Package 'POPdemog'

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Description Plot demographic graph for single/multiple populations based on the demographic description from scripts of simulation tools.
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R topics documented:
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POPdemog-package Plot population demographic hisotry

Description

Type Package

Title Plot population demographic hisotry

Plot demographic graph for single/multiple populations based on the demographic description from scripts of simulation tools.

POPdemog-package

Details

This package contains three main functions to visualize the population demographic history from simulation commands. This package can support ms, msa, msHot, MaCS, msprime, and cosi. Please check the tutorial file for the update of supporting list. Users could copy the simulation script (for ms(a), msHot, MaCS, and msprime(need a translation)) into one file, or into a R string, or just use the parameter file (for Cosi) as the input command file. Each input should only contain one demographic model.

Once the command file is prepared, function PlotMS could be used to generate the demographic graph or to give the parameters to PlotMMig plot the multiple migrations. More adjustments on migration graph would be achieved by using function PlotMig, which is allowed to add the migration patterns to other plots, such as geological map.

Author(s)

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References

ms: Hudson, R. R. "Generating Samples under a Wright-Fisher Neutral Model of Genetic Variation." Bioinformatics 18.2 (2002): 337-38.

msHot: Hellenthal, G., and M. Stephens. "MsHOT: Modifying Hudson's Ms Simulator to Incorporate Crossover and Gene Conversion Hotspots." Bioinformatics 23.4 (2006): 520-21.

MaCS: Chen, G. K., P. Marjoram, and J. D. Wall. "Fast and Flexible Simulation of DNA Sequence Data." Genome Research 19.1 (2008): 136-42.

Cosi: Shlyakhter, Ilya, Pardis C. Sabeti, and Stephen F. Schaffner. "Cosi2: An Efficient Simulator of Exact and Approximate Coalescent with Selection | Bioinformatics | Oxford Academic." OUP Academic. Oxford University Press, 22 Aug. 2014.

msprime: Jerome Kelleher, Alison M Etheridge and Gilean McVean, "Efficient Coalescent Simulation and Genealogical Analysis for Large Sample Sizes", PLoS Comput Biol 12(5): e1004842. doi: 10.1371/journal.pcbi.1004842. 2016.

Examples

```
###Tennessen's standard model
cat("macs 2025 15000000 -i 10 -r 3.0e-04 -t 0.00069 -T -I 4 10 1006 1008 1 0
-n 4 0.205 -n 1 58.00274 -n 2 70.041 -n 3 187.55 -eg 0.9e-10 1 482.46
-eg 1.0e-10 2 570.18 -eg 1.1e-10 3 720.23 -em 1.2e-10 1 2 0.731
-em 1.3e-10 2 1 0.731 -em 1.4e-10 3 1 0.2281 -em 1.5e-10 1 3 0.2281
-em 1.6e-10 2 3 0.9094 -em 1.7e-10 3 2 0.9094 -eg 0.007 1 0
-en 0.007001 1 1.98 -eg 0.007002 2 89.7668 -eg 0.007003 3 113.3896
-eG 0.031456 0 -en 0.031457 2 0.1412 -en 0.031458 3 0.07579
-eM 0.031459 0 -ej 0.03146 3 2 -en 0.0314601 2 0.2546
-em 0.0314602 2 1 4.386 -em 0.0314603 1 2 4.386 -eM 0.0697669 0
-ej 0.069767 2 1 -en 0.0697671 1 1.98 -en 0.2025 1 1 -ej 0.9575923 4 1
-em 0.06765 2 4 32 -em 0.06840 2 4 0", file="std-model-Tennessen.cmd")
#plot the demographic graph
par(mfrow=c(1,2))
PlotMS(input.file="std-model-Tennessen.cmd", type="macs", N4=10000, pop.scale="log",
log.base =50, inpos = c(1,4,7,9), time.scale = "log10year",
col.pop=c("brown", "blue", "gold3", "forestgreen"),
pops=c("AFR", "EUR", "ASIA", "ARC"),
```

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```
cex.lab=1, cex.axis = 1, xlab="", length.arrow=0.1)
title("Demographic histoy")
PlotMS(input.file="std-model-Tennessen.cmd", type="macs", N4=10000,
time.scale = "log10year", plot.out=FALSE, demo.out=TRUE)->out;
PlotMig(time_pt=2, demograph_out=out$demograph_out,mig_par=out$mig_par,
col.pop=c("brown", "blue", "gold3", "forestgreen"), pop.scale="topology");
legend("topleft", legend=c("AFR", "EUR", "ASIA", "ARC"),
col=c("brown", "blue", "gold3", "forestgreen"), pch=20, bty="n")
title("Migrations at 200 years ago");unlink("std-model-Tennessen.cmd")
###Archaic introgrssion model
cat("./ms 44 1 -r 20000 50000000 -t 30000 -I 6 20 20 1 1 1 1 1 -en 0 1 1
-en 0 2 1 -en 0 3 1e-10 -en 0 4 1e-10 -en 0 5 1e-10 -en 0 6 1e-10
-es 0.0125 2 0.97 -en 0.02500025 7 0.25 -en 0.02500025 2 1 -ej 0.05 4 3
-ej 0.05 6 5 -en 0.05000025 3 0.25 -en 0.05000025 5 0.25 -ej 0.0500025 5 3
-en 0.050005 3 0.25 -ej 0.075 2 1 -en 0.0750025 1 1 -ej 0.1 7 3
-en 0.1000025 3 0.25 -ej 0.3 3 1 -en 0.3000025 1 1", file="test.1.ms.cmd")
PlotMS(input.file = "test.1.ms.cmd", type="ms", N4=10000,
time.scale = "kyear", length.arrow=0.1, inpos=c(1,2,5,4.5,5.5,6,3),
col.pop=c("brown", "blue", "forestgreen", rainbow(10)[6:9]));\\
unlink("test.1.ms.cmd")
###Migration model from ms.
cat("./ms 15 100 -t 3.0 -I 6 0 7 0 0 8 0 -m 1 2 2.5 -m 2 1 2.5 -m 2 3 2.5
 -m \ 3 \ 2 \ 2.5 \ -m \ 4 \ 5 \ 2.5 \ -m \ 5 \ 4 \ 2.5 \ -m \ 5 \ 6 \ 2.5 \ -m \ 6 \ 5 \ 2.5 \ -em \ 2.0 \ 3 \ 4 \ 2.5 
-em 2.0 4 3 2.5", file="test.2.ms.cmd")
PlotMS(input.file = "test.2.ms.cmd", type="ms", N4=10000, col.pop="gray",
col.arrow="black", length.arrow=0.1, lwd.arrow=2);unlink("test.2.ms.cmd")
```

NRuler

add population size ruler

Description

This function works like the function legend,

Usage

```
NRuler(x, y = NULL, Nsize, Nlab = N.szie, N4, pop.scale,
linear.scale = 0.2, log.base = 10, ...)
```

Arguments

x, y	Position of the population size ruler, if y does not have numeric value, x will support the keywords inputs from the list "bottomright", "bottom", "bottom-left", "topleft", "topleft", "topright", "right" and "center".
Nsize	The population sizes of the ticks on the ruler
Nlab	The labels on the ticks of the ruler, default as the population sizes
N4	Scalar to scale real population size
pop.scale	A keyword to define the way to reflect population size to lineage width. It supports "log" and "linear".

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linear.scale	A numeric number used to reflect population size linearly. This value will be applied when pop.scale="linear".
log.base	The base of logarithm. This value will be applied when pop.scale="log".
	Additional arguments can be passed, such as col, lwd, lty.

PlotMig

Plot migration event(s) at a particular time

Description

This function is used to plot migration events at a particular time point based on the output of PlotMS with demo.out=T. The parameters of add and map.pos allow the migration graph to be added to any other background. User can use time_pt or event to define the time to plot migration(s).

Usage

```
PlotMig(time_pt = NULL, event = 1, mig_par, demograph_out,
   pop.scale = mig_par$pop.scale, linear.scale = mig_par$linear.scale,
   log.base = mig_par$log.base, col.pop = mig_par$col.pop,
   col.arrow = mig_par$col.arrow, xlim = mig_par$xlim, ylim = mig_par$ylim,
   lwd.arrow = mig_par$lwd.arrow, length.arrow = mig_par$length.arrow,
   angle.arrow = mig_par$angle.arrow, topology.scale = 1, add = FALSE,
   map.pos = NULL, m.adjust = 0)
```

Arguments

time_pt	A numeric value define at which time to plot migration(s). time_pt should be in the same scale as the setting of time.scale. For example, time_pt = 3 when time.scale = "log10year" gives us the migrations at 10^3 years ago.
event	An index to define at which time to plot migration(s) out. Every demographic event has an index in the order of time. Any demographic changes at the same time is defined as the same event and share the same index.
mig_par	A list contained all settings for plot demographic graph, refer PlotMS.
demograph_out	A list contained all demographic information, refer PlotMS.
pop.scale	A keyword to define the way to adjust the lineage width. "topology" returns only topology structure among simulated populations, ignoring either the population sizes and the time duration difference between any demographic events. "linear" linearly reflects the population sizes to lineage widths, the scale magnitude is defined in variable linear.scale; "log" is to use logarithm function to reflect the population sizes, the base is defined in log.base.
linear.scale	Linear scale magnitude, be applied when pop.scale="linear".
log.base	The base of logarithm, be applied when pop.scale="log".
col.pop	Population lineage color.

col.arrow Migration arrow color.

xlim The range of x-axis.

ylim The range of y-axis.

1wd.arrow The line width of arrow, defined by 0.5+migration strength*1wd.arrow.

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length.arrow The size of arrow end
angle.arrow The angle of the arrow end, between 0 and 90.
topology.scale Used for scale the size of circle when the pop.scale= "topology".

add A logical value allowing to add the migrations to other background (2D only).
Positions for every population dime should be well defined in map.pos.

map.pos A matrix, each row is the position for each population.

m.adjust Migration threshold for plotting the migration events. Migration event with strength higher than m.adjust will be shown. Default value is 0.

See Also

PlotMS

Examples

```
\mathtt{cat("./ms\ 15\ 100\ -t\ 3.0\ -I\ 6\ 0\ 7\ 0\ 0\ 8\ 0\ -m\ 1\ 2\ 2.5\ -m\ 2\ 1\ 2.5\ -m\ 2\ 3\ 2.5}
-m 3 2 2.5 -m 4 5 2.5 -m 5 4 2.5 -m 5 6 2.5 -m 6 5 2.5 -em 2.0 3 4 2.5
-em 2.0 4 3 2.5", file = "test.mig.cmd")
out <- PlotMS(input.file = "test.mig.cmd", type = "ms", N4 = 10000, plot.out = FALSE, demo.out = TRUE);</pre>
#check all migration events
events <- out$mig_par$events</pre>
print(events)
#check the time for those migration events
timeofevents <- out$mig_par$time[events]</pre>
print(timeofevents)
#plot event by event
par(mfrow = c(1, 2))
PlotMig(event = 1, demograph_out = out$demograph_out, mig_par = out$mig_par)
title("Event-1");
PlotMig(event = 2, demograph_out = out$demograph_out, mig_par = out$mig_par,
col.pop = 1:6, xlim = c(-5,4)
title("Event-2", cex.main = 3);
legend("topleft", col = 1:6, pch = 20, bty = "n", cex = 2,
legend = c("pop-1", "pop-2", "pop-3", "pop-4", "pop-5", "pop-6"))
unlink("test.mig.cmd")
```

 ${\tt PlotMMig}$

Plot Multiple Migrations

Description

This function is used to plot all the migration evens. It based on the output of PlotMS with demo.out=FALSE. Plot settings should be customized in PlotMS. Use function PlotMig to customize the plot of single migration.

Usage

```
PlotMMig(demograph_out, mig_par, m.adjust = 0)
```

Arguments

demograph_out A list of all demographic information, more descriptions will be found in the return value description of PlotMS.

mig_par A list of all settings for plotting demographic graph, more descriptions will be found in the return value description of PlotMS.

m.adjust Migration threshold for plotting the migration events. Migration event with strength higher than m.adjust will be shown. Default value is 0.

See Also

PlotMS

Examples

```
cat("./ms 15 100 -t 3.0 -I 6 0 7 0 0 8 0 -m 1 2 2.5 -m 2 1 2.5 -m 2 3 2.5
-m 3 2 2.5 -m 4 5 2.5 -m 5 4 2.5 -m 5 6 2.5 -m 6 5 2.5 -em 2.0 3 4 2.5
-em 2.0 4 3 2.5", file = "test.mig.cmd")
out<-PlotMS(input.file = "test.mig.cmd", type = "ms", N4 = 10000,
plot.out = FALSE, demo.out = TRUE, col.pop=1:6, cex.lab=1.5);
PlotMMig(demograph_out = out$demograph_out, mig_par = out$mig_par)
unlink("test.mig.cmd")</pre>
```

PlotMS

Plot population demographic graph and generate demographic parameters

Description

This is the main function to plot demographic graph for single/multiple populations, which is named after Hudson's ms. It can read the simulation script from ms, msa, msHot, MaCS, scrm, and cosi.

The input.file or imput.cmd and command type are required to give a quick plot of demographic history. Principally, every component in the output graph could be directly customized according to need.

In the demographic graph, each population has a lineage vertically stretch back in time (inpos for positions, col.pop for the colors, pops for the names, and time.scale for the time axis). The width of the lineage reflects the relative population size (pop.scale for lineage widths). Population splits and migrations are in arrows (col.arrow, length.arrow, lwd.arrow, angle.arrow).

Usage

```
PlotMS(input.cmd = NULL, input.file = NULL, type, inpos = NULL,
  N4 = 10000, pop.scale = "linear", linear.scale = 0.2, log.base = 10,
  time.scale = "4Ne", gen = 25, m.adjust = 0, col.pop = "gray45",
  col.arrow = col.pop, length.arrow = 0.15, lwd.arrow = 1,
  angle.arrow = 15, pops = NULL, xlab = "Population",
  ylab = paste("Time before present (", time.scale, ")", sep = ""),
  xlim = NULL, ylim = NULL, plot.out = T, demo.out = F, cex.lab = 1,
  cex.axis = 1, axes = T)
```

Arguments

input.cmd A string input for demographic events. User can paste simulation scripts for the

convenient of debugging.

input.file A file for demographic events. It can be simulation scripts (for ms, msa, MaCS)

or parameter files (for cosi).

type A keyword indicates the type of simulation command, currently supports "ms"

for ms, "msa" for msa, "macs" for MaCS, "scrm" for scrm, "cosi" for cosi, "msprime" for msprime. Please check tutorial file to see more support simu-

lators.

inpos Population positions at time 0.

N4 A numeric for the effective population size of 4Ne, exact the same definition as

4N0 in the well-known simulation tool ms.

pop.scale A keyword to define the way to adjust the lineage width. "topology" returns only

topology structure among simulated populations, ignoring either the population sizes and the time duration difference between any demographic events. "linear" linearly reflects the population sizes to lineage widths, the scale magnitude is defined in variable linear.scale; "log" is to use logarithm function to reflect

the population sizes, the base is defined in log.base.

linear.scale Linear scale magnitude, be applied when pop.scale="linear".

log.base The base of logarithm, be applied when pop.scale="log".

time.scale A keyword to define the time unit used in the plot, it can be "4Ne", "genera-

tion", "year", "kyear", and "log10year". When the pop.scale="topology", this

parameter will be ignored.

gen Years per generation. Default value is 25.

m. adjust Threshold for migration strength. Only arrows with migration strength bigger

than m. adjust are plotted. This value should be stay between 0 and 1.

col.pop Color for each population.

col.arrow Color for each migration arrow

length.arrow Size of arrow ends

lwd.arrow Arrow width, which is determined as 0.5+migration strength*lwd.arrow.

angle.arrow Arrow end angle, between 0 and 90.

pops Population name labels. Default as 1:total population number.

xlab Title for the x-axis
ylab Title for the y-axis
xlim Range of x-axis
ylim Range of y-axis

plot.out A logical variable to give out the demographic plot or not. If TRUE, the demo-

graphic plot will be given.

demo.out A logical variable to output the demographic parameters. If TRUE, all demo-

graphic parameters that used for the graph will output as a list.

cex.lab The magnification to be used for x and y labels relative to the current setting of

cex.

cex.axis The magnification to be used for axis annotation relative to the current setting

of cex.

axes A logical value to plot the axes or not.

Value

if the parameter plot=F/FALSE, following three lists will be returned:

demograph_out

This list contains all demographic details from the input command file: time.series is a vector of time; Pos is a numeric matrix of positions for each population at every demographic event; N is a numeric matrix of population size for each population at every demographic event; m is 3-D numeric matrix of migration rates between populations at every demographic event; survive is a matrix recording the begin and end for each population according to the demographic events; g.rate is a matrix of exponential growth rates at every demographic event; pop.pos is a numeric vector of the population positions at time 0; pop.lab is a vector of population names; mscmd is the ms command for demographic plot. All simulation script will turn to ms command for extract the demographic information; present.pop.num is the number of populations at present; total.pop.num is the number of total populations exist in the plot; N4 is 4Ne, please refers to Hudson's ms for more description; gen is the years per generation.

evo_par

This list contains all parameters used to draw the demographic graph, including: pop.scale define the way to scale population size; linear.scale is used for scale population size linearly; log.base is the base of logarithm of the population size; time.scale defines the time unit used in the plot; time is the scaled time vector according to the time.scale; col.pop specifies the plotting color for each population; col.arrow specifies the plotting color for each migration arrow; length.arrow is the size of arrow ends; lwd.arrow is the width of arrow; angle.arrow is the angle of the arrow end; lab.pop is a vector of population names; lab.pos is a vector of positions to plot the population names; xlim is the range of x-axis; ylim is the range of y-axis; xlab is the title of x-axis; ylab it the title of y-axis; cex.lab is the magnification to be used for x and y labels; cex.axis is the magnification to be used for axis annotation; axes is a logical value to plot the axes or not.

mig_par

This list contains all parameters used to draw the migrations, including: pop.scale defines the way to scale population size; linear.scale is used for scaling population size linearly; log.base is the base of logarithm of the population size; time.scale defines the time unit used in the plot; time is the scaled time vector according to the time.scale; lab.pop is a vector of population names; col.pop specifies the plotting color for each population; col.arrow specifies the plotting color for each migration arrow; length.arrow is the size of arrow ends; lwd.arrow is the width of arrow; angle.arrow is the angle of the arrow end; xlim specifies the range of x-axis; ylim specifies the range of y-axis; mfrow sets the layout of multiple migrations in one plot; events records the events when migration state changes; cex.lab is the magnification to be used for x and y labels;

References

4Ne: http://home.uchicago.edu/rhudson1/source/mksamples.html

Examples

```
#example 1
cat("./ms 44 1 -r 20000 50000000 -t 30000 -I 6 20 20 1 1 1 1 -en 0 1 1
-en 0 2 1 -en 0 3 1e-10 -en 0 4 1e-10 -en 0 5 1e-10 -en 0 6 1e-10
```

```
-es 0.0125 2 0.97 -en 0.02500025 7 0.25 -en 0.02500025 2 1 -ej 0.05 4 3
-ej 0.05 6 5 -en 0.05000025 3 0.25 -en 0.05000025 5 0.25 -ej 0.0500025 5 3
-en 0.050005 3 0.25 -ej 0.075 2 1 -en 0.0750025 1 1 -ej 0.1 7 3
-en 0.1000025 3 0.25 -ej 0.3 3 1 -en 0.3000025 1 1", file="test.1.ms.cmd")
PlotMS(input.file = "test.1.ms.cmd", type="ms", N4=10000, time.scale = "kyear")
#adjust the population position
PlotMS(input.file = "test.1.ms.cmd", type="ms", N4=10000, time.scale = "kyear",
inpos=c(1,2,5,4,6,7,3))
#add color for each population
PlotMS(input.file = "test.1.ms.cmd", type="ms", N4=10000, time.scale = "kyear",
inpos=c(1,2,5,4,6,7,3), col.pop=rainbow(10)[3:9])
#add population names
unlink("test.1.ms.cmd")
#example 2
cat("./ms 15 100 -t 3.0 -I 6 0 7 0 0 8 0 -m 1 2 2.5 -m 2 1 2.5 -m 2 3 2.5
 -m \ \ 3 \ \ 2 \ \ 2.5 \ \ -m \ \ 4 \ \ 5 \ \ 2.5 \ \ -m \ \ 5 \ \ 6 \ \ 2.5 \ \ -m \ \ 6 \ \ 5 \ \ 2.5 \ \ -em \ \ 2.0 \ \ 3 \ \ 4 \ \ 2.5 
-em 2.0 4 3 2.5", file="test.2.ms.cmd")
PlotMS(input.file = "test.2.ms.cmd", type="ms", N4=10000, col.pop=3,
col.arrow=1, lwd.arrow=1.5)
unlink("test.2.ms.cmd")
#example 3
cat("./ms 1 1 -t 1.0 -I 3 10 10 10 -n 1 1.682020 -n 2 3.736830
-n 3 7.292050 -eg 0 2 116.010723 -eg 0 3 160.246047
-ma 0 0.881098 0.561966 0.881098 0 2.797460 0.561966 2.797460 0
-ej 0.028985 3 2 -en 0.028985 2 0.287184
-ema 0.028985 3 0 7.293140 0 7.293140 0 0 0 0 0 -ej 0.197963 2 1
-en 0.303501 1 1", file="Ryan2009.cmd")
PlotMS(input.file = "Ryan2009.cmd", type="ms", N4=10000, pop.scale="log",
log.base=200, pops=c("AFR", "EUR", "ESA"),
col.pop=c("brown", "blue", "yellow"));
unlink("Ryan2009.cmd")
#example 4
cat("ms 4 1 -t 7156.0000000 -r 2000.0000 10000000 -eN 0 5
-eG 0.000582262 1318.18 -eG 0.00232905 -329.546 -eG 0.00931619 82.3865
-eG 0.0372648 -20.5966 -eG 0.149059 5.14916 -eN 0.596236 0.5 -T",
file="zigzag.cmd")
par(mfrow=c(1,2))
PlotMS(input.file = "zigzag.cmd", type="ms", N4=10000)
#change the time unit
PlotMS(input.file = "zigzag.cmd", type="ms", N4=10000, time.scale="log10year")
unlink("zigzag.cmd")
```

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