Package ‘waves’

April 21, 2021

Title Vis-NIR Spectral Analysis Wrapper

Version 0.1.1

Description Originally designed application in the context of resource-limited plant research and breeding programs, ‘waves’ provides an open-source solution to spectral data processing and model development by bringing useful packages together into a streamlined pipeline. This package is wrapper for functions related to the analysis of point visible and near-infrared reflectance measurements. It includes visualization, filtering, aggregation, preprocessing, cross-validation set formation, model training, and prediction functions to enable open-source association of spectral and reference data. This package is documented in a peer-reviewed manuscript in the Plant Phenome Journal <doi:10.1002/ppj2.20012>. Specialized cross-validation schemes are described in detail in Jarquín et al. (2017) <doi:10.3835/plantgenome2016.12.0130>. Example data is from Ikeogu et al. (2017) <doi:10.1371/journal.pone.0188918>.

URL https://github.com/GoreLab/waves

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BugReports https://github.com/GoreLab/waves/issues

Depends R (>= 3.5)

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Encoding UTF-8

LazyData true

Imports dplyr, prospectr, spectacles, caret, pls, randomForest, wesanderson, magrittr, tidyselect, ggplot2, tidyr (>= 1.0), stringr, rlang

RoxygenNote 7.1.1

Suggests testthat (>= 2.1.0)

NeedsCompilation no

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AggregateSpectra

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| AggregateSpectra | Aggregate data based on grouping variables and a user-provided function |

Description

Use grouping variables to collapse spectral data.frame by mean or median. Recommended for use after FilterSpectra.

Usage

AggregateSpectra(df, grouping.colnames, reference.value.colname, agg.function)

Arguments

- df: data.frame object containing one or multiple columns of grouping variables (must be consistent within each group), column of reference values (optional), and columns of spectra. Spectral column names must start with "X".

- grouping.colnames: Names of columns to be used as grouping variables. Minimum 2 variables required. Default is c("trial", "plot").

- reference.value.colname: Name of reference column to be aggregated along with spectra. Default is "reference".

- agg.function: Name of function (string format) to be used for sample aggregation. Must be either "mean" or "median". Default is "mean".

Value

data.frame object df aggregated based on grouping column by agg.function

Author(s)

Jenna Hershberger <jmh579@cornell.edu>
### Examples

```r
library(magrittr)
aggregated.test <- ikeogu.2017 %>%
dplyr::select(-TCC) %>%
ao.omit() %>%
AggregateSpectra(grouping.colnames = c("study.name"),
                   reference.value.colname = "DMC.oven",
                   agg.function = "mean")
aggregated.test[1:5, 1:5]
```

### Description

Preprocessing, also known as pretreatment, is often used to increase the signal to noise ratio in vis-NIR datasets. The `waves` function `DoPreprocessing` applies common spectral preprocessing methods such as standard normal variate and the Savitzky–Golay filter.

### Usage

```r
DoPreprocessing(
  df,  
  test.data = NULL,  
  preprocessing.method = 1,  
  wavelengths = 740:1070
)
```

### Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>df</code></td>
<td>data.frame object containing spectral data. First column(s) (optional) include metadata (with or without reference value column) followed by spectral columns. Spectral column names must be formatted as &quot;X&quot; followed by wavelength Include no other columns to right of spectra! No missing values permitted.</td>
</tr>
<tr>
<td><code>test.data</code></td>
<td>data.frame object with same format as <code>train.data</code>. Will be appended to <code>df</code> during preprocessing so that the same transformations are applied to each row. Default is <code>NULL</code>.</td>
</tr>
<tr>
<td><code>preprocessing.method</code></td>
<td>Number or list of numbers 1:13 corresponding to desired pretreatment method(s):</td>
</tr>
<tr>
<td></td>
<td>• 1 = raw data (default)</td>
</tr>
<tr>
<td></td>
<td>• 2 = standard normal variate (SNV)</td>
</tr>
<tr>
<td></td>
<td>• 3 = SNV and first derivative</td>
</tr>
<tr>
<td></td>
<td>• 4 = SNV and second derivative</td>
</tr>
<tr>
<td></td>
<td>• 5 = first derivative</td>
</tr>
<tr>
<td></td>
<td>• 6 = second derivative</td>
</tr>
<tr>
<td></td>
<td>• 7 = Savitzky–Golay filter (SG)</td>
</tr>
</tbody>
</table>
FilterSpectra

• 8 = SNV and SG
• 9 = gap segment derivative (window size = 11)
• 10 = SG and first derivative (window size = 5)
• 11 = SG and first derivative (window size = 11)
• 12 = SG and second derivative (window size = 5)
• 13 = SG and second derivative (window size = 11)

wavelengths List of wavelengths represented by each column in df. Default is 740:1070.

Value
Preprocessed df' (or list of data.frames) with reference column intact

Author(s)
Jenna Hershberger <jmh579@cornell.edu>

Examples

---

FilterSpectra  Filter spectral data frame based on Mahalanobis distance

Description
Determine Mahalanobis distances of observations (rows) within a given data.frame with spectral data. Option to filter out observations based on these distances.

Usage
FilterSpectra(df, filter, return.distances, num.col.before.spectra, window.size, verbose)

Arguments

  df a data.frame object containing columns of spectra and rows of observations. May also contain columns of metadata to the left of the spectra.

  filter boolean that determines whether or not the input data.frame will be filtered. If TRUE, df will be filtered according to squared Mahalanobis distance with a 95% cutoff from a chi-square distribution with degrees of freedom = number of spectral columns. If FALSE, a column of squared Mahalanobis distances h.distance will be added to the right side of df and all rows will be returned. Default is TRUE.

  return.distances boolean that determines whether a column of squared Mahalanobis distances will be included in output data.frame. If TRUE, a column of Mahalanobis distances for each row will be added to the right side of df. Default is FALSE.
**num.col.before.spectra**

number of columns to the left of the spectral matrix in df. Default is 4.

**window.size**

number defining the size of window to use when calculating the covariance of the spectra (required to calculate Mahalanobis distance). Default is 10.

**verbose**

If TRUE, the number of rows removed through filtering will be printed to the console. Default is TRUE.

**Details**

This function uses a chi-square distribution with 95% cutoff where degrees of freedom = number of wavelengths (columns) in the input data.frame.

**Value**

If filter is TRUE, returns filtered data frame df and reports the number of rows removed. The Mahalanobis distance with a cutoff of 95% of chi-square distribution (degrees of freedom = number of wavelengths) is used as filtering criteria. If filter is FALSE, returns full input df with column h.distances containing the Mahalanobis distance for each row.

**Author(s)**

Jenna Hershberger <jmh579@cornell.edu>

**References**


**Examples**

```r
library(magrittr)
filtered.test <- ikeogu.2017 %>%
dplyr::select(-TCC) %>%
na.omit() %>%
FilterSpectra(df = ., filter = TRUE, return.distances = TRUE, num.col.before.spectra = 5, window.size = 15)
filtered.test[1:5, c(1:5, (ncol(filtered.test)-5):ncol(filtered.test))]
```

**FormatCV**

*Format multiple trials with or without overlapping genotypes into training and test sets according to user-provided cross validation scheme*
**Description**

Standalone function that is also used within `TrainSpectralModel` to divide trials or studies into training and test sets based on overlap in trial environments and genotype entries.

**Usage**

```r
FormatCV(
  trial1,
  trial2,
  trial3 = NULL,
  cv.scheme,
  seed = NULL,
  remove.genotype = FALSE
)
```

**Arguments**

- **trial1**  
  data.frame object that is for use only when `cv.scheme` is provided. Contains the trial to be tested in subsequent model training functions. The first column contains unique identifiers, second contains genotypes, third contains reference values, followed by spectral columns. Include no other columns to right of spectra! Column names of spectra must start with "X", reference column must be named "reference", and genotype column must be named "genotype".

- **trial2**  
  data.frame object that is for use only when `cv.scheme` is provided. This data.frame contains a trial that has overlapping genotypes with `trial1` but that were grown in a different site/year (different environment). Formatting must be consistent with `trial1`.

- **trial3**  
  data.frame object that is for use only when `cv.scheme` is provided. This data.frame contains a trial that may or may not contain genotypes that overlap with `trial1`. Formatting must be consistent with `trial1`.

- **cv.scheme**  
  A cross validation (CV) scheme from Jarquín et al., 2017. Options for cv.scheme include:
  - "CV1": untested lines in tested environments
  - "CV2": tested lines in tested environments
  - "CV0": tested lines in untested environments
  - "CV00": untested lines in untested environments

- **seed**  
  Number used in the function `set.seed()` for reproducible randomization. If `NULL`, no seed is set. Default is `NULL`.

- **remove.genotype**  
  boolean that, if TRUE, removes the "genotype" column is removed from the output data.frame. Default is FALSE.

**Details**

Use of a cross-validation scheme requires a column in the input data.frame named "genotype" to ensure proper sorting of training and test sets. Variables `trial1` and `trial2` are required, while `trial3` is optional.
Value

List of data.frames (training set, test set) compiled according to user-provided cross validation scheme.

Author(s)

Jenna Hershberger <jmh579@cornell.edu>

References


Examples

# Must have a column called "genotype", so we'll create a fake one for now
# We will use CV00, which does not require any overlap in genotypes
# In real scenarios, CV schemes that rely on genotypes should not be applied when
# genotypes are unknown, as in this case.
library(magrittr)
trials <- ikeogu.2017 %>%
  dplyr::mutate(genotype = 1:nrow(ikeogu.2017)) %>% # fake for this example
  dplyr::rename(reference = DMC.oven) %>%
  dplyr::select(study.name, sample.id, genotype, reference,
                dplyr::starts_with("X"))
trial1 <- trials %>%
  dplyr::filter(study.name == "C16Mcal") %>%
  dplyr::select(-study.name)
trial2 <- trials %>%
  dplyr::filter(study.name == "C16Mval") %>%
  dplyr::select(-study.name)
cv.list <- FormatCV(trial1 = trial1, trial2 = trial2, cv.scheme = "CV00",
                    remove.genotype = TRUE)
cv.list[[1]][1:5, 1:5]

ikeogu.2017

Example vis-NIRS and reference dataset

Description

The ‘ikeogu.2017’ data set contains raw vis-NIRS scans, total carotenoid content, and cassava root dry matter content (using the oven method) from the 2017 PLOS One paper by Ikeogu et al. This dataset contains a subset of the original scans and reference values from the supplementary files of the paper. ‘ikeogu.2017’ is a ‘data.frame’ that contains the following columns:

- study.name = Name of the study as described in Ikeogu et al. (2017).
- sample.id = Unique identifier for each individual root sample
• DMC.oven = Cassava root dry matter content, the percentage of dry weight relative to fresh weight of a sample after oven drying.
• TCC = Total carotenoid content (µg/g, unknown whether on a fresh or dry weight basis) as measured by high performance liquid chromatography
• X350:X2500 = spectral reflectance measured with the QualitySpec Trek: S-10016 vis-NIR spectrometer. Each cell represents the mean of 150 scans on a single root at a single wavelength.

Usage
ikeogu.2017

Format
An object of class tbl_df (inherits from tbl, data.frame) with 175 rows and 2155 columns.

Author(s)

References

Examples
library(magrittr)
library(ggplot2)
data(ikeogu.2017)
ikeogu.2017[1:10,1:10] %>%
dplyr::select(-starts_with("X")) %>%
dplyr::group_by(study.name) %>%
tidyr::gather(trait, value, c(DMC.oven:TCC), na.rm = TRUE) %>%
ggplot2::ggplot(aes(x = study.name, y = value, fill = study.name)) +
  facet_wrap(~ trait, scales = 'free_y', nrow = 2) +
geom_boxplot()

---

PlotSpectra

Plot spectral data, highlighting outliers as identified using Mahalanobis distance

Description
Generates a ggplot object of given spectra, with wavelength on the x axis and given spectral values on the y. Mahalanobis distance is used to calculate outliers, which are both identified on the plot. Rows from the original dataframe are printed to the console for each outlier that is identified.
PlotSpectra

Usage

PlotSpectra(
  input.df,
  wavelengths,
  num.col.before.spectra = 1,
  window.size = 10,
  verbose = TRUE
)

Arguments

input.df  data.frame object containing columns of spectra. Spectral columns must be labeled with an "X" and then the wavelength (example: "X740" = 740nm). Leftmost column must be unique ID. May also contain columns of metadata between the unique ID and spectral columns. Cannot contain any missing values

wavelengths  List of wavelengths (numerical format) represented by each spectral column in input.df

num.col.before.spectra  Number of columns to the left of the spectral matrix (including unique ID). Default is 1.

window.size  number defining the size of window to use when calculating the covariance of the spectra (required to calculate Mahalanobis distance). Default is 10.

verbose  If TRUE, the number of rows removed through filtering will be printed to the console. Default is TRUE.

Value

If verbose, prints unique ID and metadata for rows identified as outliers. Returns plot of spectral data with non-outliers in blue and outliers in red. X-axis is wavelengths and y-axis is spectral values.

Author(s)

Jenna Hershberger <jmh579@cornell.edu>

Examples

library(magrittr)
ikeogu.2017 %>%
dplyr::rename(unique.id = sample.id) %>%
dplyr::select(unique.id, dplyr::everything(), -TCC) %>%
na.omit() %>%
PlotSpectra(input.df = .,
  wavelengths = 350:2500,
  num.col.before.spectra = 5,
  window.size = 15)
PredictFromSavedModel  

Use provided model object to predict trait values with input dataset

Description

Loads an existing model and cross-validation performance statistics (created with SaveModel) and makes predictions based on new spectra.

Usage

PredictFromSavedModel(
  input.data,  
  model.stats.location,  
  model.location,  
  wavelengths = 740:1070,  
  model.method = "pls"  
)

Arguments

input.data  
data.frame object of spectral data for input into a spectral prediction model. First column contains unique identifiers followed by spectral columns. Include no other columns to right of spectra! Column names of spectra must start with "X".

model.stats.location  
String containing file path (including file name) to save location of 
"(model.name)_stats.csv" as output from the SaveModel function.

model.location  
String containing file path (including file name) to location where the trained model 
"(model.name).Rds") was saved as output by the SaveModel function.

wavelengths  
List of wavelengths represented by each column in input.data

model.method  
Model type to use for training. Valid options include:
  • "pls": Partial least squares regression (Default)
  • "rf": Random forest
  • "svmLinear": Support vector machine with linear kernel
  • "svmRadial": Support vector machine with radial kernel

Value

data.frame object of predictions for each sample (row). First column is unique identifier supplied by input.data and second is predicted values

Author(s)

Jenna Hershberger <jmh579@cornell.edu>
SaveModel

Examples

```r
## Not run:
ikeogu.2017 %>%
dplyr::select(sample.id, dplyr::starts_with("X")) %>%
PredictFromSavedModel(input.data = .,
  model.stats.location = paste0(getwd(),
                              "/my_model_stats.csv"),
  model.location = paste0(getwd(), "/my_model.Rds"),
  wavelengths = 350:2500)

## End(Not run)
```

SaveModel

Save spectral prediction model and model performance statistics

Description

Saves spectral prediction model and model statistics to `model.save.folder` as `model.name.Rds` and `model.name_stats.csv` respectively

Usage

```r
SaveModel(
  df,
  save.model = TRUE,
  autoselect.preprocessing = TRUE,
  preprocessing.method = NULL,
  model.save.folder = NULL,
  model.name = "PredictionModel",
  best.model.metric = "RMSE",
  tune.length = 50,
  model.method = "pls",
  num.iterations = 10,
  wavelengths = 740:1070,
  stratified.sampling = TRUE,
  cv.scheme = NULL,
  trial1 = NULL,
  trial2 = NULL,
  trial3 = NULL,
  verbose = TRUE
)
```

Arguments

df  
data.frame object. First column contains unique identifiers, second contains reference values, followed by spectral columns. Include no other columns to right of spectra! Column names of spectra must start with "X" and reference column must be named "reference"
save.model If TRUE, the trained model will be saved in .Rds format to the location specified by model.save.folder. If FALSE, the model will be output by function but will not save to file. Default is TRUE.

autoselect.preprocessing Boolean that, if TRUE, will choose the preprocessing method for the saved model using the best.model.metric. If FALSE, the user must supply the preprocessing method (1-12, see DoPreprocessing() documentation for more information) of the saved model. Default is TRUE.

preprocessing.method Number or list of numbers 1:13 corresponding to desired pretreatment method(s):
- 1 = raw data (default)
- 2 = standard normal variate (SNV)
- 3 = SNV and first derivative
- 4 = SNV and second derivative
- 5 = first derivative
- 6 = second derivative
- 7 = Savitzky–Golay filter (SG)
- 8 = SNV and SG
- 9 = gap segment derivative (window size = 11)
- 10 = SG and first derivative (window size = 5)
- 11 = SG and first derivative (window size = 11)
- 12 = SG and second derivative (window size = 5)
- 13 = SG and second derivative (window size = 11)

model.save.folder Path to folder where model will be saved. If not provided, will save to working directory.

model.name Name that model will be saved as in model.save.folder. Default is “PredictionModel”.

best.model.metric Metric used to decide which model is best. Must be either "RMSE" or "Rsquared"

tune.length Number delineating search space for tuning of the PLSR hyperparameter ncomp. Default is 50.

model.method Model type to use for training. Valid options include:
- "pls": Partial least squares regression (Default)
- "rf": Random forest
- "svmLinear": Support vector machine with linear kernel
- "svmRadial": Support vector machine with radial kernel

num.iterations Number of training iterations to perform

wavelengths List of wavelengths represented by each column in df

stratified.sampling If TRUE, training and test sets will be selected using stratified random sampling. This term is only used if test.data == NULL. Default is TRUE.

cv.scheme A cross validation (CV) scheme from Jarquin et al., 2017. Options for cv.scheme include:
SaveModel 13

- "CV1": untested lines in tested environments
- "CV2": tested lines in tested environments
- "CV0": tested lines in untested environments
- "CV00": untested lines in untested environments

**trial1** data.frame object that is for use only when cv.scheme is provided. Contains the trial to be tested in subsequent model training functions. The first column contains unique identifiers, second contains genotypes, third contains reference values, followed by spectral columns. Include no other columns to right of spectra! Column names of spectra must start with "X", reference column must be named "reference", and genotype column must be named "genotype".

**trial2** data.frame object that is for use only when cv.scheme is provided. This data.frame contains a trial that has overlapping genotypes with trial1 but that were grown in a different site/year (different environment). Formatting must be consistent with trial1.

**trial3** data.frame object that is for use only when cv.scheme is provided. This data.frame contains a trial that may or may not contain genotypes that overlap with trial1. Formatting must be consistent with trial1.

**verbose** If TRUE, the number of rows removed through filtering will be printed to the console. Default is TRUE.

### Details

Wrapper that uses `DoPreprocessing`, `FormatCV`, and `TrainSpectralModel` functions.

### Value

List of model stats (in data.frame) and trained model object. Saves both to model.save.folder as well. To use optimally trained model for predictions, use tuned parameters from $bestTune

### Author(s)

Jenna Hershberger <jmh579@cornell.edu>

### Examples

```r
library(magrittr)

test.model <- ikeogu.2017 %>%
  dplyr::filter(study.name == "C10Mcal") %>%
  dplyr::rename(reference = DMC.oven) %>%
  dplyr::select(sample.id, reference, dplyr::starts_with("X")) %>%
  na.omit() %>%
  SaveModel(df = ., save.model = FALSE,
             autoselect.preprocessing = TRUE,
             model.name = "my_prediction_model",
             tune.length = 50, num.iterations = 10,
             wavelengths = 350:2500)

summary(test.model[1])
test.model[2]
```
TestModelPerformance

Test the performance of spectral models

Description

Wrapper that trains models based spectral data to predict reference values and reports model performance statistics

Usage

TestModelPerformance(
  train.data,  # data.frame object of spectral data for input into a spectral prediction model. First column contains unique identifiers, second contains reference values, followed by spectral columns. Include no other columns to right of spectra! Column names of spectra must start with "X" and reference column must be named "reference".
  num.iterations,  # Number of training iterations to perform
  test.data = NULL,  # data.frame with same specifications as df. Use if specific test set is desired for hyperparameter tuning. If NULL, function will automatically train with a stratified sample of 70%. Default is NULL.
  preprocessing = TRUE,  # If TRUE, 12 preprocessing methods will be applied and their performance analyzed. If FALSE, input data is analyzed as is (raw). Default is FALSE.
  wavelengths = 740:1070,  # List of wavelengths represented by each column in train.data
  tune.length = 50,  # Number delineating search space for tuning of the PLSR hyperparameter ncomp. Default is 50.
  model.method = "pls",  # "pls", "plsda", "spls", etc.
  output.summary = TRUE,  # Boolean, if TRUE output model performance summary
  rf.variable.importance = FALSE,  # Boolean, if TRUE output variable importance
  stratified.sampling = TRUE,  # Boolean, if TRUE use stratified sampling
  cv.scheme = NULL,  # data.frame of cross-validation indices
  trial1 = NULL,  # data.frame if you want to split the trial
  trial2 = NULL,  # data.frame if you want to split the trial
  trial3 = NULL,  # data.frame if you want to split the trial
  split.test = FALSE,  # Boolean, if TRUE split test data
  verbose = TRUE  # Boolean, if TRUE print additional information
)

Arguments

test.data  # Number of training iterations to perform

train.data  # data.frame object of spectral data for input into a spectral prediction model. First column contains unique identifiers, second contains reference values, followed by spectral columns. Include no other columns to right of spectra! Column names of spectra must start with "X" and reference column must be named "reference".

num.iterations  # Number of training iterations to perform

test.data  # data.frame with same specifications as df. Use if specific test set is desired for hyperparameter tuning. If NULL, function will automatically train with a stratified sample of 70%. Default is NULL.

preprocessing  # If TRUE, 12 preprocessing methods will be applied and their performance analyzed. If FALSE, input data is analyzed as is (raw). Default is FALSE.

wavelengths  # List of wavelengths represented by each column in train.data

tune.length  # Number delineating search space for tuning of the PLSR hyperparameter ncomp. Default is 50.
model.method  Model type to use for training. Valid options include:

- "pls": Partial least squares regression (Default)
- "rf": Random forest
- "svmLinear": Support vector machine with linear kernel
- "svmRadial": Support vector machine with radial kernel

output.summary  boolean that controls function output.

- If TRUE, a summary df will be output (1st row = means, 2nd row = standard deviations). Default is TRUE.
- If FALSE, entire results data frame will be output

rf.variable.importance  boolean that:

- If TRUE, model.method must be set to "rf". Returns a list with a model performance data.frame and a second data.frame with variable importance values for each wavelength for each training iteration. If return.model is also TRUE, returns list of three elements with trained model first, model performance second, and variable importance last. Dimensions are nrow = num.iterations, ncol = length(wavelengths).
- If FALSE, no variable importance is returned. Default is FALSE.

stratified.sampling  If TRUE, training and test sets will be selected using stratified random sampling. This term is only used if test.data == NULL. Default is TRUE.

cv.scheme  A cross validation (CV) scheme from Jarquín et al., 2017. Options for cv.scheme include:

- "CV1": untested lines in tested environments
- "CV2": tested lines in tested environments
- "CV0": tested lines in untested environments
- "CV00": untested lines in untested environments

trial1  data.frame object that is for use only when cv.scheme is provided. Contains the trial to be tested in subsequent model training functions. The first column contains unique identifiers, second contains genotypes, third contains reference values, followed by spectral columns. Include no other columns to right of spectra! Column names of spectra must start with "X", reference column must be named "reference", and genotype column must be named "genotype".

trial2  data.frame object that is for use only when cv.scheme is provided. This data.frame contains a trial that has overlapping genotypes with trial1 but that were grown in a different site/year (different environment). Formatting must be consistent with trial1.

trial3  data.frame object that is for use only when cv.scheme is provided. This data.frame contains a trial that may or may not contain genotypes that overlap with trial1. Formatting must be consistent with trial1.

split.test  boolean that allows for a fixed training set and a split test set. Example// train model on data from two breeding programs and a stratified subset (70%) of a third and test on the remaining samples (30%) of the third. If FALSE, the entire provided test set test.data will remain as a testing set or if none is provided, 30% of the provided train.data will be used for testing. Default is FALSE.
verbose If TRUE, the number of rows removed through filtering will be printed to the console. Default is TRUE.

Details

Calls `DoPreprocessing`, `FormatCV`, and `TrainSpectralModel` functions.

Value
data.frame with model performance statistics in summary format (2 rows, one with mean and one with standard deviation of all training iterations) or in long format (number of rows = num.iterations).

Note if preprocessing = TRUE, only the first mean of summary statistics for all iterations of training are provided for each technique. Included summary statistics:

- Tuned parameters depending on the model algorithm:
  - **Best.n.comp**, the best number of components
  - **Best.ntree**, the best number of trees in an RF model
  - **Best.mtry**, the best number of variables to include at every decision point in an RF model
- **RMSECV**, the root mean squared error of cross-validation
- **R2cv**, the coefficient of multiple determination of cross-validation for PLSR models
- **RMSEP**, the root mean squared error of prediction
- **R2p**, the squared Pearson’s correlation between predicted and observed test set values
- **RPD**, the ratio of standard deviation of observed test set values to RMSEP
- **RPIQ**, the ratio of performance to interquartile difference
- **CCC**, the concordance correlation coefficient
- **Bias**, the average difference between the predicted and observed values
- **SEP**, the standard error of prediction
- **R2sp**, the squared Spearman’s rank correlation between predicted and observed test set values

Author(s)

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Examples

```r
library(magrittr)
ikeogu.2017 %>%
dplyr::rename(reference = DMC.oven) %>%
dplyr::rename(unique.id = sample.id) %>%
dplyr::select(unique.id, reference, dplyr::starts_with("X")) %>%
na.omit() %>%
TestModelPerformance(train.data = .,
  tune.length = 3,
  num.iterations = 3,
  preprocessing = FALSE,
  wavelengths = 350:2500)
```
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