Spectra-trait PLSR example using leaf-level spectra and leaf mass per area (LMA) data from CONUS NEON sites

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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

Load libraries

```
list.of.packages <- c("pls", "dplyr", "here", "plotrix", "ggplot2", "gridExtra", "spectratrait")
invisible(lapply(list.of.packages, library, character.only = TRUE))
##
## 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/spectratrait
##
## Attaching package: 'gridExtra'
## The following object is masked from 'package:dplyr':
##
##
       combine
```

Setup other functions and options

```
### Setup other functions and options
# 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_gDW_m2"</pre>
# What is the source dataset from EcoSIS?
ecosis_id <- "5617da17-c925-49fb-b395-45a51291bd2d"
# Specify output directory, output_dir
# Options:
# tempdir - use a OS-specified temporary directory
# user defined PATH - e.q. "~/scratch/PLSR"
output_dir <- "tempdir"</pre>
```

Set working directory (scratch space)

[1] "/private/var/folders/xp/h3k9vf3n2jx181ts786_yjrn9c2gjq/T/RtmpOAOtX1"

Grab data from EcoSIS

```
print(paste0("Output directory: ",getwd())) # check wd
```

 $\label{lem:ura} \begin{tabular}{ll} URL: & https://ecosis.org/package/fresh-leaf-spectra-to-estimate-lma-over-neon-domains-ineastern-united-states \end{tabular}$

```
## [1] "Output directory: /Users/sserbin/Data/GitHub/spectratrait/vignettes"
### Get source dataset from EcoSIS
dat_raw <- spectratrait::get_ecosis_data(ecosis_id = ecosis_id)</pre>
## [1] "**** Downloading Ecosis data ****"
## Downloading data...
.default = col_double(),
##
    Affiliation = col_character(),
##
    `Common Name` = col_character(),
##
## Domain = col_character(),
##
    Functional_type = col_character(),
##
    `Latin Genus` = col_character(),
   `Latin Species` = col_character(),
##
```

```
##
     Project = col_character(),
     Sample ID = col character(),
##
     `USDA Symbol` = col_character()
##
## )
## i Use `spec()` for the full column specifications.
## Download complete!
head(dat raw)
## # A tibble: 6 x 2,162
##
     Affiliation
                            `Common Name` Domain Functional_type
                                                                   LMA `Latin Genus`
##
     <chr>>
                            <chr>
                                          <chr> <chr>
                                                                  <dbl> <chr>
## 1 University of Wiscon~ black walnut DO2
                                                                   72.9 Juglans
                                                 broadleaf
## 2 University of Wiscon~ black walnut DO2
                                                 broadleaf
                                                                   72.9 Juglans
## 3 University of Wiscon~ black walnut DO2
                                                 broadleaf
                                                                   60.8 Juglans
## 4 University of Wiscon~ black walnut DO2
                                                 broadleaf
                                                                   60.8 Juglans
## 5 University of Wiscon~ black walnut DO2
                                                 broadleaf
                                                                   85.9 Juglans
## 6 University of Wiscon~ black walnut DO2
                                                 broadleaf
                                                                   85.9 Juglans
## # ... with 2,156 more variables: Latin Species <chr>, PI <chr>, Project <chr>,
       Sample_ID <chr>, USDA Symbol <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>, ...
names(dat_raw)[1:40]
## [1] "Affiliation"
                           "Common Name"
                                             "Domain"
                                                                "Functional_type"
                                                                "PI"
   [5] "LMA"
                           "Latin Genus"
                                             "Latin Species"
  [9] "Project"
                           "Sample_ID"
                                                                "350"
##
                                             "USDA Symbol"
## [13] "351"
                           "352"
                                             "353"
                                                                "354"
## [17] "355"
                           "356"
                                             "357"
                                                                "358"
## [21] "359"
                           "360"
                                             "361"
                                                                "362"
## [25] "363"
                           "364"
                                             "365"
                                                                "366"
## [29] "367"
                           "368"
                                             "369"
                                                                "370"
## [33] "371"
                                             "373"
                                                                "374"
                           "372"
## [37] "375"
                           "376"
                                             "377"
                                                                "378"
Create full plsr dataset
### Create plsr dataset
Start.wave <- 500
End.wave <- 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 11
##
     Affiliation
                            `Common Name` Domain Functional_type
                                                                    LMA `Latin Genus`
```

##

##

<chr>>

PI = col_character(),

<chr> <chr>

<dbl> <chr>

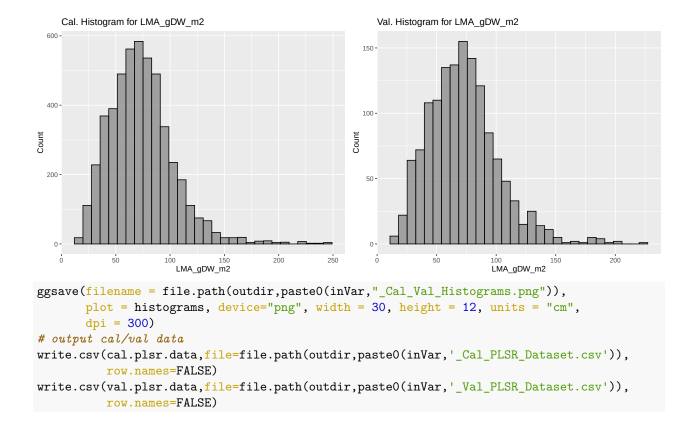
<chr>

```
## 1 University of Wiscon~ black walnut DO2
                                                 broadleaf
                                                                   72.9 Juglans
## 2 University of Wiscon~ black walnut DO2
                                                 broadleaf
                                                                   72.9 Juglans
                                                                   60.8 Juglans
## 3 University of Wiscon~ black walnut DO2
                                                 broadleaf
## 4 University of Wiscon~ black walnut DO2
                                                                   60.8 Juglans
                                                 broadleaf
## 5 University of Wiscon~ black walnut DO2
                                                 broadleaf
                                                                   85.9 Juglans
## 6 University of Wiscon~ black walnut DO2
                                                 broadleaf
                                                                   85.9 Juglans
## # ... with 5 more variables: Latin Species <chr>, PI <chr>, Project <chr>,
       Sample_ID <chr>, USDA Symbol <chr>
sample info2 <- sample info %>%
  select(Domain, Functional_type, Sample_ID, USDA_Species_Code=`USDA_Symbol`, LMA_gDW_m2=LMA)
head(sample_info2)
## # A tibble: 6 x 5
     Domain Functional_type Sample_ID USDA_Species_Code LMA_gDW_m2
##
                             <chr>
##
     <chr> <chr>
                                       <chr>
                                                               <dbl>
## 1 D02
            broadleaf
                            P0001
                                       JUNI
                                                                72.9
## 2 D02
            broadleaf
                            L0001
                                       JUNI
                                                                72.9
## 3 D02
            broadleaf
                            P0002
                                       JUNI
                                                                60.8
## 4 D02
            broadleaf
                                       JUNI
                            L0002
                                                                60.8
## 5 D02
            broadleaf
                            P0003
                                       JUNI
                                                                85.9
## 6 D02
            broadleaf
                            L0003
                                       JUNI
                                                                85.9
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 <- "dplyr" #base/dplyr
# base R - a bit slow
# dplyr - much faster
split data <- spectratrait::create data split(dataset=plsr data,approach=method, split seed=2356812,
                                               prop=0.8, group variables=c("USDA Species Code", "Domain")
names(split_data)
## [1] "cal data" "val data"
cal.plsr.data <- split_data$cal_data</pre>
head(cal.plsr.data)[1:8]
     Domain Functional_type Sample_ID USDA_Species_Code LMA_gDW_m2
                                                                       Wave 500
## 1
        D08
                  broadleaf
                                                               44.18 0.04170800
                                L2644
                                                    ACBA
        D08
## 2
                  broadleaf
                                L2646
                                                    ACBA
                                                               41.71 0.05067800
## 3
        D08
                  broadleaf
                                L2645
                                                    ACBA
                                                               40.66 0.04701700
## 4
        D08
                  broadleaf
                                P2639
                                                    ACBA
                                                               44.18 0.04125300
## 5
        D03
                  broadleaf
                                P0614
                                                    ACFL
                                                               52.91 0.03895800
## 6
        D03
                  broadleaf
                                L0609
                                                    ACFL
                                                               81.67 0.04186169
       Wave_501
                  Wave 502
## 1 0.04208700 0.04283700
## 2 0.05087600 0.05153500
## 3 0.04718200 0.04766500
## 4 0.04150300 0.04247100
```

```
## 5 0.03915100 0.03956200
## 6 0.04217802 0.04258768
val.plsr.data <- split_data$val_data</pre>
head(val.plsr.data)[1:8]
      Domain Functional_type Sample_ID USDA_Species_Code LMA_gDW_m2 Wave_500
## 3
         D02
                   broadleaf
                                 P0002
                                                     JUNI
                                                               60.77 0.043758
## 12
         D02
                   broadleaf
                                 L0006
                                                     JUNI
                                                               42.54 0.044338
## 13
        D02
                   broadleaf
                                 P0007
                                                     QUVE
                                                              106.57 0.015643
         D02
                                                               78.82 0.033019
## 19
                   broadleaf
                                 P0010
                                                     PRSE
## 21
         D02
                   broadleaf
                                 P0011
                                                     PRSE
                                                               86.09 0.024819
## 28
         D02
                   broadleaf
                                 L0014
                                                     PRSE
                                                               67.11 0.040095
     Wave_501 Wave_502
##
## 3 0.044171 0.044869
## 12 0.044748 0.045294
## 13 0.015579 0.015431
## 19 0.033102 0.033245
## 21 0.024826 0.025045
## 28 0.040397 0.040864
rm(split_data)
# Datasets:
print(paste("Cal observations: ",dim(cal.plsr.data)[1],sep=""))
## [1] "Cal observations: 4922"
print(paste("Val observations: ",dim(val.plsr.data)[1],sep=""))
## [1] "Val observations: 1390"
cal_hist_plot <- qplot(cal.plsr.data[,paste0(inVar)],geom="histogram",</pre>
                       main = paste0("Cal. 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("Val. Histogram for ",inVar),
                       xlab = pasteO(inVar), ylab = "Count", fill=I("grey50"),
                       col=I("black"),alpha=I(.7))
histograms <- grid.arrange(cal_hist_plot, val_hist_plot, ncol=2)</pre>
## `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))],</pre>
                              Spectra=I(cal_spec))
head(cal.plsr.data)[1:5]
##
     Domain Functional_type Sample_ID USDA_Species_Code LMA_gDW_m2
## 1
        D08
                   broadleaf
                                  L2644
                                                                 44.18
                                                      ACBA
        D08
                   broadleaf
                                                                 41.71
## 2
                                  L2646
                                                      ACBA
## 3
                   broadleaf
                                                                 40.66
        D08
                                  L2645
                                                      ACBA
## 4
        D08
                   broadleaf
                                  P2639
                                                      ACBA
                                                                 44.18
## 5
        D03
                   broadleaf
                                  P0614
                                                      ACFL
                                                                 52.91
## 6
        D03
                   broadleaf
                                  L0609
                                                      ACFL
                                                                 81.67
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]
      Domain Functional_type Sample_ID USDA_Species_Code LMA_gDW_m2
##
## 3
         D02
                    broadleaf
                                   P0002
                                                       JUNI
                                                                  60.77
         D02
                    broadleaf
                                                                  42.54
## 12
                                   L0006
                                                       JUNI
## 13
         D02
                    broadleaf
                                   P0007
                                                       QUVE
                                                                 106.57
                    broadleaf
## 19
         D02
                                   P0010
                                                       PRSE
                                                                  78.82
## 21
         D02
                    broadleaf
                                   P0011
                                                       PRSE
                                                                  86.09
## 28
         D02
                    broadleaf
                                   L0014
                                                       PRSE
                                                                  67.11
```

plot cal and val spectra

```
par(mfrow=c(1,2)) # B, L, T, R
spectratrait::f.plot.spec(Z=cal.plsr.data$Spectra,wv=wv,plot_label="Calibration")
spectratrait::f.plot.spec(Z=val.plsr.data$Spectra,wv=wv,plot_label="Validation")
                         Calibration
                                                                                  Validation
    100
                                                             100
                                       Mean Reflectance
                                                                                               Mean Reflectance
                                       Min/Max
                                                                                               Min/Max
                                       95% CI
                                                                                               95% CI
    80
                                                             80
                                                         Reflectance (%)
Reflectance (%)
    9
                                                            9
    4
                                                             4
    20
                                                            20
                  1000
                             1500
                                                                           1000
                                                                                      1500
        500
                                        2000
                                                                500
                                                                                                 2000
                        Wavelength (nm)
                                                                                Wavelength (nm)
dev.copy(png,file.path(outdir,paste0(inVar,'_Cal_Val_Spectra.png')),
           height=2500, width=4900, res=340)
## quartz_off_screen
##
dev.off();
## pdf
##
par(mfrow=c(1,1))
```

Use Jackknife permutation to determine optimal number of components

```
random_seed=random_seed)
  print(paste0("*** Optimal number of components: ", nComps))
} else {
 nComps <- spectratrait::find_optimal_components(dataset=cal.plsr.data, targetVariable=inVar,
                                                  method=method,
                                                  maxComps=maxComps, iterations=iterations,
                                                  seg=seg, prop=prop,
                                                  random seed=random seed)
}
## [1] "*** Identifying optimal number of PLSR components ***"
## [1] "*** Running permutation test. Please hang tight, this can take awhile ***"
## [1] "Options:"
## [1] "Max Components: 20 Iterations: 40 Data Proportion (percent): 70"
## [1] "*** Providing PRESS and coefficient array output ***"
## No id variables; using all as measure variables
## [1] "*** Optimal number of components based on t.test: 12"
                    ╛╇╅╇╇╇
 2e+05
dev.copy(png,file.path(outdir,paste0(paste0(inVar,"_PLSR_Component_Selection.png"))),
        height=2800, width=3400, res=340)
## quartz_off_screen
dev.off();
## pdf
##
     2
Fit final model
### Fit final model
segs <- 100
plsr.out <- plsr(as.formula(paste(inVar,"~","Spectra")),scale=FALSE,ncomp=nComps,</pre>
```

segments=segs, segment.type="interleaved",trace=FALSE,

validation="CV",

```
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
pls::RMSEP(plsr.out, newdata = val.plsr.data)
## (Intercept)
                                                  3 comps
                                                                               5 comps
                      1 comps
                                    2 comps
                                                                 4 comps
                                                    16.187
        29.372
                       18.664
                                     18.166
                                                                  12.760
                                                                                12.149
##
##
       6 comps
                      7 comps
                                    8 comps
                                                  9 comps
                                                                10 comps
                                                                              11 comps
##
        12.004
                       11.465
                                     11.144
                                                    10.389
                                                                  10.063
                                                                                 9.732
##
      12 comps
          9.633
##
plot(pls::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)
pls::R2(plsr.out, newdata = val.plsr.data)
## (Intercept)
                      1 comps
                                    2 comps
                                                  3 comps
                                                                 4 comps
                                                                               5 comps
##
     -0.001908
                     0.595475
                                   0.616770
                                                 0.695732
                                                                0.810908
                                                                              0.828593
##
       6 comps
                      7 comps
                                    8 comps
                                                  9 comps
                                                                10 comps
                                                                              11 comps
                     0.847338
                                   0.855775
##
      0.832656
                                                 0.874647
                                                                0.882410
                                                                              0.890000
##
      12 comps
##
      0.892247
plot(pls::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
   30
                                                       0.8
   25
Model Validation RMSEP
                                                   Model Validation R2
                                                       9.0
   20
                                                       0.4
   15
                                                       0.2
   9
                                                       0.0
              2
                          6
                                                                 2
                                                                              6
                                 8
                                       10
                                             12
                                                                                    8
                                                                                          10
                                                                                                12
```

Number of Components

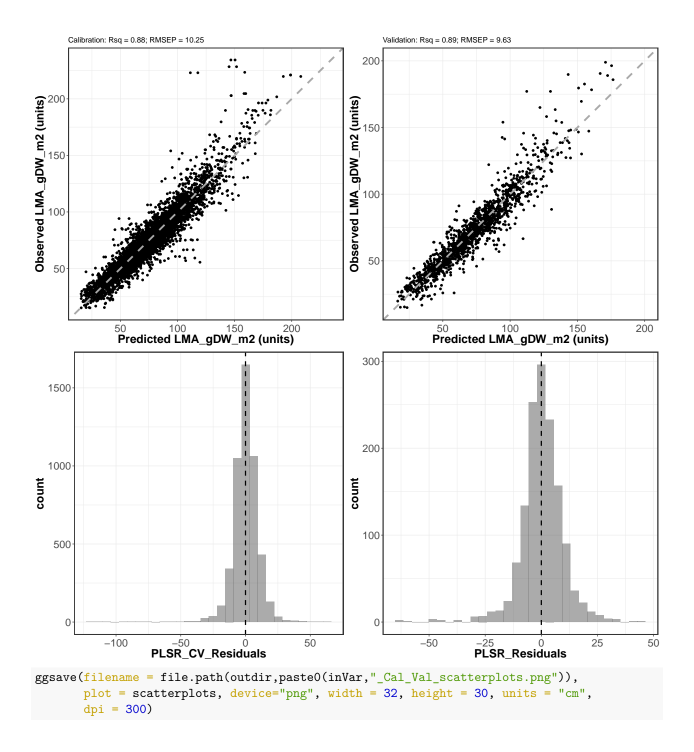
Number of Components

par(opar)

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)
     Domain Functional_type Sample_ID USDA_Species_Code LMA_gDW_m2 PLSR_Predicted
##
## 1
        D08
                  broadleaf
                                 L2644
                                                     ACBA
                                                                44.18
                                                                             53.00773
## 2
        D08
                  broadleaf
                                 L2646
                                                     ACBA
                                                                41.71
                                                                             44.02712
## 3
        D08
                  broadleaf
                                                                             46.76972
                                 L2645
                                                     ACBA
                                                                40.66
## 4
        D08
                  broadleaf
                                 P2639
                                                     ACBA
                                                                44.18
                                                                             49.62804
## 5
        D03
                  broadleaf
                                 P0614
                                                     ACFL
                                                                52.91
                                                                             64.78900
## 6
        D03
                  broadleaf
                                 L0609
                                                     ACFL
                                                                81.67
                                                                             96.85291
    PLSR CV Predicted PLSR CV Residuals
##
## 1
              53.03952
                                 8.859523
## 2
              44.03282
                                 2.322823
              46.90528
## 3
                                 6.245280
## 4
              49.80084
                                 5.620843
## 5
              64.98964
                                12.079641
              96.86039
                                15.190390
cal.R2 <- round(pls::R2(plsr.out,intercept=F)[[1]][nComps],2)
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)
      Domain Functional_type Sample_ID USDA_Species_Code LMA_gDW_m2 PLSR_Predicted
##
                   broadleaf
## 3
         D02
                                  P0002
                                                       JUNI
                                                                 60.77
                                                                              63.90905
         D02
                   broadleaf
                                  L0006
                                                      JUNI
                                                                 42.54
                                                                              41.54133
## 12
         D02
## 13
                   broadleaf
                                  P0007
                                                      QUVE
                                                                106.57
                                                                              99.99662
         D02
                   broadleaf
                                                      PRSE
                                                                 78.82
                                                                              89.03078
## 19
                                  P0010
## 21
         D02
                   broadleaf
                                  P0011
                                                      PRSE
                                                                 86.09
                                                                              85.17273
                   broadleaf
                                                      PRSE
                                                                 67.11
                                                                              67.95549
## 28
         D02
                                  L0014
##
      PLSR_Residuals
## 3
           3.1390459
## 12
          -0.9986720
## 13
          -6.5733831
## 19
          10.2107788
## 21
          -0.9172668
## 28
           0.8454930
val.R2 <- round(pls::R2(plsr.out,newdata=val.plsr.data,intercept=F)[[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>
```

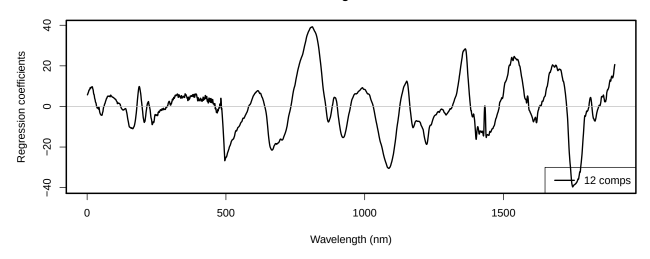
```
cal_scatter_plot <- ggplot(cal.plsr.output, aes(x=PLSR_CV_Predicted,</pre>
                                                 y=get(inVar))) +
  theme_bw() + geom_point() + geom_abline(intercept = 0, slope = 1,
                                           color="dark grey".
                                           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,</pre>
                              aes(x=PLSR_CV_Residuals)) +
  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],</pre>
                      probs = c(0.001, 0.999))
val_scatter_plot <- ggplot(val.plsr.output,</pre>
                           aes(x=PLSR_Predicted, y=get(inVar))) +
  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",
```

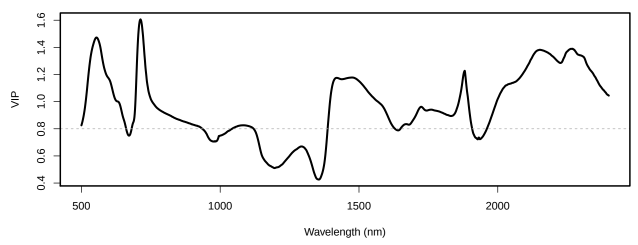


Generate Coefficient and VIP plots

```
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_gDW_m2





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

pdf ## 2

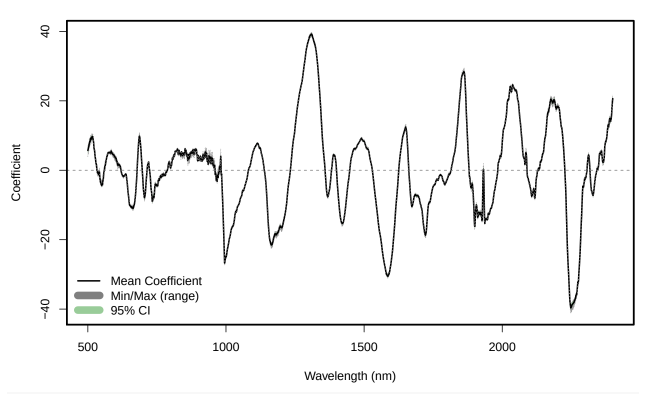
par(opar)

Jackknife validation

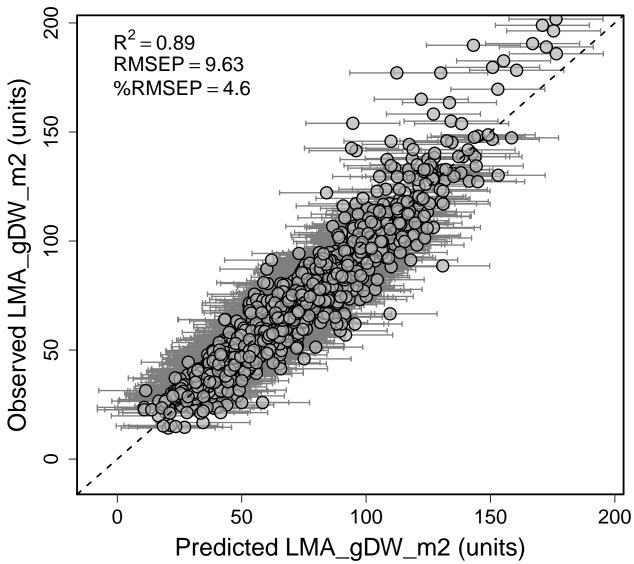
```
if(grepl("Windows", sessionInfo()$running)){
  pls.options(parallel =NULL)
} else {
  pls.options(parallel = parallel::detectCores()-1)
seg <- 100
jk.plsr.out <- pls::plsr(as.formula(paste(inVar,"~","Spectra")), scale=FALSE,
                          center=TRUE, ncomp=nComps,
                          validation="CV", segments = seg,
                          segment.type="interleaved", trace=FALSE,
                          jackknife=TRUE, data=cal.plsr.data)
pls.options(parallel = NULL)
Jackknife_coef <- spectratrait::f.coef.valid(plsr.out = jk.plsr.out, data_plsr = cal.plsr.data,</pre>
                                ncomp = nComps, inVar=inVar)
Jackknife intercept <- Jackknife coef[1,,,]</pre>
Jackknife_coef <- Jackknife_coef[2:dim(Jackknife_coef)[1],,,]</pre>
interval <-c(0.025, 0.975)
Jackknife_Pred <- val.plsr.data$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,</pre>
                        FUN = quantile,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_Conf[1,]</pre>
val.plsr.output$UCI <- Interval_Conf[2,]</pre>
val.plsr.output$LPI <- val.plsr.output$PLSR_Predicted-1.96*sd_tot</pre>
val.plsr.output$UPI <- val.plsr.output$PLSR_Predicted+1.96*sd_tot</pre>
head(val.plsr.output)
##
      Domain Functional_type Sample_ID USDA_Species_Code LMA_gDW_m2 PLSR_Predicted
## 3
         D02
                   broadleaf
                                  P0002
                                                      JUNI
                                                                 60.77
                                                                             63.90905
## 12
         D02
                   broadleaf
                                  1.0006
                                                      JUNI
                                                                 42.54
                                                                             41.54133
## 13
         D02
                   broadleaf
                                  P0007
                                                      QUVE
                                                                106.57
                                                                             99.99662
## 19
         D02
                   broadleaf
                                  P0010
                                                      PRSE
                                                                 78.82
                                                                             89.03078
## 21
         D02
                   broadleaf
                                  P0011
                                                      PRSE
                                                                 86.09
                                                                             85.17273
## 28
         D02
                   broadleaf
                                  L0014
                                                      PRSE
                                                                 67.11
                                                                             67.95549
##
      PLSR_Residuals
                           LCI
                                               LPI
                                                          UPI
                                     UCI
## 3
           3.1390459 63.75673 64.12043 45.02836 82.78973
## 12
          -0.9986720 41.42248 41.69728 22.66069 60.42196
          -6.5733831 99.88029 100.11962 81.11612 118.87712
## 13
## 19
          10.2107788 88.83274 89.21623 70.14949 107.91207
## 21
          -0.9172668 85.02330 85.32067 66.29194 104.05353
## 28
           0.8454930 67.82558 68.15298 49.07457 86.83642
```

Jackknife coefficient plot

Jackknife regression coefficients



Jackknife validation plot



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

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
##
                1 69.44171 5.796399 6.112194 6.482897 6.814518
## Seg 1
                 2 69.36719 6.316698 6.662910 7.031691 7.382632
## Seg 2
## Seg 3
                3 68.73866 6.729135 7.031912 7.372056 7.650510
                4 69.09350 5.899933 6.199538 6.614751 6.975426
## Seg 4
## Seg 5
                5 69.12959 5.355288 5.701339 6.043622 6.383870
                 6 68.67202 5.615024 5.968986 6.358249 6.654510
## Seg 6
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/spectratrait/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,
                                          pasteO(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
```

Confirm files were written to temp space

```
print("**** PLSR output files: ")
## [1] "**** PLSR output files: "
print(list.files(outdir)[grep(pattern = inVar, list.files(outdir))])
    [1] "LMA_gDW_m2_Cal_PLSR_Dataset.csv"
##
##
   [2] "LMA_gDW_m2_Cal_Val_Histograms.png"
##
   [3] "LMA_gDW_m2_Cal_Val_scatterplots.png"
##
   [4] "LMA_gDW_m2_Cal_Val_Spectra.png"
##
   [5] "LMA gDW m2 Coefficient VIP plot.png"
   [6] "LMA_gDW_m2_Jackkife_PLSR_Coefficients.csv"
##
   [7] "LMA_gDW_m2_Jackknife_Regression_Coefficients.png"
##
## [8] "LMA_gDW_m2_Observed_PLSR_CV_Pred_12comp.csv"
## [9] "LMA_gDW_m2_PLSR_Coefficients_12comp.csv"
## [10] "LMA_gDW_m2_PLSR_Component_Selection.png"
## [11] "LMA gDW m2 PLSR Validation Scatterplot.png"
## [12] "LMA_gDW_m2_PLSR_VIPs_12comp.csv"
## [13] "LMA_gDW_m2_Val_PLSR_Dataset.csv"
## [14] "LMA_gDW_m2_Validation_PLSR_Pred_12comp.csv"
```