

 Open access • Journal Article • DOI:10.1093/BIOINFORMATICS/BTP696

## **pegas: an R package for population genetics with an integrated-modular approach.**

— [Source link](#) 

Emmanuel Paradis

**Institutions:** Institut de recherche pour le développement

**Published on:** 01 Feb 2010 - Bioinformatics (Oxford University Press)

**Topics:** Population

Related papers:

- [adegenet: a R package for the multivariate analysis of genetic markers](#)
- [Inference of population structure using multilocus genotype data](#)
- [R: A language and environment for statistical computing.](#)
- [APE: Analyses of Phylogenetics and Evolution in R language](#)
- [Detecting the number of clusters of individuals using the software STRUCTURE: a simulation study.](#)

Share this paper:    

View more about this paper here: <https://typeset.io/papers/pegas-an-r-package-for-population-genetics-with-an-135vipnci2>



**HAL**  
open science

## pegas: an R package for population genetics with an integrated-modular approach

E. Paradis

► **To cite this version:**

E. Paradis. pegas: an R package for population genetics with an integrated-modular approach. *Bioinformatics*, Oxford University Press (OUP), 2010, 26 (3), pp.419 - 420. 10.1093/bioinformatics/btp696 . hal-01822130

**HAL Id: hal-01822130**

**<https://hal.archives-ouvertes.fr/hal-01822130>**

Submitted on 23 Jun 2018

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

# PEGAS: an R package for population genetics with an integrated–modular approach

Emmanuel Paradis\*

Institut de Recherche pour le Développement, UR175 CAVIAR, BP 5095, 361 rue Jean-François Breton, F-34196 Montpellier Cédex 5, France

Received on XXXXX; revised on XXXXX; accepted on XXXXX

Associate Editor: Jeffrey Barrett

## ABSTRACT

**Summary:** *pegas* is a new package for the analysis of population genetic data. It is written in R and is integrated with two other existing R packages (*ape* and *adegenet*). *pegas* provides functions for standard population genetic methods, as well as low-level functions for developing new methods. The flexible and efficient graphical capabilities of R are used for plotting haplotype networks as well as for other functionalities. *pegas* emphasises the need to further develop an integrated–modular approach for software dedicated to the analysis of population genetic data.

**Availability:** *pegas* is distributed through the Comprehensive R Archive Network (CRAN):

<http://cran.r-project.org/web/packages/pegas/index.html>

Further information may be found at: <http://ape.mpl.ird.fr/pegas/>

**Contact:** [Emmanuel.Paradis@ird.fr](mailto:Emmanuel.Paradis@ird.fr)

Population genetics has a strong mathematical background, and therefore genetic data analyses heavily relies on computer programs. Currently, no unified framework for these programs exists making the use of the many different population genetics programs a complicated task (Excoffier and Heckel, 2006). On the other hand, R (R Development Core Team, 2009) appeared as a unified framework for analysing data in bioinformatics (Gentleman, 2008) and phylogenetics (Paradis, 2006). In this note, I introduce *pegas* (Population and Evolutionary Genetics Analysis System) which aims to fill the gap between the existing R packages and the current need in population genetic data analyses.

*pegas* is entirely written in R to insure maximum portability among operating systems. It requires a standard R installation as well as two packages: *ape* (Paradis *et al.*, 2004) and *adegenet* (Jombart, 2008). These three packages make an integrated environment for population genetic data analysis.

One of the strengths of R is the flexibility of its data structures, so that it is easy to adapt them for a particular field, resulting in much easier data input/output and manipulation (one of the most time-consuming step in data analysis). Two such structures have been derived for *pegas*. The main one is the class "loci" which is a simple data frame where the rows are individuals and the columns are loci and optional variables. The latter may be of any kind (continuous, integer, logical, etc). Loci are coded with R

factors (i.e., categorical variables) which are vectors of integers. Since computing on integers is fast, counting genotypic frequencies can be done efficiently. The convention is to code a genotype with the alleles separated with a forward slash, so that once genotypes are counted, allelic frequencies are easily obtained by counting the number of alleles in each genotype. Consequently, any level of ploidy may be handled. Since most operations on allelic data require counting allele and genotype frequencies, this operation has been optimised in the function `summary.loci` which is used by other functions in *pegas*.

The class "loci" inherits the class "data.frame" (a standard class in R) so all functionalities available for the latter (manipulation, subsetting, etc.) can be used for the former (Chambers, 2008). The typical example is in the use of the indexing operator `[, ]`. A special version of this operator has been written for the class "loci" so that the loci columns are correctly identified even after deletion of some rows and/or columns. Another advantage of the data frame structure is that each row is identified by a unique label (the `rownames`) so that these data may easily be matched with other data that have similar labels (e.g., a distance matrix, a DNA sequence alignment, or a genealogy).

R has many tools to analyse data frames and these may be used directly for the class "loci" and functions in *pegas*. A common task in population genetics is to perform some analyses for different subsets of the data (populations, breeds, plots, etc). For instance, if a data set `x` has a variable `population`, the R function `by` may be used to compute some statistics for each level of this variable, e.g., `by(x, x$population, hw.test)` to perform Hardy–Weinberg test. The second argument may be a list of variables in which case the analyses will be done for each combination of them, e.g., `by(x, list(x$plot, x$treatment), summary)` will compute allele and genotype frequencies for every combination of `plot` and `treatment`.

The second main special data structure of *pegas* is the class "haplotype" which inherits the class "DNABin" of *ape*. In addition to the set of unique DNA sequences, this class includes a vector of indices identifying the individuals belonging to each haplotype. This makes possible to link a set of haplotypes with individual data (phenotypes, geographic locations, etc).

Missing data are coded explicitly in *pegas*. In the class "loci" the standard NA is used and these are handled in the R standard way. In the class "haplotype", the IUPAC code for ambiguous nucleotides is used and this is taken into account during calculations.

\*[Emmanuel.Paradis@ird.fr](mailto:Emmanuel.Paradis@ird.fr)

	row_names	Fca25	Fca23	Fca43	Fca45	Fca77	Fca78	Fca90	Fca96	Fca37
1	0215	000/000	136/146	139/139	116/120	156/156	142/148	199/199	113/113	208/208
2	0216	000/000	146/146	138/145	120/126	156/156	140/148	185/199	113/113	208/208
3	0217	136/143	136/146	141/141	116/116	152/156	142/142	197/197	113/113	210/210
4	0218	133/135	138/138	139/141	116/126	150/150	142/148	199/199	091/106	208/208
5	0219	133/135	140/146	141/145	126/126	152/152	142/148	193/199	113/113	208/208
6	0220	135/143	136/146	145/149	120/126	150/156	148/148	193/199	091/113	208/208
7	0221	136/135	136/146	139/145	116/126	152/152	142/148	199/199	106/113	208/208
8	0222	136/143	136/146	136/149	120/126	154/158	142/148	193/197	091/091	208/212
9	0223	137/143	136/146	139/139	116/126	150/160	142/142	197/197	106/113	208/212
10	0224	136/135	132/132	141/145	120/126	150/156	148/148	197/197	091/106	208/208
11	07	137/141	120/136	137/145	128/128	152/152	142/150	193/199	091/091	182/182
12	041	129/133	120/126	135/145	126/126	144/150	140/140	197/199	091/113	182/208
13	042	129/133	120/120	135/145	120/120	152/156	142/142	197/199	091/091	202/202
14	043	133/133	120/126	135/135	120/120	156/156	142/142	199/199	091/091	182/206
15	044	131/135	136/136	137/137	126/120	152/152	140/142	199/199	091/091	202/202
16	045	129/135	136/146	135/135	120/120	144/144	142/142	193/199	091/091	182/182
17	046	129/133	120/144	133/133	126/126	144/144	140/140	191/191	091/091	182/182
18	047	129/135	138/138	135/135	120/126	146/158	140/148	191/199	091/113	182/182
19	048	136/135	136/144	135/145	126/126	146/156	140/150	185/191	091/091	182/182
20	049	131/135	130/136	135/137	120/126	152/158	140/150	191/199	101/121	208/208
21	051	129/133	136/136	135/145	128/120	146/158	142/142	193/199	091/113	182/206
22	053	131/135	136/136	135/145	128/128	146/158	140/148	193/199	113/113	182/208
23	054	133/135	120/120	137/145	128/128	148/160	142/142	193/199	091/091	208/208
24	056	131/133	136/146	136/135	126/120	144/148	142/142	191/191	113/113	182/182

Fig. 1. The R data editor showing microsatellite data. Note the first column allowing to edit the individual (row) labels.

The current version of *pegas* includes standard tools for population genetic analyses: genotypic and allelic frequencies, Hardy–Weinberg equilibrium,  $F_{ST}$ , analysis of molecular variance (AMOVA), haplotype network, mismatch distribution, Tajima's  $D$  and  $R_2$  tests for population stability, nucleotide diversity ( $\pi$ ), the population parameter  $\theta$  ( $= 4N_e\mu$ ), the site frequency spectrum, as well as several functions for reading and writing data files. *pegas* has also several low-level functions for manipulating its data structures (extracting observed alleles and genotypes, their frequencies, ploidy levels, building tables of all possible genotypes), making possible to extend its functionalities in a straightforward way. *pegas* includes basic tools for the coalescent such a computing likelihoods for a given tree, and estimating  $\theta$  by maximum likelihood. All functions are accompanied with a help page describing how the calculations are done and giving the relevant literature references. These pages are compiled in a reference manual in PDF available from the above website.

There are already a large number of computer programs for handling genetic data, so it is crucial for R users to be able to analyse the data files from the most widely used ones. *pegas* provides a function `read.loci` to read allelic data in tabular form from a text file. This function has several options to specify the loci and allele separator as well as which columns should be considered as loci or as additional variables. *pegas* has several functions to convert the data structures from *adegenet* (class "genind") to the class "loci". Since *adegenet* has functions to read files created for the programs *STRUCTURE*, *FSTAT*, *GENETIX*, and *GENEPOP* (`read.structure`, `read.fstat`, `read.genetix`, `read.genepop`), these file formats may also be readily used into *pegas* as well. Once read into R, and possibly converted into the class "loci", data can be edited by hand with the R spreadsheet data editor (Fig. 1). *pegas* is distributed with a tutorial explaining, step-by-step, how to input data from different file formats (in R type `vignette("ReadingFiles")`).

*pegas* uses the efficient and flexible graphical capabilities of R (Murrell, 2006). For instance, the options for plotting a haplotype network (Templeton *et al.*, 1992) are:

```
plot.haploNet(x, size = 1, col = "black",
  bg = "white", col.link = "black",
  lwd = 1, lty = 1, pie = NULL,
  labels = TRUE, scale.ratio = 1,
  legend = FALSE, fast = FALSE, ...)
```

These options make possible to plot symbols of different sizes, colours (contour and background), and the links between them may be of different colours, widths, or line types (solid, dotted, dashed, etc). These may be controlled by variables computed from the original data set with R's standard statistical and computing tools.

*pegas* illustrates the use of an integrated–modular approach for the development of software data analysis. The three packages *adegenet*, *ape*, and *pegas*, complement each other for population genetics as they provide functions for spatial and multivariate analyses (*adegenet*), trees structures and DNA sequences manipulation (*ape*), and basic population genetic analyses (*pegas*). Furthermore, each package has enough functionalities to be used on its own, independently of the others.

## ACKNOWLEDGEMENT

I am grateful to two anonymous referees for their comments on a previous version of this paper, and to Thibaut Jombart for useful discussions.

*Funding:* IRD program "Spirales", and European Science Foundation (ESF) through the MolArch (Molecular Archives of Past Climatic Changes) project.

## REFERENCES

- Chambers, J. M. (2008). *Software for Data Analysis: Programming with R*. Springer, New York.
- Excoffier, L. and Heckel, G. (2006). Computer programs for population genetics data analysis: a survival guide. *Nature Rev. Genet.* **7**, 745–758.
- Gentleman, R. (2008). *R Programming for Bioinformatics*. Chapman & Hall/CRC, Boca Raton.
- Jombart, T. (2008). *adegenet*: a R package for the multivariate analysis of genetic markers. *Bioinformatics*, **24**, 1403–1405.
- Murrell, P. (2006). *R Graphics*. Chapman & Hall/CRC, Boca Raton, FL.
- Paradis, E. (2006). *Analysis of Phylogenetics and Evolution with R*. Springer, New York.
- Paradis, E., Claude, J. and Strimmer, K. (2004). APE: analyses of phylogenetics and evolution in R language. *Bioinformatics*, **20**, 289–290.
- R Development Core Team (2009). *R: A Language and Environment for Statistical Computing*. R Foundation for Statistical Computing Vienna, Austria.
- Templeton, A. R., Crandall, K. A. and Sing, C. F. (1992). A cladistic analysis of phenotypic association with haplotypes inferred from restriction endonuclease mapping and DNA sequence data. III. Cladogram estimation. *Genetics*, **132**, 619–635.