Example application of multi-biome LMA PLSR model

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Overview

This is an R Markdown Notebook to illustrate how to apply the multi-biome LMA PLSR model to leaf reflectance spectra.

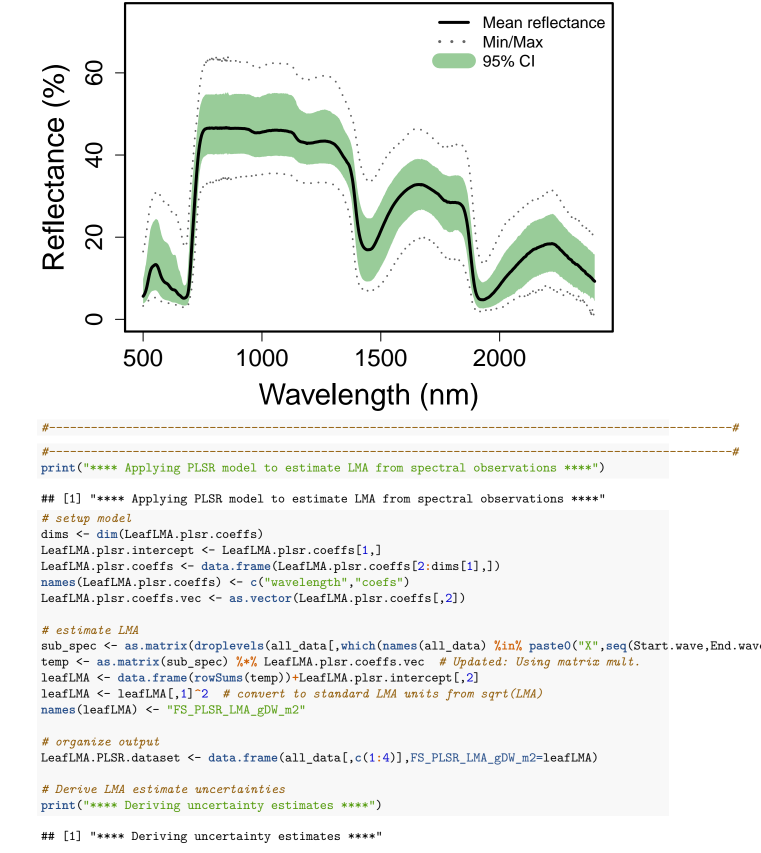
Try executing this chunk by clicking the Run button within the chunk or by placing your cursor inside it and pressing Cmd+Shift+Enter.

```
# get all required libraries
list.of.packages <- c("readr", "scales", "plotrix", "httr", "devtools") # packages needed for script
# check for dependencies and install if needed
new.packages <- list.of.packages[!(list.of.packages %in% installed.packages()[,"Package"])]
if(length(new.packages)) install.packages(new.packages)
# load libraries needed for script
library(readr) # readr - read_csv function to pull data from EcoSIS
library(plotrix) # plotCI - to generate obsvered vs predicted plot with CIs
library(scales) # alpha() - for applying a transparency to data points
##
## Attaching package: 'scales'
## The following object is masked from 'package:plotrix':
##
##
       rescale
## The following object is masked from 'package:readr':
##
       col_factor
library(devtools)
## Loading required package: usethis
library(httr)
# define function to grab PLSR model from GitHub
\#devtools::source\_gist("gist.github.com/christophergandrud/4466237")
source_GitHubData <-function(url, sep = ",", header = TRUE) {</pre>
 require(httr)
 request <- GET(url)</pre>
  stop_for_status(request)
 handle <- textConnection(content(request, as = 'text'))</pre>
  on.exit(close(handle))
  read.table(handle, sep = sep, header = header)
### PLSR Coefficients - Grab 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 <- 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 <- source_GitHubData(githubURL)</pre>
Example datasets
URL: https://ecosis.org/#result/13aef0ce-dd6f-4b35-91d9-28932e506c41 (Lopex)
URL: https://ecosis.org/#result/2231d4f6-981e-4408-bf23-1b2b303f475e (Angers)
#-----
### Grab data
print("**** Downloading Ecosis data ****")
## [1] "**** Downloading Ecosis data ****"
ecosis_id <- "13aef0ce-dd6f-4b35-91d9-28932e506c41" # lopex
ecosis_file <- sprintf(</pre>
  "https://ecosis.org/package/export?package_id=%s&filters=&metadata=true",
  ecosis_id
message("Downloading data...")
## Downloading data...
dat_raw <- read_csv(ecosis_file)</pre>
## Parsed with column specification:
## cols(
##
     .default = col_double(),
##
     `English Name` = col_character(),
    `Latin Name` = col_character(),
##
    Measurement_type = col_character(),
##
##
    Refl_file = col_character(),
##
    Trans_file = col_character()
## )
```

```
## See spec(...) for full column specifications.
message("Download complete!")
## Download complete!
# keep just fresh leaf refl obs. remove dried leaves from sample set
remove \leftarrow c(176,177,178,179,180,196,197,198,199,200,321,322,323,324,325)
remove <- which(dat_raw$Measurement_type=="transmittance" | dat_raw$`Sample_#` %in% remove)
lopex_dat_clean <- dat_raw[-remove,]</pre>
ecosis_id <- "2231d4f6-981e-4408-bf23-1b2b303f475e" # angers
ecosis file <- sprintf(</pre>
  "https://ecosis.org/package/export?package id=%s&filters=&metadata=true",
  ecosis_id
message("Downloading data...")
## Downloading data...
dat_raw <- read_csv(ecosis_file)</pre>
## Parsed with column specification:
## cols(
     .default = col_double(),
##
     `English Name` = col_character(),
##
    `Latin Name` = col_character(),
##
     Measurement_type = col_character(),
    Refl_file = col_character(),
    Trans_file = col_character()
##
## )
## See spec(...) for full column specifications.
message("Download complete!")
## Download complete!
# cleanup and remove dried leaves from dataset
remove <- c(178,179,184,185,196,197,241,250,254,257,258,269)
remove <- which(dat_raw$Measurement_type=="transmittance" | dat_raw$`Sample_#` %in% remove)
angers_dat_clean <- dat_raw[-remove,]</pre>
rm(dat raw)
## Concatenate data
Start.wave <- 500
End.wave <- 2400
wv <- seq(Start.wave, End.wave, 1)</pre>
lopex_spectra_sub <- lopex_dat_clean[,names(lopex_dat_clean)[match(seq(Start.wave,End.wave,1),names(lop
lopex_info <- data.frame(Sample_Num=lopex_dat_clean$`Sample_#`, Common_Species_Name=lopex_dat_clean$`En
                         LMA_gDW_m2=(lopex_dat_clean$`Leaf mass per area (g/cm²)`)*10000)
angers_spectra_sub <- angers_dat_clean[,names(angers_dat_clean)[match(seq(Start.wave,End.wave,1),names(
angers_spectra_sub <- na.omit(angers_spectra_sub)</pre>
angers_info <- data.frame(Sample_Num=angers_dat_clean$`Sample_#`, Common_Species_Name=angers_dat_clean$
```

```
LMA_gDW_m2=(angers_dat_clean$`Leaf mass per area (g/cm )`)*10000)
angers_info <- na.omit(angers_info)</pre>
all_data <- rbind(data.frame(Dataset=rep("Lopex",dim(lopex_info)[1]), lopex_info, lopex_spectra_sub),
                  data.frame(Dataset=rep("Angers",dim(angers_info)[1]), angers_info, angers_spectra_sub
## cleanup
rm(angers_dat_clean,lopex_dat_clean, angers_info, angers_spectra_sub, lopex_info, lopex_spectra_sub)
## Plot data
waves <- seq(500, 2400, 1)
cexaxis <- 1.5
cexlab <- 1.8
vlim <- 74
ylim2 <- 80
mean_spec <- colMeans(all_data[,which(names(all_data) %in% paste0("X",seq(Start.wave,End.wave,1)))])
spectra_quantiles <- apply(all_data[,which(names(all_data) %in% paste0("X",seq(Start.wave,End.wave,1)));</pre>
                           2,quantile,na.rm=T,probs=c(0,0.025,0.05,0.5,0.95,0.975,1))
print("**** Plotting Ecosis data. Writing to scratch space ****")
## [1] "**** Plotting Ecosis data. Writing to scratch space ****"
png(file=file.path("~",wd,'Angers_Lopex_spectra_summary_plot.png'),height=3000,
    width=3900, res=340)
par(mfrow=c(1,1), mar=c(4.5,5.7,0.3,0.4), om=c(0.3,0.9,0.3,0.1)) # B, L, T, R
plot(waves, mean spec*100, ylim=c(0, ylim), cex=0.00001, col="white", xlab="Wavelength (nm)",
     ylab="Reflectance (%)",cex.axis=cexaxis, cex.lab=cexlab)
polygon(c(waves ,rev(waves)),c(spectra_quantiles[6,]*100, rev(spectra_quantiles[2,]*100)),
        col="#99CC99",border=NA)
lines(waves,mean_spec*100,lwd=3, lty=1, col="black")
lines(waves, spectra quantiles[1,]*100,lwd=1.85, lty=3, col="grey40")
lines(waves, spectra_quantiles[7,]*100,lwd=1.85, lty=3, col="grey40")
legend("topright",legend=c("Mean reflectance","Min/Max", "95% CI"),lty=c(1,3,1),
       lwd=c(3,3,15),col=c("black","grey40","#99CC99"),bty="n", cex=1.7)
box(1wd=2.2)
dev.off()
## pdf
par(mfrow=c(1,1), mar=c(4.5,5.7,0.3,0.4), om=c(0.3,0.9,0.3,0.1)) # B, L, T, R
plot(waves, mean_spec*100, ylim=c(0, ylim), cex=0.00001, col="white", xlab="Wavelength (nm)",
     ylab="Reflectance (%)",cex.axis=cexaxis, cex.lab=cexlab)
polygon(c(waves ,rev(waves)),c(spectra_quantiles[6,]*100, rev(spectra_quantiles[2,]*100)),
        col="#99CC99",border=NA)
lines(waves,mean_spec*100,lwd=3, lty=1, col="black")
lines(waves, spectra_quantiles[1,]*100,lwd=1.85, lty=3, col="grey40")
lines(waves, spectra quantiles[7,]*100,lwd=1.85, lty=3, col="grey40")
legend("topright",legend=c("Mean reflectance","Min/Max", "95% CI"),lty=c(1,3,1),
       lwd=c(3,3,15),col=c("black","grey40","#99CC99"),bty="n", cex=1)
box(1wd=2.2)
```



```
dims <- dim(LeafLMA.plsr.jk.coeffs)</pre>
intercepts <- LeafLMA.plsr.jk.coeffs[,2]</pre>
jk.leaf.lma.est <- array(data=NA,dim=c(dim(sub_spec)[1],dims[1]))
for (i in 1:length(intercepts)){
  coefs <- unlist(as.vector(LeafLMA.plsr.jk.coeffs[i,3:dims[2]]))</pre>
  temp <- sub_spec %*% coefs</pre>
  values <- data.frame(rowSums(temp))+intercepts[i]</pre>
  jk.leaf.lma.est[,i] <- values[,1]^2</pre>
 rm(temp)
jk.leaf.lma.est.quant <- apply(jk.leaf.lma.est,1,quantile,probs=c(0.025,0.975))
jk.leaf.lma.est.quant2 <- data.frame(t(jk.leaf.lma.est.quant))</pre>
names(jk.leaf.lma.est.quant2) <- c("FS_PLSR_Leaf_LMA_L5","FS_PLSR_Leaf_LMA_U95")</pre>
jk.leaf.lma.est.sd <- apply(jk.leaf.lma.est,1,sd)</pre>
names(jk.leaf.lma.est.sd) <- "FS_PLSR_Leaf_LMA_Sdev"</pre>
## Combine into final dataset
stats <- data.frame(jk.leaf.lma.est.sd,jk.leaf.lma.est.quant2)
names(stats) <- c("FS_PLSR_Leaf_LMA_Sdev","FS_PLSR_Leaf_LMA_L5","FS_PLSR_Leaf_LMA_U95")</pre>
LeafLMA.PLSR.dataset.out <- data.frame(LeafLMA.PLSR.dataset,stats,</pre>
                                        residual=(LeafLMA.PLSR.dataset$FS_PLSR_LMA_gDW_m2-LeafLMA.PLSR.d
# output results
write.csv(x = LeafLMA.PLSR.dataset.out, file = file.path("~",wd,"Angers_Lopex_PLSR_estimated_LMA_data.c
          row.names = F)
# calculate error stats
rmse <- sqrt(mean(LeafLMA.PLSR.dataset.out$residual^2))</pre>
# calculate fit stats
reg <- lm(LeafLMA.PLSR.dataset.out$FS_PLSR_LMA_gDW_m2~LeafLMA.PLSR.dataset.out$LMA_gDW_m2)
## Plot up results
ptcex <- 1.8
cexaxis <- 1.3
cexlab <- 1.8
print("**** Plotting Lopex/Angers LMA validation plot. Writing to scratch space ****")
## [1] "**** Plotting Lopex/Angers LMA validation plot. Writing to scratch space ****"
png(file=file.path("~",wd,'Angers_Lopex_LMA_validation_plot.png'),height=3000,
    width=3900, res=340)
par(mfrow=c(1,1), mar=c(4.5,5.4,1,1), oma=c(0.3,0.9,0.3,0.1)) # B, L, T, R
plotCI(LeafLMA.PLSR.dataset.out$FS_PLSR_LMA_gDW_m2,LeafLMA.PLSR.dataset.out$LMA_gDW_m2,
       li=LeafLMA.PLSR.dataset.out$FS_PLSR_Leaf_LMA_L5,gap=0.009,sfrac=0.004,lwd=1.6,
       ui=LeafLMA.PLSR.dataset.out$FS_PLSR_Leaf_LMA_U95,err="x",pch=21,col="black",
       pt.bg=alpha("grey70",0.7),scol="grey30",xlim=c(0,340),cex=ptcex,
       ylim=c(0,340),xlab="",
       vlab=expression(paste("Observed LMA (",g~m^{-2},")")),main="",
       cex.axis=cexaxis,cex.lab=cexlab)
mtext(side = 1, text = expression(paste(Predicted~LMA," (",g~m^{-2},")")), line = 3.5,
      cex=2.2)
```

```
abline(0,1,lty=2,lw=2)
legend("topleft",legend = c(paste0("RMSE = ",round(rmse)),
                            paste0("R2 = ",round(summary(reg)$r.squared,2))), bty="n", cex=1.5)
box(1wd=2.2)
dev.off()
## pdf
##
par(mfrow=c(1,1), mar=c(4.5,5.4,1,1), oma=c(0.3,0.9,0.3,0.1)) # B, L, T, R
plotCI(LeafLMA.PLSR.dataset.out$FS_PLSR_LMA_gDW_m2,LeafLMA.PLSR.dataset.out$LMA_gDW_m2,
       li=LeafLMA.PLSR.dataset.out$FS_PLSR_Leaf_LMA_L5,gap=0.009,sfrac=0.004,lwd=1.6,
       ui=LeafLMA.PLSR.dataset.out$FS_PLSR_Leaf_LMA_U95,err="x",pch=21,col="black",
       pt.bg=alpha("grey70",0.7),scol="grey30",xlim=c(0,340),cex=ptcex,
      ylim=c(0,340),xlab="",
      ylab=expression(paste("Observed LMA (",g~m^{-2},")")),main="",
       cex.axis=cexaxis,cex.lab=cexlab)
mtext(side = 1, text = expression(paste(Predicted~LMA," (",g~m^{-2},")")), line = 3.5,
      cex=2.2)
abline(0,1,lty=2,lw=2)
legend("topleft",legend = c(paste0("RMSE = ",round(rmse)),
                            paste0("R2 = ",round(summary(reg)$r.squared,2))), bty="n", cex=1.5)
box(lwd=2.2)
        350
                                                      00
Observed LMA (g \mathsf{m}^{-2}
                    RMSE = 21
                    R2 = 0.66
        250
        50
        50
                                          150
                                                   200
                                                            250
               0
                        50
                                100
                                                                      300
                                                                               350
                        Predicted LMA (g m<sup>-2</sup>)
```

clear workspace

rm(list=ls(all=TRUE))

EOF