# Package 'GenTag'

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Type Package

Title Generate Color Tag Sequences

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Author Carlos Biagolini-Jr.
Maintainer Carlos Biagolini-Jr. <c.biagolini@gmail.com></c.biagolini@gmail.com>
<b>Description</b> Implement a coherent and flexible protocol for animal color tagging. 'GenTag' provides a simple computational routine with low CPU usage to create color sequences for animal tag. First, a single-color tag sequence is created from an algorithm selected by the user, followed by verification of the combination uniqueness. Three methods to produce color tag sequences are provided. Users can modify the main function core to allow a wide range of applications.
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GenTag-package Generate Color Tag Sequences

### Description

Implement a coherent and flexible protocol for animal color tagging. 'GenTag' provides a simple computational routine with low CPU usage to create color sequences for animal tag. First, a single-color tag sequence is created from an algorithm selected by the user, followed by verification of the combination uniqueness. Three methods to produce color tag sequences are provided. Users can modify the main function core to allow a wide range of applications.

### **Details**

#### The DESCRIPTION file:

Package: GenTag Type: Package

Title: Generate Color Tag Sequences

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Description: Implement a coherent and flexible protocol for animal color tagging. 'GenTag' provides a simple computational

License: GPL (>= 2)

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pre\_used Pre-used combinations combinations

scy Summary color year

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Implement a coherent and flexible protocol for animal color tagging. 'GenTag' provides a simple computational routine with low CPU usage to create color sequences for animal tag. First, a single-color tag sequence is created from an algorithm selected by the user, followed by verification of the combination uniqueness. Three methods to produce color tag sequences are provided. Users can modify the main function core to allow a wide range of applications.

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#### Author(s)

Carlos Biagolini-Jr.

Maintainer: Carlos Biagolini-Jr.<c.biagolini@gmail.com>

#### References

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```
genseq(30, 4,c("Black","Blue","Brown","Gray","Green"))
```

4 allequal

	allequal	All equal tag sequence sample	
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#### **Description**

Create combinations with equal sample probability to all colors.

### Usage

```
allequal(ntag, colorsname, nspecial = 0, name1 = "Metal", name2 = "EMPTY", location1 = 1,
location2 = 2, nspecial1 = 1, nspecial2 = 1)
```

## Arguments

ntag	Number of tag to be used in each animal.
colorsname	Names/Code of color tags to be sample.
nspecial	Number of special tags/codes, such as metallic, "EMPTY", or flag (min 0, max 2).
name1	Name of special tag 1.
name2	Name of special tag 2.
location1	Position (or group of positions) to special band 1.
location2	Position (or group of positions) to special band 2.
nspecial1	Number of special tag 1 that will be present in all sequences genetated.
nspecial2	Number of special tag 2 that will be present in all sequences genetated.

### Value

A sequencie of tags

#### Author(s)

Biagolini-Jr.

#### See Also

genseq vfrequency lifexp

```
# Create an object contain the name/code of tag colors
tcol<-c("Black","Blue","Brown","Gray","Green","Pink","Purple","Red","White","Yellow")
# Generate color tag combination without especial tags
genseq(30, 4, colorsname= tcol)
#Generate color tag combination with especial color (ex metallic tag for numeric identification)
genseq(30, 4, tcol, nspecial=1, name1="Metal",location1=c(2,4))
# For ongoing works, use the argument usedcombinations to informe the previus used combinations
data(pre_used) # Data example
genseq(100, 4, tcol, usedcombinations=pre_used[,1:4])</pre>
```

erc 5

erc

Estimates remaining color

## Description

Estimates number of remaining color tags in the field

### Usage

```
erc(usedcombinations, yearusedcombinations, currentyear = NA,
  yearsurvival = NA, lifespan = NA, hide_color = NA)
```

# Arguments

usedcombinations

Pre used combinations

yearusedcombinations

The year in which the combination was used.

currentyear Current year.

yearsurvival An estimation of the proportion of animals that survive between years.

lifespan Combinations older the lifespan will be automatically disregard.

hide\_color Color(s) to be hide in the estimation of remain colors tags present in nature.

#### Value

A estimation of the number of remaining color tags in the field.

### Author(s)

Biagolini-Jr.

#### See Also

scy

```
## The function is currently defined as
data(pre_used) # Data example
erc(pre_used[,1:4],pre_used[,5],2019,0.85, hide_color="EMPTY")
```

6 escombination

escode

Empty synonym code

## Description

Find synonyms in a group of colors with code for empty tag.

#### Usage

```
escode(row_under_review, emptyname = NA, columns_set = NA)
```

### **Arguments**

row\_under\_review

Color sequence to be analyzed.

emptyname

Code used to define empty code.

columns\_set

Range of codes which belong to the same tag region.

#### Value

Matrix with all synonym

### **Examples**

```
# Example of a full sequence
combination<- c("EMPTY","Red","Yellow","Red","Blue","Green")
# See synonym for the code group 1 to 3
escode(combination,"EMPTY",columns_set=1:3)</pre>
```

escombination

Empty synonym combination

## Description

Find synonyms of a sequence with code for empty tag.

### Usage

```
escombination(row_under_review, emptyname, g1 = NA, g2 = NA, g3 = NA, g4 = NA, g5 = NA, g6 = NA)
```

esdataset 7

### **Arguments**

row_under_revie	ew .
	Color sequence to be analyzed.
emptyname	Code used to define empty code.
g1	Group of tags from group 1.
g2	Group of tags from group 2.
g3	Group of tags from group 3.
g4	Group of tags from group 4.
g5	Group of tags from group 5.
g6	Group of tags from group 6.

## Value

Matrix with all synonyms

## Author(s)

Biagolini-Jr

### See Also

escode esdataset

## **Examples**

```
# Example of a full sequence
combination<- c("EMPTY","Red","Yellow","Red","Blue","Green") # See synonym for the full sequence
escombination(combination,"EMPTY",g1=1:3,g2=4:6)</pre>
```

## Description

Find synonyms in a dataset contain combination with code for empty tag.

# Usage

```
esdataset(tag_sheet, emptyname, g1 = NA, g2 = NA, g3 = NA, g4 = NA, g5 = NA, g6 = NA)
```

8 genseq

## Arguments

tag_sheet	Dataset contain sequences to be analyzed.
emptyname	Code used to define empty code.
g1	Group of tags from group 1. For example, in a code of 6 colors for bird tag, tags from 1 to 3 belongs to the left leg, and tags from 1 to 3 belongs to the right. g1 must to address the left leg, and g2 must to address right leg. Thus: "g1=1:3" and "g2=4:6".
g2	Group of tags from group 2. For example, in a code of 6 colors for bird tag, tags from 1 to 3 belongs to the left leg, and tags from 1 to 3 belongs to the right. g1 must to address the left leg, and g2 must to address right leg. Thus: "g1=1:3" and "g2=4:6".
g3	Group of tags from group 3.
g4	Group of tags from group 4.
g5	Group of tags from group 5.
g6	Group of tags from group 6.

### Value

Matrix with all synonyms for from entire dataset

# Author(s)

Biagolini-Jr.

## See Also

escode escombination

# Examples

```
(combination<- matrix(c("EMPTY","Red","Blue","Green"),4,6,TRUE)) \# Example of a dataset esdataset(combination,"EMPTY",g1=1:3,g2=4:6)
```

genseq	Color tag combination generator	

# Description

Create sequences of color tags for animal identification.

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#### Usage

```
genseq(ncombinations = 100, ntag = 4, colorsname, gen_method = "allequal", usedcombinations = NA, colorsf = NA, nspecial = 0, name1 = "Metal", name2 = "EMPTY", location1 = 1, location2 = 2, nspecial1 = 1, nspecial2 = 1, emptyused = FALSE, emptyname = "EMPTY", currentyear = NA, yearsurvival = 1, lifespan = NA, iotf = FALSE, yearusedcombinations = NA, speed = 1, ignorecolor = NA, g1 = NA, g2 = NA, g3 = NA, g4 = NA, g5 = NA, g6 = NA, parameterslist = NA)
```

#### **Arguments**

ncombinations Number of combinations to be generated.

ntag Number of tag to be used in each animal.

colorsname Names/Code of color tags to be sample.

gen\_method method used for sample colors for tag sequence.

usedcombinations

Pre used combinations.

colorsf Frequencies/ratio for color sample.

nspecial Number of special tags/codes, such as metallic, "EMPTY", or flag (min 0, max

2).

name1 Name of special tag 1.

Name of special tag 2.

location1 Position (or group of positions) to special band 1.
location2 Position (or group of positions) to special band 2.

nspecial 1 Number of special tag 1 that will be present in all sequences genetated.

Number of special tag 2 that will be present in all sequences genetated.

Prepresent in all sequences genetated.

If pre-used combination has code for empty set as TRUE, otherwise FALSE

emptyname Code used to define empty code.

currentyear Current year.

yearsurvival An estimation of the proportion of animals that survive between years.

lifespan Combinations older the lifespan will be automatically disregard.

iotf Ignore older than lifespan. If TRUE ignore pre-used combinations older than

lifespan

yearusedcombinations

The year in which the combination was used.

speed Speed for color frequency adjustment.

ignorecolor Color to be ignored on lifexp.

g1 Group of tags from group 1. For example, in a code of 6 colors for bird tag, tags

from 1 to 3 belongs to the left leg, and tags from 1 to 3 belongs to the right. g1 must to address the left leg, and g2 must to address right leg. Thus: "g1=1:3"

and "g2=4:6".

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g2	Group of tags from group 2. For example, in a code of 6 colors for bird tag, tags
	from 1 to 3 belongs to the left leg, and tags from 1 to 3 belongs to the right. g1
	must to address the left leg, and g2 must to address right leg. Thus: "g1=1:3"
	and "g2=4:6".
g3	Group of tags from group 3.
g4	Group of tags from group 4.
g5	Group of tags from group 5.
g6	Group of tags from group 6.
parameterslist	parameters for methods not provide by 'GenTag'

#### Value

A list of combinations

#### Author(s)

Biagolini-Jr.

#### See Also

allequal vfrequency lifexp

#### **Examples**

```
# Create an object contain the name/code of tag colors
tcol<-c("Black", "Blue", "Brown", "Gray", "Green", "Pink", "Purple", "Red", "White", "Yellow")
# Generate color tag combination without especial tags
genseq(30, 4, colorsname= tcol)
#Generate color tag combination with especial color (ex metallic tag for numeric identification)
genseq(30, 4, tcol, nspecial=1, name1="Metal", location1=c(2,4))
# For ongoing works, use the argument usedcombinations to informe the previus used combinations
data(pre_used) # Data example
genseq(30, 4, colorsname= tcol, usedcombinations=pre_used[,1:4])
combinations<-genseq(100, 4, tcol) # save combinations into an object</pre>
```

lifexp

Life expectancy tag sequence sample

### **Description**

Create combinations with variable sample probability.

# Usage

```
lifexp(ntag, colorsname, nspecial = 0, name1 = "Metal", name2 = "EMPTY",
location1 = 1, location2 = 2, nspecial1 = 1, nspecial2 = 1, currentyear = NA,
yearsurvival = 1, lifespan = NA, yearusedcombinations,
usedcombinations, speed = 1, ignorecolor = NA)
```

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#### **Arguments**

ntag Number of tag to be used in each animal. colorsname Names/Code of color tags to be sample.

nspecial Number of special tags/codes, such as metallic, "EMPTY", or flag (min 0, max

2).

name1 Name of special tag 1. name2 Name of special tag 2.

location1 Position (or group of positions) to special band 1.
location2 Position (or group of positions) to special band 2.

nspecial 1 Number of special tag 1 that will be present in all sequences genetated.

Number of special tag 2 that will be present in all sequences genetated.

Number of special tag 2 that will be present in all sequences genetated.

currentyear Current year.

yearsurvival An estimation of the proportion of animals that survive between years.

lifespan Combinations older the lifespan will be automatically disregard.

yearusedcombinations

The year in which the combination was used.

usedcombinations

Pre used combinations.

speed Speed for color frequency adjustment.

ignorecolor Color to be ignored on lifexp.

#### Value

A sequencie of tags

#### Author(s)

Biagolini-Jr.

#### See Also

allequal genseq vfrequency

```
data(pre_used) # Data example
# Create an object contain the name/code of tag colors
tcol<-c("Black","Blue","Brown","Gray","Green","Pink","Purple","Red","White","Yellow")
genseq(30, 4,tcol, "lifexp", pre_used[,1:4],
yearusedcombinations=pre_used[,5], yearsurvival= 0.8, lifespan=5, currentyear=2019)</pre>
```

pre\_used

pre\_used

Pre-used combinations combinations

### **Description**

Simulated data of pre-used combinations

### Usage

```
data("pre_used")
```

#### **Format**

A data frame with 1200 observations on the following 5 variables.

- Tag\_1 a factor with levels Black Brown Dark\_Blue EMPTY Gray Green Light\_Blue Orange Pink Red White Yellow
- Tag\_2 a factor with levels Black Brown Dark\_Blue Gray Green Light\_Blue Metal Orange Pink Red White Yellow
- Tag\_3 a factor with levels Black Brown Dark\_Blue EMPTY Gray Green Light\_Blue Orange Pink Red White Yellow
- Tag\_4 a factor with levels Black Brown Dark\_Blue Gray Green Light\_Blue Metal Orange Pink Red White Yellow

Year a numeric vector

### **Details**

Simulated database to example of registers of color tag sequences usage. Columns 1, 2, 3 and 4 represent the tag code, and column 5 is the date of tagging.

#### Source

Simulated database

```
data(pre_used)
```

scy 13

scy

Summary color year

## Description

Summary the number of each color tag used per year.

## Usage

```
scy(usedcombinations, yearusedcombinations, hide_color = NA)
```

### **Arguments**

```
usedcombinations
```

Pre used combinations.

yearusedcombinations

The year in which the combination was used.

hide\_color

Color(s) to be hide in the estimation of remain colors tags present in nature.

### Value

A summary of the numeber of tag colors used by year

### Author(s)

Biagolini-Jr.

#### See Also

erc

```
data(pre_used) # Data example
scy(pre_used[,1:4],pre_used[,5], hide_color="EMPTY")
```

14 vfrequency

vfrequency Variable frequency tag sequence sample
---

# Description

Create combinations with defined sample probability to each colors.

### Usage

```
vfrequency(ntag, colorsname, colorsf, nspecial = 0, name1 = "Metal",
name2 = "EMPTY", location1 = 1, location2 = 2, nspecial1 = 1, nspecial2 = 1)
```

# Arguments

ntag	Number of tag to be used in each animal.
colorsname	Names/Code of color tags to be sample.
colorsf	Frequencies/ratio for color sample.
nspecial	Number of special tags/codes, such as metallic, "EMPTY", or flag (min 0, max 2).
name1	Name of special tag 1.
name2	Name of special tag 2.
location1	Position (or group of positions) to special band 1.
location2	Position (or group of positions) to special band 2.
nspecial1	Number of special tag 1 that will be present in all sequences genetated.

Number of special tag 2 that will be present in all sequences genetated.

#### Value

A sequencie of tags

#### Author(s)

Biagolini-Jr

nspecial2

### See Also

allequal genseq lifexp

```
tcol<-c("Black","Blue","Brown","Gray","Green","Pink","Purple","Red","White","Yellow")
p<-c(1,2,5,1,2,2,4,5,8,5)
genseq(30, 4, tcol, gen_method="vfrequency", colorsf=p)</pre>
```

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