Package 'HMLET'

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Author Alireza Kazemi
Maintainer Alireza Kazemi <alireza.kzmi@gmail.com></alireza.kzmi@gmail.com>
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AddClusterInfotoData_HMLET ClusterStats_HMLET ComputeSamplingInterval_HMLET ComputeSubjectLevelPermLabels_HMLET

	ComputeTValues_HMLET	4
	CreateTimeBinData_HMLET	5
	ExportDataForGUI_HMLET	6
	FilterClusterstimePoints_HMLET	7
	FindClusters_HMLET	7
	GenerateRandomData_HMLET	8
	PermutationTestDataPrep_HMLET	8
	PermutationTestSingleParticipant_HMLET	9
	PermutationTest_HMLET	10
	PlotNullDistribution_HMLET	10
	PlotTimeSeries_HMLET	11
	Preprocess_CheckData_HMLET	13
	Preprocess_FixSampling_HMLET	14
	Preprocess_Interpolate_HMLET	15
	RelativeViewingProportionLabels_HMLET	16
	RemoveIncompletetimePoints_HMLET	17
	SubjectLevelPermutationTestWithin_HMLET	17
	TrialLevelPermutationTestWithin_HMLET	18
	UniquePermutations_HMLET	18
Index		19

AddClusterInfotoData_HMLET

Add cluster information and stats to the corresponding time points in the initial dataframe.

Description

Add cluster information and stats to the corresponding time points in the initial dataframe.

Usage

AddClusterInfotoData_HMLET(data, clusterInf)

Arguments

data frame with temporal data for permutation tests routine, output from Permu-

tationTestDataPrep_HMLET.R.

clusterInf data frame with cluster information, output from ClusterStats_HMLET or Per-

mutationTest_HMLET

Value

data frame compatible for permutation tests and other HMLET modules

ClusterStats_HMLET 3

ClusterStats_HMLET	Adds cluster statistics to t value dataframe, includes all info for all
	tests.

Description

Adds cluster statistics to t value dataframe, includes all info for all tests.

Usage

```
ClusterStats_HMLET(data, paired = T, detailed = F, threshold_t = NULL)
```

Arguments

data	data frame with temporal data for permutation tests routine, output from PermutationTestDataPrep_HMLET.R.
paired	optional boolean to control T-tests. True is for statistical comparisons in paired mode, False for a two-sample T-Test; defaults to True.
detailed	optional boolean to include cluster statistics and t value statistics within one dataframe. Defaults to False to populate dataframe only with cluster statistics.
threshold_t	optional probability threshold for statistical comparison computed based on number of subjects. alpha = 0.025 .

Value

returns data frame with cluster statistics

```
ComputeSamplingInterval_HMLET

Compute the sampling time interval
```

Description

This function estimates the sampling interval for each trial based on the time differences between each time point and uses mode to find the interval. Note that your dataset has to have a unique consistent time interval between all samples. In case, several time intervals is estimated for your dataset it use "Preprocess_FixSampling_HMLET" function and pass the highest sampling interval to fix the inconsistency.

Usage

```
ComputeSamplingInterval_HMLET(
  data,
  ID = "ID",
  trial,
  timePoint,
  appendFlag = T,
  userCallFlag = T
)
```

Arguments

data long format dataframe containing temporal data.

ID string for column name that represents IDs within data frame, defaults to "ID".

trial string for column name that represents trials within data frame. Can be numeri-

cal or categorical.

timePoint string for column name that represents time stamps of each sample.

appendFlag optional flag specifies whether the interval(s) being appended to the data or re-

turned separately, defaults to TRUE.

userCallFlag optional flag to print out data when users are calling this function, defaults to

TRUE.

ComputeSubjectLevelPermLabels_HMLET

Provides unique permutation indices within each subject.

Description

Provides unique permutation indices within each subject.

Usage

ComputeSubjectLevelPermLabels_HMLET(labels, n = 1)

Arguments

labels labels from the data frame.

n optional integer for number of permutations to resample data.

Value

labels of unique permutation indices to use for subject level permutation test.

 ${\tt Compute t\ values_HMLET} \quad \textit{Compute t\ values for\ permutation\ tests}.$

Description

Compute t values for permutation tests.

Usage

ComputeTValues_HMLET(respTime, paired = TRUE)

Arguments

respTime dataframe for responses in temporal order.

paired optional boolean to control T-tests. True is for statistical comparisons in paired

mode, False for a two-sample T-Test; defaults to True.

Value

t values in the format of a data frame.

CreateTimeBinData_HMLET

Collapse time points in specified time bins.

Description

Collapse time points in specified time bins.

Usage

```
CreateTimeBinData_HMLET(
  data,
  groupingColumns = NULL,
  timeBinWidth = 250,
  timeMax = 3000,
  FixatedOn,
  timePoint = "timeStamp",
  AOIs = NULL,
  timeForward = T,
  aggregateFun = mean
)
```

Arguments

data dataframe containing temporal data.

groupingColumns

optional array of strings for column names of variables to be grouped, defaults

to NULL.

timeBinWidth optional integer specified time interval for time bins, defaults to 250.

timeMax optional integer for maximum time in temporal order, samples are left out if

duration is longer. Defaults to 3000.

Fixated0n string for column name of specific AOI in dataframe to be fixated on.

timePoint optional string representing column name for time Stamps, defaults to "timeS-

tamp".

AOIs optional array of strings representing column names for areas of interest, de-

faults to NULL.

timeForward optional boolean to sort timebins, defaults to True for ascending order.

aggregateFun optional function for aggregation, defaults to mean.

Value

temporal data with sorted time points into time bins.

ExportDataForGUI_HMLET

ExportDataForGUI_HMLET Export samples of trial with maximum time point to CSV file for MATLAB GUI.

Description

ExportDataForGUI_HMLET Export samples of trial with maximum time point to CSV file for MATLAB GUI.

Usage

```
ExportDataForGUI_HMLET(
  data,
  ID = "ID",
  trial = "trial",
  timePoint = "timePoint",
  timeMax = NULL,
  dataPointDuration,
  response,
  fixation,
  condition,
  testName = NULL,
  gazeX,
  gazeY,
  gazeXRelative = NULL,
  gazeYRelative = NULL,
  miscVars = NULL,
  fileName = "HMLET_DataforGUI.csv",
  path = getwd()
)
```

Arguments

data	dataframe containing temporal data.	
ID	optional string for column name that represents IDs within data frame, defaults to "ID." $$	
trial	string for column name that represents trials within data frame, for example "TrialNum."	
timePoint	string for column name that represents time intervals, for example "timeStamp."	
timeMax	optional integer for maximum time in temporal order, samples are left out if duration is longer. Default is NULL	
dataPointDuration		
	optional integer or character for specified duration of samples, used for time bins in CreateTimeBinData_HMLET.	
response	the column in which participants' responses are stored e.g., HIT, Miss, FA, and CR.	
fixation	data for specific AOI in dataframe to be fixated on. #TODO: ASK	
condition	string for column name that specifies a condition within the data frame.	

testName	optional string	for name of	data – used	as condition name	e or test names to com-

pare permutation test results between different tests/conditions later, defaults to

NULL.

gazeX integer X-coordinate of gaze point on the screen according to top left corner. integer Y-coordinate of gaze point on the screen according to top left corner. gazeY optional integer X-coordinate of gaze point relative to an arbitrary center. gazeXRelative optional integer Y-coordinate of gaze point relative to an arbitrary center. gazeYRelative miscVars

optional list of strings containing column names of additional variables to be

included in exported file

fileName optional string for output csv file, defaults to "HMLET_DataforGUI.csv."

optional path to identify the destination location default: current working direcpath

FilterClusterstimePoints_HMLET

not being used.

Description

not being used.

Usage

FilterClusterstimePoints_HMLET(data, clusterInf = clusterInf)

FindClusters_HMLET Cluster all samples whose t value is greater than threshold value by

temporal adjacency.

Description

Cluster all samples whose t value is greater than threshold value by temporal adjacency.

Usage

FindClusters_HMLET(tValues, threshold_t = threshold_t)

Arguments

tValues dataframe of t values calculated from ComputeTValues HMLET.

threshold_t optional probability threshold for statistical comparison, defaults to NA and will

> be computed based on number of trials when "between trials permutation" is called or will be computed based on number of subjects when "between subjects

permutation" is called. Alpha = 0.025.

Value

t value dataframe with appended cluster information.

GenerateRandomData_HMLET

Generate unique random permutations To be completed

Description

Generate unique random permutations To be completed

Usage

```
GenerateRandomData_HMLET(
   tMax = 20,
   effectOffset = 5,
   trialNum = 40,
   subjNum = 20,
   effectSize = 0.2
)
```

PermutationTestDataPrep_HMLET

Return a data frame compatible for permutation tests routine.

Description

Return a data frame compatible for permutation tests routine.

Usage

```
PermutationTestDataPrep_HMLET(
  data,
  ID = "ID",
  trial = NULL,
  timePoint,
  condition,
  gazeMeasure,
  conditionLevels = NULL,
  targetAOI = NULL,
  testName = NULL,
  timeBinName = NULL
)
```

Arguments

data frame with temporal data for permutation tests routine.

ID optional string for column name that represents IDs within data frame, defaults

to "ID".

trial string for column name that represents trials within data frame.

timePoint string for column name that includes time point or index values in numeric form.

condition string for column name that specifies a condition within the data frame.

gazeMeasure string for column name which includes numerical value of the gaze measure at

each time point such as gazeproportion.

conditionLevels

optional array of string column names to denote all the values of conditions,

defaults to NULL.

targetAOI optional string specify the name of the target AOI which should be set as 1 in

the gazeMeasure variable, defaults to NULL.

testName optional string for name of data – used as condition name or test names to com-

pare permutation test results between different tests/conditions later, defaults to

NULL.

timeBinName optional string for the column name that includes time bin names, defaults to

NULL.

Value

data frame compatible for permutation tests and other HMLET modules

PermutationTestSingleParticipant_HMLET

Conduct trial level permutation test for a single participant.

Description

Conduct trial level permutation test for a single participant.

Usage

PermutationTestSingleParticipant_HMLET(data, samples = 2000, threshold_t = NA)

Arguments

data frame with temporal data for permutation tests routine, output from Permu-

tationTestDataPrep_HMLET.R.

samples optional integer for number of samples, defaults to 2000.

threshold_t optional probability threshold for statistical comparison, defaults to NA and will

be computed based on number of trials when "between trials permutation" is called or will be computed based on number of subjects when "between subjects

permutation" is called. Alpha = 0.025.

Value

a list of permutation tests in which distribution of desired statistic under the null hypothesis is estimated in a large number of permutations of the original data

PermutationTest_HMLET Permutation Tests General Routine.

Description

Permutation Tests General Routine.

Usage

```
PermutationTest_HMLET(
  data,
  samples = 2000,
  paired = T,
  permuteTrialsWithinSubject = F,
  threshold_t = NA
)
```

Arguments

data frame with temporal data for permutation tests routine, output from Permu-

tationTestDataPrep_HMLET.R.

samples optional integer for number of samples, defaults to 2000.

paired optional boolean to control T-tests. True is for statistical comparisons in paired

mode, False for a two-sample T-Test; defaults to True.

permuteTrialsWithinSubject

optional boolean to run permutation test pipeline for conducting permutation

tests between trials or between subjects, defaults to False.

threshold_t optional probability threshold for statistical comparison, defaults to NA and will

be computed based on number of trials when "between trials permutation" is called or will be computed based on number of subjects when "between subjects

permutation" is called. Alpha = 0.025.

Value

a list of permutation tests in which distribution of desired statistic under the null hypothesis is estimated in a large number of permutations of the original data

PlotNullDistribution_HMLET

Plot Null Distribution.

Description

Plot Null Distribution.

Usage

PlotNullDistribution_HMLET(resultList, smoothingBandWidth = NULL)

Arguments

resultList list of the dataframes that is already prepared by PrepareMLETData_HMLET or a list that is the result of PermutationTest_HMLET. smoothingBandWidth

The smoothing bandwidth which determines the granularity of distribution estimation, defaults to sd/3

Value

null distribution plot handle.

PlotTimeSeries_HMLET Plot temporal trends for time series.

Description

Plot temporal trends for time series.

Usage

```
PlotTimeSeries_HMLET(
  resultList,
  showDataPointProp = T,
  showOverallMean = "none",
  gazePropRibbonAlpha = 0.1,
  clusterFillColor = "#CC9933",
  clusterFillAlpha = 0.5,
  pointSize = 1,
  pointAlpha = 0.7,
  pointFatten = 3,
  testNameOrder = NULL,
  conditionOrder = NULL,
  onlySignificantClusters = T,
  clusterData = NULL,
  yLabel = "Gaze Proportion",
  dataAxisLabel = "Available Observations (%)",
  xLabel = "Time (ms)",
  lineWidthGazeProp = 1,
  lineWidthDataPointProp = 1,
  lineWidthOverallMean = 1,
  lineTypeOverallMean = "dotted",
  alphaOverallMean = 0.5,
  shapeCodeOverallMean = 1,
  tickSizeOverallMean = NULL
)
```

Arguments

resultList dataframe of the data that is already prepared by PrepareMLETData_HMLET or a list that is the result of PermutationTest_HMLET.

showDataPointProp

optional boolean to display data points on the axes, defaults to True.

showOverallMean

optional string to specify the overall mean. "Line" = to display a horizontal line on the y = average. "Point" to display overall average and range in a pointRange format. "none" to not display it, defaults to "none".

gazePropRibbonAlpha

optional float to change opacity of gaze ribbon, defaults to 0.1.

clusterFillColor

optional string to change fill color of clustered data points, defaults to "#CC9933".

clusterFillAlpha

optional float to change opacity of clustered data points, defaults to 0.5.

pointSize optional integer to change size of points on plot, defaults to 1.

pointAlpha optional float to change opacity of points on plot, defaults to 0.7.

pointFatten optional integer to change size of lines on plot, defaults to 3.

testNameOrder specify order of test name labels, defaults to NULL.

conditionOrder specify order of condition levels, defaults to NULL.

only Significant Clusters

optional Boolean to plot only the significant clusters, defaults to True.

clusterData data containing all clusters generated from ClusterStats_HMLET to display on

plot, defaults to NULL.

yLabel Proportion" specify label for the y axis, defaults to "Gaze"

dataAxisLabel Specify label for the second y axis on the right, defaults to "Data Points \((%\)\".

xLabel Specify label for the x axis, defaults to "Time $\mbox{\mbox{$\mbox{$\mbox{$w$}$}}}$ "

lineWidthGazeProp

Specify line width for gaze proportion, defaults to 1.

lineWidthDataPointProp

Specify line width for data point numbers, defaults to 1.

lineWidthOverallMean

Specify line width for overall mean, defaults to 1.

lineTypeOverallMean

Specify line type for overall mean, defaults to "dotted".

alphaOverallMean

Specify alpha for overall mean, defaults to 0.5.

shapeCodeOverallMean

Specify the shape of the overall mean points when showOverallMean = "Point", defaults to $1 \cdot (circle)$.

tickSizeOverallMean

Optional to specify where to plot the overall means when showOverallMean = "Point", defaults to NULL.

Value

a plot handle that visualizes the data from PrepareMLETData_HMLET or the list from Permuation-Test_HMLET.

```
Preprocess_CheckData_HMLET
```

Check Data for Critical Issues

Description

Raw eye-tracking data comes with time stamps of each sample and Gaze Coordinates. These time stamps has to have consistent and unique time interval across all datapoints. Gaze Coordinates may have missing values due to blinks, head movement, etc. This function reports issues about inconsistent sampling intervals and missing Gaze coordinates.

Usage

```
Preprocess_CheckData_HMLET(
  data,
  ID = "ID",
  trial = "trial",
  timePoint = "timePoint",
  GazeX,
  GazeY,
  groupingColumns = NULL,
  missRateCap = 50,
  shortTrialsThreshold = 2
)
```

Arguments

data	long format dataframe	containing temporal data.

ID string for column name that represents IDs within data frame, defaults to "ID".

trial string for column name that identify unique trials within data frame. Can be

numerical or categorical.

timePoint string for column name that represents time stamps of each sample.

GazeX string for column name that includes X coordinate of gaze points

GazeY string for column name that includes Y coordinate of gaze points

groupingColumns

optional array of strings for column names of variables to be grouped for the

final miss Rate Summary report, defaults to NULL.

missRateCap Specify the percentage of missing gaze points cap to be counted as high Miss

Rate

shortTrialsThreshold

Specify the minimum number of samples that should be available within a trial to be counted as a valid trial. defaults to 2

Details

This function returns a dataframe with summary of missing gaze points The summary dataframe has 4 columns: ID(with the original column name): indicate participant's ID nTrials: number of trials nTrialsMoreThanCap: number of trials in which, more than missRateCap (defaults to 50 averageMissRate: Average percentage of missing gaze points across all trials of each participant.

Data Cleaning Note: You may want to remove trials with more than missRateCap (defaults to 50 participants that most of their trials has high rate of missing gazepoints.

```
Preprocess_FixSampling_HMLET
Fix Sampling Inconsistencies
```

Description

Raw eye-tracking data comes with time stamps of each sample. These time stamps has to have consistent and unique time interval across all datapoints. If across participants there are different sampling intervals this function downsample all datapoint to the biggest sampling interval. This function also relabel all time stamps with a consistent time stamps to be comparable across trials and participants. If there is a gap between samples of a trial this function fill in the gap with NA values for Gaze coordinates. difference between each two consecutive samples within a trial. Note that your dataset has to have a unique consistent time interval between all samples. In case, several time intervals is estimated for your dataset it use "Preprocess_FixSampling_HMLET" function and pass the highest sampling interval to fix the inconsistency.

Usage

```
Preprocess_FixSampling_HMLET(
   data,
   ID = "ID",
   trial,
   timePoint,
   GazeX,
   GazeY,
   samplingInterval = NULL,
   fillGenratedRows = T,
   ignoreColumns = NULL,
   removeShortTrials = T,
   shortTrialsThreshold = 2
)
```

Arguments

data long format dataframe containing temporal data.

ID string for column name that represents IDs within data frame, defaults to "ID".

trial string for column name that identify unique trials within data frame. Can be

numerical or categorical.

timePoint string for column name that represents time stamps of each sample.

GazeX string for column name that includes X coordinate of gazepoints

GazeY string for column name that includes Y coordinate of gazepoints

samplingInterval

specifies working directory of project.

fillGenratedRows

Optional flag to specify whether the new rows generated to fill temporal jumps should be filled or not. This function uses a 'down-up' order (each value filled with its preceding value; otherwise, with its following value) to fill these columns. Defaults to TRUE.

 $ignore {\tt Columns} \quad Optional\ string (s)\ to\ specify\ columns\ that\ shouldn't\ be\ filled,\ defaults\ to\ NULL.$ $remove {\tt ShortTrials}$

Optional flag to specify whether short trials has to be removed or not, defaults to T.

shortTrialsThreshold

Specify the minimum number of samples that should be available within a trial to be counted as a valid trial. defaults to 2

Details

The output is the same dataframe as input with three new columns: "time": includes new time stamps restarting to (0+sampling interval) in each trial "sampleIdx": includes index for each sample restarting to 1 in each trial "interval": includes the unique sampling interval which is equal to the "time" Note: If your dataframe has columns with same names make sure to rename them before using this function.

Preprocess_Interpolate_HMLET

Interpolate missing Gaze Points

Description

Gaze Coordinates may have missing values due to blinks, head movement, etc. This function fill the NA Gaze points with linear interpolation. You can specify how many consecutive NAs should be interpolated by passing maxNAToFill parameter.

Usage

```
Preprocess_Interpolate_HMLET(
  data,
  ID = "ID",
  trial,
  timePoint,
  GazeX,
  GazeY,
  maxNAToFill = Inf
)
```

Arguments

data	long format dataframe containing temporal data.
ID	string for column name that represents IDs within data frame, defaults to "ID".
trial	string for column name that identify unique trials within data frame. Can be numerical or categorical.
timePoint	string for column name that represents time stamps of each sample.
GazeX	string for column name that includes X coordinate of gazepoints
GazeY	string for column name that includes Y coordinate of gazepoints
maxNAToFill	numeric value to denote the max number of consecutive NAs that should be interpolated, defaults to Inf

Details

The output is the same dataframe as input with three new columns: "GazeX_Interpolated": includes all original GazeX coordinates and interpolated values "GazeY_Interpolated": includes all original GazeY coordinates and interpolated values "Interpolated": is a flag that is 1 for samples that are interpolated and 0 otherwise. Note: If your dataframe has columns with same names make sure to rename them before using this function.

 $Relative Viewing Proportion Labels_HMLET$

Label trials whether participant was disproportionately or equally looking toward items

Description

Label trials whether participant was disproportionately or equally looking toward items

Usage

```
RelativeViewingProportionLabels_HMLET(
  data,
  ID = "ID"
  trial,
  AOIName,
  comparingAOI = c("Target", "Lure"),
  targetAOI = "Target",
  dispropotionCriterion = 10
)
```

Arguments

data data frame with temporal data for permutation tests routine.

ID optional string for column name that represents IDs within data frame, defaults

trial string for column name that represents trials within data frame.

string for column name which includes name of the AOI that the subject is look-**AOIName**

ing at each timePoint.

comparingAOI names of the two AOIs that are their looking proportion should be compared. names of the two AOIs that are their looking proportion should be compared.

dispropotionCriterion

targetA0I

optional string for column name to focus on specific AOI, defaults to NULL.

testName Name of this data – can be used as condition name or test names to compare

permutation test results between different tests/conditions later

Value

data frame as original data with an extracolumn with labels

RemoveIncompletetimePoints_HMLET

Remove Time points with missing conditions. This function is currently only applicable for within participant manipulations.

Description

Remove Time points with missing conditions. This function is currently only applicable for within participant manipulations.

Usage

RemoveIncompletetimePoints_HMLET(data)

Arguments

data dataframe containing temporal data.

Value

data with no missing conditions.

SubjectLevelPermutationTestWithin_HMLET

Conduct subject level permutation tests between trials.

Description

Conduct subject level permutation tests between trials.

Usage

```
SubjectLevelPermutationTestWithin_HMLET(
  data,
  samples = 2000,
  paired = T,
  threshold_t = NULL
)
```

Arguments

data frame with temporal data for permutation tests routine, output from Permu-

tationTestDataPrep_HMLET.R.

samples optional number of resampling data, how many permutations needed.

paired optional boolean to control T-tests. True is for statistical comparisons in paired

mode, False for a two-sample T-Test; defaults to True.

threshold_t optional probability threshold for statistical comparison computed based on num-

ber of subjects. alpha = 0.025.

Value

returns t value distribution presented in a data frame.

TrialLevelPermutationTestWithin_HMLET

Conduct trial level permutation tests between trials.

Description

Conduct trial level permutation tests between trials.

Usage

```
TrialLevelPermutationTestWithin_HMLET(
  data,
  samples = 2000,
  paired = T,
  threshold_t = NULL
)
```

Arguments

data frame with temporal data for permutation tests routine, output from Permu-

tationTestDataPrep_HMLET.R.

samples optional number of resampling data, how many permutations needed.

paired optional boolean to control T-tests. True is for statistical comparisons in paired

mode, False for a two-sample T-Test; defaults to True.

threshold_t optional probability threshold for statistical comparison computed based on num-

ber of trials. alpha = 0.025.

Value

returns t value distribution presented in a data frame.

UniquePermutations_HMLET

Generate unique random permutations.

Description

Generate unique random permutations.

Usage

```
UniquePermutations_HMLET(listInput, uniqueLabels, n = 1)
```

Arguments

listInput Arbitrary list.
uniqueLabels All possible Labels.

n An integer number indicating the number of permutations.

Index

```
AddClusterInfotoData_HMLET, 2
ClusterStats_HMLET, 3
ComputeSamplingInterval_HMLET, 3
ComputeSubjectLevelPermLabels_HMLET, 4
{\tt ComputeTValues\_HMLET, 4}
CreateTimeBinData_HMLET, 5
ExportDataForGUI_HMLET, 6
FilterClusterstimePoints_HMLET, 7
FindClusters_HMLET, 7
{\tt GenerateRandomData\_HMLET,\,8}
{\tt PermutationTest\_HMLET},\, 10
PermutationTestDataPrep_HMLET, 8
PermutationTestSingleParticipant_HMLET,
{\tt PlotNullDistribution\_HMLET}, 10
PlotTimeSeries_HMLET, 11
Preprocess_CheckData_HMLET, 13
Preprocess_FixSampling_HMLET, 14
Preprocess_Interpolate_HMLET, 15
Relative Viewing Proportion Labels\_HMLET,
RemoveIncompletetimePoints_HMLET, 17
{\tt SubjectLevelPermutationTestWithin\_HMLET},
{\tt TrialLevelPermutationTestWithin\_HMLET},
UniquePermutations_HMLET, 18
```