# Package 'GridOptimizer'

## January 17, 2025

| Title Op  | otimize Randomized Sample Placement in Grid-like spaces   |
|---|---|
| Version   | 0.0.1.1   |
| ce  | cion This script generates ``grids" of samples under specific conditions while resolving adjacet conflicts and calculating entropy metrics. The primary purpose is to optimize same placement in experimental setups to balance diversity and randomness. |
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|   | calculate_box_metrics   |

2 calculate\_box\_metrics

```
      is_adjacent
      6

      resolve_conflicts
      7

      scoreGrids
      7

      validate_conflicts
      8

Index
```

calculate\_box\_metrics Calculate Metrics for Multiple Boxes

## Description

Computes aggregated metrics (entropy, unique genotypes, Simpson's Diversity Index, randomness percentage) across multiple boxes.

#### Usage

```
calculate_box_metrics(boxes, genotypes)
```

#### **Arguments**

boxes A list of boxes (matrices) to analyze.
genotypes A vector of all possible genotypes.

#### Value

A list containing average and standard deviation of entropy, genotype counts, and Simpson's Diversity Index.

## Examples

```
boxes <- list(
   list(box = matrix(c("cond1_G1_1", "cond1_G2_1", NA, NA), nrow = 2, byrow = TRUE)),
   list(box = matrix(c("cond1_G1_2", "cond1_G3_1", NA, NA), nrow = 2, byrow = TRUE))
)
genotypes <- c("G1", "G2", "G3")
calculate_box_metrics(boxes, genotypes)</pre>
```

compute\_box\_metrics 3

#### **Description**

Computes entropy, unique genotype count, and Simpson's Diversity Index for a box.

#### Usage

```
compute_box_metrics(box)
```

#### **Arguments**

box

A matrix representing the box.

#### Value

A list containing entropy, unique genotype count, and Simpson's Diversity Index.

## **Examples**

```
box <- matrix(c("cond1_G1_1", "cond1_G2_1", "cond1_G1_2", NA), nrow = 2, ncol = 2)
compute_box_metrics(box)</pre>
```

exportGrid

Export Box Results

#### **Description**

Exports box results to Excel and optionally generates PDF for visualization.

## Usage

```
exportGrid(experiment, boxes, file_name = "experiment_output")
```

## Arguments

experiment The experiment number to export.

boxes A list of box generation results.

file\_name Base name for output files.

#### Value

Generates an Excel file with the results and optionally a PDF file.

4 generateGrids

#### **Examples**

```
samples <- data.frame(Genotype = c("G1", "G2", "G3"), Reps = c(2, 2, 1))
conditions <- c("cond1", "cond2")
boxes <- generateGrids(samples, "Genotype", "Reps", conditions, 2, 2, 10, 1)
temp_file <- file.path(tempdir(), "experiment_results")
exportGrid(1, boxes, file_name = temp_file)</pre>
```

generateGrids

Generate Boxes

#### Description

Generates boxes of genotypic samples based on given conditions and parameters.

#### Usage

```
generateGrids(
  samples,
  sample_labels,
  reps,
  conditions,
  gridRows,
  gridCols,
  nIterations,
  nCores,
  availableGrids = NULL
)
```

#### **Arguments**

samples Data frame with genotype and replicate information.

sample\_labels Column name for genotype data.

reps Column name for replicate data.

conditions Experimental conditions to consider.

gridRows Number of rows in each box.
gridCols Number of columns in each box.

nIterations Number of iterations to attempt box generation.

nCores Number of cores for parallel execution.

availableGrids Optional; maximum number of grids allowed.

#### Value

A list of generated boxes with conflict resolution applied.

generateMaps 5

#### **Examples**

```
samples <- data.frame(Genotype = c("G1", "G2", "G3"), Reps = c(2, 2, 1)) conditions <- c("cond1", "cond2") boxes <- generateGrids(samples, "Genotype", "Reps", conditions, 2, 2, 10, 1)
```

generateMaps

Generate Maps from Updated Sample Placement

## Description

This function reads a modified "Sample Placement" sheet and generates updated PDF maps for each box. It processes the user's modifications to the box layout and creates visualizations for all the updated boxes.

## Usage

```
generateMaps(
  data,
  samples,
  rep,
  cond,
  box,
  row,
  col,
  file_name = "updated_maps",
  font_size = NULL
)
```

#### **Arguments**

| data      | A data frame (e.g., read from the "Sample Placement" sheet). |
|-----------|--|
| samples   | The column name containing sample names.                     |
| rep       | The column name containing replicate numbers.                |
| cond      | The column name containing condition labels.                 |
| box       | The column name containing box identifiers.                  |
| row       | The column name containing row numbers.                      |
| col       | The column name containing column numbers.                   |
| file_name | The base name for the output PDF file.                       |
| font_size | Optional font size for the labels on the maps.               |

### Value

A PDF file with visualizations for all boxes.

6 is\_adjacent

#### **Examples**

```
mock_data <- data.frame(</pre>
Sample = c("G1", "G2", "G3", "G4", "G5"),
Replicate = c(1, 2, 1, 3, 1),
Condition = c("cond1", "cond1", "cond2", "cond2", "cond3"),
Box = c(1, 1, 1, 2, 2),
Row = c(1, 2, 3, 1, 2),
Column = c(1, 2, 3, 1, 2)
temp_file <- file.path(tempdir(), "updated_map")</pre>
generateMaps(
  data = mock_data,
  samples = "Sample",
  rep = "Replicate",
  cond = "Condition",
  box = "Box",
  row = "Row",
  col = "Column",
  file_name = temp_file
```

is\_adjacent

Check for Adjacency Conflicts

### **Description**

Checks if a specific cell in a box matrix has adjacency conflicts based on the given genotype label.

#### Usage

```
is_adjacent(box, row, col, label)
```

#### **Arguments**

box A matrix representing the box.

row Row index of the cell to check.

col Column index of the cell to check.

label The genotype label in the cell.

#### Value

Logical value: TRUE if there is a conflict, FALSE otherwise.

resolve\_conflicts 7

#### **Examples**

```
box <- matrix(NA, nrow = 5, ncol = 5)
box[1, 1] <- "cond1_G1_1"
is_adjacent(box, 1, 2, "cond1_G1_2")</pre>
```

resolve\_conflicts

Resolve Adjacency Conflicts

#### **Description**

Resolves adjacency conflicts in a box by swapping labels between cells.

## Usage

```
resolve_conflicts(box, max_iterations = 1000)
```

## Arguments

box

A matrix representing the box.

 ${\tt max\_iterations}$  Maximum number of iterations to attempt resolution.

#### Value

A matrix with resolved conflicts.

## **Examples**

```
box <- matrix(c(NA, "cond1_G1_1", "cond1_G1_2", NA), nrow = 2, ncol = 2)
resolve_conflicts(box, max_iterations = 100)</pre>
```

scoreGrids

Score Boxes

## Description

Evaluates generated boxes and computes metrics for each experiment.

## Usage

```
scoreGrids(box_results)
```

#### **Arguments**

box\_results

A list of box generation results.

8 validate\_conflicts

#### Value

A data frame with metrics for each experiment, including entropy, unique genotype counts, and Simpson's Diversity Index.

## **Examples**

```
box_results <- list(list(
  iteration = 1,
  boxes = list(
    cond1 = list(
      list(box = matrix(c("cond1_G1_1", "cond1_G2_1", NA, NA), nrow = 2, ncol = 2))
    )
    )
)
scoreGrids(box_results)</pre>
```

validate\_conflicts

Validate Adjacency Conflicts

## Description

Validates if any adjacency conflicts exist in a box.

## Usage

```
validate_conflicts(box)
```

#### **Arguments**

box

A matrix representing the box.

#### Value

Logical value: TRUE if no conflicts exist, FALSE otherwise.

#### **Examples**

```
box <- matrix(c(NA, "cond1_G1_1", "cond1_G1_2", NA), nrow = 2, ncol = 2) validate_conflicts(box)
```

## **Index**

```
calculate_box_metrics, 2
compute_box_metrics, 3

exportGrid, 3

generateGrids, 4
generateMaps, 5

is_adjacent, 6

resolve_conflicts, 7

scoreGrids, 7

validate_conflicts, 8
```