

Initialization

```
> library(NCStats)
> library(plotrix) #histStack()
```

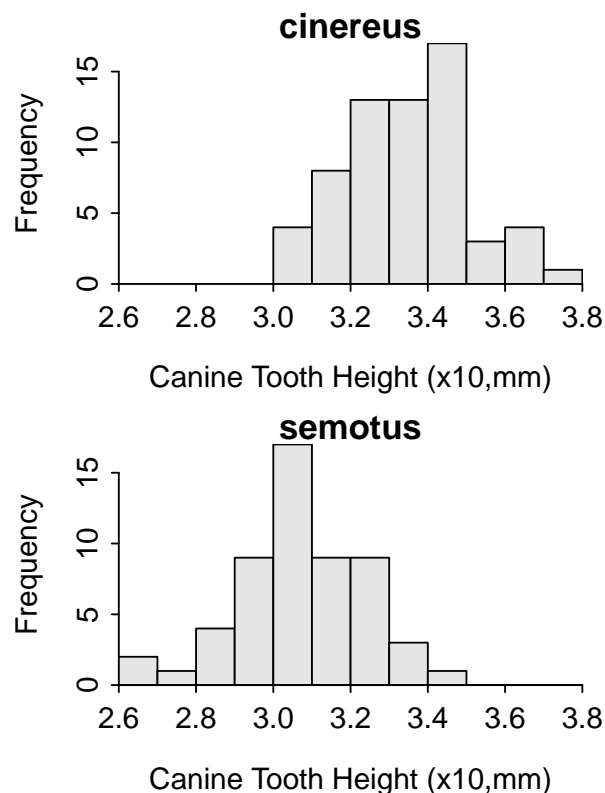
1 Bat Subspecies Example

1.1 Data Preparation and Examination

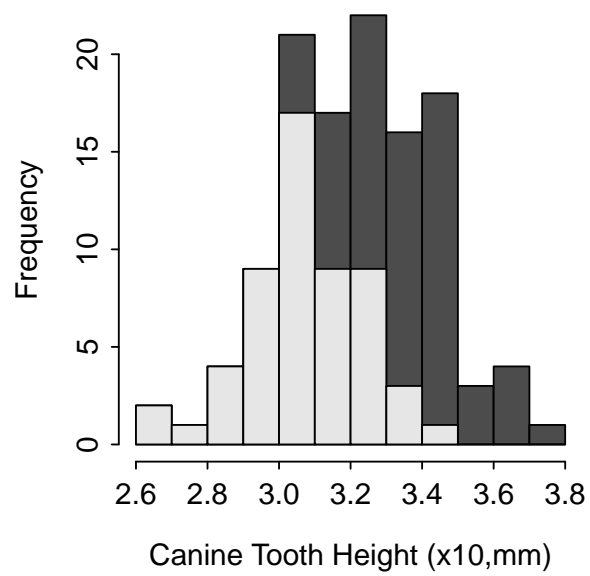
```
> bat <- read.csv("https://raw.githubusercontent.com/droglenc/NCData/master/Batmorph.csv")
> str(bat)

'data.frame': 118 obs. of 7 variables:
 $ subsp      : Factor w/ 2 levels "cinereus","semotus": 2 2 2 2 2 2 2 2 2 2 ...
 $ bodymass   : num  19.5 16.2 17 16.5 14.3 ...
 $ skulllength: num  1.6 1.55 1.56 1.56 1.53 ...
 $ canine     : num  0.326 0.308 0.291 0.287 0.301 0.305 0.277 0.313 0.289 0.293 ...
 $ coronoid   : num  0.303 0.282 0.292 0.303 0.279 0.284 0.286 0.281 0.278 0.28 ...
 $ wingspan   : num  0.358 0.358 0.359 0.353 0.351 0.361 0.351 0.363 0.34 0.365 ...
 $ hab       : Factor w/ 3 levels "A","B","C": 1 1 1 1 1 1 1 1 1 2 ...

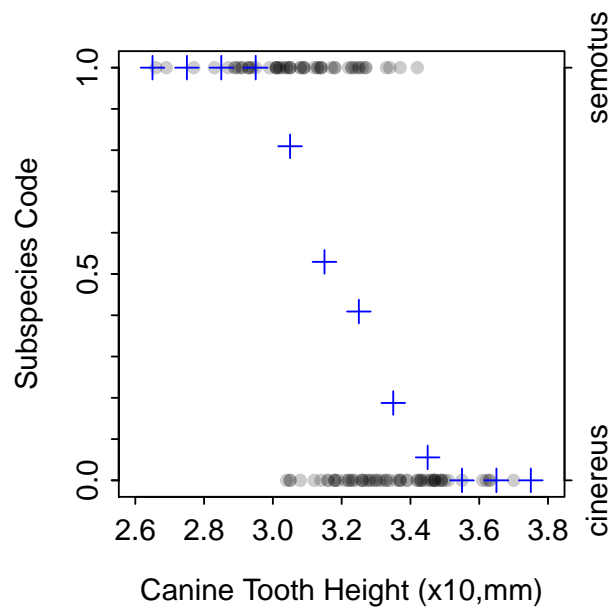
> bat$canine10 <- bat$canine*10
> xlbl <- "Canine Tooth Height (x10,mm)"
> ylbl <- "Subspecies Code"
> hist(canine10~subsp,data=bat,breaks=seq(2.6,3.8,0.1),xlim=c(2.6,3.8),
      xlab=xlbl,nrow=2,ncol=1)
```



```
> histStack(canine10~subsp,data=bat,breaks=seq(2.6,3.8,0.1),xlim=c(2.6,3.8),
  col="gray.colors",xlab=xlbl,right=FALSE)
```

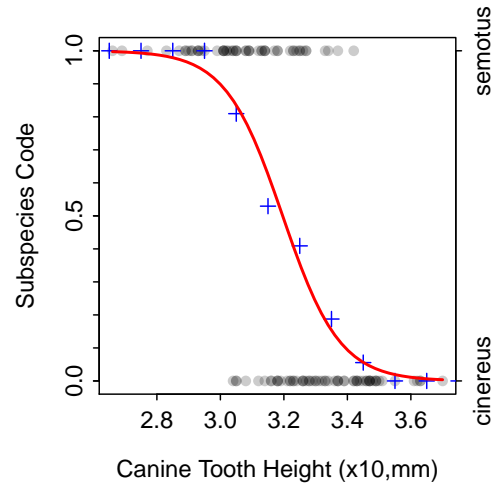


```
> plotBinResp(subsp~canine10,data=bat,breaks=seq(2.6,3.8,0.1),xlim=c(2.6,3.8),
  xlab=xlbl,ylab=ylbl)
```



1.2 Model Fitting and Examination

```
> glm1 <- glm(subsp~canine10,data=bat,family=binomial)
> fitPlot(glm1,breaks=seq(2.6,3.8,0.1),xlab=xlbl,ylab=ylbl,main="")
```



```
> summary(glm1)
```

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	35.516	6.428	5.525	3.29e-08
canine10	-11.112	2.005	-5.543	2.97e-08

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 163.040 on 117 degrees of freedom
Residual deviance: 97.178 on 116 degrees of freedom
AIC: 101.18

Number of Fisher Scoring iterations: 5

```
> confint(glm1)
```

Waiting for profiling to be done...

	2.5 %	97.5 %
(Intercept)	24.21685	49.66132
canine10	-15.52430	-7.58941

1.3 Interpretation of Slope

```
> x1 <- c(3,4) # purposely pick two canine10 values 1 unit apart
> ( p1 <- predict(glm1,data.frame(canine10=x1)) )
      1      2
2.179940 -8.931994
> p1[2]-p1[1]
      2
-11.11193
```

```

> exp(-11.112)                                # back-transformed 'slope' from summary() above
[1] 1.493206e-05
> ( bp1 <- exp(p1) )
      1      2
8.8457728416 0.0001320944
> bp1[2]/bp1[1]
      2
1.493306e-05

```

1.4 Predicting Probabilities

```

> ( p2 <- predict(glm1,data.frame(canine10=c(3,3.4))) )
      1      2
2.179940 -2.264834
> exp(p2)/(1+exp(p2))
      1      2
0.89843357 0.09407761
> predict(glm1,data.frame(canine10=c(3,3.4)),type="response")
      1      2
0.89843357 0.09407761

```

1.5 X for a Certain Proportion

```

> ( cfs <- coef(glm1) )
(Intercept)    canine10
    35.51574    -11.11193
> p <- 0.5    # canine tooth height where subspecies ratio is 50/50
> ( x <- (log(p/(1-p))-cfs[1])/cfs[2] )
(Intercept)
    3.19618
> predict(glm1,data.frame(canine10=x),type="response")    # test the answer
(Intercept)
    0.5
> p <- 0.9    # length where 90% are semotus, 10% are cinereus
> (log(p/(1-p))-cfs[1])/cfs[2]
(Intercept)
    2.998444

```

2 Bootstrapping

```
> bc1 <- bootCase(glm1)      # bootstrapping, be patient!
> head(bc1)

      (Intercept)  canine10
[1,]    44.47487 -14.05668
[2,]    36.54978 -11.43092
[3,]    39.62964 -12.24824
[4,]    44.05369 -13.72758
[5,]    45.15736 -14.29452
[6,]    35.57576 -11.21272

> confint(bc1)

      95% LCI   95% UCI
(Intercept) 26.61595 52.169407
canine10    -16.38357 -8.331562

> hist(bc1,breaks=15)
Error in int_abline(a = a, b = b, h = h, v = v, untf = untf, ...): plot.new has not been called yet
> predProb <- function(x,alpha,beta1) exp(alpha+beta1*x)/(1+exp(alpha+beta1*x))
> predProb(3,coef(glm1)[1],coef(glm1)[2])
(Intercept)
 0.8984336

> p3 <- predProb(3,bc1[,1],bc1[,2])
> head(p3)
[1] 0.9092764 0.9052549 0.9470960 0.9463911 0.9066831 0.8740872

> quantile(p3,c(0.025,0.975))

      2.5%      97.5%
0.8166296 0.9622341

> predX <- function(p,alpha,beta1) (log(p/(1-p))-alpha)/beta1
> x50 <- predX(0.5,bc1[,1],bc1[,2])
> head(x50)
[1] 3.163967 3.197449 3.235538 3.209137 3.159067 3.172803

> quantile(x50,c(0.025,0.975))

      2.5%      97.5%
3.150743 3.240848
```

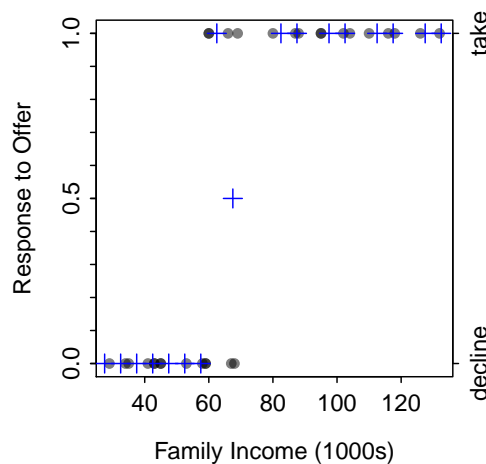
3 Solar Panel Offer Data

```
> sp <- read.csv("https://raw.githubusercontent.com/droglenc/NCDData/master/SolarOffer.csv")
> str(sp)

'data.frame': 30 obs. of 5 variables:
 $ income   : int  80 60 35 45 29 43 34 104 102 59 ...
 $ age      : int  30 34 25 27 23 28 24 43 46 36 ...
 $ takeover: Factor w/ 2 levels "decline","take": 2 2 1 1 1 1 1 2 2 1 ...
 $ mortgage : int  2000 2100 1500 1800 1900 1600 1500 2400 2700 2600 ...
 $ famsize  : int   4 3 2 4 2 3 1 5 3 2 ...

> xlbl <- "Family Income (1000s)"
> ylbl <- "Response to Offer"
```

```
> plotBinResp(takeoffer~income,data=sp,xlab=xlbl,ylab=ylbl,breaks=seq(25,135,5))
```



```
> glm2 <- glm(takeoffer~income,data=sp,family=binomial)
> summary(glm2)
```

Coefficients:

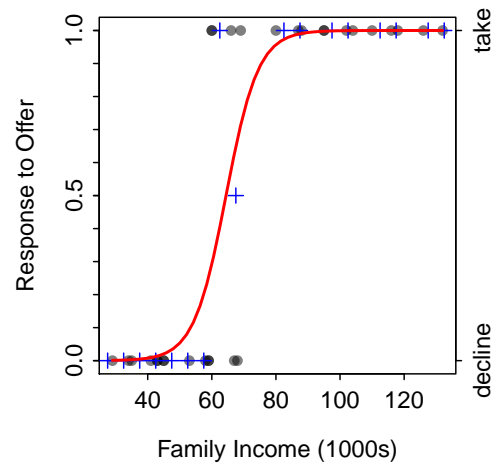
	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	-12.84503	6.16398	-2.084	0.0372
income	0.19934	0.09774	2.039	0.0414

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 41.455 on 29 degrees of freedom
Residual deviance: 13.035 on 28 degrees of freedom
AIC: 17.035

Number of Fisher Scoring iterations: 8

```
> fitPlot(glm2,xlab=xlbl,ylab=ylbl,breaks=seq(25,135,5),main="")
```



```
> p <- 0.25
> (log(p/(1-p))-coef(glm2)[1])/coef(glm2)[2]
(Intercept)
  58.92646
> predict(glm2,data.frame(income=80),type="response")
      1
0.9569831
```