# Package 'PopGenHelpR'

August 2, 2024

```
Version 1.3.2Description Estimate commonly used population genomic statistics and generate publication quality figures. 'PopGenHelpR' uses vcf, 'geno' (012), and csv files to generate output.
```

Title Streamline Population Genomic and Genetic Analyses

```
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```

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Ancestry\_barchart

Plot an ancestry matrix for individuals and(or) populations.

# Description

Plot an ancestry matrix for individuals and(or) populations.

# Usage

```
Ancestry_barchart(
  anc.mat,
  pops,
  K,
  plot.type = "all",
  col,
  ind.order = NULL,
  pop.order = NULL
)
```

# Arguments

anc.mat

Data frame or character string that supplies the input data. If it is a character string, the file should be a csv. The first column should be the names of each sample/population, followed by the estimated contribution of each cluster to that individual/pop.

Differentiation 3

pops	Data frame or character string that supplies the input data. If it is a character string, the file should be a csv. The columns should be named Sample, containing the sample IDs; Population indicating the population assignment of the individual, population and sample names must be the same type (i.e., both numeric or both characters); Long, indicating the longitude of the sample; Lat, indicating the latitude of the sample.
K	Numeric. The number of genetic clusters in your data set, please contact the package authors if you need help doing this.
plot.type	Character string. Options are all, individual, and population. All is default and recommended, this will plot a barchart for both the individuals and populations.
col	Character vector indicating the colors you wish to use for plotting.
ind.order	Character vector indicating the order to plot the individuals in the individual ancestry bar chart.
pop.order	Chracter vector indicating the order to plot the populations in the population ancesyry bar chart.

## Value

A list containing your plots and the data frames used to generate the plots.

## Author(s)

Keaka Farleigh

# **Examples**

```
data(Q_dat)
Qmat <- Q_dat[[1]]
rownames(Qmat) <- Qmat[,1]
Loc <- Q_dat[[2]]
Test_all <- Ancestry_barchart(anc.mat = Qmat, pops = Loc, K = 5,
plot.type = 'all',col = c('#d73027', '#fc8d59', '#e0f3f8', '#91bfdb', '#4575b4'))</pre>
```

Differentiation

A function to estimate three measures of genetic differentiation using geno files, vcf files, or vcfR objects. Data is assumed to be bi-allelic.

# Description

A function to estimate three measures of genetic differentiation using geno files, vcf files, or vcfR objects. Data is assumed to be bi-allelic.

4 Differentiation

#### Usage

```
Differentiation(
  data,
  pops,
  statistic = "all",
  missing_value = NA,
  write = FALSE,
  prefix = NULL,
  population_col = NULL,
  individual_col = NULL
)
```

#### **Arguments**

data Character. String indicating the name of the vcf file, geno file or vcfR object to

be used in the analysis.

pops Character. String indicating the name of the population assignment file or dataframe

containing the population assignment information for each individual in the data. This file must be in the same order as the vcf file and include columns specifying the individual and the population that individual belongs to. The first column should contain individual names and the second column should indicate the population assignment of each individual. Alternatively, you can indicate the column containing the individual and population information using the indi-

vidual\_col and population\_col arguments.

statistic Character. String or vector indicating the statistic to calculate. Options are any

of: all; all of the statistics; Fst, Weir and Cockerham (1984) Fst; NeisD, Nei's D

statistic; JostsD, Jost's D.

missing\_value Character. String indicating missing data in the input data. It is assumed to be

NA, but that may not be true (is likely not) in the case of geno files.

write Boolean. Whether or not to write the output to files in the current working

directory. There will be one or two files for each statistic. Files will be named

based on their statistic such as Fst\_perpop.csv.

prefix Character. Optional argument. String that will be appended to file output. Please

provide a prefix if write is set to TRUE.

population\_col Numeric. Optional argument (a number) indicating the column that contains the

population assignment information.

individual\_col Numeric. Optional argument (a number) indicating the column that contains the

individuals (i.e., sample name) in the data.

#### Value

A list containing the estimated heterozygosity statistics. The per pop values are calculated by taking the average of the per locus estimates.

## Author(s)

Keaka Farleigh

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#### References

#### Fst:

Pembleton, L. W., Cogan, N. O., & Forster, J. W. (2013). StAMPP: An R package for calculation of genetic differentiation and structure of mixed-ploidy level populations. Molecular ecology resources, 13(5), 946-952.doi:10.1111/17550998.12129

Weir, B. S., & Cockerham, C. C. (1984). Estimating F-statistics for the analysis of population structure. evolution, 1358-1370.

#### Nei's D:

Nei, M. (1972). Genetic distance between populations. The American Naturalist, 106(949), 283-292.doi:10.1086/282771

doi:10.1111/17550998.12129 Pembleton, L. W., Cogan, N. O., & Forster, J. W. (2013). StAMPP: An R package for calculation of genetic differentiation and structure of mixed-ploidy level populations. Molecular ecology resources, 13(5), 946-952.

#### Jost's D:

Jost L (2008). GST and its relatives do not measure differentiation. Molecular Ecology, 17, 4015-4026.doi:10.1111/j.1365294X.2008.03887.x

## **Examples**

```
data("HornedLizard_Pop")
data("HornedLizard_VCF")
Test <- Differentiation(data = HornedLizard_VCF, pops = HornedLizard_Pop, write = FALSE)</pre>
```

Fst\_dat

A genetic differentiation matrix and locality information for each population. This data was generated by subsetting data of Farleigh et al., 2021.

# Description

A symmetric matrix with estimated genetic differentiation (Fst) between 3 populations.

#### Usage

```
data(Fst_dat)
```

#### Format

A list with two elements:

**Fst\_dat** Data frame with three rows and three columns

**Loc\_dat** Data frame containing the locality information for each population

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6 Heterozygosity

#### **Source**

Farleigh, K., Vladimirova, S. A., Blair, C., Bracken, J. T., Koochekian, N., Schield, D. R., ... & Jezkova, T. (2021). The effects of climate and demographic history in shaping genomic variation across populations of the Desert Horned Lizard (Phrynosoma platyrhinos). Molecular Ecology, 30(18), 4481-4496.

#### **Examples**

```
data(Fst_dat)
Fst <- Fst_dat[[1]]
Loc <- Fst_dat[[2]]

Test <- Network_map(dat = Fst, pops = Loc,
neighbors = 2,col = c('#4575b4', '#91bfdb', '#e0f3f8','#fd8d3c','#fc4e2a'),
statistic = "Fst", Lat_buffer = 1, Long_buffer = 1)

Fstat_plot <- Pairwise_heatmap(dat = Fst, statistic = 'FST')</pre>
```

Heterozygosity

A function to estimate seven measures of heterozygosity using geno files, vcf files, or vcfR objects. Data is assumed to be bi-allelic.

# Description

A function to estimate seven measures of heterozygosity using geno files, vcf files, or vcfR objects. Data is assumed to be bi-allelic.

#### Usage

```
Heterozygosity(
  data,
  pops,
  statistic = "all",
  missing_value = NA,
  write = FALSE,
  prefix = NULL,
  population_col = NULL,
  individual_col = NULL
)
```

## **Arguments**

data

Character. String indicating the name of the vcf file, geno file or vcfR object to be used in the analysis.

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pops Character. String indicating the name of the population assignment file or dataframe

containing the population assignment information for each individual in the data. This file must be in the same order as the vcf file and include columns specifying the individual and the population that individual belongs to. The first column should contain individual names and the second column should indicate the population assignment of each individual. Alternatively, you can indicate the column containing the individual and population information using the indi-

vidual col and population col arguments.

statistic Character. String or vector indicating the statistic to calculate. Options are

any of: all; all of the statistics; Ho, observed heterozygosity; He, expected heterozygosity; PHt, proportion of heterozygous loci; Hs\_exp, heterozygosity standardized by the average expected heterozygosity; Hs\_obs, heterozygosity standardized by the average observed heterozygosity; IR, internal relatedness;

HL, homozygosity by locus.

missing\_value Character. String indicating missing data in the input data. It is assumed to be

NA, but that may not be true (is likely not) in the case of geno files.

write Boolean. Whether or not to write the output to files in the current working

directory. There will be one or two files for each statistic. Files will be named

based on their statistic such as Ho\_perpop.csv or Ho\_perloc.csv.

prefix Character. Optional argument. String that will be appended to file output. Please

provide a prefix if write is set to TRUE.

population\_col Numeric. Optional argument (a number) indicating the column that contains the

population assignment information.

individual\_col Numeric. Optional argument (a number) indicating the column that contains the

individuals (i.e., sample name) in the data.

#### Value

A list containing the estimated heterozygosity statistics. The per pop values are calculated by taking the average of the per locus estimates.

#### Author(s)

Keaka Farleigh

#### References

# Expected (He) and observed heterozygosity (Ho):

Nei, M. (1987) Molecular Evolutionary Genetics. Columbia University Press

#### Homozygosity by locus (HL) and internal relatedness (IR):

Alho, J. S., Välimäki, K., & Merilä, J. (2010). Rhh: an R extension for estimating multilocus heterozygosity and heterozygosity—heterozygosity correlation. Molecular ecology resources, 10(4), 720-722.

Amos, W., Worthington Wilmer, J., Fullard, K., Burg, T. M., Croxall, J. P., Bloch, D., & Coulson, T. (2001). The influence of parental relatedness on reproductive success. Proceedings of the Royal Society of London. Series B: Biological Sciences, 268(1480), 2021-2027.doi:10.1098/rspb.2001.1751

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Aparicio, J. M., Ortego, J., & Cordero, P. J. (2006). What should we weigh to estimate heterozygosity, alleles or loci?. Molecular Ecology, 15(14), 4659-4665.

Heterozygosity standardized by expected (Hs\_exp) and observed heterozygosity (Hs\_obs):

Coltman, D. W., Pilkington, J. G., Smith, J. A., & Pemberton, J. M. (1999). Parasite-mediated selection against Inbred Soay sheep in a free-living island populaton. Evolution, 53(4), 1259-1267.

## **Examples**

```
data("HornedLizard_Pop")
data("HornedLizard_VCF")
Test <- Heterozygosity(data = HornedLizard_VCF, pops = HornedLizard_Pop, write = FALSE)</pre>
```

Het\_dat

A data frame of hypothetical heterozygosity data produced by Heterozygosity.

#### **Description**

Data frame containing 5 columns and 3 rows

#### Usage

```
data(Het_dat)
```

#### **Format**

A data frame with 5 columns and 3 rows:

**Heterozygosity** Estimated heterozygosity

Pop Population assignment

Standard.Deviation standard deviation

Longitude Longitude Latitude Latitude

...

#### Source

Coordinates and population names taken from Farleigh, K., Vladimirova, S. A., Blair, C., Bracken, J. T., Koochekian, N., Schield, D. R., ... & Jezkova, T. (2021). The effects of climate and demographic history in shaping genomic variation across populations of the Desert Horned Lizard (Phrynosoma platyrhinos). Molecular Ecology, 30(18), 4481-4496.

```
data(Het_dat)
Test <- Point_map(Het_dat, statistic = "Heterozygosity")</pre>
```

HornedLizard\_Pop 9

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HornedLizard_Pop	A population assignment data frame to be used in Heterozygosity
	and Differentiation.

# Description

Data frame containing 4 columns and 72 rows

#### Usage

```
data(HornedLizard_Pop)
```

#### **Format**

A data frame with 4 columns and 72 rows:

Sample Sample Name

Population Population assignment according to sNMF results (see citation)

Longitude Longitude

Latitude Latitude

...

#### **Source**

Coordinates and population names taken from Farleigh, K., Vladimirova, S. A., Blair, C., Bracken, J. T., Koochekian, N., Schield, D. R., ... & Jezkova, T. (2021). The effects of climate and demographic history in shaping genomic variation across populations of the Desert Horned Lizard (Phrynosoma platyrhinos). Molecular Ecology, 30(18), 4481-4496.

```
data("HornedLizard_Pop")
data("HornedLizard_VCF")
Test <- Differentiation(data = HornedLizard_VCF, pops = HornedLizard_Pop, write = FALSE)</pre>
```

Network\_map

HornedLizard\_VCF

A vcfR object to be used in Heterozygosity and Differentiation.

## **Description**

Data frame containing 4 columns and 72 rows

## Usage

```
data(HornedLizard_Pop)
```

#### **Format**

A vcfR object

vcfR object A vcfR object containing genotype and sample information for 72 individuals.

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#### **Source**

Farleigh, K., Vladimirova, S. A., Blair, C., Bracken, J. T., Koochekian, N., Schield, D. R., ... & Jezkova, T. (2021). The effects of climate and demographic history in shaping genomic variation across populations of the Desert Horned Lizard (Phrynosoma platyrhinos). Molecular Ecology, 30(18), 4481-4496.

## **Examples**

```
data("HornedLizard_Pop")
data("HornedLizard_VCF")
Test <- Heterozygosity(data = HornedLizard_VCF, pops = HornedLizard_Pop, write = FALSE)</pre>
```

Network\_map

A function to map statistics (i.e., genetic differentiation) between points as a network on a map.

## **Description**

A function to map statistics (i.e., genetic differentiation) between points as a network on a map.

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#### Usage

```
Network_map(
  dat,
  pops,
  neighbors,
  col,
  statistic = NULL,
  breaks = NULL,
  Lat_buffer = 1,
  Long_buffer = 1,
  Latitude_col = NULL,
  Longitude_col = NULL)
)
```

#### **Arguments**

dat Data frame or character string that supplies the input data. If it is a character

string, the file should be a csv. If it is a csv, the 1st row should contain the individual/population names. The columns should also be named in this fashion.

pops Data frame or character string that supplies the input data. If it is a character

string, the file should be a csv. The columns should be named Sample, containing the sample IDs; Population indicating the population assignment of the individual; Long, indicating the longitude of the sample; Lat, indicating the latitude of the sample. Alternatively, see the Longitude\_col and Latitude\_col

arguments.

neighbors Numeric or character. The number of neighbors to plot connections with, or

the specific relationship that you want to visualize. Names should match those in the population assignment file and be seperated by an underscore. If I want to visualize the relationship between East and West, for example, I would set

neighbors = "East\_West".

col Character vector indicating the colors you wish to use for plotting.

statistic Character indicating the statistic being plotted. This will be used to title the

legend. The legend title will be blank if left as NULL.

breaks Numeric. The breaks used to generate the color ramp when plotting. Users

should supply 3 values if custom breaks are desired.

Lat\_buffer Numeric. A buffer to customize visualization.

Long\_buffer Numeric. A buffer to customize visualization.

Latitude\_col Numeric. The number of the column indicating the latitude for each sample. If

this is not null, PopGenHelpR will use this column instead of looking for the

Lat column.

Longitude\_col Numeric. The number of the column indicating the longitude for each sample.

If this is not null, PopGenHelpR will use this column instead of looking for the

Long column.

#### Value

A list containing the map and the matrix used to plot the map.

Pairwise\_heatmap

#### Author(s)

Keaka Farleigh

# **Examples**

```
data(Fst_dat)
Fst <- Fst_dat[[1]]
Loc <- Fst_dat[[2]]
Test <- Network_map(dat = Fst, pops = Loc,
neighbors = 2,col = c('#4575b4', '#91bfdb', '#e0f3f8','#fd8d3c','#fc4e2a'),
statistic = "Fst", Lat_buffer = 1, Long_buffer = 1)</pre>
```

Pairwise\_heatmap

A function to plot a heatmap from a symmetric matrix.

# Description

A function to plot a heatmap from a symmetric matrix.

# Usage

```
Pairwise_heatmap(dat, statistic, col = NULL)
```

## **Arguments**

dat	Data frame or character string that supplies the input data. If it is a character string, the file should be a csv. If it is a csv, the 1st row should contain the individual/population names. The columns should also be named in this fashion.
statistic	Character indicating the statistic represented in the matrix, this will be used to label the plot.
col	Character vector indicating the colors to be used in plotting. The vector should

contain two colors, the first will be the low value, the second will be the high value.

## Value

A heatmap plot

```
#' data(Fst_dat)
Fst <- Fst_dat[[1]]
Fstat_plot <- Pairwise_heatmap(dat = Fst, statistic = 'FST')</pre>
```

PCA 13

PCA	A function to perform principal component analysis (PCA) on genetic data. Loci with missing data will be removed prior to PCA.
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# Description

A function to perform principal component analysis (PCA) on genetic data. Loci with missing data will be removed prior to PCA.

# Usage

```
PCA(
   data,
   center = TRUE,
   scale = FALSE,
   missing_value = NA,
   write = FALSE,
   prefix = NULL
)
```

# Arguments

data	Character. String indicating the name of the vcf file, geno file or vcfR object to be used in the analysis.
center	Boolean. Whether or not to center the data before principal component analysis.
scale	Boolean. Whether or not to scale the data before principal component analysis.
missing_value	Character. String indicating missing data in the input data. It is assumed to be NA, but that may not be true (is likely not) in the case of geno files.
write	Boolean. Whether or not to write the output to files in the current working directory. There will be two files, one for the individual loadings and the other for the percent variance explained by each axis.
prefix	Character. Optional argument. String that will be appended to file output. Please provide a prefix if write is set to TRUE.

## Value

A list containing two elements: the loadings of individuals on each principal component and the variance explained by each principal component.

## Author(s)

Keaka Farleigh

```
data("HornedLizard_VCF")
Test <- PCA(data = HornedLizard_VCF)</pre>
```

14 Piechart\_map

Piechart\_map

Plot a map of ancestry pie charts.

#### **Description**

Plot a map of ancestry pie charts.

#### Usage

```
Piechart_map(
   anc.mat,
   pops,
   K,
   plot.type = "all",
   col,
   piesize = 0.35,
   Lat_buffer,
   Long_buffer,
   Latitude_col = NULL,
   Longitude_col = NULL)
```

#### **Arguments**

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Data frame or character string that supplies the input data. If it is a character string, the file should be a csv. The first column should be the names of each sample/population, followed by the estimated contribution of each cluster to that individual/pop.

pops

Data frame or character string that supplies the input data. If it is a character string, the file should be a csv. The columns should be named Sample, containing the sample IDs; Population indicating the population assignment of the individual, population and sample names must be the same type (i.e., both numeric or both characters); Long, indicating the longitude of the sample; Lat, indicating the latitude of the sample. Alternatively, see the Longitude\_col and Latitude\_col arguments.

Κ

Numeric. The number of genetic clusters in your data set, please contact the package authors if you need help doing this.

plot.type

Character string. Options are all, individual, and population. All is default and recommended, this will plot a piechart map for both the individuals and

populations.

col

Character vector indicating the colors you wish to use for plotting.

piesize

Numeric. The radius of the pie chart for ancestry mapping.

Lat\_buffer

Numeric. A buffer to customize visualization.

Long\_buffer

Numeric. A buffer to customize visualization.

Plot\_coordinates 15

Latitude\_col Numeric. The number of the column indicating the latitude for each sample. If this is not null, PopGenHelpR will use this column instead of looking for the Lat column.

Longitude\_col Numeric. The number of the column indicating the longitude for each sample. If this is not null, PopGenHelpR will use this column instead of looking for the Long column.

#### Value

A list containing your plots and the data frames used to generate the plots.

#### Author(s)

Keaka Farleigh

## **Examples**

```
data(Q_dat)
Qmat <- Q_dat[[1]]
rownames(Qmat) <- Qmat[,1]
Loc <- Q_dat[[2]]
Test_all <- Piechart_map(anc.mat = Qmat, pops = Loc, K = 5,
plot.type = 'all', col = c('#d73027', '#fc8d59', '#e0f3f8', '#91bfdb', '#4575b4'), piesize = 0.35,
Lat_buffer = 1, Long_buffer = 1)</pre>
```

Plot\_coordinates

A function to plot coordinates on a map.

## **Description**

A function to plot coordinates on a map.

## Usage

```
Plot_coordinates(
  dat,
  col = c("#A9A9A9", "#000000"),
  size = 3,
  Lat_buffer = 1,
  Long_buffer = 1,
  Latitude_col = NULL,
  Longitude_col = NULL
)
```

Point\_map

## **Arguments**

dat Data frame or character string that supplies the input data. If it is a character

string, the file should be a csv. The coordinates of each row should be indicated by columns named Longitude and Latitude. Alternatively, see the Latitude\_col

and Longitude\_col arugments.

col Character vector indicating the colors you wish to use for plotting, two colors

are allowed. The first color will be the fill color, the second is the outline color. For example, if I want red points with a black outline I would set col to col =

c("#FF0000", "#000000").

size Numeric. The size of the points to plot.

Lat\_buffer Numeric. A buffer to customize visualization.

Long\_buffer Numeric. A buffer to customize visualization.

Latitude\_col Numeric. The number of the column indicating the latitude for each sample. If

this is not null, PopGenHelpR will use this column instead of looking for the

Latitude column.

Longitude\_col Numeric. The number of the column indicating the longitude for each sample.

If this is not null, PopGenHelpR will use this column instead of looking for the

Longitude column.

#### Value

A ggplot object.

#### Author(s)

Keaka Farleigh

#### **Examples**

```
data("HornedLizard_Pop")
Test <- Plot_coordinates(HornedLizard_Pop)</pre>
```

Point\_map

A function to map statistics as colored points on a map.

## **Description**

A function to map statistics as colored points on a map.

Point\_map 17

#### Usage

```
Point_map(
  dat,
  statistic,
  size = 3,
  breaks = NULL,
  col,
  out.col = NULL,
  Lat_buffer = 1,
  Long_buffer = 1,
  Latitude_col = NULL,
  Longitude_col = NULL)
```

## Arguments

dat	Data frame or character string that supplies the input data. If it is a character
	string, the file should be a csv. The first column should be the statistic to be
	plotted. The coordinates of each row should be indicated by columns named
	Longitude and Latitude. Alternatively, see the Longitude_col and Latitude_col
	arguments.

statistic Character string. The statistic to be plotted.
size Numeric. The size of the points to plot.

should supply 3 values if custom breaks are desired.

col Character vector indicating the colors you wish to use for plotting, three colors

are allowed (low, mid, high). The first color will be the low color, the second

the middle, the third the high.

out.col Character. A color for outlining points on the map. There will be no visible

outline if left as NULL.

Lat\_buffer Numeric. A buffer to customize visualization.

Long\_buffer Numeric. A buffer to customize visualization.

Latitude\_col Numeric. The number of the column indicating the latitude for each sample. If

this is not null, PopGenHelpR will use this column instead of looking for the

Latitude column.

Longitude\_col Numeric. The number of the column indicating the longitude for each sample.

If this is not null, PopGenHelpR will use this column instead of looking for the

Longitude column.

#### Value

A list containing maps and the data frames used to generate them.

#### Author(s)

Keaka Farleigh

18 Private.alleles

#### **Examples**

```
data(Het_dat)
Test <- Point_map(Het_dat, statistic = "Heterozygosity")</pre>
```

Private.alleles

A function to estimate the number of private alleles in each population.

#### **Description**

A function to estimate the number of private alleles in each population.

# Usage

```
Private.alleles(
  data,
  pops,
  write = FALSE,
  prefix = NULL,
  population_col = NULL,
  individual_col = NULL
)
```

#### **Arguments**

data Character. String indicating the name of the vcf file or vcfR object to be used	in
--	----

the analysis.

pops Character. String indicating the name of the population assignment file or dataframe

containing the population assignment information for each individual in the data. This file must be in the same order as the vcf file and include columns specifying the individual and the population that individual belongs to. The first column should contain individual names and the second column should indicate the population assignment of each individual. Alternatively, you can indicate the column containing the individual and population information using the indi-

vidual col and population col arguments.

write Boolean. Optional argument indicating Whether or not to write the output to

a file in the current working directory. This will output to files; 1) the table of private allele counts per population (named prefix\_PrivateAlleles\_countperpop)

and 2) metadata associated with the private alleles (named prefix\_PrivateAlleles\_metadata).

Please supply a prefix it you write files to your working directory as a best prac-

tice.

prefix Character. Optional argument indicating a string that will be appended to file

output. Please set a prefix if write is TRUE.

population\_col Numeric. Optional argument (a number) indicating the column that contains the

population assignment information.

individual\_col Numeric. Optional argument (a number) indicating the column that contains the

individuals (i.e., sample name) in the data.

Q\_dat

#### Value

A list containing the count of private alleles in each population and the metadata for those alleles. The metadata is a list that contains the private allele and locus name for each population.

#### Author(s)

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## **Examples**

```
data("HornedLizard_Pop")
data("HornedLizard_VCF")
Test <- Private.alleles(data = HornedLizard_VCF, pops = HornedLizard_Pop, write = FALSE)</pre>
```

Q\_dat

A list representing a q-matrix and the locality information associated with the qmatrix

#### **Description**

List with two elements

#### Usage

```
data(Q_dat)
```

#### **Format**

A list with two elements:

**Qmat** A q-matrix with 6 columns and 30 rows, the first column lists the sample name and the remaining 5 represent the contribution a genetic cluster to that individuals ancestry

Loc\_dat The locality information for each individual in the q-matrix

## Source

Data was generated by package authors.

```
data(Q_dat)
Qmat <- Q_dat[[1]]
rownames(Qmat) <- Qmat[,1]
Loc <- Q_dat[[2]]
Test_all <- Ancestry_barchart(anc.mat = Qmat, pops = Loc, K = 5,
plot.type = 'all',col = c('#d73027', '#fc8d59', '#e0f3f8', '#91bfdb', '#4575b4'))</pre>
```

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