Measuring Performance: Ours vs Mourad

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Loading Libraries	
library(caret)	
<pre>## Warning: package 'caret' was built under R version 3.4.4 ## Loading required package: lattice ## Loading required package: ggplot2 ## Warning: package 'ggplot2' was built under R version 3.4.4 library(data.table)</pre>	
<pre>## Warning: package 'data.table' was built under R version 3.4.4 library(ggplot2) library(gbm)</pre>	
<pre>## Loading required package: survival ## ## Attaching package: 'survival' ## The following object is masked from 'package:caret': ## ## cluster ## Loading required package: splines ## Loading required package: parallel ## Loaded gbm 2.1.3</pre>	
library(pROC)	
## Warning: package 'pROC' was built under R version 3.4.4	

```
## Type 'citation("pROC")' for a citation.
##
## Attaching package: 'pROC'
## The following objects are masked from 'package:stats':
##
##
       cov, smooth, var
library(plyr)
library(dplyr)
## Warning: package 'dplyr' was built under R version 3.4.4
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:plyr':
##
##
       arrange, count, desc, failwith, id, mutate, rename, summarise,
##
       summarize
## The following objects are masked from 'package:data.table':
##
       between, first, last
##
## The following objects are masked from 'package:stats':
##
       filter, lag
##
## The following objects are masked from 'package:base':
       intersect, setdiff, setequal, union
##
library(DMwR)
## Loading required package: grid
##
## Attaching package: 'DMwR'
## The following object is masked from 'package:plyr':
##
##
       join
library(gridExtra)
##
## Attaching package: 'gridExtra'
## The following object is masked from 'package:dplyr':
##
       combine
library(knitr)
## Warning: package 'knitr' was built under R version 3.4.4
library(ROCR)
```

Loading required package: gplots

```
##
## Attaching package: 'gplots'
## The following object is masked from 'package:stats':
##
## lowess
```

Reading in RDS objects

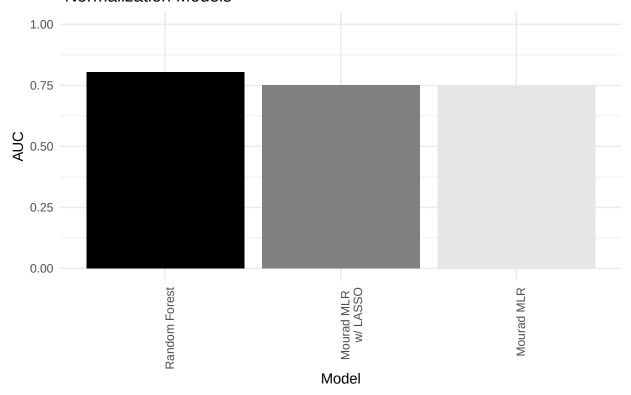
```
#mourad model
mourad.auc <- readRDS("C:/Users/Spiro Stilianoudakis/Documents/TAD_data/RData/GM12878/mourad_model/mour
mourad.roc <- readRDS("C:/Users/Spiro Stilianoudakis/Documents/TAD_data/RData/GM12878/mourad_model/mour
mourad.summary <- readRDS("C:/Users/Spiro Stilianoudakis/Documents/TAD_data/RData/GM12878/mourad_model/summard model with lasso
mourad.lasso.auc <- readRDS("C:/Users/Spiro Stilianoudakis/Documents/TAD_data/RData/GM12878/mourad_mode
mourad.lasso.roc <- readRDS("C:/Users/Spiro Stilianoudakis/Documents/TAD_data/RData/GM12878/mourad_mode
mourad.lasso.summary <- readRDS("C:/Users/Spiro Stilianoudakis/Documents/TAD_data/RData/GM12878/mourad_siterianoudakis/Documents/TAD_data/RData/GM12878/our_pipeline/rflst.rds
#random forest
rflst <- readRDS("C:/Users/Spiro Stilianoudakis/Documents/TAD_data/RData/GM12878/our_pipeline/rflst.rds
#gbm
gbmlst <- readRDS("C:/Users/Spiro Stilianoudakis/Documents/TAD_data/RData/GM12878/our_pipeline/gbmlst.re</pre>
```

Model Performance

AUCs

```
"Random Forest"),
                        auc=c(m.auc,
                              m.l.auc,
                              rf.auc))
auc.plot <- auc.plot[order(auc.plot$auc, decreasing=TRUE),]</pre>
auc.plot$Model <-factor(auc.plot$Model,</pre>
                                      levels=auc.plot$Model)
p<-ggplot(data=auc.plot, aes(x=Model, y=auc, fill=Model)) +</pre>
  xlab("Model") + ylab("AUC") +
  geom_bar(stat="identity") + ylim(0,1) +
  scale_fill_manual(values=grey(c(0,.5,.9)), guide=FALSE) +
  scale_x_discrete(labels= c("Random Forest",
                              "Mourad MLR \n w/ LASSO",
                              "Mourad MLR")) +
  theme_minimal() +
  theme(axis.text.x = element_text(angle = 90, hjust = 1)) +
  ggtitle("Model Performance for Different \n Normalization Models")
p
```

Model Performance for Different Normalization Models



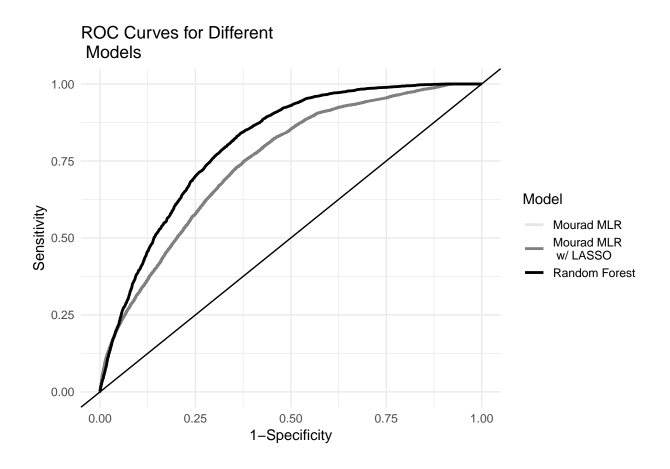
kable(auc.plot)

	Model	auc
3	Random Forest	0.8045185

	Model	auc
2	Mourad MLR w/ LASSO	0.7503714
1	Mourad MLR	0.7474979

ROC Curves

```
#mourad model
mourad.roc.fpr <- mourad.roc@ x.values[[1]]</pre>
mourad.roc.tpr <- mourad.roc@ y.values[[1]]</pre>
mourad.roc.df <- cbind.data.frame(fpr=mourad.roc.fpr,</pre>
                                    tpr=mourad.roc.tpr,
                                    Model = rep("M", length(mourad.roc.tpr)))
#mourad model w/ lasso
mourad.lasso.roc.fpr <- mourad.roc@ x.values[[1]]</pre>
mourad.lasso.roc.tpr <- mourad.roc@ y.values[[1]]</pre>
mourad.lasso.roc.df <- cbind.data.frame(fpr=mourad.lasso.roc.fpr,</pre>
                                    tpr=mourad.lasso.roc.tpr,
                                    Model = rep("MwL", length(mourad.lasso.roc.fpr)))
#random forest
rf.fpr <- rowMeans(rflst[[2]])</pre>
rf.tpr <- rowMeans(rflst[[1]])</pre>
rf.roc.df <- cbind.data.frame(fpr=rf.fpr,</pre>
                                    tpr=rf.tpr,
                                    Model = rep("RF", length(rf.fpr)))
#concatenating data frames
allrocdat <- rbind.data.frame(mourad.roc.df, mourad.lasso.roc.df, rf.roc.df)
ggplot(data=allrocdat, aes(x=fpr, y=tpr, color=Model)) +
  geom_line(size=1) +
  scale colour manual(name="Model",
    labels=c("Mourad MLR",
             "Mourad MLR \n w/ LASSO",
             "Random Forest"),
    values=grey(c(.9,.5,0))) +
  xlab("1-Specificity") +
  ylab("Sensitivity") +
  xlim(0, 1) +
  ylim(0, 1) +
  geom_abline(intercept=0, slope=1) +
  theme_minimal() +
  ggtitle("ROC Curves for Different \n Models")
```

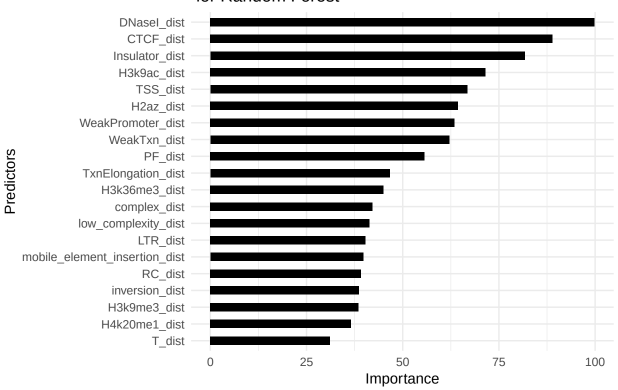


Variable Importance Plot

```
varimp.rf <- as.vector(rowMeans(rflst[[4]]))</pre>
rownames(rflst[[4]])[grep("Gm12878_", rownames(rflst[[4]]))] <- gsub("Gm12878_", "", rownames(rflst[[4]])
\#rownames(rflst[[4]])[grep("\_dist", rownames(rflst[[4]]))] \leftarrow gsub("\_dist", "", rownames(rflst[[4]])[grep("\_dist", rownames(rflst[[4]])]]
varimp.rf.df <- data.frame(Feature=rownames(rflst[[4]]),</pre>
                              Importance=varimp.rf)
varimp.rf.df <- varimp.rf.df[order(varimp.rf.df$Importance),]</pre>
numvarrf <- dim(varimp.rf.df)[1]</pre>
varimp.rf.df <- varimp.rf.df[(numvarrf-19):numvarrf,]</pre>
varimp.rf.df$Feature <- factor(varimp.rf.df$Feature,levels=varimp.rf.df$Feature)</pre>
rfp <- ggplot(varimp.rf.df, aes(x=Feature,</pre>
                                    y=Importance)) +
  xlab("Predictors") +
  ylab("Importance") +
  #ggtitle("Importance Plot for Gradient Boosting Machine") +
  geom_bar(stat="identity",
            width=.5,
            position="dodge",
```

```
fill="black") +
coord_flip() +
theme_minimal() +
ggtitle("Variable Importance Plot \n for Random Forest")
rfp
```

Variable Importance Plot for Random Forest



Estimates from Mourad Models

```
#mourad model
dim(mourad.summary)

## [1] 60 9

sig.vars <- mourad.summary[mourad.summary$^Pr(>|z|)^ < 0.05,]
dim(sig.vars)

## [1] 37 9

sig.vars <- sig.vars[order(abs(sig.vars$Estimate), decreasing = TRUE),]
rownames(sig.vars) <- NULL
sig.vars <- sig.vars[1:20, which(colnames(sig.vars) %in% c("GenomicFeature", "Estimate", "Pr(>|z|)"))]
kable(sig.vars)
```

Estimate	$\Pr(> z)$
3.8435655	0.0000000
3.0004561	0.0000000
1.8631419	0.0000000
1.8180323	0.0000000
1.5380977	0.0000000
1.3605898	0.0000001
1.2286235	0.0000006
1.1905517	0.0000011
1.1758725	0.0000019
1.1510972	0.0000458
1.1367741	0.0000071
1.0690906	0.0000000
1.0399866	0.0000254
1.0286227	0.0000242
1.0038337	0.0001420
0.9625891	0.0002621
0.7478179	0.0034025
-0.4638145	0.0056090
0.4612106	0.0016588
-0.4475249	0.0036293
	3.8435655 3.0004561 1.8631419 1.8180323 1.5380977 1.3605898 1.2286235 1.1905517 1.1758725 1.1510972 1.1367741 1.0690906 1.0399866 1.0286227 1.0038337 0.9625891 0.7478179 -0.4638145 0.4612106

```
#mourad model w/ lasso
dim(mourad.lasso.summary)
```

```
## [1] 60 3
```

```
mourad.lasso.summary <- as.data.frame(mourad.lasso.summary)
mourad.lasso.summary$Estimate <- as.numeric(as.character(mourad.lasso.summary$Estimate))

sig.vars.lasso <- mourad.lasso.summary[order(abs(mourad.lasso.summary$Estimate), decreasing = TRUE),]
rownames(sig.vars.lasso) <- NULL
sig.vars.lasso <- sig.vars.lasso[1:20, which(colnames(sig.vars.lasso) %in% c("GenomicFeature", "Estimate
kable(sig.vars.lasso)</pre>
```

GenomicFeature	Estimate
srpRNA	-4.24869
A	3.43288
В	2.58396
RC	-1.29609
$novel_sequence_insertion$	-1.26161
CTCF	1.14327
Insulator	1.01586
ActivePromoter	0.92045
WeakPromoter	0.66831
DNaseI	0.57307
WeakEnhancer6	0.53279
satellite	0.44559
RepetitiveCNV15	-0.40150
H3k27me3	0.36100
H2az	0.32995
$sequence_alteration$	-0.31331

GenomicFeature	Estimate
WE	-0.30484
H3k9me3	-0.26855
TxnElongation	0.25109
se	-0.24898

Comparing Results

Feature	Random Forest	Mourad	Mourad w/ LASSO
DNaseI	1	19	10
CTCF	2	12	6
Insulator	3	3	7
H3k9ac	4	NA	NA
TSS	5	NA	NA
H2az	6	NA	15
WeakPromoter	7	5	9
WeakTxn	8	8	NA
PF	9	NA	NA
TxnElongation	10	7	19
H3k36me3	11	NA	NA
complex	12	NA	NA
low_complexity	13	NA	NA
LTR	14	NA	NA
$mobile_element_insertion$	15	NA	NA
RC	16	NA	4
inversion	17	NA	NA
H3k9me3	18	NA	18
H4k20me1	19	NA	NA
T	20	NA	NA