An example showing how to apply an existing PLSR model to new data. In this case applying the LMA model from Serbin et al., (2019; DOI - 10.1111/nph.16123) to a dataset collected at CONUS NEON field sites

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Getting Started

Load libraries

```
list.of.packages <- c("pls","dplyr","reshape2","here","plotrix","ggplot2","gridExtra",</pre>
                      "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 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?
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.g. "~/scratch/PLSR"
output_dir <- "tempdir"</pre>
Set working directory (scratch space)
## [1] "/private/var/folders/tq/tydmhlwn1bdf_Opmpcq70r2c0000gn/T/RtmpwSX0i9"
Grab PLSR Coefficients from GitHub
git_repo <- "https://raw.githubusercontent.com/serbinsh/SSerbin_etal_2019_NewPhytologist/master/"</pre>
print("**** Downloading PLSR coefficients ****")
## [1] "**** Downloading PLSR coefficients ****"
githubURL <- paste0(git_repo, "SSerbin_multibiome_lma_plsr_model/sqrt_LMA_gDW_m2_PLSR_Coefficients_10com
LeafLMA.plsr.coeffs <- spectratrait::source_GitHubData(githubURL)</pre>
rm(githubURL)
githubURL <- paste0(git_repo, "SSerbin_multibiome_lma_plsr_model/sqrt_LMA_gDW_m2_Jackkife_PLSR_Coefficie.
LeafLMA.plsr.jk.coeffs <- spectratrait::source_GitHubData(githubURL)</pre>
rm(githubURL)
Get source dataset from EcoSIS
dat_raw <- spectratrait::get_ecosis_data(ecosis_id = ecosis_id)</pre>
## [1] "**** Downloading Ecosis data ****"
## Downloading data...
## Rows: 6312 Columns: 2162
## -- Column specification -------
## Delimiter: ","
       (10): Affiliation, Common Name, Domain, Functional type, Latin Genus, ...
## dbl (2152): LMA, 350, 351, 352, 353, 354, 355, 356, 357, 358, 359, 360, 361,...
## i Use `spec()` to retrieve the full column specification for this data.
## i Specify the column types or set `show_col_types = FALSE` to quiet this message.
```

Download complete!

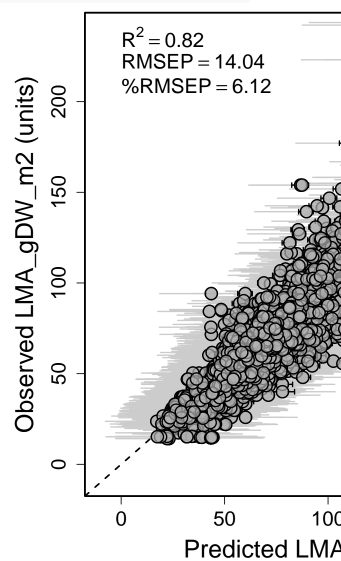
```
head(dat_raw)
## # A tibble: 6 x 2,162
     Affiliation
                           `Common Name` Domain Functional_type
                                                                  LMA `Latin Genus`
     <chr>
                           <chr>
                                          <chr>
                                                 <chr>>
                                                                  <dbl> <chr>
## 1 University of Wiscon~ black walnut DO2
                                                 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
                                                                   60.8 Juglans
## 4 University of Wiscon~ black walnut DO2
                                                 broadleaf
## 5 University of Wiscon~ black walnut DO2
                                                                   85.9 Juglans
                                                 broadleaf
## 6 University of Wiscon~ black walnut DO2
                                                 broadleaf
                                                                   85.9 Juglans
## # i 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>, ...
## #
names(dat raw)[1:40]
##
   [1] "Affiliation"
                          "Common Name"
                                             "Domain"
                                                                "Functional_type"
   [5] "LMA"
                          "Latin Genus"
                                             "Latin Species"
                                                                "PI"
                                                                "350"
   [9] "Project"
                          "Sample_ID"
                                             "USDA Symbol"
##
## [13] "351"
                          "352"
                                             "353"
                                                                "354"
                          "356"
                                             "357"
## [17] "355"
                                                                "358"
## [21] "359"
                          "360"
                                             "361"
                                                                "362"
                          "364"
## [25] "363"
                                             "365"
                                                                "366"
## [29] "367"
                          "368"
                                             "369"
                                                                "370"
                          "372"
                                             "373"
                                                                "374"
## [33] "371"
## [37] "375"
                          "376"
                                             "377"
                                                                "378"
Prepare new data for estimation
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>
head(Spectra)[1:6,1:10]
        Wave_500 Wave_501 Wave_502 Wave_503 Wave_504 Wave_505 Wave_506 Wave_507
##
## [1,] 0.044226 0.044605 0.044927 0.045473 0.046241 0.046878 0.047826 0.049090
## [2,] 0.046855 0.047601 0.047944 0.048478 0.049381 0.050235 0.051161 0.052191
## [3,] 0.043758 0.044171 0.044869 0.045465 0.045984 0.046933 0.047993 0.049090
## [4,] 0.041154 0.041603 0.042088 0.042408 0.042639 0.043260 0.044140 0.045058
## [5,] 0.037296 0.037944 0.038209 0.038677 0.039388 0.039948 0.040630 0.041501
## [6,] 0.043878 0.044257 0.044723 0.045295 0.045949 0.046575 0.047378 0.048357
        Wave_508 Wave_509
## [1,] 0.050268 0.051525
## [2,] 0.053322 0.054357
## [3,] 0.050168 0.051441
## [4,] 0.045700 0.046476
## [5,] 0.042613 0.043731
## [6,] 0.049392 0.050387
```

```
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>>
                           <chr>>
                                          <chr> <chr>
                                                                 <dbl> <chr>
## 1 University of Wiscon~ black walnut DO2
                                                 broadleaf
                                                                  72.9 Juglans
## 2 University of Wiscon~ black walnut DO2
                                                                  72.9 Juglans
                                                broadleaf
## 3 University of Wiscon~ black walnut DO2 broadleaf
                                                                  60.8 Juglans
                                                                  60.8 Juglans
## 4 University of Wiscon~ black walnut DO2
                                                broadleaf
## 5 University of Wiscon~ black walnut DO2 broadleaf
                                                                  85.9 Juglans
## 6 University of Wiscon~ black walnut DO2
                                                broadleaf
                                                                  85.9 Juglans
## # i 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 DO2 broadleaf
                            L0002
                                      JUNI
                                                               60.8
## 5 DO2 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)
#### 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[complete.cases(plsr_data[,names(plsr_data) %in%
                                                   c(inVar,paste0("Wave_",wv))]),]
Example data cleaning.
print("**** Applying PLSR model to estimate LMA from spectral observations ****")
Prepare PLSR model
## [1] "**** Applying PLSR model to estimate LMA from spectral observations ****"
# setup model
dims <- dim(LeafLMA.plsr.coeffs)</pre>
LeafLMA.plsr.intercept <- LeafLMA.plsr.coeffs[1,]</pre>
LeafLMA.plsr.coeffs <- data.frame(LeafLMA.plsr.coeffs[2:dims[1],])</pre>
names(LeafLMA.plsr.coeffs) <- c("wavelength", "coefs")</pre>
LeafLMA.plsr.coeffs.vec <- as.vector(LeafLMA.plsr.coeffs[,2])</pre>
sub_spec <- droplevels(plsr_data[,which(names(plsr_data) %in%</pre>
                                                    paste0("Wave_",seq(Start.wave,End.wave,1)))])
```

```
plsr_pred <- as.matrix(sub_spec) %*% LeafLMA.plsr.coeffs.vec + LeafLMA.plsr.intercept[,2]
leafLMA <- plsr pred[,1]^2 # convert to standard LMA units from sqrt(LMA)</pre>
names(leafLMA) <- "PLSR LMA gDW m2"</pre>
# organize output
LeafLMA.PLSR.dataset <- data.frame(plsr_data[,which(names(plsr_data) %notin%
                                                       paste0("Wave_",seq(Start.wave,End.wave,1)))],
                                    PLSR LMA gDW m2=leafLMA, PLSR Residuals=leafLMA-plsr data[,inVar])
head(LeafLMA.PLSR.dataset)
Apply PLSR model
     Domain Functional_type Sample_ID USDA_Species_Code LMA_gDW_m2 PLSR_LMA_gDW_m2
## 1
        D02
                  broadleaf
                                 P0001
                                                    JUNI
                                                               72.87
                                                                            96.26243
## 2
        D02
                  broadleaf
                                                    JUNI
                                                               72.87
                                                                            90.09453
                                 L0001
## 3
        D02
                  broadleaf
                                 P0002
                                                    JUNI
                                                               60.77
                                                                            77.16475
## 4
        D02
                  broadleaf
                                 L0002
                                                    JUNI
                                                               60.77
                                                                            60.99039
## 5
        D02
                  broadleaf
                                                               85.92
                                                                           101.22709
                                 P0003
                                                    JUNI
## 6
        D02
                  broadleaf
                                 L0003
                                                    JUNI
                                                               85.92
                                                                            97.13018
    PLSR_Residuals
##
         23.3924343
## 1
         17.2245326
## 2
## 3
         16.3947533
## 4
         0.2203913
## 5
         15.3070857
## 6
         11.2101840
print("**** Generate PLSR uncertainty estimates ****")
Generate PLSR uncertainty estimates
## [1] "**** Generate PLSR uncertainty estimates ****"
jk_coef <- data.frame(LeafLMA.plsr.jk.coeffs[,3:dim(LeafLMA.plsr.jk.coeffs)[2]])</pre>
jk coef <- t(jk coef)</pre>
head(jk_coef)[,1:6]
                                      [,3]
##
                 [,1]
                            [,2]
                                                [,4]
                                                           [,5]
                                                                     [.6]
## Wave 500 1.0005875 0.9952840 0.5652908 0.9793160 1.1052207 0.9370473
## Wave_501 0.9584235 0.9631434 0.5230544 0.9330803 1.0477469 0.9042780
## Wave 502 0.8960202 0.9065954 0.4597413 0.8710298 0.9658130 0.8628370
## Wave_503 0.8722135 0.8936197 0.4420696 0.8456098 0.9272967 0.8513741
## Wave_504 0.8452831 0.8644923 0.4159567 0.8110004 0.8903192 0.8320347
## Wave_505 0.8240743 0.8378399 0.3902871 0.7829891 0.8570048 0.8150339
jk_int <- t(LeafLMA.plsr.jk.coeffs[,2])</pre>
head(jk int)[,1:6]
## [1] 7.787098 7.959443 8.015161 8.018586 7.658080 7.998432
jk pred <- as.matrix(sub spec) ** jk coef + matrix(rep(jk int, length(plsr data[,inVar])),
                                                     byrow=TRUE, ncol=length(jk_int))
jk_pred <- jk_pred^2
head(jk_pred)[,1:6]
```

```
[,2]
                              [,3]
         [,1]
                                       [,4]
                                                  [,5]
                                                            [,6]
## 1 94.28721 96.77712 96.44452 95.11992 96.72830 95.33877
## 2 90.36051 90.57120 90.77562 89.77821 90.24826 89.61806
## 3 75.71088 77.91861 76.42730 76.11473 77.67179 76.68756
## 4 61.37001 61.30963 60.56606 60.72330 61.63712 60.69649
## 5 99.24456 101.75948 101.22916 99.96305 101.70397 100.16758
## 6 97.40414 97.65463 97.52687 97.00817 97.33677 96.08535
dim(jk pred)
## [1] 6312 1000
interval <-c(0.025,0.975)
Interval_Conf <- apply(X = jk_pred, MARGIN = 1, FUN = quantile,</pre>
                       probs=c(interval[1], interval[2]))
sd_mean <- apply(X = jk_pred, MARGIN = 1, FUN =sd)</pre>
sd_res <- sd(LeafLMA.PLSR.dataset$PLSR_Residuals)</pre>
sd tot <- sqrt(sd mean^2+sd res^2)</pre>
LeafLMA.PLSR.dataset$LCI <- Interval Conf[1,]</pre>
LeafLMA.PLSR.dataset$UCI <- Interval_Conf[2,]</pre>
LeafLMA.PLSR.dataset$LPI <- LeafLMA.PLSR.dataset$PLSR_LMA_gDW_m2-1.96*sd_tot
LeafLMA.PLSR.dataset$UPI <- LeafLMA.PLSR.dataset$PLSR_LMA_gDW_m2+1.96*sd_tot
head(LeafLMA.PLSR.dataset)
     Domain Functional_type Sample_ID USDA_Species_Code LMA_gDW_m2 PLSR_LMA_gDW_m2
##
## 1
        D02
                  broadleaf
                                 P0001
                                                     JUNI
                                                               72.87
                                                                            96.26243
                                                                            90.09453
## 2
        D02
                  broadleaf
                                 L0001
                                                     JUNI
                                                               72.87
## 3
        D02
                  broadleaf
                                 P0002
                                                     JUNI
                                                               60.77
                                                                            77.16475
## 4
        D02
                  broadleaf
                                                               60.77
                                                                            60.99039
                                 L0002
                                                     JUNI
## 5
        D02
                  broadleaf
                                 P0003
                                                     JUNI
                                                               85.92
                                                                           101.22709
                                                               85.92
## 6
        D02
                  broadleaf
                                 L0003
                                                     JUNI
                                                                            97.13018
    PLSR Residuals
                         LCI
                                    UCI
                                             LPI
                                                       UPT
         23.3924343 93.95423 99.03625 71.30476 121.2201
## 1
## 2
         17.2245326 88.81329 92.00078 65.21071 114.9784
## 3
        16.3947533 74.79509 79.85715 52.19722 102.1323
## 4
         0.2203913 59.80058 62.29402 36.12678 85.8540
## 5
         15.3070857 98.86570 103.97701 76.26586 126.1883
## 6
         11.2101840 95.50843 99.66865 72.20971 122.0507
rmsep_percrmsep <- spectratrait::percent_rmse(plsr_dataset = LeafLMA.PLSR.dataset,</pre>
                                               inVar = inVar,
                                               residuals = LeafLMA.PLSR.dataset$PLSR_Residuals,
                                               range="full")
RMSEP <- rmsep_percrmsep$rmse
perc_RMSEP <- rmsep_percrmsep$perc_rmse</pre>
r2 <- round(summary(lm(LeafLMA.PLSR.dataset$PLSR_LMA_gDW_m2~
                         LeafLMA.PLSR.dataset[,inVar]))$adj.r.squared,2)
expr <- vector("expression", 3)</pre>
expr[[1]] \leftarrow 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(LeafLMA.PLSR.dataset$LPI), max(LeafLMA.PLSR.dataset$UPI))</pre>
par(mfrow=c(1,1), mar=c(4,5.3,1,0.4), oma=c(0.1, 0.1, 0.1, 0.2))
plotrix::plotCI(LeafLMA.PLSR.dataset$PLSR_LMA_gDW_m2,LeafLMA.PLSR.dataset[,inVar],
```

```
li=LeafLMA.PLSR.dataset$LPI, ui=LeafLMA.PLSR.dataset$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(LeafLMA.PLSR.dataset$PLSR LMA gDW m2,LeafLMA.PLSR.dataset[,inVar],
                li=LeafLMA.PLSR.dataset$LCI, ui=LeafLMA.PLSR.dataset$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(lwd=2.2)
```



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_PLSR_Estimates.csv"

## [2] "LMA_gDW_m2_PLSR_Validation_Scatterplot.png"
```