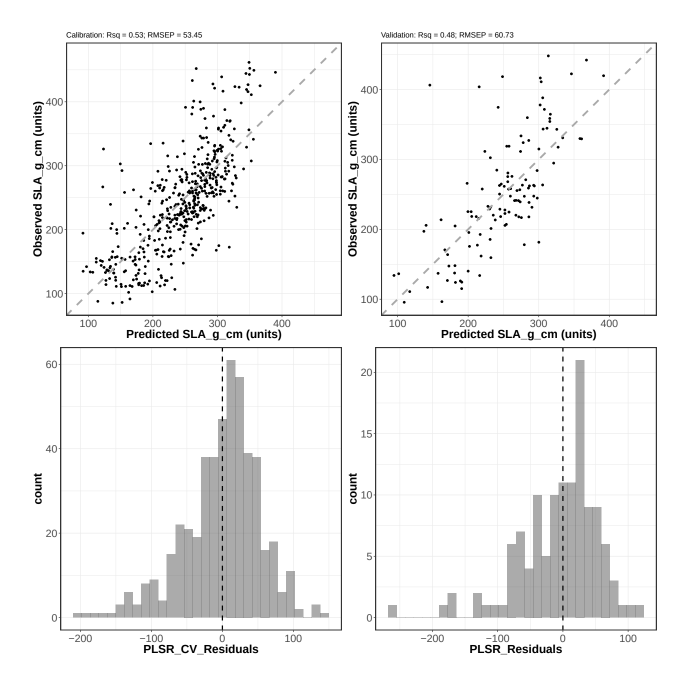


#### PLSR fit observed vs. predicted plot data

```
#calibration
cal.plsr.output <- data.frame(cal.plsr.data[, which(names(cal.plsr.data) %notin% "Spectra")],</pre>
                               PLSR Predicted=fit,
                               PLSR CV Predicted=as.vector(plsr.out$validation$pred[,,nComps]))
cal.plsr.output <- cal.plsr.output %>%
  mutate(PLSR_CV_Residuals = PLSR_CV_Predicted-get(inVar))
head(cal.plsr.output)
##
              Plant_Species Growth_Form
                                               timestamp SLA_g_cm CalVal
## 1 Calamagrostis epigejos
                               graminoid 5/25/2016 12:20 106.6500
                                                                      Cal
## 2
      Anthoxanthum odoratum
                               graminoid 5/27/2016 8:40 293.3565
                                                                      Cal
                                                                      Cal
## 3
       Alopecurus pratensis
                               graminoid 5/27/2016 9:23 220.2703
## 4
              Festuca ovina
                               graminoid 5/27/2016 9:23 137.1220
                                                                      Cal
## 5
        Agrostis capillaris
                               graminoid 5/27/2016 9:42 237.4237
                                                                      Cal
     Aegopodium podagraria
                                    forb 5/25/2016 12:20 388.2384
                                                                      Cal
## 6
##
     PLSR_Predicted PLSR_CV_Predicted PLSR_CV_Residuals
## 1
           231.9307
                              234.1193
                                              127.469378
## 2
           237.6749
                              236.7755
                                              -56.581079
## 3
           262.8365
                              263.8336
                                               43.563272
## 4
                                               -8.283722
           126.5863
                              128.8382
## 5
           251.2489
                              251.3030
                                               13.879308
## 6
           277.2292
                              274.2644
                                             -113.974044
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")],</pre>
                               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)
```

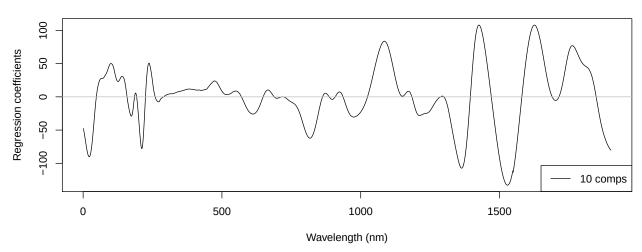
```
##
             Plant_Species Growth_Form
                                             timestamp SLA_g_cm CalVal
## 1
             Urtica dioica
                                  forb 5/25/2016 12:37 284.6788
                                                                    Val
## 2
           Stellaria media
                                  forb 5/25/2016 13:21 418.4284
                                                                    Val
## 3 Alopecurus pratensis graminoid 6/1/2016 11:32 218.2117
                                                                    Val
## 4 Alopecurus pratensis graminoid
                                        6/8/2016 8:37 216.7568
                                                                    Val
       Agrostis capillaris
                             graminoid 6/8/2016 9:05 231.5292
                                                                    Val
## 5
## 6 Aegopodium podagraria
                                  forb 6/7/2016 9:05 311.4018
                                                                    Val
     PLSR Predicted PLSR Residuals
##
## 1
           240.6023
                        -44.076512
## 2
           248.6923
                       -169.736117
## 3
           211.4638
                         -6.747881
           275.4544
                         58.697587
## 4
## 5
           290.4019
                         58.872672
## 6
           274.2311
                        -37.170622
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"),
```

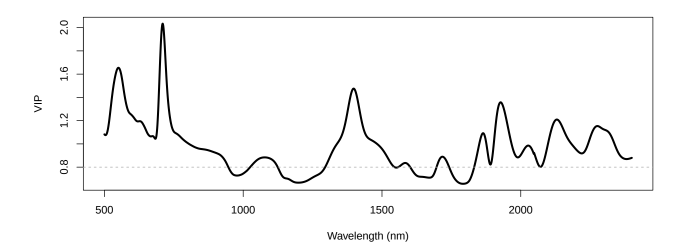
```
axis.text.x = element_text(angle = 0, vjust = 0.5),
        panel.border = element_rect(linetype = "solid", fill = NA, size=1.5))
val_resid_histogram <- ggplot(val.plsr.output, aes(x=PLSR_Residuals)) +</pre>
  geom_histogram(alpha=.5, position="identity") +
  geom_vline(xintercept = 0, color="black",
             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
grid.arrange(cal_scatter_plot, val_scatter_plot, cal_resid_histogram, val_resid_histogram,
            nrow=2,ncol=2)
## Warning: Removed 7 rows containing missing values (geom_point).
## Warning: Removed 3 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







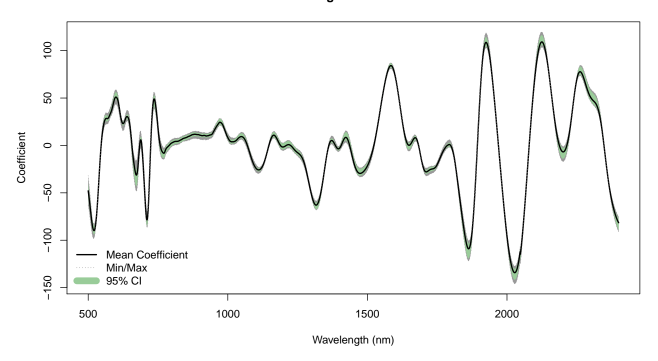
## Jackknife validation

```
Jackknife_coef <- Jackknife_coef[2:dim(Jackknife_coef)[1],,,]</pre>
#interval <- c(0.025,0.975)
interval <-c(0.05,0.95)
Jackknife_Pred <- val.plsr.data$Spectra%*%Jackknife_coef+Jackknife_intercept</pre>
Interval_Conf <- apply(X = Jackknife_Pred, MARGIN = 1, FUN = quantile,</pre>
                        probs=c(interval[1], interval[2]))
Interval Pred <- apply(X = Jackknife Pred, MARGIN = 1, FUN = quantile,</pre>
                        probs=c(interval[1], interval[2]))
sd_mean <- apply(X = Jackknife_Pred, MARGIN = 1, FUN =sd)</pre>
sd_res <- sd(val.plsr.output$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 Growth_Form
                                            timestamp SLA_g_cm CalVal
## 1
                                forb 5/25/2016 12:37 284.6788
            Urtica dioica
## 2
                                 forb 5/25/2016 13:21 418.4284
          Stellaria media
                                                                 Val
## 3 Alopecurus pratensis
                           graminoid 6/1/2016 11:32 218.2117
                                                                 Val
## 4 Alopecurus pratensis graminoid
                                      6/8/2016 8:37 216.7568
                                                                 Val
      Agrostis capillaris graminoid
                                      6/8/2016 9:05 231.5292
                                                                 Val
## 6 Aegopodium podagraria
                                       6/7/2016 9:05 311.4018
                                                                 Val
                                 forb
## PLSR Predicted PLSR Residuals
                                       LCI
                                                UCI
                                                         LPI
                                                                  UPI
## 1
          240.6023
                       -44.076512 231.2077 253.7969 120.83400 360.3705
## 2
                      -169.736117 239.6933 260.5583 129.04378 368.3409
          248.6923
## 3
          211.4638
                        -6.747881 199.3682 221.3377 91.78604 331.1416
## 4
                        58.697587 262.5270 290.0982 155.52083 395.3880
          275.4544
## 5
          290.4019
                        58.872672 280.1643 298.9968 170.77762 410.0261
## 6
          274.2311
                       -37.170622 261.9091 285.7860 154.51259 393.9497
```

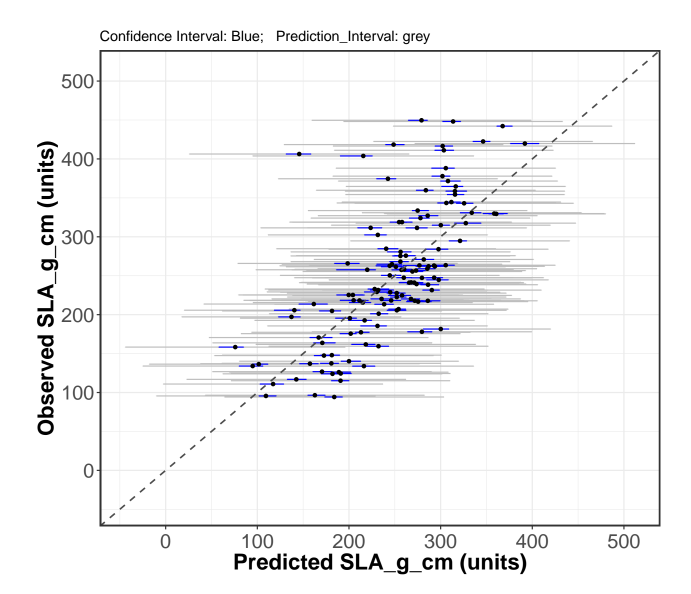
# Jackknife coefficient plot

### Jackknife regression coefficients



### Jackknife validation plot

```
\#rng\_vals \leftarrow quantile(val.plsr.output[,inVar], probs = c(0.001, 0.999))
rng vals <- c(min(val.plsr.output$LPI), max(val.plsr.output$UPI))</pre>
jk_val_scatterplot <- ggplot(val.plsr.output, aes(x=PLSR_Predicted,</pre>
                                                   y=get(inVar))) +
  theme_bw()+ geom_errorbar(aes(xmin = LPI,xmax = UPI),color='grey',
                            width=0.2) +
  geom errorbar(aes(xmin = LCI,xmax = UCI),color='blue',width=0.2) +
  geom point(size=1.3) +
  geom_abline(intercept = 0, slope = 1, color="grey30",
              linetype="dashed", size=0.7) +
  xlim(rng_vals[1], rng_vals[2]) +
  ylim(rng_vals[1], rng_vals[2]) +
  labs(x=paste0("Predicted ", paste(inVar), " (units)"),
       y=paste0("Observed ", paste(inVar), " (units)"),
       title=paste0("Confidence Interval: Blue; Prediction_Interval: grey")) +
  theme(axis.text=element_text(size=18),legend.position = 'right',
        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))
print(jk_val_scatterplot)
```



# Output jackknife results

```
out.jk.coefs <- data.frame(Iteration=seq(1,seg,1),</pre>
                           Intercept=Jackknife_intercept,t(Jackknife_coef))
head(out.jk.coefs)[1:6]
         Iteration Intercept Wave_500 Wave_501 Wave_502 Wave_503
## Seg 1
                 1 246.6837 -49.80782 -52.32289 -54.88084 -57.63716
                 2 254.8287 -52.24947 -54.31513 -56.41444 -58.71748
## Seg 2
## Seg 3
                 3 246.2546 -54.91885 -57.12727 -59.35903 -61.78247
                 4 249.9940 -49.37912 -51.77580 -54.22486 -56.87922
## Seg 4
                 5 257.4183 -45.54171 -47.92949 -50.36257 -53.01337
## Seg 5
## Seg 6
                 6 247.2549 -40.72975 -42.81360 -44.93902 -47.28299
write.csv(out.jk.coefs,file=file.path(outdir,
                                      paste0(inVar,
                                             '_Jackkife_PLSR_Coefficients.csv')),
          row.names=FALSE)
```

#### Create core PLSR outputs

```
print(paste("Output directory: ", getwd()))
## [1] "Output directory: /Users/sserbin/Data/GitHub/PLSR_for_plant_trait_prediction/vignettes"
# 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,
                               paste0(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] "SLA_g_cm_Jackkife_PLSR_Coefficients.csv"

## [2] "SLA_g_cm_Observed_PLSR_CV_Pred_10comp.csv"

## [3] "SLA_g_cm_PLSR_Coefficients_10comp.csv"

## [4] "SLA_g_cm_PLSR_VIPs_10comp.csv"

## [5] "SLA_g_cm_Validation_PLSR_Pred_10comp.csv"
```