## Powerlifting Strength Comparison

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#### Data Prep for males >=20y

```
data <- read.csv("openpowerlifting.csv", header=TRUE)</pre>
d2 <- data[(data$Sex=="M")&(data$Equipment=="Raw")</pre>
            &(data$Event=="SBD")&(data$Age>=20),]
rm("data")
d2 <- d2[!is.na(d2$Best3BenchKg),]</pre>
d2 <- d2[!is.na(d2$Best3SquatKg),]</pre>
d2 <- d2[!is.na(d2$Best3DeadliftKg),]</pre>
d2 <- d2[!is.na(d2$TotalKg),]</pre>
d2 <- d2[!is.na(d2$BodyweightKg),]</pre>
d2 <- d2[(d2$Best3BenchKg>0)&(d2$Best3DeadliftKg>0)
          &(d2$Best3SquatKg>0)&(d2$TotalKg>0),]
d2$logBW <- log(d2$BodyweightKg)</pre>
d2$logBench <- log(d2$Best3BenchKg)</pre>
d2$logSquat <- log(d2$Best3SquatKg)</pre>
d2$logTotal <- log(d2$TotalKg)</pre>
d2$logDead <- log(d2$Best3DeadliftKg)</pre>
nrow(d2)
## [1] 111028
d2 <- d2[!duplicated(d2[,c('Name')]),]</pre>
nrow(d2)
## [1] 47003
d2$Rel_Total <- d2$TotalKg / d2$BodyweightKg</pre>
indices = sort(sample(nrow(d2), nrow(d2)*.8))
d2_train <- d2[indices,]</pre>
d2_test <- d2[-indices,]</pre>
```

```
library(MASS)
library(entropy)
```

#### Coloured Density Curves

If the density curves moves as the colour changes, that means there are different distributions of the score as the bodyweight changes. In other words, it isn't a fair score.

#### Wandering Schematic Plot

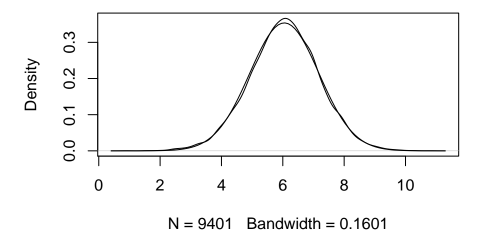
whuber, Measures of residuals heteroscedasticity, URL (version: 2021-06-09): https://stats.stackexchange. com/q/33033

### Relative Bodyweight

```
Relative_model <- quap(
  alist(
    Rel_Total ~ dnorm( mu , sigma ) ,</pre>
```

```
mu <- dnorm( 1.5 , 1 ) ,</pre>
    sigma ~ dexp( 0.1 )
  ) , data=d2_train )
precis(Relative_model)
##
              mean
                             sd
                                     5.5%
                                             94.5%
## mu
         6.049543 0.005815522 6.040249 6.058837
## sigma 1.127721 0.004112346 1.121148 1.134293
post <- extract.samples( Relative_model )</pre>
map_mu <- mean(post$mu)</pre>
map_sigma <- mean(post$sigma)</pre>
plot(density(d2_test$Rel_Total))
curve(dnorm(x,map_mu,map_sigma),add=TRUE)
```

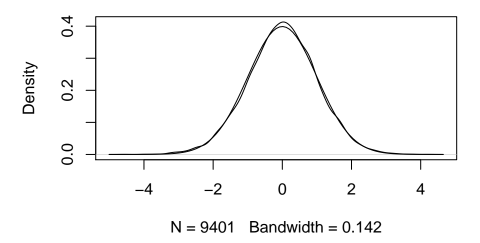
### density.default(x = d2\_test\$Rel\_Total)



#### Normalize to Z Score

```
d2_test$Rel_Total <- ( d2_test$Rel_Total - map_mu) / map_sigma
plot(density(d2_test$Rel_Total))
curve(dnorm(x,0,1),add=TRUE)</pre>
```

## density.default(x = d2\_test\$Rel\_Total)



### Log Log / Allometric model

$$\ln(y) = \alpha + \beta \ln(x)$$

$$y = e^{\alpha + \beta \ln(x)}$$

$$y = e^{\alpha} e^{\beta \ln(x)}$$

$$y = e^{\alpha} x^{\beta}$$

Hence, the log transform of both variables is known as power law or allometric scaling

```
xbar <- mean(d2_train$logTotal)
loglogTotal_model <- quap(
   alist(
    logTotal ~ dnorm( mu , sigma ) ,
    mu <- a + b*( logBW - xbar ) ,
    a <- dnorm( 7 , 0.5) ,
    b <- dexp( 2 ) ,
    sigma ~ dexp( 3 )

) , data=d2_train )
precis(loglogTotal_model)</pre>
```

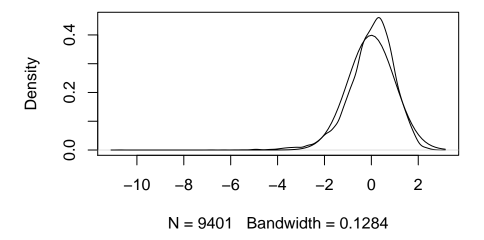
```
## a mean sd 5.5% 94.5%
## a 7.2657587 0.0083006921 7.2524926 7.2790249
## b 0.5531140 0.0046313499 0.5457122 0.5605157
## sigma 0.1802518 0.0006572385 0.1792014 0.1813022
```

```
xbar <- mean(d2_train$logTotal)
post <- extract.samples( loglogTotal_model )
map_a <- mean(post$a)
map_b <- mean(post$b)
map_sigma <- mean(post$sigma)
map_means <- map_a + (map_b * (d2_test$logBW - xbar) )</pre>
```

#### Normalize to Z Score

```
d2_test$LogLogScore <- ( d2_test$logTotal - map_means ) / map_sigma
plot(density(d2_test$LogLogScore))
curve(dnorm(x,0,1),add=TRUE)</pre>
```

### density.default(x = d2\_test\$LogLogScore)



# Varying Variance Model

```
xbar <- mean(d2_train$BodyweightKg)

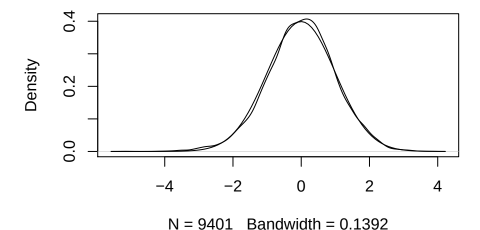
varying_variance <- quap(
   alist(
    TotalKg ~ dnorm( mu , sigma ) ,
    mu <- a + b*( BodyweightKg - xbar ) ,
    a ~ dnorm( 500 , 40 ) ,
   b ~ dnorm( 3, 2 ) ,
   sigma <- d + c*(BodyweightKg - xbar) ,
   d ~ dnorm( 80, 5 ) ,
   c ~ dexp( 2 )</pre>
```

```
) , data=d2_train )
precis(varying_variance)
##
             mean
                                      5.5%
                                                  94.5%
                           sd
## a 546.0363968 0.47588089 545.2758472 546.7969464
       3.3654696 0.02769424
                                3.3212089
                                             3.4097304
## d 91.3831835 0.33474153 90.8482019 91.9181651
      0.6526772 0.01732313
                                0.6249915
                                             0.6803629
xbar <- mean(d2_train$BodyweightKg)</pre>
post <- extract.samples( varying_variance )</pre>
map_a <- mean(post$a)</pre>
map_b <- mean(post$b)</pre>
map_c <- mean(post$c)</pre>
map_d <- mean(post$d)</pre>
map_sigmas <- map_d + (map_c * ( d2_test$BodyweightKg - xbar) )</pre>
map_means <- map_a + (map_b * ( d2_test$BodyweightKg - xbar) )</pre>
```

#### Normalize to Z Score

```
d2_test$VVScore <- ( d2_test$TotalKg - map_means ) / (map_sigmas)
plot(density(d2_test$VVScore))
curve(dnorm(x,0,1),add=TRUE)</pre>
```

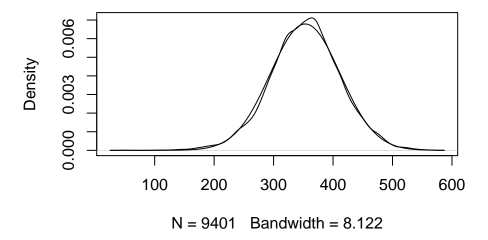
# density.default(x = d2\_test\$VVScore)



#### Wilks

```
Wilks_model <- quap(</pre>
  alist(
    Wilks ~ dnorm( mu , sigma ) ,
    mu <- dnorm( 300 , 50 ) ,
    sigma \sim dexp(0.02)
  ) , data=d2_train )
precis(Wilks_model)
##
               mean
                            sd
                                    5.5%
                                              94.5%
## mu
         353.16017 0.3028881 352.67610 353.64425
## sigma 58.73485 0.2141735 58.39256 59.07714
post <- extract.samples( Wilks_model )</pre>
map_mu <- mean(post$mu)</pre>
map_sigma <- mean(post$sigma)</pre>
plot(density(d2_test$Wilks))
curve(dnorm(x,map_mu,map_sigma),add=TRUE)
```

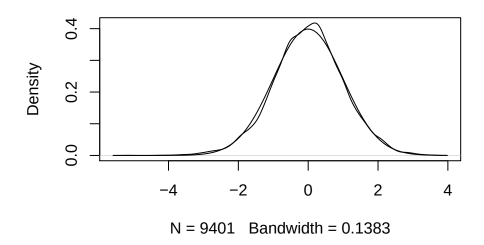
### density.default(x = d2\_test\$Wilks)



#### Normalize to Z Score

```
d2_test$Wilks_Z <- ( d2_test$Wilks - map_mu) / map_sigma
plot(density(d2_test$Wilks_Z))
curve(dnorm(x,0,1),add=TRUE)</pre>
```

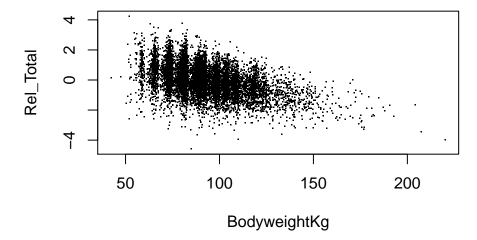
# density.default(x = d2\_test\$Wilks\_Z)

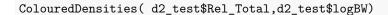


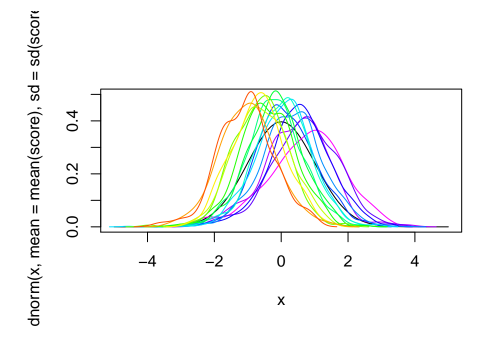
## Plots Comparing Heteroskedasticity in Scores

### Relative Bodyweight

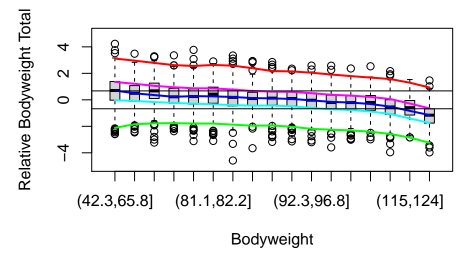
```
#par(mfrow=c(3,1))
plot(Rel_Total~BodyweightKg,data=d2_test,pch=".",cex=0.001)
```



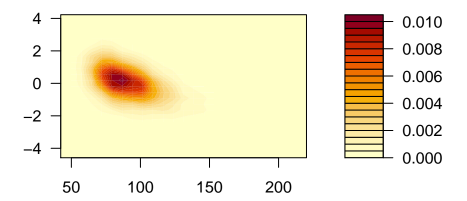




## Relative Bodyweight Z Score vs Bodyweight

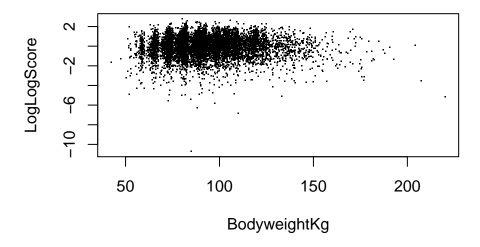


filled.contour(kde2d(d2\_test\$BodyweightKg,d2\_test\$Rel\_Total))

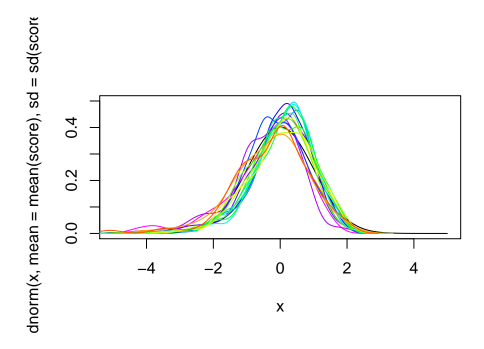


### Log Log

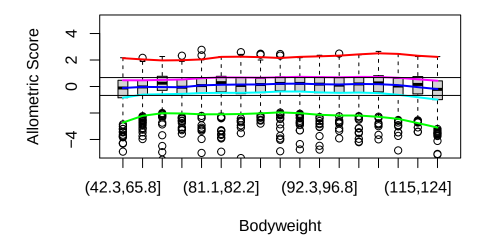
```
#par(mfrow=c(2,2))
plot(LogLogScore~BodyweightKg,data=d2_test,pch=".",cex=0.001)
```



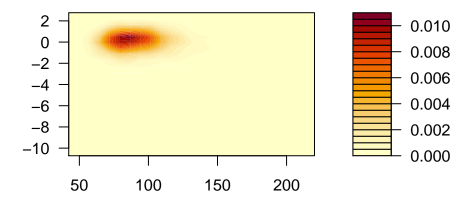
ColouredDensities( d2\_test\$LogLogScore,d2\_test\$logBW)



## Log Log Z Score vs Bodyweight

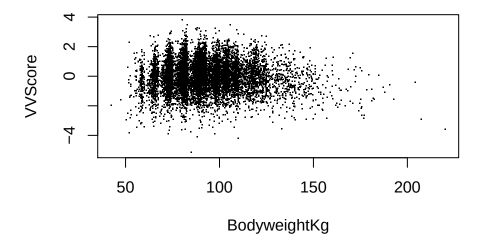


filled.contour(kde2d(d2\_test\$BodyweightKg,d2\_test\$LogLogScore))

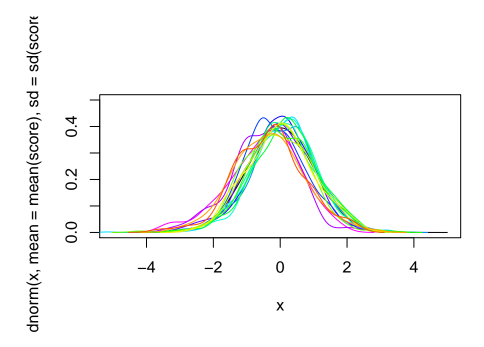


### Varying Variance

```
#par(mfrow=c(2,2))
plot(VVScore~BodyweightKg,data=d2_test,pch=".",cex=0.001)
```



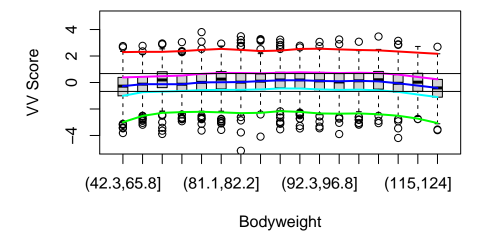
ColouredDensities( d2\_test\$VVScore,d2\_test\$logBW)



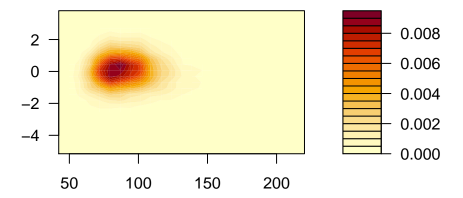
WanderingSchematic(17,d2\_test\$VVScore, d2\_test\$BodyweightKg ,

"Varying Variance Z Score vs Bodyweight", "VV Score")

## **Varying Variance Z Score vs Bodyweight**

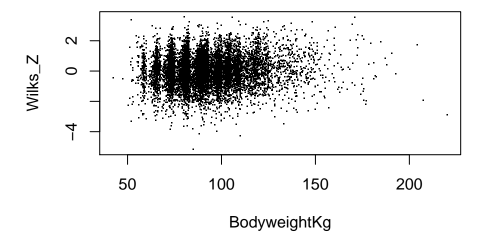


filled.contour(kde2d(d2\_test\$BodyweightKg,d2\_test\$VVScore))

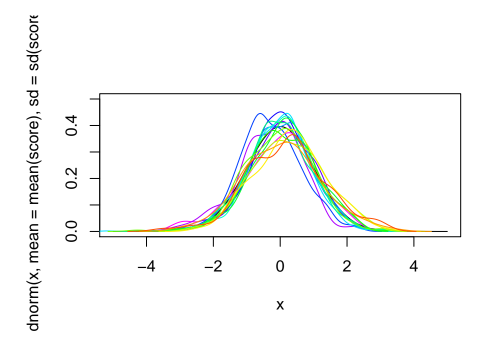


### Wilks

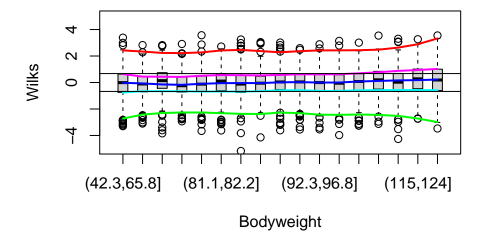
```
#par(mfrow=c(2,2))
plot(Wilks_Z~BodyweightKg,data=d2_test,pch=".",cex=0.001)
```



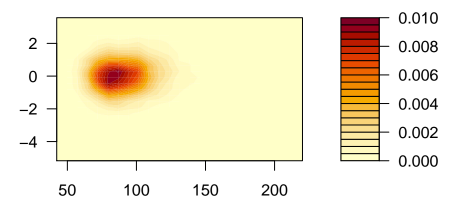
ColouredDensities( d2\_test\$Wilks\_Z,d2\_test\$logBW)



## Wilks Z Score vs Bodyweight



filled.contour(kde2d(d2\_test\$BodyweightKg,d2\_test\$Wilks\_Z))



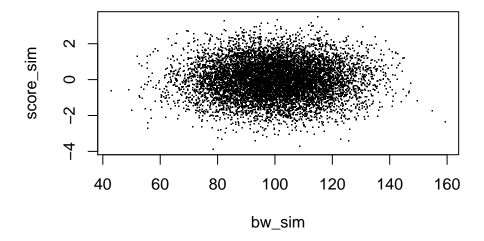
### Fair Score

Question: what does a truly fair score look like?

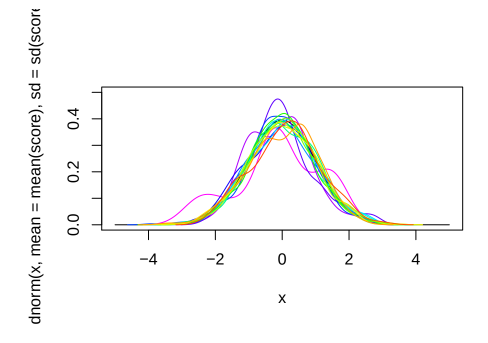
Simulation:

```
n_sim <- nrow(d2_test)
bw_sim <- rnorm(n_sim,100,15)
score_sim <- rnorm(n_sim,0,1) #standard normal independent of bodyweight</pre>
```

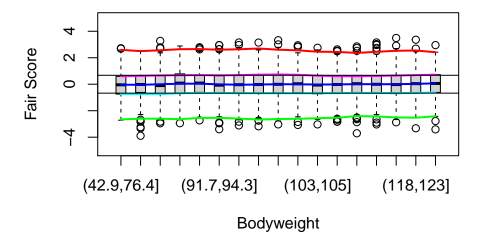
```
#par(mfrow=c(2,2))
plot(score_sim~bw_sim,pch=".",cex=0.001)
```



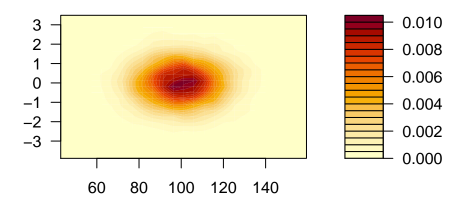
ColouredDensities( score\_sim,log(bw\_sim))



## Fair Score vs Bodyweight



filled.contour(kde2d(bw\_sim,score\_sim))



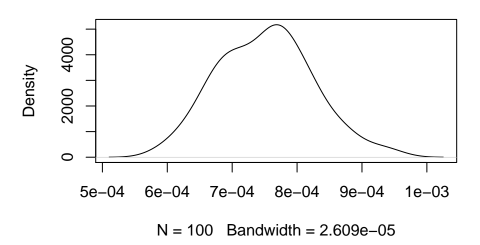
The most telling graph is probably the Wandering Schematic which is much better than any of the real scores used.

Under the null hypothesis that score is independent of bodyweight, what is the distribution of mutual information and information quality ratio?

```
sim_trials <- 100
mutInfo <- c()
joints <- c()</pre>
```

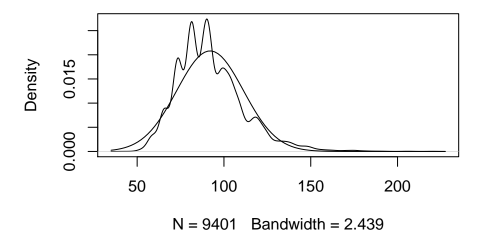
```
for(i in 1:sim_trials){
    #if(i%/floor(sim_trials/10)==0)    print(i/sim_trials)
    n_sim <- nrow(d2_test)
    bw_sim <- rnorm(n_sim,mean(d2_test$BodyweightKg),sd(d2_test$BodyweightKg))
    score_sim <- rnorm(n_sim,0,1)
    bins <- floor(sqrt(n_sim))
    joint <- kde2d(bw_sim,score_sim,n=95)$z
    mutInfo <- append(mutInfo,mi.plugin(joint))
    joints <- append(joints,entropy(joint))
}
plot(density(mutInfo/joints))</pre>
```

## density.default(x = mutInfo/joints)



curve(dnorm(x,mean(d2\_test\$BodyweightKg)),sd(d2\_test\$BodyweightKg)),add=TRUE)

### density.default(x = d2\_test\$BodyweightKg, adjust =



#### Score comparison using mutual information

This normalized version also known as Information Quality Ratio (IQR) which quantifies the amount of information of a variable based on another variable against total uncertainty: [26]

$$IQR(X,Y) = \mathrm{E}[\mathrm{I}(X;Y)] = \frac{\mathrm{I}(X;Y)}{\mathrm{H}(X,Y)} = \frac{\sum_{x \in X} \sum_{y \in Y} p(x,y) \log p(x) p(y)}{\sum_{x \in X} \sum_{y \in Y} p(x,y) \log p(x,y)} - 1$$

Figure 1: According to wikipedia, dividing by the joint entropy is a normalized version of MI known as IQR

As shown by the simulation and a priori reasoning, if the score is fair then the MI or IQR should approach 0

```
rel_joint <- kde2d(d2_test$BodyweightKg,d2_test$Rel_Total,n=95)$z
LL_joint <- kde2d(d2_test$BodyweightKg,d2_test$LogLogScore,n=95)$z
VV_joint <- kde2d(d2_test$BodyweightKg,d2_test$VVScore,n=95)$z
Wilks_joint <- kde2d(d2_test$BodyweightKg,d2_test$Wilks_Z,n=95)$z
mi.plugin(rel_joint) / entropy(rel_joint)</pre>
```

## [1] 0.01732757

mi.plugin(LL\_joint) / entropy(LL\_joint)

## [1] 0.002687419

mi.plugin(VV\_joint) / entropy(VV\_joint)

## [1] 0.002826842

```
mi.plugin(Wilks_joint) / entropy(Wilks_joint)
```

## [1] 0.002226846

HPDI(mutInfo/joints)

```
## | 0.89 | 0.89 | ## 0.0006485940 | 0.0008738527
```

All scores are well above the 89% interval for the information quality ratio. It appears that Wilks is slightly better than Allometric or varying variance scores. Relative Bodyweight Score is of course horrendous.