# Spectra-trait PLSR example using NEON AOP pixel spectra and field-sampled leaf nitrogen content from CONUS NEON sites

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#### Overview

This is an R Markdown Notebook to illustrate how to develop pixel-scale spectra-trait PLSR models. This example uses image data from NEON AOP and associated field measurements of leaf nitrogen content collected across a range of CONUS NEON sites. For more information refer to the dataset EcoSIS page: https://ecosis.org/package/canopy-spectra-to-map-foliar-functional-traits-over-neon-domains-in-eastern-united-states

#### **Getting Started**

#### Load libraries

```
list.of.packages <- c("pls", "dplyr", "here", "plotrix", "ggplot2", "gridExtra", "spectratrait")
invisible(lapply(list.of.packages, library, character.only = TRUE))
## Warning: package 'pls' was built under R version 4.3.1
##
## Attaching package: 'pls'
## The following object is masked from 'package:stats':
##
##
       loadings
## Warning: package 'dplyr' was built under R version 4.3.1
##
## 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/Library/CloudStorage/OneDrive-NASA/Data/Github/spectratrait
## Warning: package 'plotrix' was built under R version 4.3.1
## Warning: package 'ggplot2' was built under R version 4.3.1
## Attaching package: 'gridExtra'
```

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

#### Setup other functions and options

```
### Setup options
# 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? What is the variable name in the input dataset?
inVar <- "Nitrogen"</pre>
# What is the source dataset from EcoSIS?
ecosis id <- "b9dbf3db-5b9c-4ab2-88c2-26c8b39d0903"
# 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/th/fpt\_z3417gn8xgply92pvy6r0000gq/T/RtmpY2MT01"

## Grab data from EcoSIS

## Download complete!

```
head(dat_raw)
## # A tibble: 6 x 459
     Affiliation
                   Boron Calcium Carbon Carotenoids_area Carotenoids_mass Cellulose
                   <dbl>
                            <dbl>
                                   <dbl>
                                                     <dbl>
     <chr>
                                                                       <dbl>
                                                                                 <dbl>
## 1 University ~ 0.0420
                            24.2
                                    463.
                                                      9.19
                                                                        1.18
                                                                                  221.
## 2 University ~ 0.0361
                             6.90
                                    558.
                                                     10.8
                                                                        1.17
                                                                                  183.
## 3 University ~ 0.0407
                                                                        1.52
                            16.7
                                    532.
                                                     12.2
                                                                                  133.
## 4 University ~ 0.0461
                            13.9
                                    461.
                                                      9.16
                                                                        1.50
                                                                                  220.
## 5 University ~ 0.0401
                            13.7
                                    510.
                                                     11.0
                                                                        1.53
                                                                                  101.
## 6 University ~ 0.0456
                            14.5
                                    557.
                                                      8.90
                                                                        1.24
                                                                                  214.
## # i 452 more variables: Chlorophylls_area <dbl>, Chlorophylls_mass <dbl>,
       Copper <dbl>, EWT <dbl>, Fiber <dbl>, Flavonoids <dbl>, LMA <dbl>,
       Lignin <dbl>, Magnesium <dbl>, Manganese <dbl>, NSC <dbl>, Nitrogen <dbl>,
## #
## #
       PI <chr>, Phenolics <dbl>, Phosphorus <dbl>, Plot_ID <chr>,
## #
       Potassium <dbl>, Project <chr>, SLA <dbl>, Sample Year <dbl>, Starch <dbl>,
## #
       Sugar <dbl>, Sulfur <dbl>, Water <dbl>, d13C <dbl>, d15N <dbl>,
## #
       `384` <dbl>, `389` <dbl>, `394` <dbl>, `399` <dbl>, `404` <dbl>, ...
names(dat raw)[1:40]
    [1] "Affiliation"
                             "Boron"
                                                  "Calcium"
##
   [4] "Carbon"
                             "Carotenoids area"
                                                  "Carotenoids mass"
   [7] "Cellulose"
                                                  "Chlorophylls_mass"
##
                             "Chlorophylls_area"
## [10] "Copper"
                             "EWT"
                                                  "Fiber"
                             "LMA"
## [13] "Flavonoids"
                                                  "Lignin"
                             "Manganese"
                                                  "NSC"
## [16] "Magnesium"
## [19] "Nitrogen"
                             "PI"
                                                  "Phenolics"
                             "Plot_ID"
## [22]
       "Phosphorus"
                                                  "Potassium"
## [25] "Project"
                             "SLA"
                                                  "Sample_Year"
## [28] "Starch"
                             "Sugar"
                                                  "Sulfur"
## [31] "Water"
                             "d13C"
                                                  "d15N"
## [34]
       "384"
                             "389"
                                                  "394"
                             "404"
## [37] "399"
                                                  "409"
## [40] "414"
Create full plsr dataset
# identify the trait data and other metadata
sample_info <- dat_raw[,names(dat_raw) %notin% seq(300,2600,1)]</pre>
head(sample_info)
## # A tibble: 6 x 33
##
     Affiliation
                   Boron Calcium Carbon Carotenoids_area Carotenoids_mass Cellulose
##
     <chr>>
                   dbl>
                            <dbl>
                                  <dbl>
                                                     <dbl>
                                                                       <dbl>
                                                                                 <dbl>
## 1 University ~ 0.0420
                            24.2
                                    463.
                                                      9.19
                                                                        1.18
                                                                                  221.
## 2 University ~ 0.0361
                             6.90
                                    558.
                                                     10.8
                                                                        1.17
                                                                                  183.
## 3 University ~ 0.0407
                            16.7
                                    532.
                                                     12.2
                                                                        1.52
                                                                                  133.
## 4 University ~ 0.0461
                            13.9
                                    461.
                                                      9.16
                                                                        1.50
                                                                                  220.
## 5 University ~ 0.0401
                            13.7
                                    510.
                                                     11.0
                                                                        1.53
                                                                                  101.
## 6 University ~ 0.0456
                            14.5
                                    557.
                                                      8.90
                                                                        1.24
                                                                                  214.
## # i 26 more variables: Chlorophylls_area <dbl>, Chlorophylls_mass <dbl>,
       Copper <dbl>, EWT <dbl>, Fiber <dbl>, Flavonoids <dbl>, LMA <dbl>,
## #
       Lignin <dbl>, Magnesium <dbl>, Manganese <dbl>, NSC <dbl>, Nitrogen <dbl>,
       PI <chr>, Phenolics <dbl>, Phosphorus <dbl>, Plot_ID <chr>,
## #
```

```
Potassium <dbl>, Project <chr>, SLA <dbl>, Sample_Year <dbl>, Starch <dbl>,
      Sugar <dbl>, Sulfur <dbl>, Water <dbl>, d13C <dbl>, d15N <dbl>
# spectra matrix
Spectra <- as.matrix(dat_raw[,names(dat_raw) %notin% names(sample_info)])</pre>
# set the desired spectra wavelength range to include
Start.wave <- 500
End.wave <- 2400
wv <- seq(Start.wave, End.wave, 1)</pre>
final_spec <- Spectra[,round(as.numeric(colnames(Spectra))) %in% wv]</pre>
colnames(final_spec) <- c(paste0("Wave_",colnames(final_spec)))</pre>
## Drop bad spectra data - for canopy-scale reflectance, often the "water band" wavelengths
## are too noisy to use for trait estimation. Its possible to remove these wavelengths
## prior to model fitting. Its best to first identify which wavelengths to drop
## before attempting PLSR, as these ranges may need to be considered on a case-by-case
## basis or generalized for multiple datasets
dropwaves <- c(1350:1440, 1826:1946)
final_spec <- final_spec[,colnames(final_spec) %notin% paste0("Wave_",dropwaves)]</pre>
wv <- as.numeric(gsub(pattern = "Wave_",replacement = "", x = colnames(final_spec)))</pre>
## Drop bad spectra data - for canopy-scale reflectance, often the "water band" wavelengths
## are too noisy to use for trait estimation. Its possible to remove these wavelengths
## prior to model fitting. Its best to first identify which wavelengths to drop
## before attempting PLSR, as these ranges may need to be considered on a case-by-case
## basis or generalized for multiple datasets
dropwaves <- c(1350:1440, 1826:1946)
final_spec <- final_spec[,colnames(final_spec) %notin% paste0("Wave_",dropwaves)]</pre>
wv <- as.numeric(gsub(pattern = "Wave_",replacement = "", x = colnames(final_spec)))</pre>
# assemble example dataset
sample_info2 <- sample_info %>%
  select(Plot_ID,Sample_Year,SLA,Nitrogen)
site_plot <- data.frame(matrix(unlist(strsplit(sample_info2$Plot_ID, "_")),</pre>
                               ncol=2, byrow=TRUE))
colnames(site_plot) <- c("Plot_Num", "SampleID")</pre>
sample_info3 <- data.frame(site_plot,sample_info2)</pre>
plsr data <- data.frame(sample info3,final spec*0.01)</pre>
rm(sample_info,sample_info2,sample_info3,Spectra, site_plot)
# Example data cleaning. End user needs to do what's appropriate for their
# data. This may be an iterative process.
# Keep only complete rows of inVar and spec data before fitting
plsr_data <- plsr_data %>% # remove erroneously high values, or "bad spectra"
  filter(Nitrogen<50) %>%
  filter(Wave_859<80) %>%
  filter(Wave_859>15)
plsr_data <- plsr_data[complete.cases(plsr_data[,names(plsr_data) %in%
                                                   c(inVar,paste0("Wave_",wv))]),]
```

### Example data cleaning.

## Create cal/val datasets

## [1] "Cal observations: 517"

```
## 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 <- spectratrait::create_data_split(dataset=plsr_data, approach=method, split_seed=2356326,
                                              prop=0.8, group_variables="Plot_Num")
## D02
         Cal: 80.46%
## D03
         Cal: 80.328%
## D05
         Cal: 80%
## D06
         Cal: 79.73%
## D07
         Cal: 79.245%
## D08
         Cal: 79.817%
         Cal: 79.63%
## D09
names(split_data)
## [1] "cal_data" "val_data"
cal.plsr.data <- split_data$cal_data</pre>
head(cal.plsr.data)[1:8]
     Plot_Num SampleID Plot_ID Sample_Year
                                                 SLA Nitrogen Wave_504 Wave_509
##
## 2
          D02
                  0002 D02 0002
                                       2017 10.77861 27.70598 1.2909576 1.4075910
          D02
                  0003 D02 0003
                                       2017 12.46154 34.63999 1.2976806 1.4257559
## 3
                                       2017 17.27620 26.64623 1.7735714 1.9423405
## 5
          D02
                  0005 D02 0005
                                       2017 12.92806 20.69437 1.7786337 1.9621929
## 6
          D02
                  0006 D02 0006
                                       2017 10.21521 28.87526 1.7981043 1.9359032
## 7
          D02
                  0007 D02_0007
                                       2017 20.87397 33.63137 0.8780127 0.9454703
## 8
          D02
                  0008 D02_0008
val.plsr.data <- split_data$val_data</pre>
head(val.plsr.data)[1:8]
##
      Plot_Num SampleID Plot_ID Sample_Year
                                                  SLA Nitrogen Wave_504 Wave_509
## 1
                   0001 D02_0001
           D02
                                        2017 13.66366 31.18030 1.467240 1.654816
## 4
           D02
                   0004 D02_0004
                                        2017 16.63205 34.54034 1.551933 1.764580
## 16
           D02
                   0016 D02_0016
                                        2017 14.44765 22.87740 2.198174 2.403996
           D02
                   0019 D02_0019
                                        2017 14.47103 17.73126 1.961911 2.175771
## 18
                   0020 D02 0020
                                        2017 18.98522 21.32929 1.546430 1.873175
## 19
           D02
           D02
                   0021 D02 0021
                                        2017 12.12731 29.50256 1.936263 2.065204
## 20
rm(split_data)
# Datasets:
print(paste("Cal observations: ",dim(cal.plsr.data)[1],sep=""))
```

```
print(paste("Val observations: ",dim(val.plsr.data)[1],sep=""))
## [1] "Val observations: 130"
cal_hist_plot <- ggplot(data = cal.plsr.data,</pre>
                         aes(x = cal.plsr.data[,paste0(inVar)])) +
  geom_histogram(fill=I("grey50"),col=I("black"),alpha=I(.7)) +
  labs(title=paste0("Calibration Histogram for ",inVar), x = paste0(inVar),
       y = "Count")
val_hist_plot <- ggplot(data = val.plsr.data,</pre>
                         aes(x = val.plsr.data[,paste0(inVar)])) +
  geom histogram(fill=I("grey50"),col=I("black"),alpha=I(.7)) +
  labs(title=paste0("Validation Histogram for ",inVar), x = paste0(inVar),
       y = "Count")
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`.
   Calibration Histogram for Nitrogen
                                                    Validation Histogram for Nitrogen
                                                  10.0
 40
                                                  7.5
 30 -
Count
                                                Count
5.0-
 20
                                                  2.5 -
     10
                 20
                                                         10
                       Nitrogen
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)
```

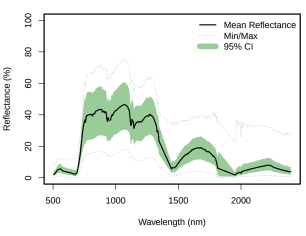
## Create calibration and validation PLSR datasets

## Plot\_Num SampleID Plot\_ID Sample\_Year SLA

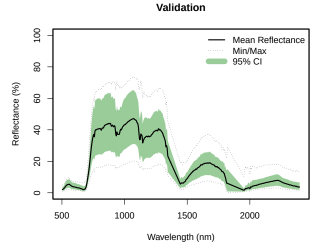
```
## 2
          D02
                   0002 D02 0002
                                         2017 10.77861
          D02
                   0003 D02 0003
## 3
                                         2017 12.46154
## 5
          D02
                  0005 D02 0005
                                         2017 17.27620
## 6
          D02
                  0006 D02_0006
                                         2017 12.92806
                   0007 D02 0007
## 7
          D02
                                         2017 10.21521
## 8
          D02
                  0008 D02 0008
                                         2017 20.87397
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]
##
      Plot_Num SampleID Plot_ID Sample_Year
                                                    SLA
## 1
           D02
                    0001 D02_0001
                                          2017 13.66366
## 4
           D02
                    0004 D02_0004
                                          2017 16.63205
## 16
           D02
                    0016 D02_0016
                                          2017 14.44765
                    0019 D02_0019
           D02
## 18
                                          2017 14.47103
           D02
## 19
                    0020 D02_0020
                                          2017 18.98522
## 20
           D02
                    0021 D02_0021
                                          2017 12.12731
```

### 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



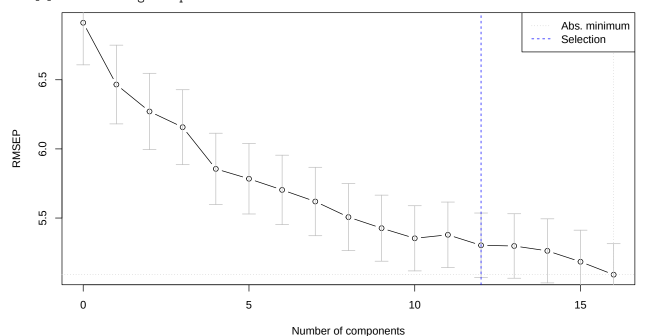
```
dev.copy(png,file.path(outdir,paste0(inVar,'_Cal_Val_Spectra.png')),
         height=2500, width=4900, res=340)
```

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

### Use permutation to determine optimal number of components

```
if(grepl("Windows", sessionInfo()$running)){
  pls.options(parallel = NULL)
} else {
  pls.options(parallel = parallel::detectCores()-1)
}
method <- "pls" #pls, firstPlateau, firstMin</pre>
random_seed <- 1245565
seg <- 50
maxComps <- 16</pre>
iterations <- 80
prop <- 0.70
if (method=="pls") {
  # pls package approach - faster but estimates more components....
  nComps <- spectratrait::find_optimal_components(dataset=cal.plsr.data, targetVariable=inVar,
                                                   method=method,
                                                   maxComps=maxComps, seg=seg,
                                                   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)
}
```

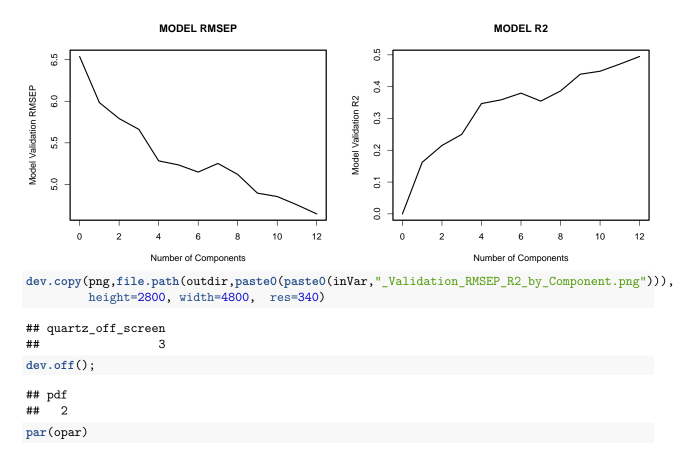
- ## [1] "\*\*\* Identifying optimal number of PLSR components \*\*\*"
- ## [1] "\*\*\* Running PLS permutation test \*\*\*"



## [1] "\*\*\* Optimal number of components: 12"

```
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
##
Fit final model
plsr.out <- plsr(as.formula(paste(inVar,"~","Spectra")),scale=FALSE,ncomp=nComps,validation="L00",
                 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
pls::RMSEP(plsr.out, newdata = val.plsr.data)
## (Intercept)
                    1 comps
                                  2 comps
                                                3 comps
                                                             4 comps
                                                                           5 comps
##
                                                                             5.235
         6.538
                      5.984
                                    5.792
                                                 5.662
                                                               5.284
##
       6 comps
                    7 comps
                                  8 comps
                                                9 comps
                                                            10 comps
                                                                          11 comps
##
         5.149
                      5.252
                                    5.121
                                                  4.896
                                                               4.855
                                                                             4.755
##
      12 comps
##
         4.646
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)
R2(plsr.out, newdata = val.plsr.data)
## (Intercept)
                                                3 comps
                                                                           5 comps
                    1 comps
                                  2 comps
                                                             4 comps
##
   -0.0001616
                  0.1621284
                                0.2150431
                                             0.2498762
                                                           0.3467097
                                                                         0.3586424
##
                                                            10 comps
                                                                         11 comps
       6 comps
                    7 comps
                                  8 comps
                                                9 comps
##
     0.3796062
                  0.3544358
                                0.3863604
                                             0.4391471
                                                           0.4484252
                                                                         0.4708911
##
      12 comps
     0.4948347
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)

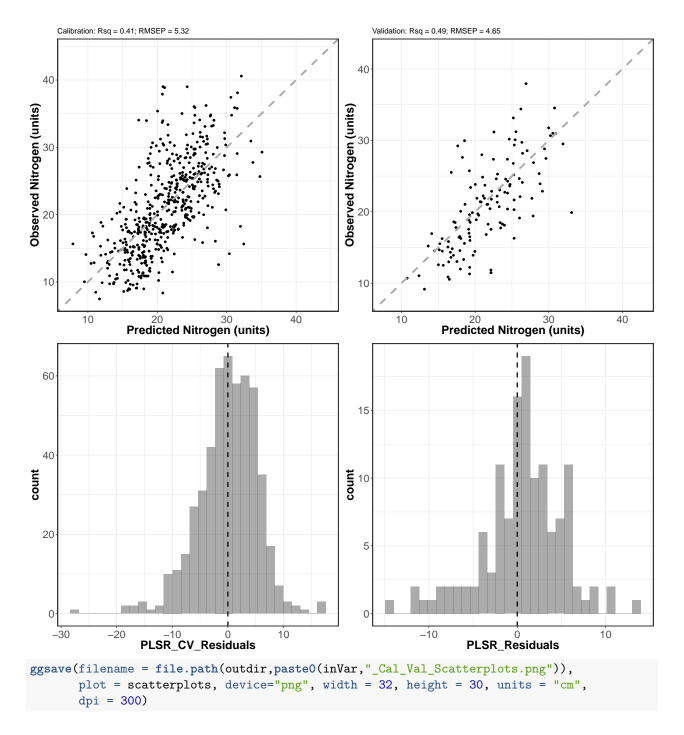


#### PLSR fit observed vs. predicted plot data

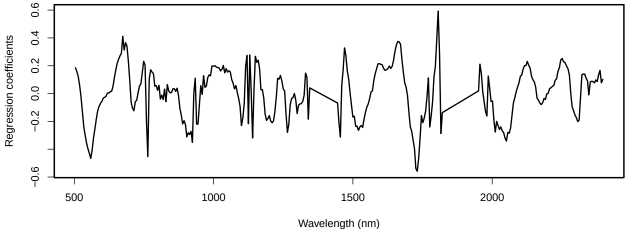
```
#calibration
cal.plsr.output <- data.frame(cal.plsr.data[, which(names(cal.plsr.data) %notin% "Spectra")],</pre>
                               PLSR_Predicted=fit,
                               PLSR_CV_Predicted=as.vector(plsr.out$validation$pred[,,nComps]))
cal.plsr.output <- cal.plsr.output %>%
  mutate(PLSR_CV_Residuals = PLSR_CV_Predicted-get(inVar))
head(cal.plsr.output)
##
     Plot_Num SampleID Plot_ID Sample_Year
                                                   SLA Nitrogen PLSR_Predicted
## 2
                  0002 D02_0002
          D02
                                        2017 10.77861 27.70598
                                                                       24.65561
## 3
          D02
                  0003 D02_0003
                                        2017 12.46154 34.63999
                                                                       27.85223
## 5
          D02
                  0005 D02_0005
                                        2017 17.27620 26.64623
                                                                       29.36467
## 6
          D02
                  0006 D02_0006
                                        2017 12.92806 20.69437
                                                                       21.66448
## 7
          D02
                  0007 D02 0007
                                        2017 10.21521 28.87526
                                                                       23.04393
                  0008 D02 0008
                                        2017 20.87397 33.63137
                                                                       25.56637
## 8
          D02
     PLSR_CV_Predicted PLSR_CV_Residuals
##
## 2
              24.59452
                               -3.1114612
## 3
              27.64033
                               -6.9996606
## 5
              29.54595
                                2.8997194
## 6
              21.68116
                                0.9867955
## 7
              22.78554
                               -6.0897138
              25.29798
## 8
                               -8.3333884
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>
```

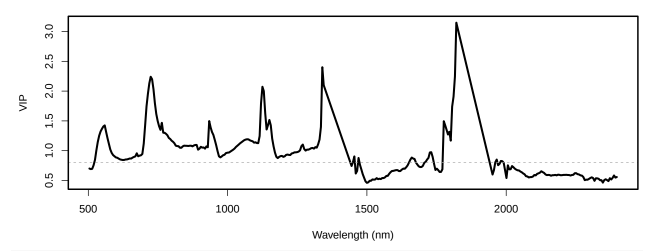
```
val.plsr.output <- data.frame(val.plsr.data[, which(names(val.plsr.data) %notin% "Spectra")],</pre>
                              PLSR_Predicted=as.vector(predict(plsr.out,
                                                                newdata = val.plsr.data,
                                                                ncomp=nComps, type="response")[,,1]))
val.plsr.output <- val.plsr.output %>%
  mutate(PLSR_Residuals = PLSR_Predicted-get(inVar))
head(val.plsr.output)
      Plot_Num SampleID Plot_ID Sample_Year
                                                   SLA Nitrogen PLSR_Predicted
## 1
           D02
                   0001 D02_0001
                                         2017 13.66366 31.18030
                                                                       22.55166
           D02
                                                                       30.79494
## 4
                   0004 D02_0004
                                         2017 16.63205 34.54034
## 16
           D02
                   0016 D02_0016
                                         2017 14.44765 22.87740
                                                                       29.14446
## 18
           D02
                   0019 D02_0019
                                         2017 14.47103 17.73126
                                                                       23,47518
                   0020 D02_0020
                                         2017 18.98522 21.32929
## 19
           D02
                                                                       23.00736
## 20
           D02
                   0021 D02 0021
                                         2017 12.12731 29.50256
                                                                       31.93483
##
     PLSR Residuals
## 1
           -8.628643
## 4
           -3.745399
## 16
           6.267060
## 18
            5.743923
## 19
            1.678070
## 20
            2.432274
val.R2 <- round(pls::R2(plsr.out,newdata=val.plsr.data,intercept=F)[[1]][nComps],2)
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", linewidth=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, linewidth=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", linewidth=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, linewidth=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", linewidth=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, linewidth=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", linewidth=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, linewidth=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 5 rows containing missing values or values outside the scale range
## (`geom_point()`).
## Warning: Removed 2 rows containing missing values or values outside the scale range
## (`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





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

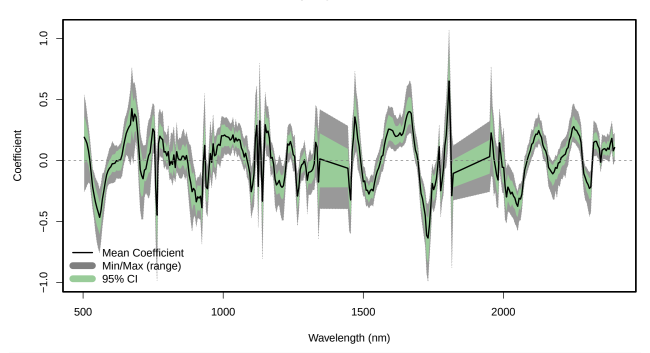
# Bootstrap validation

```
## [1] "*** Running permutation test. Please hang tight, this can take awhile ***"
## [1] "Options:"
## [1] "Max Components: 12 Iterations: 500 Data Proportion (percent): 70"
## [1] "*** Providing PRESS and coefficient array output ***"
```

```
##
      Plot_Num SampleID Plot_ID Sample_Year
                                                   SLA Nitrogen PLSR_Predicted
                                         2017 13.66366 31.18030
## 1
           D02
                   0001 D02_0001
                                                                       22.55166
## 4
           D02
                   0004 D02 0004
                                         2017 16.63205 34.54034
                                                                       30.79494
           D02
                   0016 D02_0016
                                                                       29.14446
## 16
                                         2017 14.44765 22.87740
## 18
           D02
                   0019 D02 0019
                                         2017 14.47103 17.73126
                                                                       23.47518
## 19
           D02
                   0020 D02 0020
                                         2017 18.98522 21.32929
                                                                       23.00736
## 20
           D02
                   0021 D02 0021
                                         2017 12.12731 29.50256
                                                                       31.93483
##
      PLSR Residuals
                          LCI
                                    UCI
                                             LPI
## 1
           -8.628643 21.75139 23.67919 13.44246 31.66086
## 4
           -3.745399 29.24737 32.37867 21.60577 39.98412
## 16
            6.267060 27.57462 30.82609 19.93270 38.35621
## 18
            5.743923 21.73808 24.49326 14.31158 32.63878
            1.678070 20.70321 24.57934 13.73687 32.27785
## 19
## 20
            2.432274 30.75996 34.32739 22.69357 41.17610
```

## Jackknife coefficient plot

#### **Bootstrap regression coefficients**



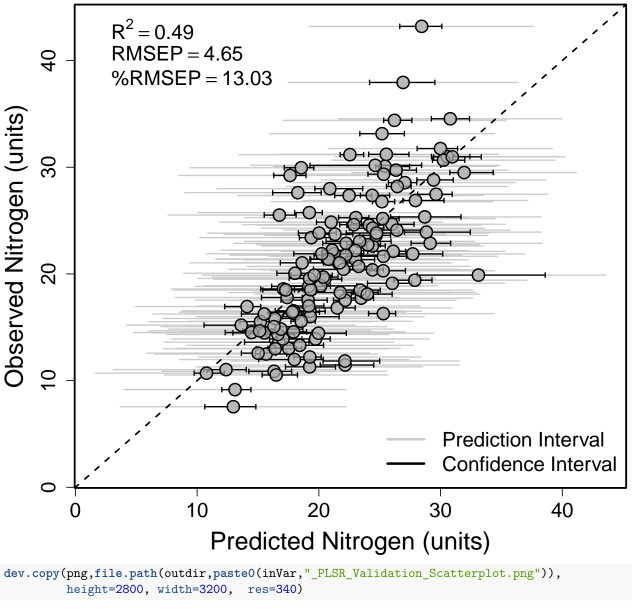
```
## quartz_off_screen
## 3
dev.off();
```

## pdf

#### ## 2

#### Bootstrap validation plot

```
rmsep_percrmsep <- spectratrait::percent_rmse(plsr_dataset = val.plsr.output,</pre>
                                               inVar = inVar,
                                               residuals = val.plsr.output$PLSR_Residuals,
                                               range="full")
RMSEP <- rmsep_percrmsep$rmse
perc_RMSEP <- rmsep_percrmsep$perc_rmse</pre>
r2 <- round(pls::R2(plsr.out, newdata = val.plsr.data, intercept=F)$val[nComps],2)
expr <- vector("expression", 3)</pre>
expr[[1]] <- bquote(R^2==.(r2))
expr[[2]] <- bquote(RMSEP==.(round(RMSEP,2)))</pre>
expr[[3]] <- bquote("%RMSEP"==.(round(perc_RMSEP,2)))</pre>
rng_vals <- c(min(val.plsr.output$LPI), max(val.plsr.output$UPI))</pre>
par(mfrow=c(1,1), mar=c(4.2,5.3,1,0.4), oma=c(0, 0.1, 0, 0.2))
plotrix::plotCI(val.plsr.output$PLSR Predicted,val.plsr.output[,inVar],
                li=val.plsr.output$LPI, ui=val.plsr.output$UPI, gap=0.009,sfrac=0.000,
                lwd=1.6, xlim=c(rng_vals[1], rng_vals[2]), ylim=c(rng_vals[1], rng_vals[2]),
                err="x", pch=21, col="black", pt.bg=scales::alpha("grey70",0.7), scol="grey80",
                cex=2, xlab=paste0("Predicted ", paste(inVar), " (units)"),
                ylab=paste0("Observed ", paste(inVar), " (units)"),
                cex.axis=1.5, cex.lab=1.8)
abline(0,1,lty=2,lw=2)
plotrix::plotCI(val.plsr.output$PLSR_Predicted,val.plsr.output[,inVar],
                li=val.plsr.output$LCI, ui=val.plsr.output$UCI, gap=0.009,sfrac=0.004,
                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=scales::alpha("grey70",0.7), scol="black",
                cex=2, xlab=paste0("Predicted ", paste(inVar), " (units)"),
                ylab=paste0("Observed ", paste(inVar), " (units)"),
                cex.axis=1.5,cex.lab=1.8, add=T)
legend("topleft", legend=expr, bty="n", cex=1.5)
legend("bottomright", legend=c("Prediction Interval", "Confidence Interval"),
       lty=c(1,1), col = c("grey80","black"), lwd=3, bty="n", cex=1.5)
box(1wd=2.2)
```



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

## pdf

#### Output bootstrap results

```
## Iteration Intercept Wave_504 Wave_509 Wave_514 Wave_519 ## 1 1 13.57171 0.2253380 0.1886856 0.1539993 0.09577521
```

## Create core PLSR outputs

```
print(paste("Output directory: ", getwd()))
## [1] "Output directory: /Users/sserbin/Library/CloudStorage/OneDrive-NASA/Data/Github/spectratrait/v
# Observed versus predicted
write.csv(cal.plsr.output,file=file.path(outdir,
                                          paste0(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,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: "
print(list.files(outdir)[grep(pattern = inVar,
                              list.files(outdir))])
##
   [1] "Nitrogen_Bootstrap_PLSR_Coefficients.csv"
  [2] "Nitrogen_Bootstrap_Regression_Coefficients.png"
##
## [3] "Nitrogen_Cal_PLSR_Dataset.csv"
## [4] "Nitrogen_Cal_Val_Histograms.png"
  [5] "Nitrogen_Cal_Val_Scatterplots.png"
##
  [6] "Nitrogen_Cal_Val_Spectra.png"
## [7] "Nitrogen_Coefficient_VIP_plot.png"
   [8] "Nitrogen_Observed_PLSR_CV_Pred_12comp.csv"
##
## [9] "Nitrogen_PLSR_Coefficients_12comp.csv"
## [10] "Nitrogen_PLSR_Component_Selection.png"
## [11] "Nitrogen_PLSR_Validation_Scatterplot.png"
```

```
## [12] "Nitrogen_PLSR_VIPs_12comp.csv"
```

- ## [13] "Nitrogen\_Val\_PLSR\_Dataset.csv"
- ## [14] "Nitrogen\_Validation\_PLSR\_Pred\_12comp.csv"
- ## [15] "Nitrogen\_Validation\_RMSEP\_R2\_by\_Component.png"