

Pondo__Allom__Params

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Introduction:

Information about allometric curves and functions used in FATES. https://fates-docs.readthedocs.io/en/latest/fates_tech_note.html#allometry-and-growth-along-allometric-curves

Install the BAAD Package:

Install the package from github:

```
#install.packages("devtools")
#devtools::install_github("richfitz/datastorr")
#devtools::install_github("traitecoevo/baad.data")
```

Review the data:

```
baad <- baad.data::baad_data()
d_baad <- baad$data
head(d_baad)
```

```
##      studyName      location latitude longitude vegetation map mat grouping
## 1  Abe1981 Japan-Oumu-46      45      143      BorF  NA  NA      <NA>
## 2  Abe1981 Japan-Oumu-46      45      143      BorF  NA  NA      <NA>
## 3  Abe1981 Japan-Oumu-46      45      143      BorF  NA  NA      <NA>
## 4  Abe1981 Japan-Oumu-46      45      143      BorF  NA  NA      <NA>
## 5  Abe1981 Japan-Oumu-46      45      143      BorF  NA  NA      <NA>
## 6  Abe1981 Japan-Oumu-46      45      143      BorF  NA  NA      <NA>
##      lai      species      speciesMatched      family pft
## 1  NA Abies sachalinensis Abies sachalinensis Pinaceae EA
## 2  NA Abies sachalinensis Abies sachalinensis Pinaceae EG
## 3  NA Abies sachalinensis Abies sachalinensis Pinaceae EG
## 4  NA Abies sachalinensis Abies sachalinensis Pinaceae EG
## 5  NA Abies sachalinensis Abies sachalinensis Pinaceae EG
## 6  NA Abies sachalinensis Abies sachalinensis Pinaceae EG
##      growingCondition status light age a.lf a.ssba a.ssbh a.ssbh a.ssbh a.shba
## 1      PM      NA      <NA>  NA  NA      NA      NA      NA      NA
## 2      PM      NA      <NA>  NA  NA      NA      NA      NA      NA
## 3      PM      NA      <NA>  NA  NA      NA      NA      NA      NA
## 4      PM      NA      <NA>  NA  NA      NA      NA      NA      NA
## 5      PM      NA      <NA>  NA  NA      NA      NA      NA      NA
## 6      PM      NA      <NA>  NA  NA      NA      NA      NA      NA
##      a.shbh a.shbh a.sbbh a.sbbh a.sbbh      a.stba      a.stbh      a.stbc
## 1      NA      NA      NA      NA      NA  0.18973040 0.13074052 0.066508302
## 2      NA      NA      NA      NA      NA  0.14286569 0.09731397 0.056410438
## 3      NA      NA      NA      NA      NA  0.19658523 0.07068583 0.035298935
## 4      NA      NA      NA      NA      NA  0.07068583 0.04523893 0.020611989
## 5      NA      NA      NA      NA      NA  0.02198274 0.01583677 0.008992024
## 6      NA      NA      NA      NA      NA  0.18957602 0.12692348 0.059828490
```

```
##   a.cp a.cs   h.t   h.c   d.ba d.bh h.bh d.cr   c.d m.lf m.ss m.sh m.sb
## 1   NA   NA 21.96  8.90 0.4915 0.408  1.3   NA 13.06 49.2   NA   NA   NA
## 2   NA   NA 19.70  8.02 0.4265 0.352  1.3   NA 11.68 28.0   NA   NA   NA
## 3   NA   NA 20.10  9.50 0.5003 0.300  1.3   NA 10.60 19.9   NA   NA   NA
## 4   NA   NA 19.23 11.60 0.3000 0.240  1.3   NA  7.63 11.1   NA   NA   NA
## 5   NA   NA 16.14  9.35 0.1673 0.142  1.3   NA  6.79  3.8   NA   NA   NA
## 6   NA   NA 22.26 11.02 0.4913 0.402  1.3   NA 11.24 33.4   NA   NA   NA
##   m.st m.so m.br m.rf m.rc m.rt m.to a.ilf ma.ilf r.st r.ss r.sb r.sh
## 1 516.8 566.0 113.8   NA   NA   NA   NA   NA   NA   NA   NA   NA   NA
## 2 427.7 455.7  84.9   NA   NA   NA   NA   NA   NA   NA   NA   NA   NA
## 3 255.0 274.9  30.2   NA   NA   NA   NA   NA   NA   NA   NA   NA   NA
## 4 172.9 184.0  14.4   NA   NA   NA   NA   NA   NA   NA   NA   NA   NA
## 5  58.6  62.4   4.6   NA   NA   NA   NA   NA   NA   NA   NA   NA   NA
## 6 561.9 595.3  61.3   NA   NA   NA   NA   NA   NA   NA   NA   NA   NA
##   n.lf n.ss n.sb n.sh n.rf n.rc
## 1   NA   NA   NA   NA   NA   NA
## 2   NA   NA   NA   NA   NA   NA
## 3   NA   NA   NA   NA   NA   NA
## 4   NA   NA   NA   NA   NA   NA
## 5   NA   NA   NA   NA   NA   NA
## 6   NA   NA   NA   NA   NA   NA
```

Refine (subset) the data for Pinus Ponderosa:

```
pipo <- d_baad[ which(d_baad$species == 'Pinus ponderosa'), ]
#head(pipo)
```

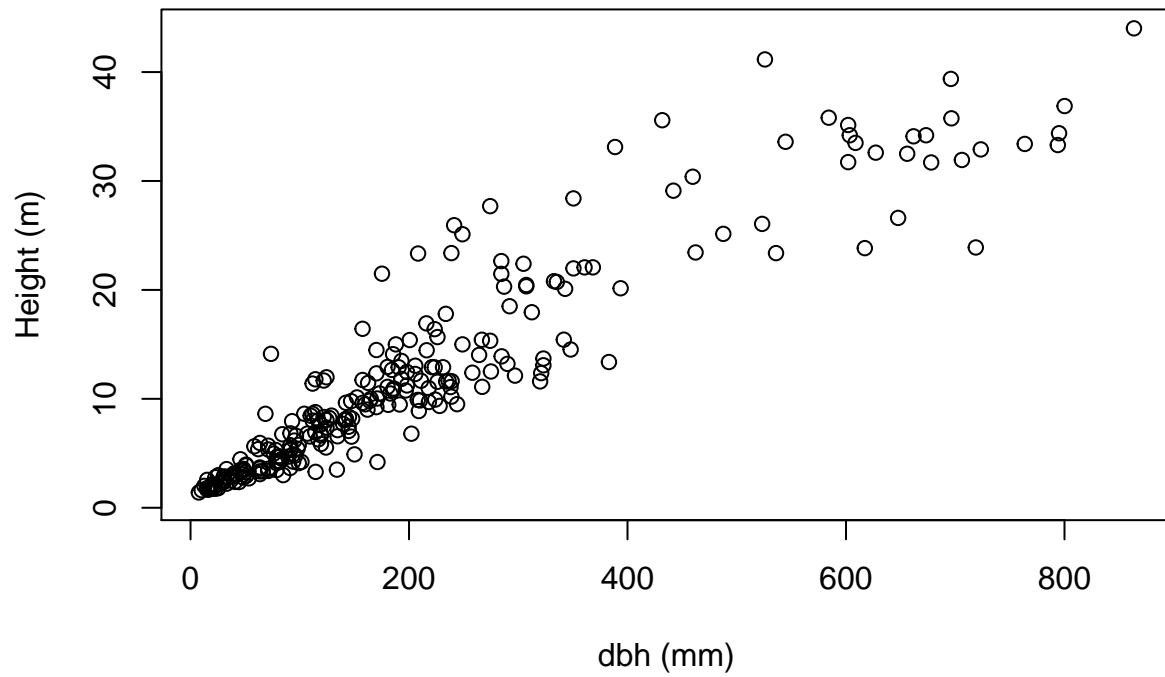
Plot raw data, dbh to height.

```
# Set variables from the data.

# Diameter at breast height (dbh) will be used for most allometric calculations
dbh_mm <- (pipo$d.bh)*1000
dbh_cm <- (pipo$d.bh)*100
# Height
h <- (pipo$h.t)

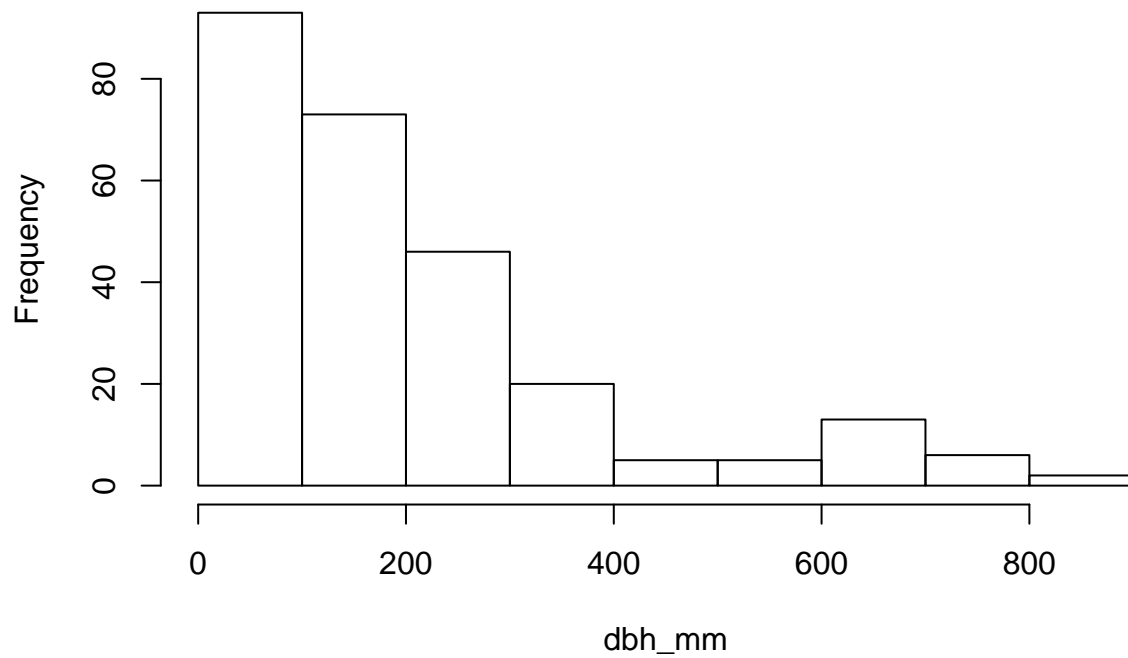
plot(h~dbh_mm,
     main="Diameter at breast height (dbh) to Height",
     xlab = "dbh (mm)",
     ylab = "Height (m)")
```

Diameter at breast height (dbh) to Height



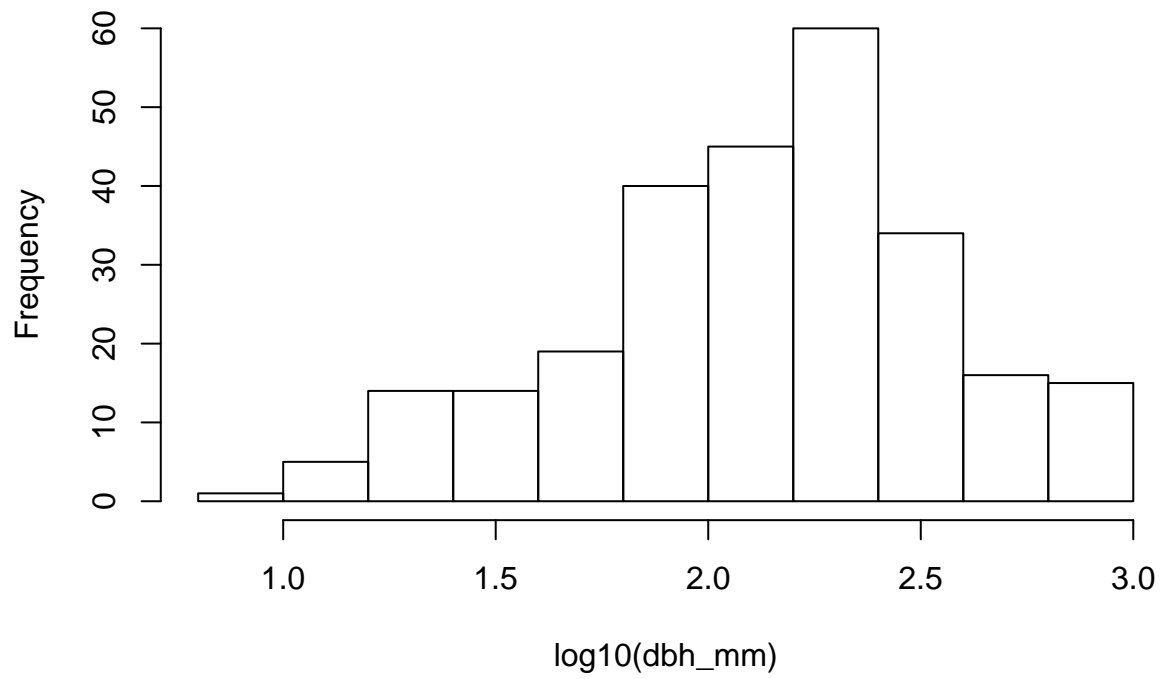
```
hist(dbh_mm)
```

Histogram of dbh_mm



```
hist(log10(dbh_mm))
```

Histogram of $\log_{10}(\text{dbh_mm})$



Diameter at breast height (dbh) to height - d2h

FATES allows for four different approaches to predicting height from dbh:

- A power function

$$h = p_1 * d^{(p_2)}$$

- O'Brien et al. (1995)

$$\log_{10}Height = \log_{10}DBH * slope + intercept$$

- Poorter et al. (2006)

$$h = p_1 * (1 - \exp(p_2 * d^{(p_3)}))$$

- Martinex Cano et al. (2019)

$$h = (p_1 * d^{(p_2)} / (p_3 + d^{(p_2)}))$$

For my research I use the dbh to height relationship from O'Brien et al., 1995. The $\log_{10}Height$ (m) is regressed on $\log_{10}DBH$ (mm). Where slope is p1 or `fates_allom_d2h1` and intercept is p2 or `fates_allom_d2h2` in the parameter file.

$$\log_{10}Height = \log_{10}DBH * slope + intercept$$

So modeled height would be. I want to solve for the best fit p1 and p2 given height and dbh data from BAAD.

$$Height = 10^{(\log_{10}(\min(d, dbh_{max})) * p1 + p2)}$$

```
# But first I will test out just a linear model
```

```
d2hmod_mm <- lm(h~dbh_mm)
coef(d2hmod_mm)
```

```
## (Intercept)      dbh_mm
##  1.88991834  0.04858541
```

```
# Then a linear model with log10(h) regressed on log10(dbh)
```

```
d2hmodlog_mm <- lm(log10(h)~log10(dbh_mm))
coef(d2hmodlog_mm)
```

```
## (Intercept) log10(dbh_mm)
##   -0.8150832    0.8181127
```

```
d2hmodlog_cm <- lm(log10(h)~log10(dbh_cm))
coef(d2hmodlog_cm)
```

```
## (Intercept) log10(dbh_cm)
##  0.003029536    0.818112721
```

Next, I will create a function to represent the O'Brien calculation used for dbh to height relationships.

```
obrien <- function(dbh, p1, p2){
  height <- 10^((log10(dbh))* p1 + p2)
  return(height)
}
```

```
dbins_mm <- c(100,200,300,400,500,600,700,800,900) # Remember in O'Brien dbh is in mm.
dbins_cm <- c(10,20,30,40,50,60,70,80,90) # But does FATES do that conversion or use cm?
```

```
# Use default parameters from O'Brien et al 1995
```

```
default_mm <- obrien(dbins_mm, 0.7, -0.2)
```

```

# Use the parameters from the log10 linear model d2hmodlog
lmfit_mm <- obrien(dbins_mm, 0.82, -0.82) # dbh in mm
lmfit_cm <- obrien(dbins_cm, 0.82, 0.003) # dbh in cm

coef(lm(default_mm~dbins_mm))

## (Intercept)      dbins_mm
## 11.67774249  0.07124032

coef(lm(lmfit_mm~dbins_mm))

## (Intercept)      dbins_mm
##  3.45859976  0.04142493

coef(lm(lmfit_cm~dbins_cm))

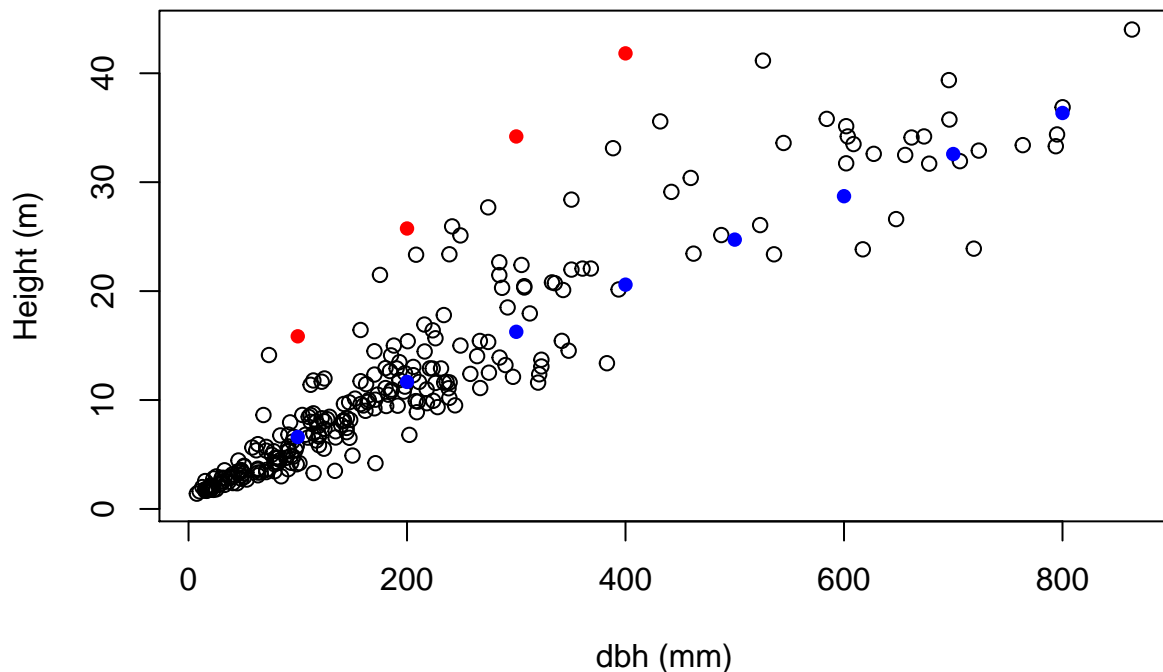
## (Intercept)      dbins_cm
##  3.4825736   0.4171207

plot(h~dbh_mm,
     main="Diameter at breast height (dbh) to Height",
     xlab = "dbh (mm)",
     ylab = "Height (m)")

points(default_mm~dbins_mm, col="red", pch=16) # default D'Brien slope (0.7) and intercept (-0.2)
points(lmfit_mm~dbins_mm, col="blue", pch=16) # BAAD data slope (0.82) and intercept (-0.82)

```

Diameter at breast height (dbh) to Height

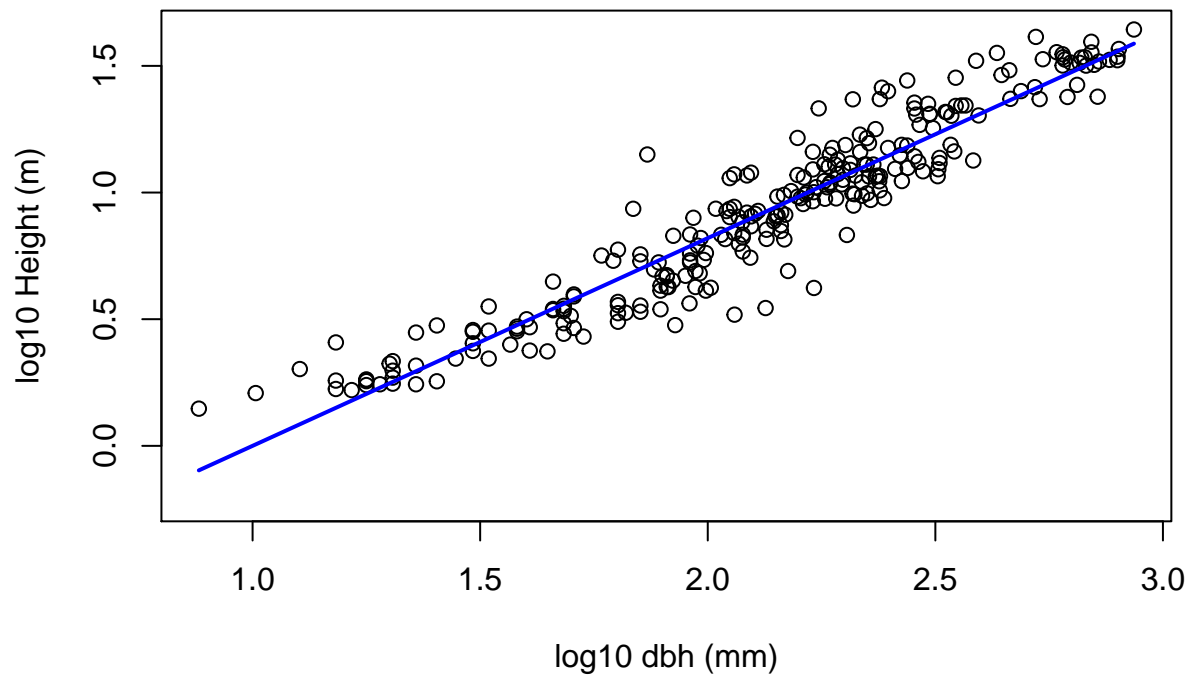


```

plot(log10(h)~log10(dbh_mm),
     main="Diameter at breast height (dbh) to Height",
     xlab = "log10 dbh (mm)",
     ylab = "log10 Height (m)")
curve(-0.82 + (0.82*x), add=T, col="blue", lwd=2) # log regression coefficients from d2hmodlog

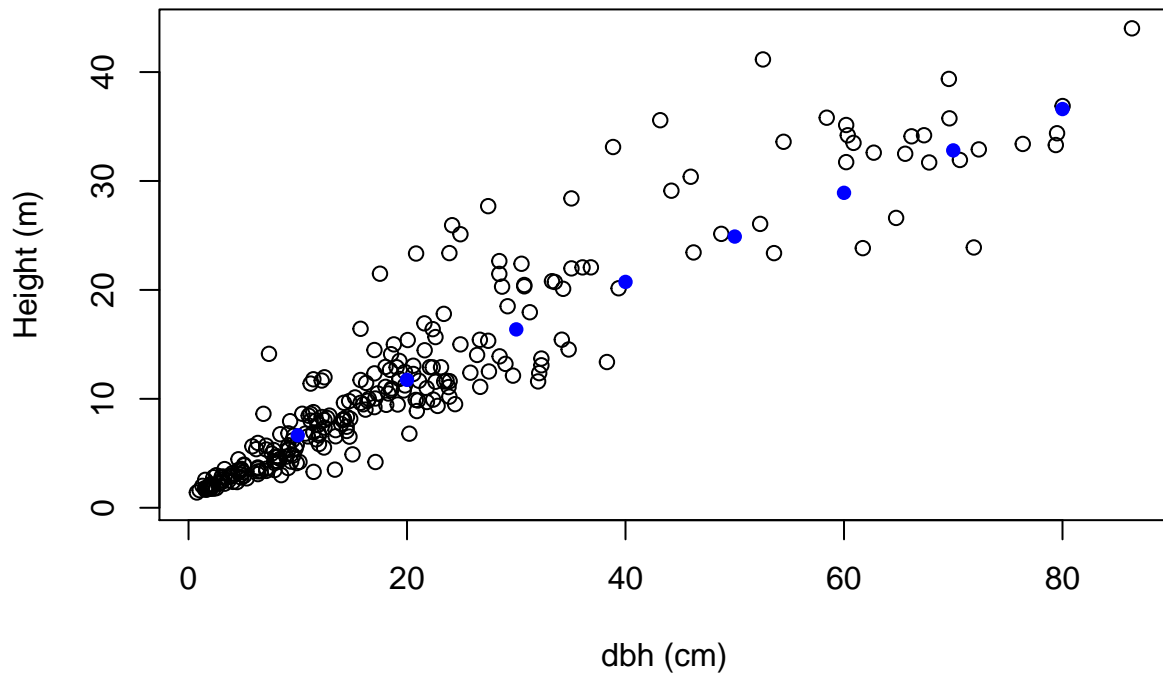
```

Diameter at breast height (dbh) to Height



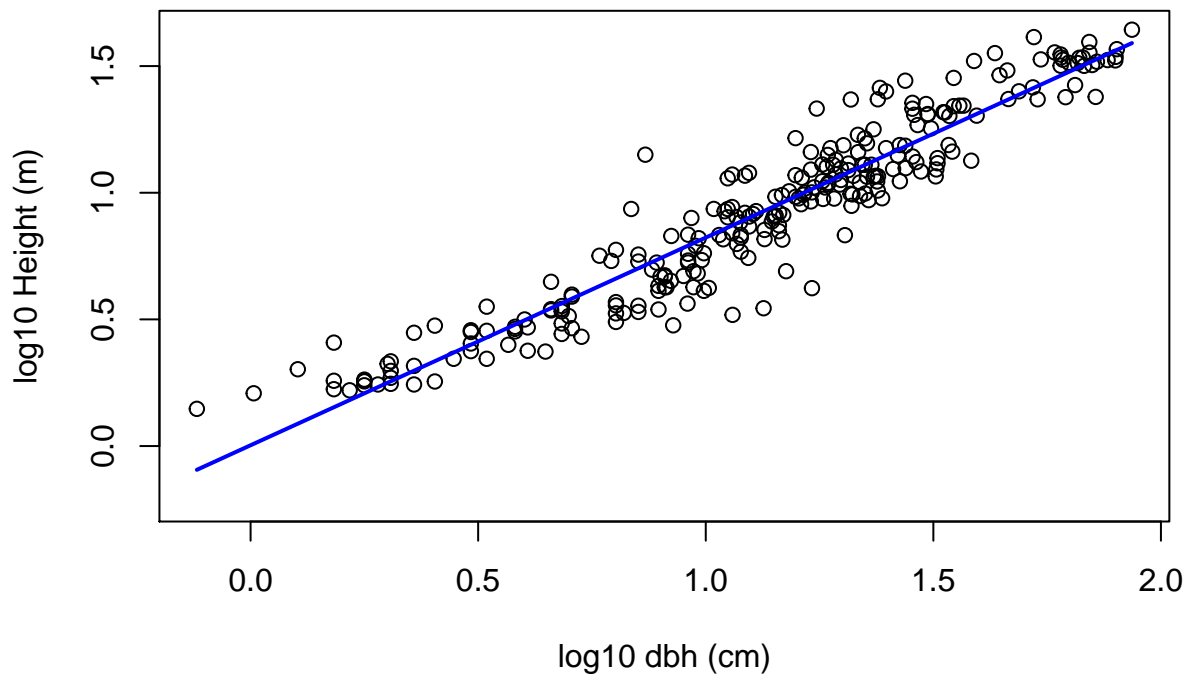
```
plot(h~dbh_cm,  
     main="Diameter at breast height (dbh) to Height",  
     xlab = "dbh (cm)",  
     ylab = "Height (m)")  
  
points(default_mm~dbins_mm, col="red", pch=16) # default O'Brien slope (0.7) and intercept (-0.2)  
points(lmfit_cm~dbins_cm, col="blue", pch=16) # BAAD data slope (0.82) and intercept (-0.82)
```

Diameter at breast height (dbh) to Height



```
plot(log10(h)~log10(dbh_cm),
     main="Diameter at breast height (dbh) to Height",
     xlab = "log10 dbh (cm)",
     ylab = "log10 Height (m)")
curve(0.003 + (0.82*x), add=T, col="blue", lwd=2) # log regression coefficients from d2hmodlog
```

Diameter at breast height (dbh) to Height



Diameter at breast height (dbh) to above ground biomass (AGB) - d2bagw

FATES provides three different options for calculating AGB from dbh:

- Saldarriaga et al. (1998)

$$(C_{agb} = f_{agb} * p_1 * h^{p_2} * d^{p_3} * rho^{p_4})$$

- 2 parameter power function

$$(C_{agb} = p_1 / c2b * d^{p_2})$$

- Chave et al. (2014)

$$(C_{agb} = p_1 / c2b * (rho * d^2 * h)^{p_2})$$

For this project I will test Saldarriaga et al. (1998) as well as the 2 parameter power function.

$$C_{agb} = f_{agb} * p_1 * h^{p_2} * d^{p_3} * rho^{p_4}$$

```
# In this case dbh and agb calculated from Chojnacky serves as the "observed data" to fit the other param

# Define the variables needed for the Saldarriaga function
rho = 0.367
f_agb = 0.6
d.agb <- c(10, 20, 30, 40, 50, 60, 70, 80, 90, 100)
h.agb <- c(2.82, 7.7, 13.9, 21.0, 29.1, 37.9, 37.9, 37.9, 37.9, 57.5)
#use heights from d2h
lmfit.h <- obrien(d.agb, 0.82, 0.003)

# Define the Saldarriaga function
sal <- function(f_agb, p1, h, p2, d, p3, rho, p4){
  agb <- f_agb * p1 * h^p2 * d^p3 * rho^p4
  return(agb)
}

# Define a 2 parameter power function (as a stand-in for Chojnacky if those param values are used)
par2_pwr <- function(p1,d,p2,c2b){
  bagw <- (p1*(d^p2))/c2b
  return(bagw)
}

choj_eq <- function(b0,b1,d){
  biom <- b0 + (b1*log(d))
  return(biom)
}
choj_p1 = -2.6177
choj_p2 = 2.4638
c2b = 2

test_2par <- par2_pwr(0.146, d.agb, 2.464, c2b)
choj <- choj_eq(choj_p1, choj_p2, d.agb)

# Parameters from earlier Jupyter Notebook. Not sure how to determine the best fit for the Sal function
#p1 = 0.131
#p2 = 0.626
#p3 = 2.46
#p4 = 2.18

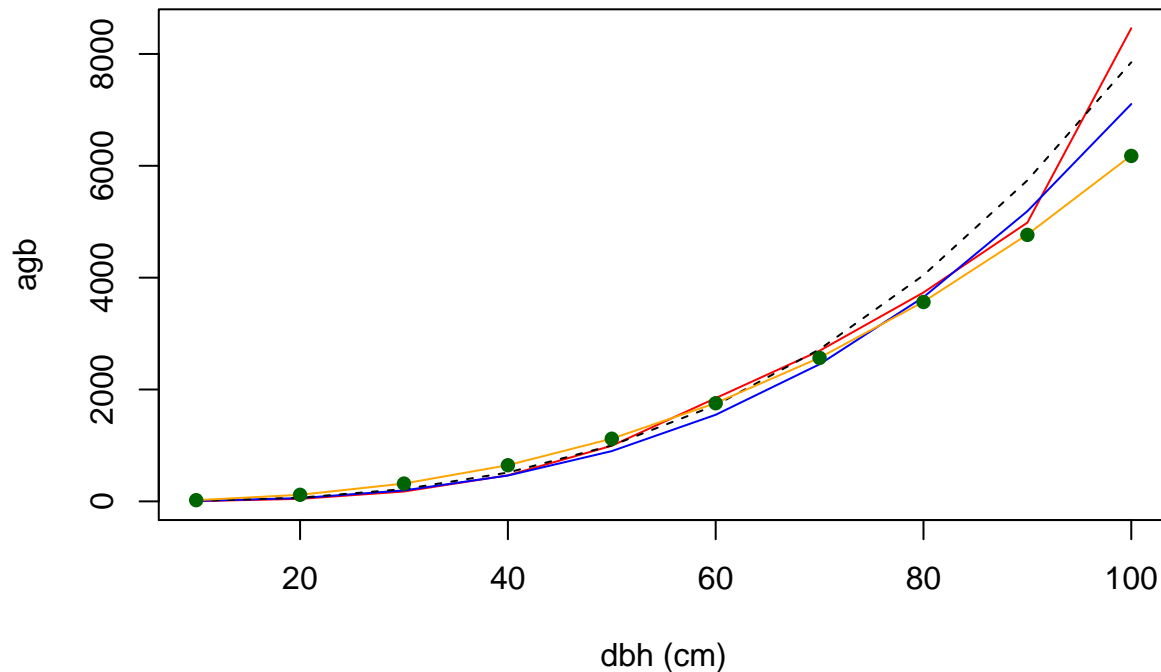
p1 = 0.11
```

```

p2 = 0.65
p3 = 2.45
p4 = 2.15

sal1 <- sal(f_agb, p1, h.agb, p2, d.agb, p3, rho, p4)
sal2 <- sal(f_agb, p1, lmfit.h, p2, d.agb, p3, rho, p4)
sal3 <- sal(f_agb, 0.131, lmfit.h, 0.626, d.agb, 2.46, rho, 2.18)
#plot(choj.h~d.agb, type="l", col="blue")
plot(sal1~d.agb, col="red", type="l", ylab="agb", xlab="dbh (cm)")
points(sal2~d.agb, col="blue", type="l")
points(sal3~d.agb, col="black", lty=2, type="l")
points(test_2par~d.agb, col="orange", type="l")
points(exp(choj)~d.agb, col="darkgreen", pch=16)

```



Here I want to get the r2 or rmse to compare the different allom functions and params to the "observe

```

rmse <- function(y_hat,y){
  return(sqrt(mean((y-y_hat)^2)))
}

```

```

r2 <- function(y_hat,y){
  RSS<-sum(((y_hat)-(y))^2)
  TSS<-sum(((y)-(mean(y)))^2)
  return(1-RSS/TSS)}

```

remember y_hat is the model predicted agb
y observed = test_choj
y_hat(s) = test (sal), test2 (sal), test_2par, py_test (sal)

```

sal1_rmse <- rmse(sal1,exp(choj))
sal1_R2 <- r2(sal1, exp(choj))

```

```

sal2_rmse <- rmse(sal2,exp(choj))
sal2_R2 <- r2(sal2,exp(choj))

sal3_rmse <- rmse(sal3, exp(choj))
sal3_R2 <- r2(sal3, exp(choj))

test_2par_rmse <- rmse(test_2par, exp(choj))
test_2par_R2 <- r2(test_2par, exp(choj))

agb_rmse <- c(sal1_rmse, sal2_rmse, sal3_rmse, test_2par_rmse)
agb_r2 <- c(sal1_R2, sal2_R2, sal3_R2, test_2par_R2)
agb_names <- c("Sal test 1", "Sal test 2", "Sal test 3", "Two Param Power")

agb_mod_fits <- data.frame(agb_rmse, agb_r2, row.names = agb_names)

agb_mod_fits

##              agb_rmse    agb_r2
## Sal test 1      734.098689 0.8675944
## Sal test 2      347.141465 0.9703919
## Sal test 3      636.566290 0.9004400
## Two Param Power    3.794937 0.9999965

```

Diameter to Leaf Biomass (d2blmax)

FATES provides three different options for calculating leaf biomass from dbh:

- Saldarriaga et al. (1998)

$$(blmax = p_1 * d^{p_2} * rho^{p_3})$$

- 2 parameter power function

$$(blmax = p_1 / c2b * d^{p_2})$$

- 2 parameter power function for large trees

$$(blmax = p_1 / c2b * \min(d, dbhmax)^{p_2})$$

Here I will test the Saldarriaga and 2 parameter power functions.

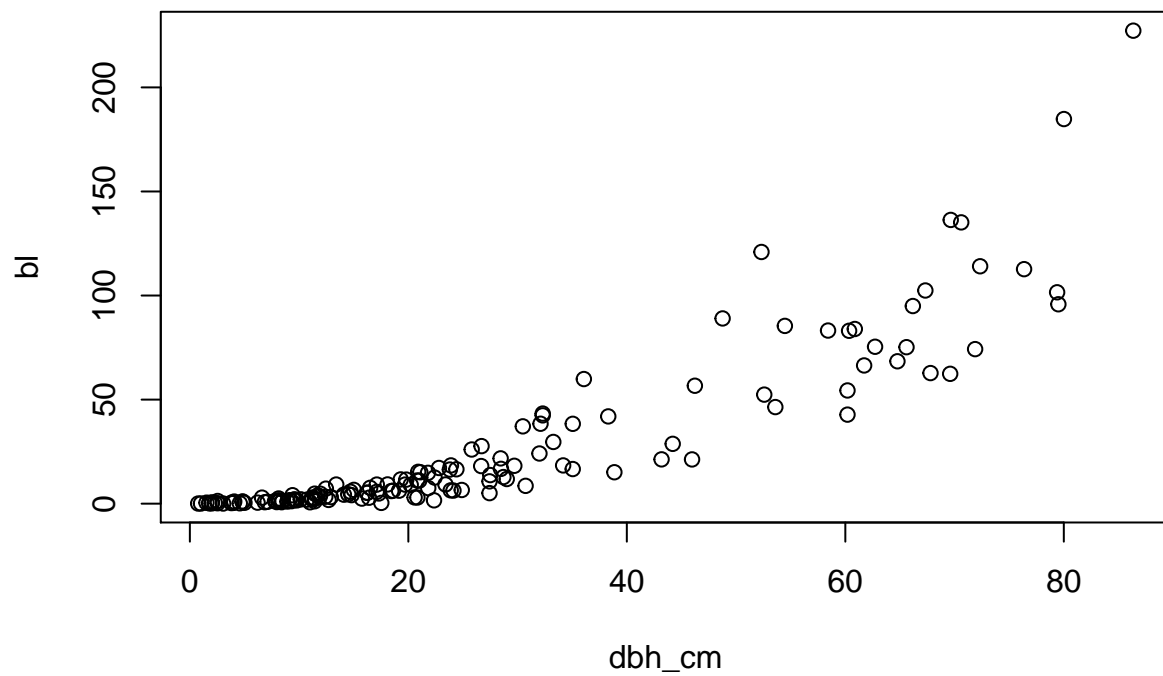
```

bl <- pipo$m.lf

bl_test <- c(0.8, 4.6, 13.2, 30, 36.2, 60, 70.5, 93) # From Polly's sheets
bl_dbh <- c(10, 20, 30, 40, 50, 60, 70, 80) #cm

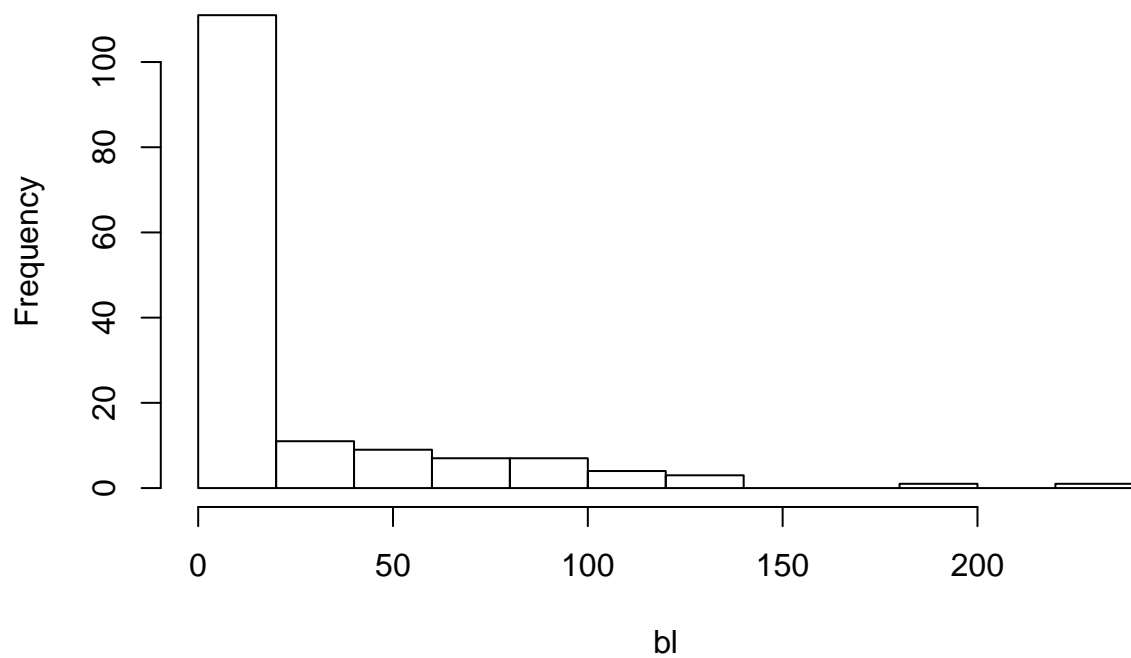
plot(bl~dbh_cm)

```

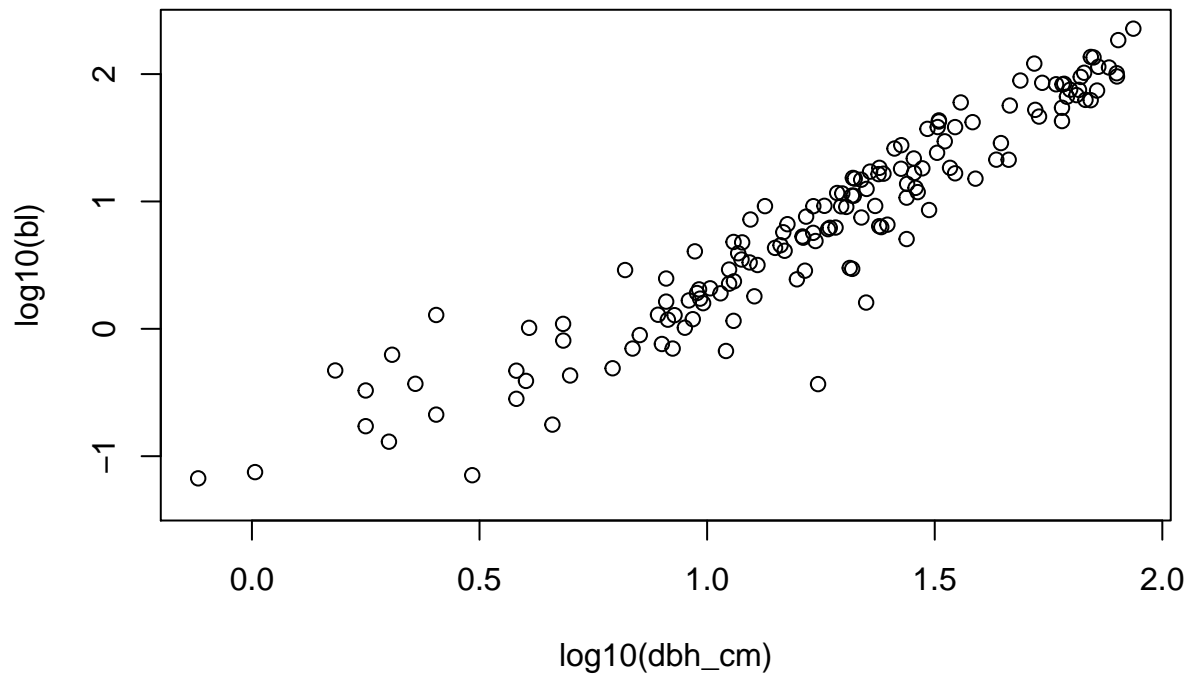


```
hist(bl)
```

Histogram of bl

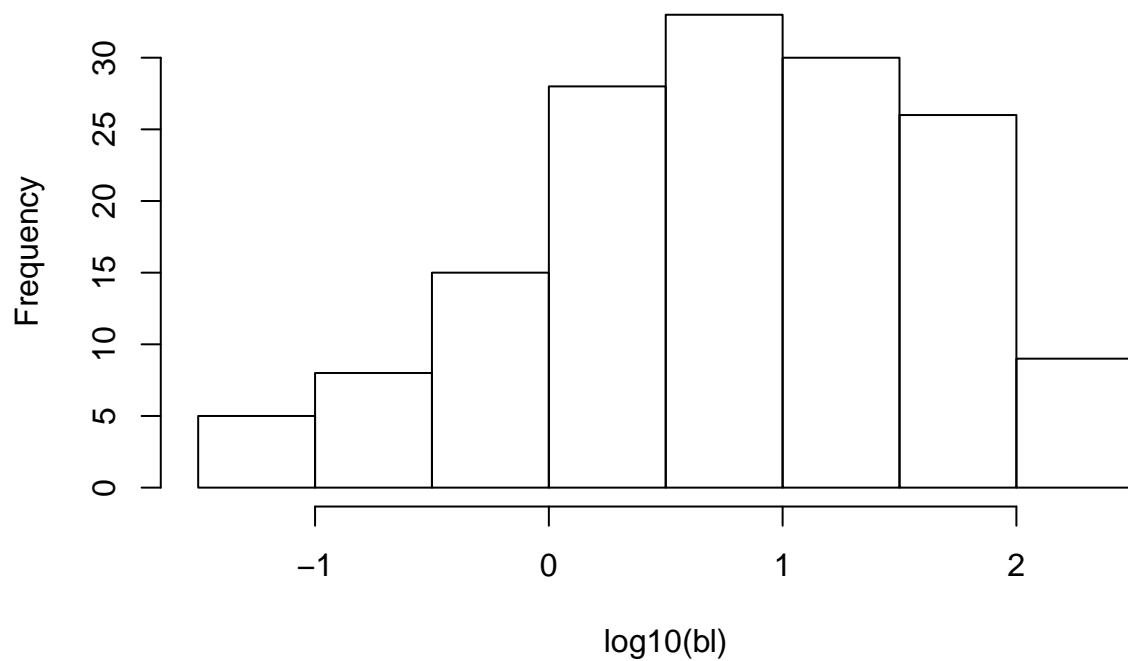


```
plot(log10(bl)~log10(dbh_cm))
```



```
hist(log10(bl))
```

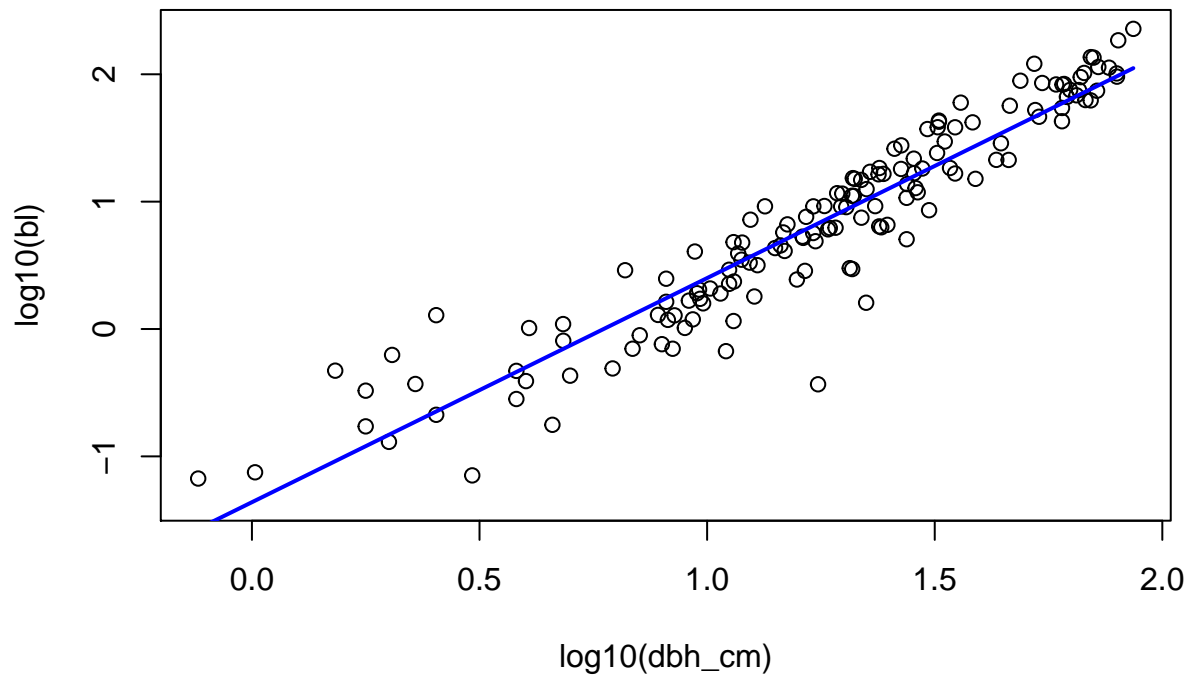
Histogram of $\log_{10}(\text{bl})$



```
d2blmod <- lm(log10(bl) ~ log10(dbh_cm))
coef(d2blmod)
```

```
## (Intercept) log10(dbh_cm)
## -1.363671 1.762904
```

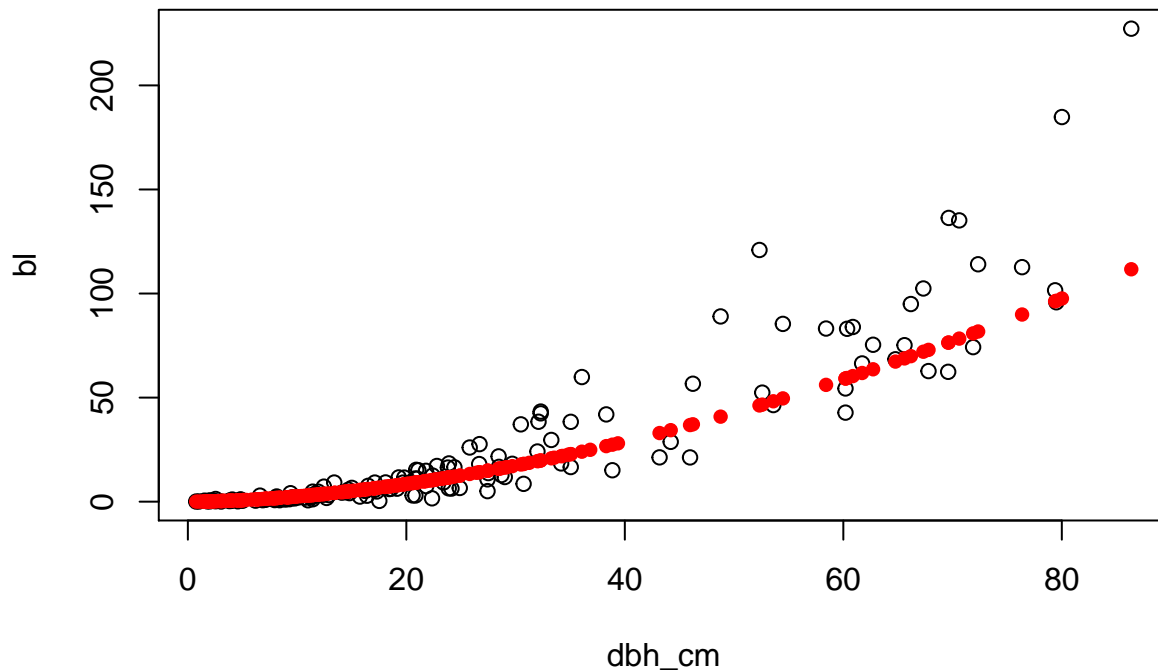
```
plot(log10(bl)~log10(dbh_cm))
curve((-1.36 + 1.76*x),add=T, col="blue", lwd=2)
```



```
bl_lm <- function(dbh, p1, p2){
  blmax <- 10^((log10(dbh))* p1 + p2)
  return(blmax)
}

bl_lm_mod <- bl_lm(sort(dbh_cm), 1.76, -1.36)

plot(bl~dbh_cm)
points(bl_lm_mod~sort(dbh_cm), col="red", pch=16)
```



```
bl_sal <- function(p1,d,p2,rho,p3){
  blmax <- (p1 * (d^p2) * (rho^p3))
  return(blmax)
}

bl_2par <- function(p1,d,p2,c2b){
  blmax <- (p1 * (d^p2))/c2b
  return(blmax)
}

# Test difference params

# Sal1: p1= 0.0169, p2= 2.592, p3= 2.251 (from Jupyter notebook curve_fit)
# Sal2: p1= 0.2, p2=1.55, p3=0.75 (from Polly)
# Sal3: p1= 0.02, p2= 2.011, p3= -0.01665 (from nls but setting p1=0.2)
# Sal4: p1= 0.115, p2= 1.834, p3= 1.33 (from nls below, p1=0.115, and using only mean bl for binned dbh)

bl_sal1_mod <- bl_sal(0.0169, sort(dbh_cm), 2.592, rho, 2.251)
#bl_sal_mod
bl_sal2_mod <- bl_sal(0.2, sort(dbh_cm), 1.55, rho, 0.75)
#bl_sal2_mod
bl_sal3_mod <- bl_sal(0.2, sort(dbh_cm), 2.011, rho, 2.28)
#bl_sal3_mod
bl_sal4_mod <- bl_sal(0.115, bl_dbh, 1.834, rho, 1.33)

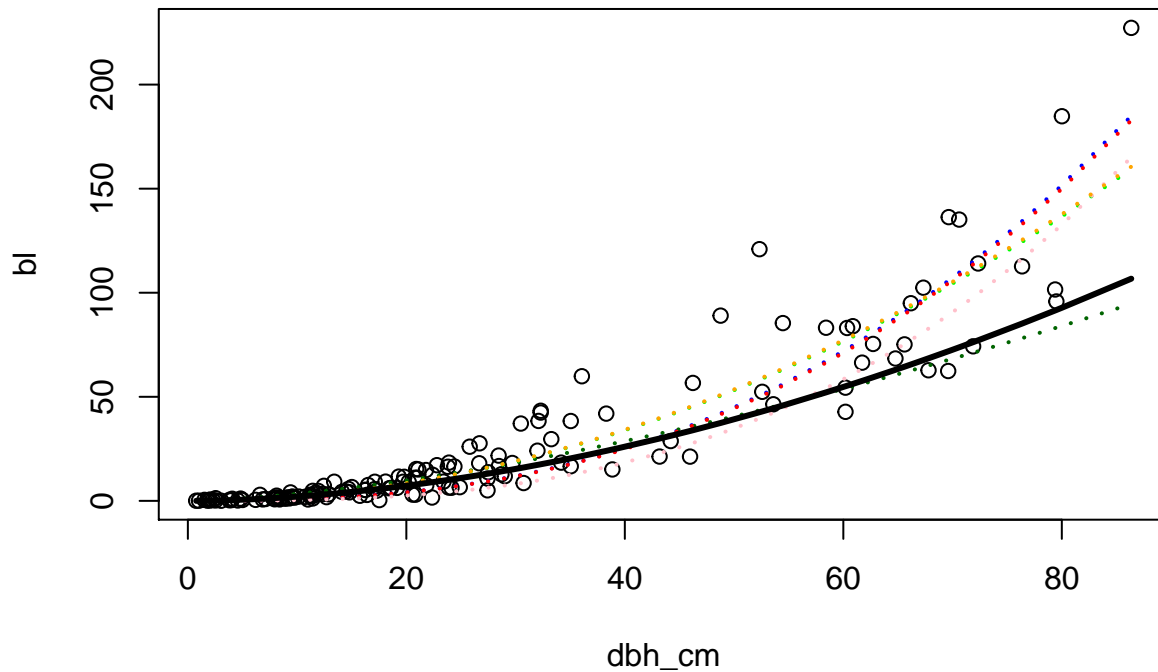
# 2 pwr1: p1= 0.0035 , p2= 2.592 (from jupyter notebook curve_fit)
# 2 pwr2: p1= 0.005, p2= 2
# 2 pwr3: p1= 0.04, p2= 2.01 (from r nls below using full bl and dbh data from BAAD)
# 2 pwr4: p1= 0.06, p2= 1.834 (from nls below using mean BL for binned dbh)

bl_2par_mod <- bl_2par(0.0035, sort(dbh_cm), 2.592, 2)
#bl_2par_mod=
```

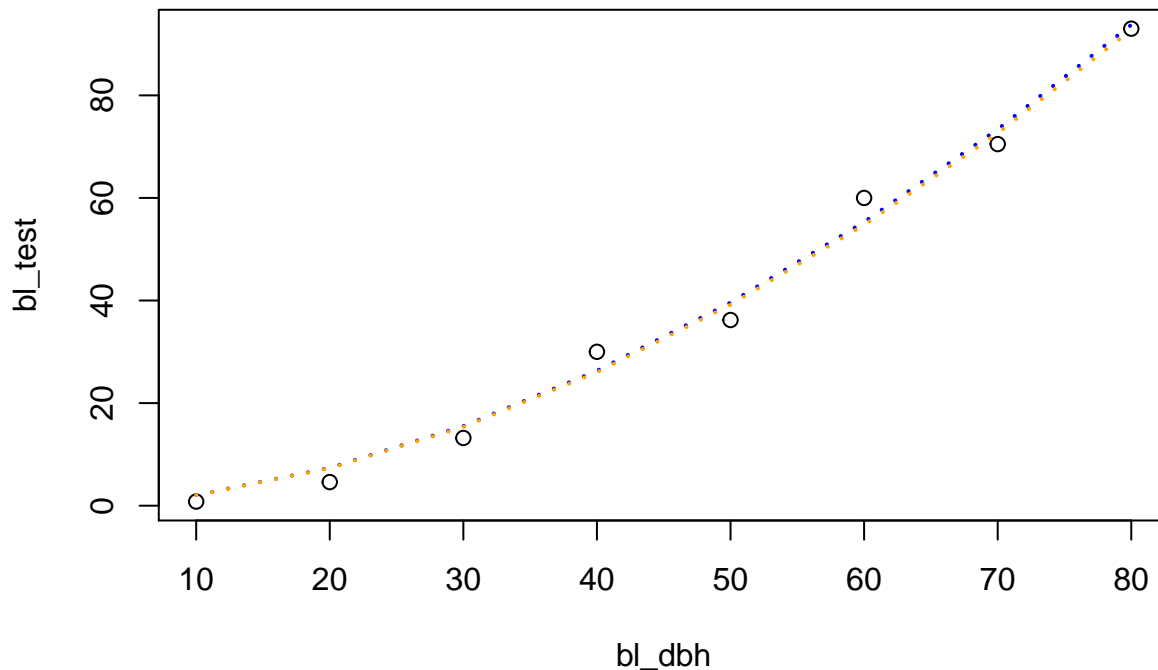
```
bl_2par2_mod <- bl_2par(0.001, sort(dbh_cm), 2.85, 2) # Brute force
#bl_2par2_mod
bl_2par3_mod <- bl_2par(0.041, sort(dbh_cm), 2.011, 2)
#bl_2par3_mod
bl_2par4_mod <- bl_2par(0.06, bl_dbh, 1.834, 2)
```

```
bl_2par5_mod <- bl_2par(0.06, sort(dbh_cm), 1.834, 2)
```

```
plot(bl~dbh_cm)
points(bl_sal1_mod~sort(dbh_cm), type="l", col="blue", lwd=2, lty=3)
points(bl_sal2_mod~sort(dbh_cm), type="l", col="darkgreen", lwd=2, lty=3)
points(bl_sal3_mod~sort(dbh_cm), type="l", col="green", lwd=2, lty=3)
points(bl_2par_mod~sort(dbh_cm), type="l", col="red", lwd=2, lty=3)
points(bl_2par2_mod~sort(dbh_cm), type="l", col="pink", lwd=2, lty=3)
points(bl_2par3_mod~sort(dbh_cm), type="l", col="orange", lwd=2, lty=3)
points(bl_2par5_mod~sort(dbh_cm), type="l", col="black", lwd=3)
```



```
plot(bl_test~bl_dbh)
points(bl_sal4_mod~bl_dbh, type="l", col="blue", lwd=2, lty=3)
points(bl_2par4_mod~bl_dbh, type="l", col="orange", lwd=2, lty=3)
```

```
bl_sal1_rmse <- rmse(bl_sal1_mod, na.omit(bl))
```

```
## Warning in y - y_hat: longer object length is not a multiple of shorter
## object length
```

```
bl_sal1_R2 <- r2(bl_sal1_mod, na.omit(bl))
```

```
## Warning in ((y_hat)) - (y): longer object length is not a multiple of
## shorter object length
```

```
bl_2par_rmse <- rmse(bl_2par_mod, na.omit(bl))
```

```
## Warning in y - y_hat: longer object length is not a multiple of shorter
## object length
```

```
bl_2par_R2 <- r2(bl_2par_mod, na.omit(bl))
```

```
## Warning in ((y_hat)) - (y): longer object length is not a multiple of
## shorter object length
```

```
bl_sal2_rmse <- rmse(bl_sal2_mod, na.omit(bl))
```

```
## Warning in y - y_hat: longer object length is not a multiple of shorter
## object length
```

```
bl_sal2_R2 <- r2(bl_sal2_mod, na.omit(bl))
```

```
## Warning in ((y_hat)) - (y): longer object length is not a multiple of
## shorter object length
```

```
bl_sal3_rmse <- rmse(bl_sal3_mod, na.omit(bl))
```

```
## Warning in y - y_hat: longer object length is not a multiple of shorter
## object length
```

```
bl_sal3_R2 <- r2(bl_sal3_mod, na.omit(bl))
```

```
## Warning in ((y_hat)) - (y): longer object length is not a multiple of
```

```

## shorter object length
bl_sal4_rmse <- rmse(bl_sal4_mod, bl_dbh)
bl_sal4_R2 <- r2(bl_sal4_mod, bl_dbh)

bl_2par2_rmse <- rmse(bl_2par2_mod, na.omit(bl))

## Warning in y - y_hat: longer object length is not a multiple of shorter
## object length
bl_2par2_R2 <- r2(bl_2par2_mod, na.omit(bl))

## Warning in ((y_hat)) - (y): longer object length is not a multiple of
## shorter object length
bl_2par3_rmse <- rmse(bl_2par3_mod, na.omit(bl))

## Warning in y - y_hat: longer object length is not a multiple of shorter
## object length
bl_2par3_R2 <- r2(bl_2par3_mod, na.omit(bl))

## Warning in ((y_hat)) - (y): longer object length is not a multiple of
## shorter object length
bl_2par4_rmse <- rmse(bl_2par4_mod, bl_test)
bl_2par4_R2 <- r2(bl_2par4_mod, bl_test)

bl_2par5_rmse <- rmse(bl_2par5_mod, na.omit(bl))

## Warning in y - y_hat: longer object length is not a multiple of shorter
## object length
bl_2par5_R2 <- r2(bl_2par5_mod, na.omit(bl))

## Warning in ((y_hat)) - (y): longer object length is not a multiple of
## shorter object length
bl_rmse <- c(bl_sal1_rmse, bl_sal2_rmse, bl_sal3_rmse, bl_2par_rmse, bl_2par2_rmse, bl_2par3_rmse, bl_2par5_rmse)
bl_r2 <- c(bl_sal1_R2, bl_sal2_R2, bl_sal3_R2, bl_2par_R2, bl_2par2_R2, bl_2par3_R2, bl_2par5_R2)
bl_names <- c("Sal test 1: BL", "Sal test 2: BL", "Sal test 3:BL", "Two Param Power: BL", "Two Param Power 2: BL", "Two Param Power 3: BL", "Two Param Power 5: BL")

bl_mod_fits <- data.frame(bl_rmse, bl_r2, row.names = bl_names)

bl_mod_fits

##              bl_rmse      bl_r2
## Sal test 1: BL      53.88777 -2.445608
## Sal test 2: BL      46.79779 -1.598580
## Sal test 3:BL      52.31379 -2.247265
## Two Param Power: BL  53.69626 -2.421160
## Two Param Power 2: BL 51.83128 -2.187640
## Two Param Power 3:BL  52.43524 -2.262360
## Two Param Power 5: BL 47.89558 -1.721926

bl2par.fit <- nls(bl_test ~ bl_2par(p1,bl_dbh, p2, 2), start=list(p1=0.0035, p2=2.5))
bl2par.fit

## Nonlinear regression model
##   model: bl_test ~ bl_2par(p1, bl_dbh, p2, 2)

```

```
## data: parent.frame()
## p1 p2
## 0.06039 1.83353
## residual sum-of-squares: 70.04
##
## Number of iterations to convergence: 20
## Achieved convergence tolerance: 6.307e-06

blsalda.fit <- nls(bl_test ~ bl_sal(0.12, bl_dbh, p2, 0.367, p3), start=list(p2=1,p3=0.5))
blsalda.fit

## Nonlinear regression model
## model: bl_test ~ bl_sal(0.12, bl_dbh, p2, 0.367, p3)
## data: parent.frame()
## p2 p3
## 1.834 1.376
## residual sum-of-squares: 70.04
##
## Number of iterations to convergence: 6
## Achieved convergence tolerance: 6.26e-06
```

Diamter to Crown Area

FATES provides one option for calculating crown area from dbh. There are actually 3 modes to choose from, but they all reference the same equation.

- A 2 power crown area function

$$carea = spreadterm * dbh^{crownareatodbhexponent}$$

where

$$spreadterm = (spread * d2ca_{max}) + ((1 - spread) * d2ca_{min})$$

and

$$crownareatodbhexponent = d2blp2 + d2blediff$$

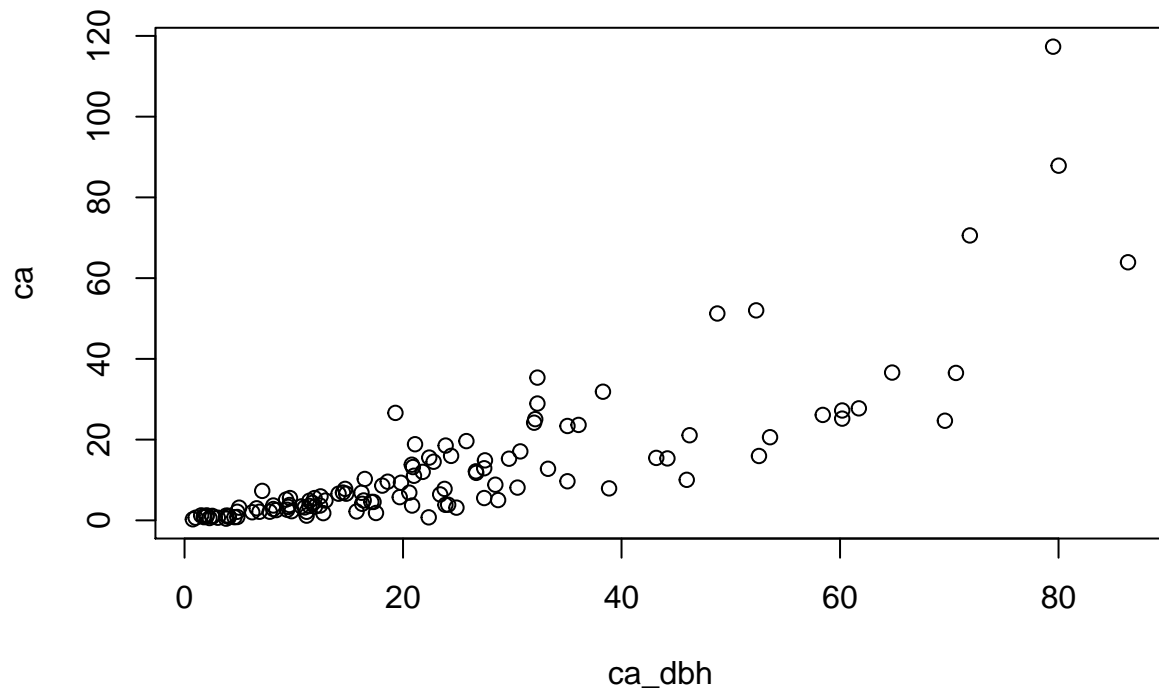
Combined all together:

$$carea = (spread * d2ca_{max}) + ((1 - spread) * d2ca_{min}) * dbh^{(d2blp2 + d2blediff)}$$

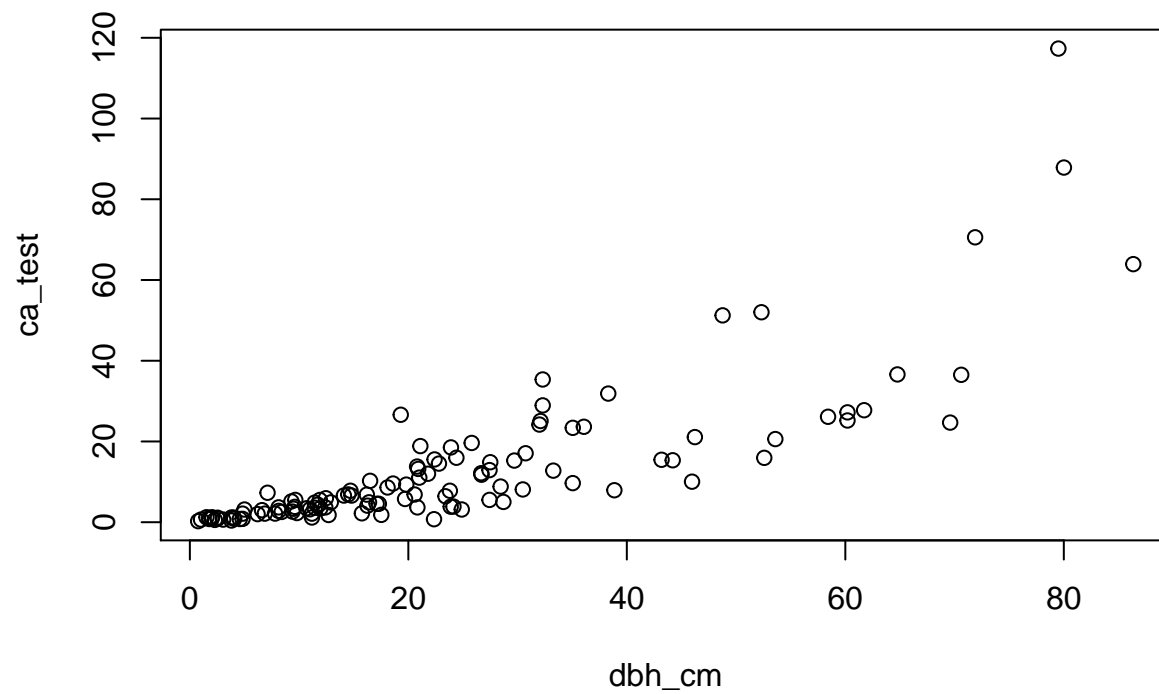
We are given dbh, d2bl p2, d2bl ediff (0), and spread? We need to fit d2camax and min? Or we should know d2ca max and min and only need to fit for spread?

```
ca <- pipo$a.cp[which(pipo$a.cp>0, pipo$dbh>0)]
ca_dbh <- pipo$d.bh[which(pipo$a.cp > 0, pipo$d.bh>0)]*100

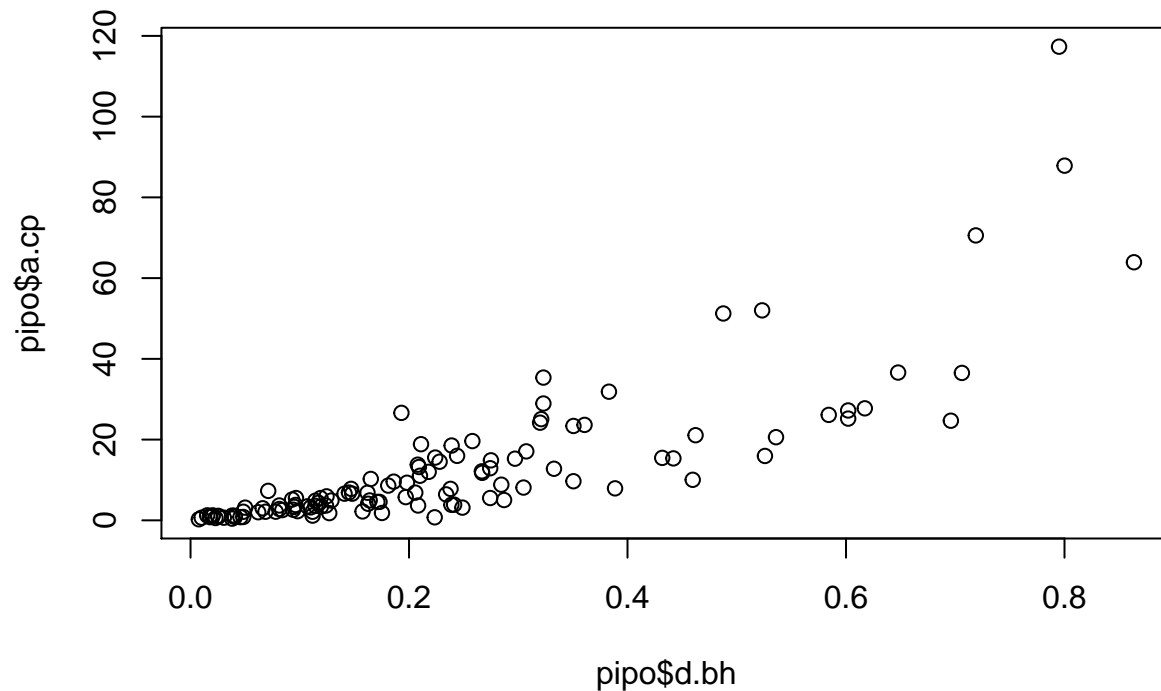
plot(ca~ca_dbh)
```



```
ca_test <- pipo$a.cp  
plot(ca_test~dbh_cm)
```



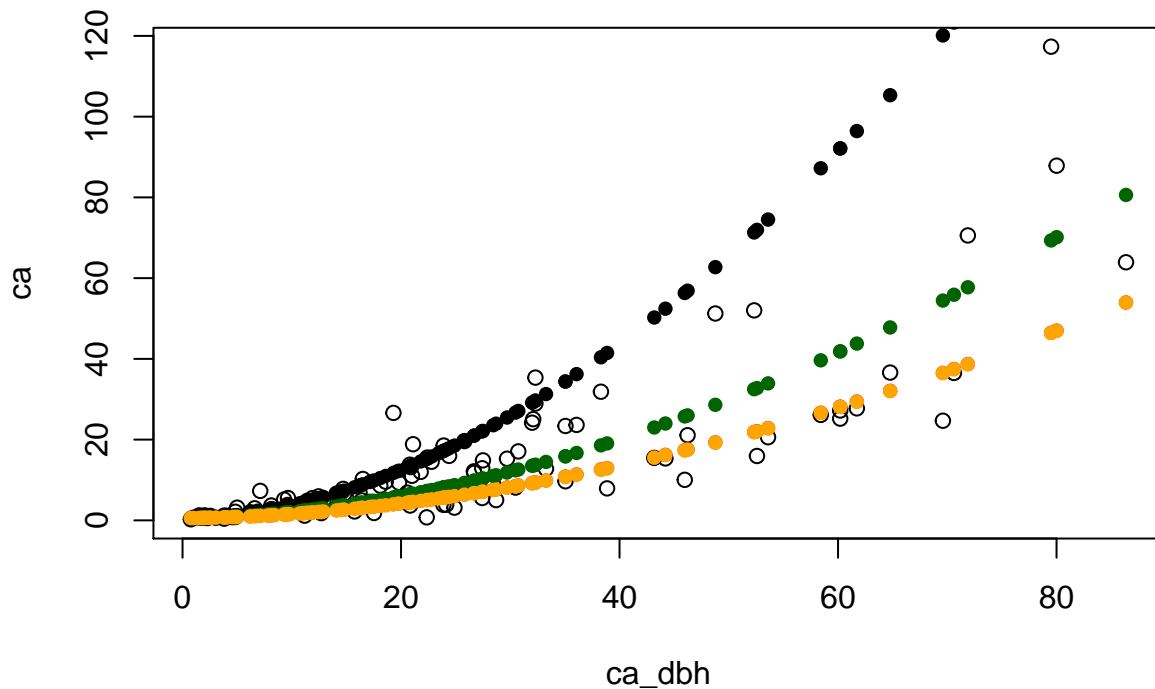
```
plot(pipo$a.cp ~ pipo$d.bh)
```



```
d2carea <- function(sprd, d2camax, d2camin, dbh, d2blp2){
  c.area <- ((sprd * d2camax) + ((1-sprd)*d2camin) * dbh^d2blp2)
  return(c.area)
}

ca_mod1 <- d2carea(0.5, 0.7, 0.1, ca_dbh, 1.834)
ca_mod2 <- d2carea(0.85, 0.7, 0.1, ca_dbh, 1.834)
ca_mod3 <- d2carea(0.85, 0.65, 0.15, ca_dbh, 1.834)
ca_mod4 <- d2carea(0.85, 0.65, 0.1, ca_dbh, 1.834)

plot(ca~ca_dbh)
points(ca_mod1 ~ ca_dbh, pch=16, col="black")
points(ca_mod2 ~ ca_dbh, pch=16, col="red")
points(ca_mod3 ~ ca_dbh, pch=16, col="darkgreen")
points(ca_mod4 ~ ca_dbh, pch=16, col="orange")
```



```
# try to fit parameters
```

```
#d2ca.fit <- nls(ca ~ d2carea(sprd, d2camax, 0.1, sort(dbh_cm), 0.1834), start=list(sprd=0.4, d2camax=0)
#d2ca.fit
```

I want to try something where I step through spread values from 0.1 to 0.9 at intervals of 0.1 and calculate the `d2camax`.

In the mean time I need to figure out the issue with NAs so that I can calculate some R^2 values.

```
#ca_mod1[is.na(ca_mod1)] <- min(ca)
#ca_mod2[is.na(ca_mod2)] <- min(ca)
#ca_mod3[is.na(ca_mod3)] <- min(ca)
#ca_mod4[is.na(ca_mod4)] <- min(ca)
```

```
ca_mod1_R2 <- r2(ca_mod1[7:125], ca[7:125])
ca_mod2_R2 <- r2(ca_mod2[7:125], ca[7:125])
ca_mod3_R2 <- r2(ca_mod3[7:125], ca[7:125])
ca_mod4_R2 <- r2(ca_mod4[7:125], ca[7:125])
```

```
ca_mod1_R2
```

```
## [1] -1.191073
```

```
ca_mod2_R2
```

```
## [1] 0.5956255
```

```
ca_mod3_R2
```

```
## [1] 0.7179147
```

```
ca_mod4_R2
```

```
## [1] 0.5942858
```