MWAS code structure:

```
mwas_analysis.r -- command line function, includes "train",
"predict", "plot", "statistics" options.
Dependencies (required packages):
1. optparse - option parsing

    biom format file operation

2. biom
3. e1071 - classifier package (SVM, cross-validation...)
4. randomForest - random forest
5. glmnet - for multi-normial regression
6. pR0C
             ROC analysis

    7. caret - variable importance analysis for feature selection
    8. xlsx - for Excel workbook output

9. beeswarm - beeswarm plot
# 10. kernlab - future version will add this function
Input Option list
Preprocessing: (universal options for all modes)
           transform_type
     -t
           suppress relative abundance conversion
     -r
     -v
           min prevalence
           collapse_table
     -е
Training:
Rscript mwas_analysis.r -w train
           otu fp
     -i
           map_fp
     -\mathsf{m}
           category
     -c
           out dir
     -0
     -M
           method (classifier type)
     -C
           classifier_parameter
           validation_type
     -a
     -n
           number of fold cv
     -f
           is_feature_selection
     -s
           feature selection paramter
Rscript bin/mwas analysis.R -w train -i
test/data/GG 100nt even10k-adults-s20.biom -m test/data/gg-
map-adults.txt -c COUNTRY -o example/train out/ -M svm
Predicting:
Rscript mwas_analysis.r -w predict
    -i
           otu_fp
           map_fp (optional)
    -m
    -c
           category (optional)
    -0
           out_dir
           trained_model_fp (classifier_parameter)
    -C
```

```
Rscript bin/mwas_analysis.R -w predict -i test/data/GG_100nt_even10k-adults-s20.biom -m test/data/gg-map-adults.txt -c COUNTRY -o example/predict_out/ -M example/train_out/trained_model.rds
```

```
Plot: (all capital for plot-only parameters)
Rscript mwas analysis.r -w plot
          otu_fp
   -i
   -0
          out_dir
    -с
          category
   -m
          map fp
   -F
          feat_stats_fp (feature_parameter)
   -M
          plot_type (method)
   −D
          distance fp
   −P
          pcoa_fp
   -T
          which taxa
   -S
          shorten taxa
   -X
          multiple axis
   -A
          Maximum false discovery rate (alpha)
   –K
          is.filter_kegg (for heatmap)
Rscript bin/mwas analysis.R -w plot -V gradients -i
test/data/taxa/merged-taxa.txt -W 'Bacteroides, Prevotella'
-o example/plot out gradient
Beeswarm with feature stats table
Rscript bin/mwas analysis.R -w plot -M beeswarm -i
test/data/taxa/merged-taxa.txt -o example/beeswarm -m
test/data/gg-map-adults.txt -c COUNTRY -F
test/data/stats/taxon-stats-table.txt -A 0.01
Beeswarm without feature stats table
Rscript bin/mwas analysis.R -w plot -M beeswarm -i
test/data/taxa/merged-taxa.txt -o example/beeswarm -m
test/data/gg-map-adults.txt -c COUNTRY -A 0.1
Scatterplot is similar to the Beeswarm plots, except
-M scatterplot
```

Statistics:

```
Rscript mwas_analysis.r -w statistics
-i otu_fp
-m map_fp
-c category
-o out dir
```

Function list

File name	Function names	Notes
import_data.R	import.train.params	
	import.predict.params	
	import.plot.params	
	import.stats.params	
util.load.R	read.qiime.table.mwas	
	read.qiime.classic.table	
	load.qiime.mapping.file	
	load.qiime.otu.table	
	otu.table.has.metadata	
	get.header.index	
	load.qiime.taxon.table	
	load.qiime.distance.matrix	
	load.qiime.pcoa.file	
	load.qiime.feature.stats	
	remove.nonoverlapping.samples	
	cluster.by.correlation	
	collapse.by.correlation	
	inspect.env	unused
	shorten.taxonomy	
	balanced.folds	unused
	is.outlier	unused
	factor.to.numeric	unused
	parse.params	updated
model.train.R	train.mwas	
	persist.model.mwas	
	cross.validation.mwas	
model.evaluation.R	model.evaluation.mwas	
	predict.knn	S3 method
preprocess_mwas.R	preprocess.mwas	
	filter.pathways	
	get.next.kegg	
visualization.R	plot.mwas	
	plot.differentiated.taxa	
	plot.beeswarm.dt	
	run.scatterplot	not used
	run.scatterplot.all.data	not used
	run.beeswarm	not used
	plot.gradients	
	qqplot.pvals	not used
export_data.R	export.mwas	

	save.xlsx	
	write.differentiation.test.results	
stats.r	differentiation.test	
	linear.test	not used
	t.test.wrapper	not used
roc_mwas.R	roc.mwas	
gradients.R	show.gradients	
	show.metadata	
	get.gradient.ixs	
	best.legend.location	
features.selection.R	feature.scores.rf.mwas	
heatmap.3.R	heatmap.3	
heatmap.mwas.R	cluster.columns	
	cluster.rows	
	shorten.taxonomy	Similar to the one in util.load.R; commented out
	create.color.bars	
	heatmap.mwas	