## Specifying prey fat content distributions for Squid diet analysis

## March 20, 2014

```
Silverside <- 2.27
Silverside.sd <- 1.02

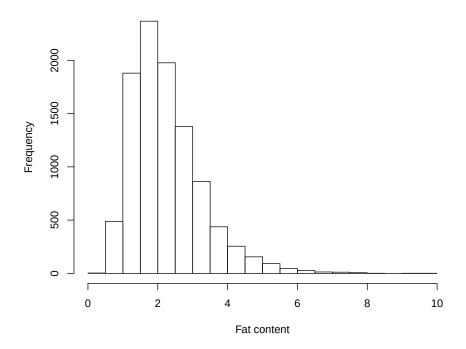
Sailfin.Molly <- 3.97
Sailfin.Molly.sd <- 1.25

Sheepshead.Minnow <- 3.59
Sheepshead.Minnow.sd <- 1.45

fish.means <- c(Silverside, Sailfin.Molly, Sheepshead.Minnow)

fish.vars <- c(Silverside.sd, Sailfin.Molly.sd, Sheepshead.Minnow.sd)^2
```

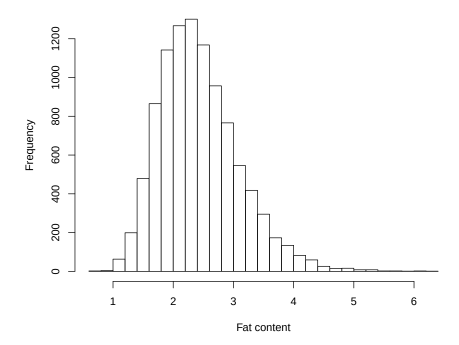
Assuming a log normal model for fat content - this is done internally in fastin-R, so the next bit is for illustrative purposes only...



```
shrimp.mean <- 2.43
shrimp.var <- 0.66^2

shrimp.ln.var = log(shrimp.var + shrimp.mean^2) - 2 * log(shrimp.mean)
shrimp.ln.mean = log(shrimp.mean) - shrimp.ln.var/2

hist(rlnorm(10000, meanlog = shrimp.ln.mean, sdlog = sqrt(shrimp.ln.var)), 30,
    xlab = "Fat content", main = "")</pre>
```



```
fat.cont <- rbind(cbind(fish.ln.means, fish.ln.vars), c(shrimp.ln.mean, shrimp.ln.var))
colnames(fat.cont) <- NULL

# add prey names
prey.ix <- t(read.csv("Prey_FA.csv", header = F, stringsAsFactors = F, row.names = 1))[,
    1]
mullets <- which(prey.ix == "Striped Mullet")
prey.ix <- prey.ix[-mullets]

rownames(fat.cont) <- unique(prey.ix)

write.table(fat.cont, file = "fat.cont.csv", col.names = F, sep = ",")</pre>
```