Main: Facial Expression Recognition Framework

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Introduction

In this project, we created a classification engine for facial emotion recognition. Our group tried out four typical machine learning algorithms, trained them on the given data set, cross-validated to find the optimized parameters, and provided fair evaluation for all four of them. The evaluation process considered the mean error (and accuracy), the area under the ROC curve (or AUC), and running time (testing and training) to give a thorough comparison of the four methods.

Models Introduction

• gbm model (baseline)

The baseline model gradient boosting is an ensemble method that uses a combination of weak learners—in this case, decision stumps—to predict classes. In each iteration of model training, the algorithm focuses on areas that the weak learners fail to properly classify and, using class weights, will focus on precisely these areas in the next iteration. This learning is done via gradient descent.

• xgboost model (proposed)

Extreme gradient boosting is a much faster version of the original gradient boosting model and uses methods faster than gradient descent, such as stochastic gradient descent in order to learn much quicker.

• PCA plus SVM model

Since the featured train data has a rather high dimension, we try PCA for data reduction as the first step. In this step we calculate all the principle components and cross validate the principle component number that gives the highest accuracy. Then we use SVM as the model since it performs well on two-group classification.

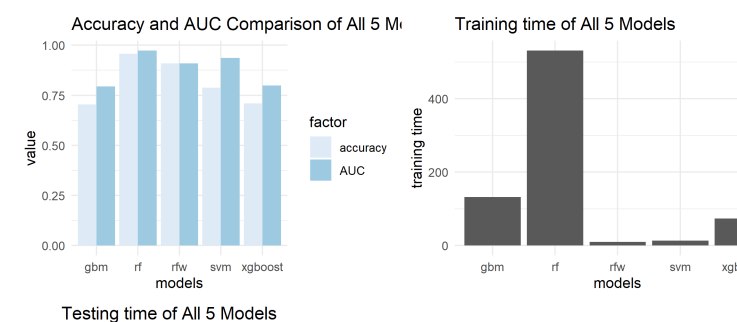
• random forest model

We try this model since random forest model is know for its high performance and powerfulness.

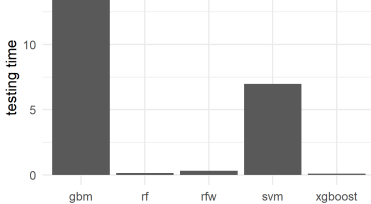
• random forest with weights model

Considering that the featured training set is highly-unbalanced, we try random forest model with weights as well.

Results







models

Analysis

Step 0 set work directories

```
set.seed(2020)
# setwd("~/GitHub/Fall2020-Project3-group1/doc") #change the working directory as needed
```

Provide directories for training images. Training images and Training fiducial points will be in different subfolders.

```
train_dir <- "../data/train_set/" #may need to be changed to local directory

train_image_dir <- paste(train_dir, "images/", sep="")
train_pt_dir <- paste(train_dir, "points/", sep="")
train_label_path <- paste(train_dir, "label.csv", sep="")</pre>
```

Step 1: set up controls for evaluation experiments.

In this chunk, we have a set of controls for the evaluation experiments.

- (T/F) cross-validation on the training set
- (T/F) reweighting the samples for training set
- (number) K, the number of CV folds
- (T/F) process features for training set
- (T/F) run evaluation on an independent test set
- (T/F) process features for test set

```
K <- 5 # number of CV folds
run.fudicial.list <- FALSE</pre>
run.feature.train <- FALSE # process features for training set
run.feature.test <- FALSE # process features for test set</pre>
sample.reweight <- TRUE # run sample reweighting in model training
run.cv.gbm <- FALSE # run cross-validation on the training set for qbm
run.train.gbm <- FALSE # run evaluation on entire train set
run.test.gbm <- TRUE # run evaluation on an independent test set
run.cv.xgboost <- FALSE # run cross-validation on the training set for xqboost
run.train.xgboost <- FALSE # run evaluation on entire train set
run.test.xgboost <- TRUE # run evaluation on an independent test set
run.cv.rforestw <- FALSE # run cross-validation on the training set for xgboost
run.train.rforestw <- FALSE # run evaluation on entire train set
run.test.rforestw <- TRUE # run evaluation on an independent test set
run.cv.RF <- FALSE # run cross-validation on the training set for xgboost
run.train.RF <- FALSE # run evaluation on entire train set
run.test.RF <- TRUE # run evaluation on an independent test set
run.cv.svm <- FALSE # run cross-validation on the training set for sum
run.train.svm <- FALSE # run evaluation on entire train set
run.test.svm <- TRUE # run evaluation on an independent test set
run.cv.pca <-FALSE # calculate pca</pre>
```

Using cross-validation or independent test set evaluation, we compare the performance of models with different specifications.

In this chunk, we set up the hyperparameters used for our models.

```
# gbm model (baseline)
hyper_grid_gbm <- expand.grid(</pre>
```

```
shrinkage = c(0.001, 0.005, 0.010, 0.050, 0.100),
  n.trees = c(600, 1200, 1800)
)
# xqboost model
hyper_grid_xgboost <- expand.grid(</pre>
  eta = c(0.01, 0.05, 0.1, 0.2, 0.3),
  lambda = c(0.001, 0.005, 0.010, 0.050, 0.100),
  gamma = c(0, 5),
  nrounds = c(600, 1200, 1800)
# sum model
hyper_grid_svm <- expand.grid(</pre>
 nprinciple = c(400, 450, 500, 550, 600, 650, 700, 750)
# random forest model
hyper_grid_rf_uw <- expand.grid(
 ntrees = c(100, 300, 500, 800, 1000),
 mtry = c(500)
# random forest with weights model
hyper_grid_rforest <- expand.grid(</pre>
 ntrees = c(100, 300, 500, 800, 1000),
 \max d = c(0, 5, 10, 15, 20, 25)
)
```

Step 2: import data and train-test split

```
info <- read.csv(train_label_path)
n <- nrow(info)
n_train <- round(n*(4/5), 0)
train_idx <- sample(info$Index, n_train, replace = F)
test_idx <- setdiff(info$Index, train_idx)</pre>
```

Fiducial points are stored in matlab format. In this step, we read them and store them in a list.

```
n_files <- length(list.files(train_image_dir))

if (run.fudicial.list){
    readMat.matrix <- function(index){
        return(round(readMat(pasteO(train_pt_dir, sprintf("%04d", index), ".mat"))[[1]],0))
    }

    fiducial_pt_list <- lapply(1:n_files, readMat.matrix)
    save(fiducial_pt_list, file="../output/fiducial_pt_list.RData")

# otherwise load the data stored for convenience
} else {</pre>
```

```
load(file="../output/fiducial_pt_list.RData")
}
```

Step 3: construct features and responses

feature.R is the wrapper for all our feature engineering functions and options. The function feature() have options that correspond to different scenarios for our project and produces an R object that contains features and responses that are required by all the models we are going to evaluate later.

- feature.R
- Input: list of images or fiducial point
- Output: an RData file that contains extracted features and corresponding responses

```
source("../lib/feature.R")

tm_feature_train <- NA

if(run.feature.train){
    tm_feature_train <- system.time(dat_train <- feature(fiducial_pt_list, train_idx))
    save(dat_train, tm_feature_train, file="../output/feature_train.RData")
}else{
    load(file="../output/feature_train.RData")
}

tm_feature_test <- NA
    if(run.feature.test){
        tm_feature_test <- system.time(dat_test <- feature(fiducial_pt_list, test_idx))
        save(dat_test, tm_feature_test, file="../output/feature_test.RData")
}else{
    load(file="../output/feature_test.RData")
}</pre>
```

Gradient Boosted Trees (gbm model) (Baseline Model)

Step 4: Train a classification model with training features and responses

Call the train_gbm model and test_gbm model from library.

 $\verb|train_gbm.R| and \verb|test_gbm.R| are wrappers for all our gbm model training steps and classification/prediction steps.$

- train_gbm.R
 - Input: a data frame containing features and labels and a parameter list.
 - Output:a trained model
- test_gbm.R
 - Input: the fitted classification model using training data and processed features from testing images
 - Input: an R object that contains a trained classifier.
 - Output: training model specification

Since other train and test documents have the same structure as the gbm model, we will omit them.

Model selection with cross-validation

• Do model selection by choosing among different values of training model parameters

Here we cross-validate parameters shrinkage and number of trees.

- n.trees: the number of trees (or decision stumps) used in the model
- shrinkage: the learning rate

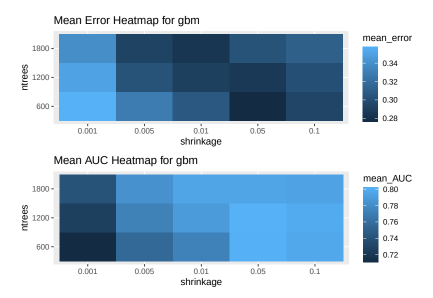
The model also uses several parameters that can be cross-validated, such as "interaction.depth" and "distribution" were set to fit the needs of this situation. For example, "interaction.depth" was set to 1 since our focus was to use decision trees and "distribution" was set to "binomial" since there are only two classes.

We cross-validated three different values of "shrinkage" and three different values of "n.trees" for a total of 15 models cross-validated. We choose to test various slower learning rates—that is, none greater than 0.1, and we chose to consider 600, 1200, and 1800 decision stumps, as it could build a model based on approximately 10%, 20%, or 30%, respectively of the 6006 fiducial features.

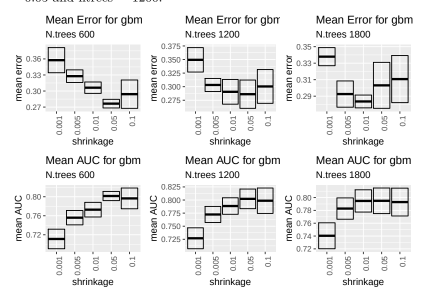
And the first 5 optimized parameters among the all 15 models we got are: (see the appendix for full table)

```
##
      shrinkage n.trees mean_error
                                      sd_error mean_AUC
## 9
          0.05
                   1200 0.2861576 0.026049925 0.8022183 0.018723658
                   600 0.2764453 0.008045855 0.8015074 0.009653744
## 4
          0.05
## 10
          0.10
                   1200 0.3004216 0.031100034 0.7986702 0.024253701
## 5
          0.10
                    600 0.2941406 0.026467746 0.7965060 0.021828465
## 14
          0.05
                   1800 0.3031199 0.027850857 0.7949472 0.019940282
```

• Visualize cross-validation results



We could also see here that the set of parameters which will give us the smallest mean error is shrinkage = 0.05 and ntrees = 600 and that the set of parameters which will give us the biggest mean AUC is shrinkage = 0.05 and ntrees = 1200.



We could see in the plots the detailed performance of each set of parameters.

• Choose the "best" parameter value

Due to the presence imbalanced data, we choose to focus out attention on highest mean AUC rather than lowest mean error. However, we notice that the second best model (model 4) has an mean AUC comparable to that of the best model (model 9) while being much simpler—model 4 has 600 trees while model 9 has 1200—we choose to select the more parsimonious model as our best baseline gbm model.

```
gbm_cv_results[order(gbm_cv_results$mean_AUC, decreasing = TRUE), ][2, ]
## shrinkage n.trees mean_error sd_error mean_AUC sd_AUC
## 4     0.05     600     0.2764453     0.008045855     0.8015074     0.009653744
```

```
par_best_gbm_ind <- 4
par_best_gbm_shrinkage <- gbm_cv_results$shrinkage[par_best_gbm_ind]
par_best_gbm_n.trees <- gbm_cv_results$n.trees[par_best_gbm_ind]</pre>
```

• Train the model with the entire training set using the selected model (model parameter) via cross-validation.

```
if (run.train.gbm) {
  # training weights
  weight_train <- rep(NA, length(label_train))</pre>
  for (v in unique(label_train)){
    weight_train[label_train == v] = 0.5 * length(label_train) /
      length(label_train[label_train == v])
  }
  if (sample.reweight){
    tm_train_gbm <- system.time(fit_train_gbm</pre>
                                 <- train(feature_train, label_train, w = weight_train,
                                          num_trees = par_best_gbm_n.trees,
                                          shrink = par_best_gbm_shrinkage))
  } else {
    tm train gbm <- system.time(fit train gbm</pre>
                                 <- train(feature_train, label_train, w = NULL,
                                          num_trees = par_best_gbm_n.trees,
                                          shrink = par_best_gbm_shrinkage))
  }
  save(fit_train_gbm, tm_train_gbm, file="../output/fit_train_gbm.RData")
} else {
  load(file="../output/fit_train_gbm.RData")
```

Step 5: Run test on test images

```
## reweight the test data to represent a balanced label distribution
weight_test <- rep(NA, length(label_test))
for (v in unique(label_test)){</pre>
```

```
weight_test[label_test == v] =
    0.5 * length(label_test) / length(label_test[label_test == v])

# convert the original 1-2 class into numeric Os and 1s
label_test <- ifelse(label_test == 2, 0, 1)

accu_gbm <- sum(weight_test * (label_pred == label_test)) / sum(weight_test)
tpr.fpr <- WeightedROC(prob_pred, label_test, weight_test)
auc_gbm<- WeightedAUC(tpr.fpr)</pre>
```

Evaluation

```
## The accuracy of the gbm model (shinkage = 0.05, n.trees = 600) is 70.48409%.
## The AUC of the gbm model (shinkage = 0.05, n.trees = 600) is 0.7935942.
```

Summarize Running Time Prediction performance matters, so does the running times for constructing features and for training the model, especially when the computation resource is limited.

```
## Time for constructing training features = 2.198 seconds
## Time for constructing testing features = 0.218 seconds
## Time for training gbm model = 132.18 seconds
## Time for testing gbm model = 14.93 seconds
```

xgboost Model (Proposed Model)

Step 4: Train a classification model with training features and responses

Call the train xgboost model and test xgboost model from library.

Model selection with cross-validation

• Do model selection by choosing among different values of training model parameters.

Here we cross-validate parameters eta, lambda, gamma and nrounds.

- nrounds: similar to the gbm model's "n.trees" parameter, which we again cross-validated the values 600, 1200, and 1800.
- eta: the learning rate, similar to the gbm model's "shrinkage" parameter. The default value is 0.3, so we cross-validate five values leading up to and including 0.3.
- gamma: allows the algorithm to decide whether to split on a certain part of the tree—the default value is 0, and other oft-used numbers include 1 and 5. Here, we choose to cross-validate with values 0 and 5.

• lambda: a regularization term for the weights. We use values ranging from 0.01 to 0.1.

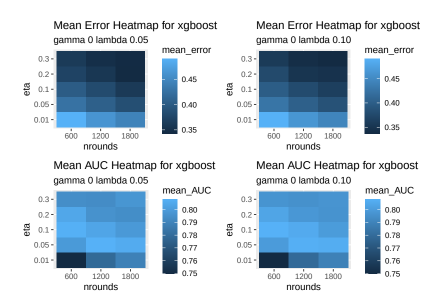
The "max_depth" parameter and have been set to 1 and the "objective" parameter was set to "binary:logistic" in order for us to compare equally with the gbm model.

In total, we have cross-validated 150 different xgboost models.

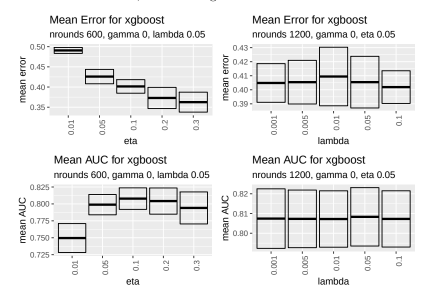
And the first 5 optimized parameters among the all 150 models we got are: (see the appendix for full table)

```
eta lambda gamma nrounds mean_error
                                           sd_error mean_AUC
##
## 67 0.05 0.050
                     0
                          1200 0.4054330 0.01838352 0.8083090 0.01479400
## 23 0.10 0.100
                           600 0.3987534 0.01527476 0.8081563 0.01500276
                     0
## 3 0.10 0.001
                     0
                           600 0.3967334 0.01385476 0.8080448 0.01802835
## 18 0.10 0.050
                     0
                           600 0.4014677 0.01679825 0.8078401 0.01590874
                          1200 0.4048485 0.01376101 0.8074209 0.01509062
## 52 0.05 0.001
                     0
```

• Visualize cross-validation results.



Since the xgboost model have 4 parameters in total, and we could only show 2 of them in a 2-D plot, we choose only to display several comparative plots to give a brief idea of visualizing the data. To actually select the best model, we are using the method to be discussed in the next section.



• Choose the "best" parameter value

Due to the presence imbalanced data, we choose to focus out attention on highest mean AUC rather than lowest mean error. However, we notice that the second best model (model 67) is has an mean AUC comparable to that of the best model (model 23) while being much simpler—model 23 has 600 trees while model 67 has 1200—we choose to select the more parsimonious model as our best proposed xgboost model.

```
par_best_res_cv_xgboost_cv_results_ind <- 23

par_best_res_cv_xgboost_cv_results_eta <-
    res_cv_xgboost_cv_results$eta[par_best_res_cv_xgboost_cv_results_ind]

par_best_res_cv_xgboost_cv_results_lambda <-
    res_cv_xgboost_cv_results$lambda[par_best_res_cv_xgboost_cv_results_ind]

par_best_res_cv_xgboost_cv_results_gamma <-
    res_cv_xgboost_cv_results$gamma[par_best_res_cv_xgboost_cv_results_ind]

par_best_res_cv_xgboost_cv_results_nrounds <-
    res_cv_xgboost_cv_results$nrounds[par_best_res_cv_xgboost_cv_results_ind]</pre>
```

• Train the model with the entire training set using the selected model (model parameters) via cross-validation.

```
if (run.train.xgboost) {
  weight_train <- rep(NA, length(label_train))</pre>
  for (v in unique(label_train)){
    weight_train[label_train == v] =
      0.5 * length(label_train) / length(label_train[label_train == v])
  }
  if (sample.reweight){
    tm_train_xgboost <- system.time(</pre>
      fit train xgboost <- train(features = feature train, labels = label train,
                                  w = weight_train,
                                  eta_val = par_best_res_cv_xgboost_cv_results_eta,
                                  lmd = par_best_res_cv_xgboost_cv_results_lambda,
                                  gam = par_best_res_cv_xgboost_cv_results_gamma,
                                  nr = par_best_res_cv_xgboost_cv_results_nrounds))
  }else{
    tm_train_xgboost <- system.time(</pre>
      fit_train_xgboost <- train(features = feature_train, labels = label_train,</pre>
                                  w = NULL,
                                  eta val = par best res cv xgboost cv results eta,
                                  lmd = par_best_res_cv_xgboost_cv_results_lambda,
                                  gam = par_best_res_cv_xgboost_cv_results_gamma,
                                  nr = par_best_res_cv_xgboost_cv_results_nrounds))
  save(fit train xgboost, tm train xgboost, file="../output/fit train xgboost.RData")
}else {
  load(file="../output/fit train xgboost.RData")
```

Step 5: Run test on test images

```
tm_test_xgboost= NA
feature_test <- as.matrix(dat_test[, -6007])
label_test <- as.integer(dat_test$label)

if(run.test.xgboost){
   load(file=".../output/fit_train_xgboost.RData")</pre>
```

```
## reweight the test data to represent a balanced label distribution
weight_test <- rep(NA, length(label_test))
for (v in unique(label_test)){
    weight_test[label_test == v] = 0.5 * length(label_test) /
        length(label_test[label_test == v])
}

# convert the original 1-2 class into numeric Os and 1s
label_test <- ifelse(label_test == 2, 0, 1)

accu_xgboost <- sum(weight_test * (label_pred == label_test)) / sum(weight_test)
tpr.fpr <- WeightedROC(prob_pred, label_test, weight_test)
auc_xgboost <- WeightedAUC(tpr.fpr)</pre>
```

Evaluation

```
## The accuracy of the xgboost model (eta = 0.1, nrounds = 600, lambda = 0.1, gamma = 0) is 70.9835%.
## The AUC of the xgboost model (eta = 0.1, nrounds = 600, lambda = 0.1, gamma = 0) is 0.7985717.
```

Summarize Running Time

```
## Time for training xgboost model = 74.03 seconds
## Time for testing xgboost model = 0.11 seconds
```

Principal Components Analysis (PCA) + Support Vector Machines (SVMs)

Step 4: Train a classification model with training features and responses

Call the train_svm model and test_svm model from library.

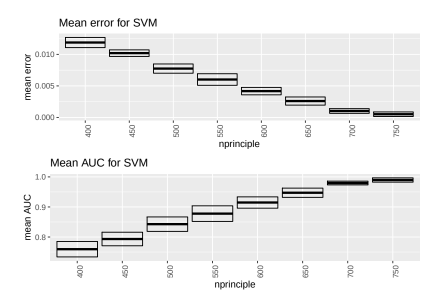
Model selection with cross-validation

• Do model selection by choosing among different values of training model parameters.

Here we cross-validate parameter nprinciple.

```
feature_train = as.matrix(dat_train[, -6007])
label_train = as.integer(dat_train$label)
if(run.cv.pca){
  pca1 <- prcomp(feature_train)
  save(pca1,file="../output/pcaCalc.RData")</pre>
```

• Visualize cross-validation results.



• Choose the "best" parameter value

```
par_best <- hyper_grid_svm$nprinciple[which.max(res_cv_svm$mean_accu)]</pre>
```

• Train the model with the entire training set using the selected model (model parameter) via cross-validation.

Step 5: Run test on test images

```
tm_test_svm = NA
feature_test <- as.matrix(dat_test[, -6007])

if(run.test.svm){
    load(file="../output/fit_train_svm.RData")
    tm_test_svm <- system.time({ prob_pred <- test(fit_train, feature_test, pca1,par_best)})
}</pre>
```

```
label_test <- as.integer(dat_test$label)
weight_test <- rep(NA, length(label_test))
for (v in unique(label_test)){
   if (as.integer(v)==2){
      weight_test[label_test == v] = 1
   }else{
      weight_test[label_test == v] = 10
   }
}
finalguess <- as.numeric(prob_pred)
accu_svm <- sum(finalguess == label_test) / sum(label_test)
tpr.fpr <- WeightedROC(as.numeric(prob_pred), label_test, weight_test)
auc_svm <- WeightedAUC(tpr.fpr)</pre>
```

Evaluation

```
## The accuracy is: 78.81706 %.

## The AUC is 0.9368997 .

Summarize Running Time

## Time for training model= 13.285 s

## Time for testing model= 6.97 s
```

Random Forest

Step 4: Train a classification model with training features and responses

Call the train_RF model and test_RF model from library.

Model selection with cross-validation

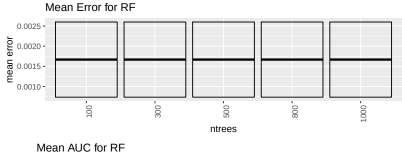
• Do model selection by choosing among different values of training model parameters.

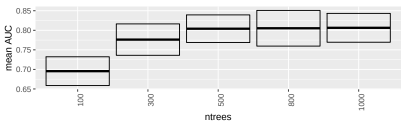
Here we cross-validate parameter ntrees and mtry (actually we use only one value for mtry)

- ntree: for random forest, the ntree parameter is usually set to be the 10% of the sample size. Since our sample has a size around 6000, we choose ntree to be 100, 300, 500, 800, and 1000.
- mtry: this parameter is more experience-related, so we set its value to be the default 500.

```
source("../lib/cross_validation_RF.R")
feature_train = as.matrix(dat_train[, -6007])
label_train = as.integer(dat_train$label)
if(run.cv.RF){
    res_cv <- matrix(0, nrow = length(ntree)*length(mtry), ncol = 4)
    for(i in 1:length(ntree)){
        cat("ntree = ", ntree[i], "\n")
        for(j in 1:length(mtry)){
        cat("mtry = ", mtry[j], "\n")
        res_cv[(i-1)*length(mtry)+j,] <- cv.function(
            features = feature_train, labels = label_train, K, ntree = ntree[i],mtry=mtry[j])
}}
save(res_cv, file="../output/res_cv_RF.RData")
}else{
    load("../output/res_cv_RF.RData")
}</pre>
```

• Visualize cross-validation results.





• Choose the "best" parameter value

```
par_best <- hyper_grid_rf_uw$ntrees[which.max(res_cv_rforest$mean_AUC)]</pre>
```

• Train the model with the entire training set using the selected model (model parameter) via cross-validation.

Step 5: Run test on test images

```
tm_test = NA
feature_test <- as.matrix(dat_test[, -6007])

if(run.test.RF){
   load(file="../output/fit_train_RF.RData")
   tm_test_RF <- system.time(label_pred <- as.integer(test_RF(fit_train_RF, feature_test)));
}</pre>
```

Evaluation

```
## Setting levels: control = 0, case = 1
## Setting direction: controls < cases
## The unweighted accuracy of the random forest model is 95.66667 %.
## The unweighted AUC of the random forest model is 0.9739479 .</pre>
```

Summarize Running Time Prediction performance matters, so does the running times for constructing features and for training the model, especially when the computation resource is limited.

```
## Time for training random forest model= 531.55 s
## Time for testing random forest model= 0.16 s
```

Random Forest with weights

Step 4: Train a classification model with training features and responses

Call the train model and test model from library.

Model selection with cross-validation

• Do model selection by choosing among different values of training model parameters.

From the experience of trying out different mtry value for the random forest model without a significant difference in the final result, we switch to another parameter worth worrying about: the max depth parameter.

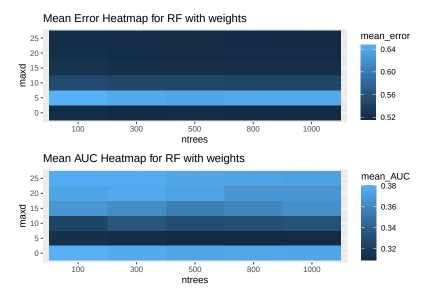
Here we cross-validate parameter ntrees and max depth.

```
sample.reweight <- T</pre>
if(run.cv.rforestw){
  res_cv <- matrix(0, nrow = nrow(hyper_grid_rforest), ncol = 4)</pre>
  for (i in 1:nrow(hyper_grid_rforest)){
    print(hyper_grid_rforest$ntree[i])
    print(hyper_grid_rforest$maxd[i])
    res_cv[i,] <- cv.function(features = feature_train,</pre>
                              labels = label_train,
                              ntree = hyper_grid_rforest$ntree[i],
                              md = hyper_grid_rforest$maxd[i],
                              K, reweight = sample.reweight
    )
  }
  save(res_cv, file="../output/res_cv_rforestw.RData")
  load("../output/res_cv_rforestw.RData")
}
```

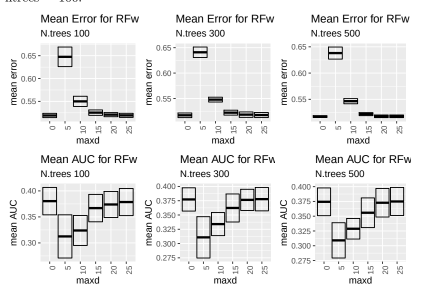
And the first 5 optimized parameters among the all 30 models we got are: (see the appendix for full table)

```
##
     ntrees maxd mean_error
                                                       sd AUC
                                sd_error mean_AUC
## 1
         100
               0 0.5191987 0.004676295 0.3803271 0.02631877
## 26
         100
              25 0.5194598 0.004461227 0.3785671 0.02605897
## 27
         300
               25 0.5173999 0.004971162 0.3777519 0.02049336
## 2
         300
               0 0.5168797 0.004553045 0.3772316 0.02038906
## 22
         300
               20 0.5181833 0.005354415 0.3764582 0.01852388
```

• Visualize cross-validation results



We could also see here that the set of parameters which will give us the smallest mean error is maxd = 25 and ntrees = 100 and that the set of parameters which will give us the biggest mean AUC is maxd = 0 and ntrees = 100.



• Choose the "best" parameter value

Due to the presence imbalanced data, we choose to focus out attention on highest mean AUC rather than lowest mean error. However, we notice that the second best model (model 67) is has an mean AUC comparable to that of the best model (model 23) while being much simpler—model 23 has 600 trees while model 67 has 1200—we choose to select the more parsimonious model as our best proposed xgboost model.

```
par_best_res_cv_rforest_cv_results_ind <- which(
   res_cv_rforest_cv_results$mean_AUC == max(res_cv_rforest_cv_results$mean_AUC))

par_best_res_cv_xgboost_cv_results_ntrees <-
   res_cv_rforest_cv_results$ntrees[par_best_res_cv_rforest_cv_results_ind]

par_best_res_cv_xgboost_cv_results_md <-
   res_cv_rforest_cv_results$maxd[par_best_res_cv_rforest_cv_results_ind]</pre>
```

• Train the model with the entire training set using the selected model (model parameter) via cross-validation.

```
if (run.train.rforestw) {
  # training weights
  weight_train <- rep(NA, length(label_train))</pre>
  for (v in unique(label train)){
    weight_train[label_train == v] = 0.5 * length(label_train) /
      length(label train[label train == v])
  }
  if (sample.reweight){
    tm_train_rforestw <- system.time(</pre>
      fit_train_rforestw <- train(features = feature_train, labels = label_train,</pre>
                                   w = weight_train,
                                   ntree = par_best_res_cv_xgboost_cv_results_ntrees,
                                   md = par_best_res_cv_xgboost_cv_results_md))
  } else {
    tm_train_rforestw <- system.time(</pre>
      fit_train_rforestw <- train(features = feature_train, labels = label_train,</pre>
                                   w = NULL,
                                   ntree = par_best_res_cv_xgboost_cv_results_ntrees,
                                   md = par_best_res_cv_xgboost_cv_results_md))
  save(fit_train_rforestw, tm_train_rforestw, file="../output/fit_train_rforestw.RData")
} else {
  load(file="../output/fit train rforestw.RData")
```

Step 5: Run test on test images

```
## reweight the test data to represent a balanced label distribution
weight_test <- rep(NA, length(label_test))
for (v in unique(label_test)){</pre>
```

```
weight_test[label_test == v] = 0.5 * length(label_test) /
    length(label_test[label_test == v])
}

# convert the original 1-2 class into numeric 0s and 1s
label_test <- ifelse(label_test == 2, 0, 1)

accu_rfw <- sum(weight_test * (label_pred == label_test)) / sum(weight_test)
tpr.fpr <- WeightedROC(prob_pred$predictions, label_test, weight_test)
auc_rfw <- WeightedAUC(tpr.fpr)</pre>
```

Evaluation

The accuracy of the random forest with weights model (ntrees = 100, max_depth = 0) is 90.99233%.

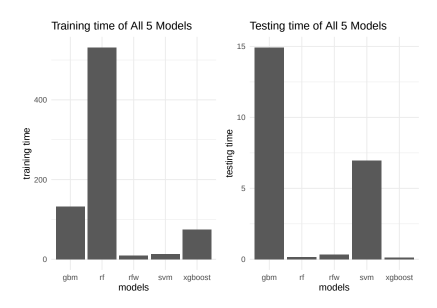
The AUC of the random forest with weights model (ntrees = 100, max_depth = 0) is 0.9099233.

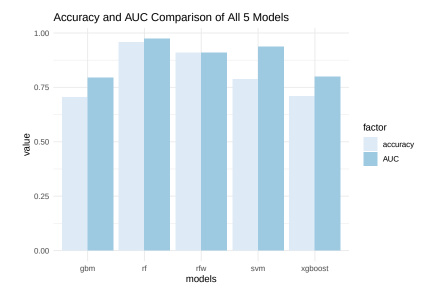
Summarize Running Time

Time for training random forest with weights model = 9.486 seconds

Time for testing random forest with weights model = 0.33 seconds

Results





Reference(s)

- Du, S., Tao, Y., & Martinez, A. M. (2014). Compound facial expressions of emotion. Proceedings of the National Academy of Sciences, 111(15), E1454-E1462.
- Revert, Félix. "Fine-Tuning XGBoost in Python like a Boss." Medium, Towards Data Science, 20 Sept. 2019, towardsdatascience.com/fine-tuning-xgboost-in-python-like-a-boss-b4543ed8b1e.
- Jain, Aarshay. "XGBoost Parameters: XGBoost Parameter Tuning." Analytics Vidhya, 18 Oct. 2020, www.analyticsvidhya.com/blog/2016/03/complete-guide-parameter-tuning-xgboost-with-codes-python/.
- "XGBoost Documentation" XGBoost Documentation Xgboost 1.3.0-SNAPSHOT Documentation, xgboost.readthedocs.io/en/latest/index.html.
- \bullet Greenwell, Brandon. "Gbm." Function | R Documentation, www.rdocumentation.org/packages /gbm/versions/2.1.8/topics/gbm.

Appendix

We provide the full output of the cross-validation table for each model below. They are ready to view after uncommenting. We did not display them in our final report because it is very space-consuming to do so.

```
# gbm_cv_results # gbm cross validation results
# res_cv_xgboost_cv_results # xgboost cross validation results
# res_cv_svm # pca and svm cross validation results
# res_cv_rforest # random forest cross validation results
# res_cv_rforest_cv_results # random forest with weights cross validation results
```