Moran's Autocorrelation Coefficient in Comparative Methods

Emmanuel Paradis

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This document clarifies the use of Moran's autocorrelation coefficient to quantify whether the distribution of a trait among a set of species is affected or not by their phylogenetic relationships.

1 Theoretical Background

Moran's autocorrelation coefficient (often denoted as I) is an extension of Pearson product-moment correlation coefficient to a univariate series [2, 5]. Recall that Pearson's correlation (denoted as ρ) between two variables x and y both of length n is:

$$\rho = \frac{\sum_{i=1}^{n} (x_i - \bar{x})(y_i - \bar{y})}{\left[\sum_{i=1}^{n} (x_i - \bar{x})^2 \sum_{i=1}^{n} (y_i - \bar{y})^2\right]^{1/2}},$$

where \bar{x} and \bar{y} are the sample means of both variables. ρ measures whether, on average, x_i and y_i are associated. For a single variable, say x, I will measure whether x_i and x_j , with $i \neq j$, are associated. Note that with ρ , x_i and x_j are not associated since the pairs (x_i, y_i) are assumed to be independent of each other.

In the study of spatial patterns and processes, we may logically expect that close observations are more likely to be similar than those far apart. It is usual to associate a weight to each pair (x_i, x_j) which quantifies this [3]. In its simplest form, these weights will take values 1 for close neighbours, and 0 otherwise. We also set $w_{ii} = 0$. These weights are sometimes referred to as a neighbouring function.

I's formula is:

$$I = \frac{n}{S_0} \frac{\sum_{i=1}^{n} \sum_{j=1}^{n} w_{ij} (x_i - \bar{x})(x_j - \bar{x})}{\sum_{i=1}^{n} (x_i - \bar{x})^2},$$
(1)

where w_{ij} is the weight between observation i and j, and S_0 is the sum of all w_{ij} 's:

$$S_0 = \sum_{i=1}^n \sum_{j=1}^n w_{ij}.$$

Quite not so intuitively, the expected value of I under the null hypothesis of no autocorrelation is not equal to zero but given by $I_0 = -1/(n-1)$. The expected variance of I_0 is also known, and so we can make a test of the null hypothesis. If the observed value of I (denoted \hat{I}) is significantly greater than I_0 , then values of x are positively autocorrelated, whereas if $\hat{I} < I_0$, this will indicate negative autocorrelation. This allows us to design one- or two-tailed tests in the standard way.

Gittleman & Kot [4] proposed to use Moran's I to test for "phylogenetic effects". They considered two ways to calculate the weights w:

- With phylogenetic distances among species, e.g., $w_{ij} = 1/d_{ij}$, where $1/d_{ij}$ are distances measured on a tree.
- With taxonomic levels where $w_{ij} = 1$ if species i and j belong to the same group, 0 otherwise.

Note that in the first situation, there are quite a lot of possibilities to set the weights. For instance, Gittleman & Kot also proposed:

$$\begin{aligned} w_{ij} &= 1/d_{ij}^{\alpha} & \text{if } d_{ij} \leq c \\ w_{ij} &= 0 & \text{if } d_{ij} > c, \end{aligned}$$

where c is a cut-off phylogenetic distance above which the species are considered to have evolved completely independently, and α is a coefficient (see [4] for details). By analogy to the use of a spatial correlogram where coefficients are calculated assuming different sizes of the "neighbourhood" and then plotted to visualize the spatial extent of autocorrelation, they proposed to calculate I at different taxonomic levels.

2 Implementation in ape

From version 1.2-6, ape has functions Moran.I and correlogram.formula implementing the approach developed by Gittleman & Kot. There was an error in the help pages of ?Moran.I (corrected in ver. 2.1) where the weights were referred to as "distance weights". This has been wrongly interpreted in my book [6, pp. 139–142]. The analyses below aim to correct this.

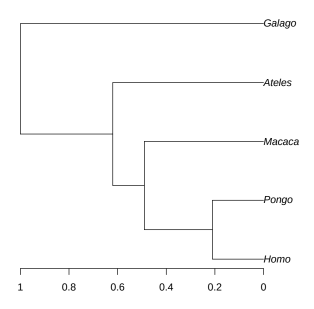
2.1 Phylogenetic Distances

The data, taken from [1], are the log-transformed body mass and longevity of five species of primates:

```
> body <- c(4.09434, 3.61092, 2.37024, 2.02815,
+     -1.46968)
> longevity <- c(4.74493, 3.3322, 3.3673, 2.89037,
+     2.30259)
> names(body) <- names(longevity) <- c("Homo", "Pongo",
+     "Macaca", "Ateles", "Galago")</pre>
```

The tree has branch lengths scaled so that the root age is one. We read the tree with <code>ape</code>, and plot it:

```
> library(ape)
> trnwk <- "((((Homo:0.21,Pongo:0.21):0.28,Macaca:0.49):0.13,Ateles:0.62)"
> trnwk[2] <- ":0.38,Galago:1.00);"
> tr <- read.tree(text = trnwk)
> plot(tr)
> axisPhylo()
```



We choose the weights as $w_{ij}=1/d_{ij}$, where the d's is the distances measured on the tree:

> w <- 1/cophenetic(tr) > w

	Homo	Pongo	Macaca	Ateles	Galago
Homo	Inf	2.3809524	1.0204082	0.8064516	0.5
Pongo	2.3809524	Inf	1.0204082	0.8064516	0.5
Macaca	1.0204082	1.0204082	Inf	0.8064516	0.5
Ateles	0.8064516	0.8064516	0.8064516	Inf	0.5
Galago	0.5000000	0.5000000	0.5000000	0.5000000	Inf

Of course, we must set the diagonal to zero:

> diag(w) <- 0

We can now perform the analysis with Moran's I:

```
> Moran.I(body, w)

$observed
[1] -0.0731218

$expected
[1] -0.25

$sd
[1] 0.08910814

$p.value
[1] 0.04714628
```

Not surprisingly, the results are opposite to those in [6] since, there, the distances (given by cophenetic(tr)) were used as weights. (Note that the argument dist has been since renamed weight.¹) We can now conclude for a slighly significant positive phylogenetic correlation among body mass values for these five species.

The new version of Moran.I gains the option alternative which specifies the alternative hypothesis ("two-sided" by default, i.e., H_1 : $I \neq I_0$). As expected from the above result, we divide the P-value be two if we define H_1 as $I > I_0$:

```
> Moran.I(body, w, alt = "greater")

$observed
[1] -0.0731218

$expected
[1] -0.25

$sd
[1] 0.08910814

$p.value
[1] 0.02357314

    The same analysis with longevity gives:
> Moran.I(longevity, w)

$observed
[1] -0.1837739

$expected
[1] -0.25
```

 $^{^{1}}$ The older code was actually correct; nevertheless, it has been rewritten, and is now much faster. The documentation has been clarified. The function correlogram.phylo, which computed Moran's I for a tree given as argument using the distances among taxa, has been removed.

\$sd

[1] 0.0911455

\$p.value

[1] 0.4674727

As for body, the results are nearly mirrored compared to [6] where a non-significant negative phylogenetic correlation was found: it is now positive but still largely not significant.

2.2 Taxonomic Levels

The function correlogram.formula provides an interface to calculate Moran's I for one or several variables giving a series of taxonomic levels. An example of its use was provided in [6, pp. 141-142]. The code of this function has been simplified, and the graphical presentation of the results have been improved.

correlogram.formula's main argument is a formula which is "sliced", and Moran.I is called for each of these elements. Two things have been changed for the end-user at this level:

- In the old version, the rhs of the formula was given in the order of the taxonomic hierarchy: e.g., Order/SuperFamily/Family/Genus. Not respecting this order resulted in an error. In the new version, any order is accepted, but the order given it is then respected when plotted the correlogram.
- Variable transformations (e.g., log) were allowed on the lhs of the formula.
 Because of the simplification of the code, this is no more possible. So it
 is the responsibility of the user to apply any tranformation before the
 analysis.

Following Gittleman & Kot [4], the autocorrelation at a higher level (e.g., family) is calculated among species belonging to the same category and to different categories at the level below (genus). To formalize this, let us write the different levels as $X^1/X^2/X^3/\ldots/X^n$ with X^n being the lowest one (Genus in the above formula):

$$w_{ij} = 1 \quad \text{if } X_i^k = X_j^k \text{ and } X_i^{k+1} \neq X_j^{k+1} \\ w_{ij} = 0 \quad \text{otherwise}$$

$$k < n$$

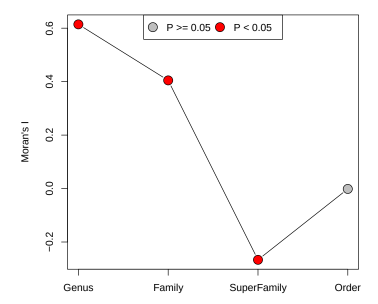
This is thus different from the idea of a "neighbourhood" of different sizes, but rather similar to the idea of partial correlation where the influence of the lowest level is removed when considering the highest ones [4].

To repeat the analyses on the carnivora data set, we first log_{10} -transform the variables mean body mass (SW) and the mean female body mass (FW):

- > data(carnivora)
- > carnivora\$log10SW <- log10(carnivora\$SW)</pre>
- > carnivora\$log10FW <- log10(carnivora\$FW)

We first consider a single variable analysis (as in [6]):

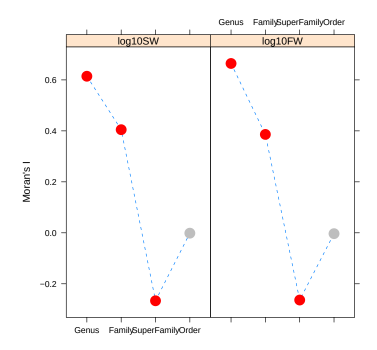
```
> fm1.carn <- log10SW ~ Order/SuperFamily/Family/Genus
> co1 <- correlogram.formula(fm1.carn, data = carnivora)
> plot(co1)
```



A legend now appears by default, but can be removed with legend = FALSE. Most of the appearance of the graph can be customized via the option of the plot method (see ?plot.correlogram for details). This is the same analysis than the one displayed on Fig. 6.3 of [6].

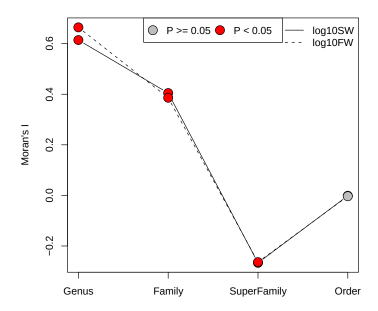
When a single variable is given in the lhs in correlogram.formula, an object of class "correlogram" is returned as above. If several variables are analysed simultaneously, the object returned is of class "correlogramList", and the correlograms can be plotted together with the appropriate plot method:

```
> fm2.carn <- log10SW + log10FW ~ Order/SuperFamily/Family/Genus
> co2 <- correlogram.formula(fm2.carn, data = carnivora)
> print(plot(co2))
```



By default, lattice is used to plot the correlograms on separate panels; using lattice = FALSE (actually the second argument, see ?plot.correlogramList) makes a standard graph superimposing the different correlograms:

> plot(co2, FALSE)



The options are roughly the same than above, but do not have always the same effect since lattice and base graphics do not have the same graphical parameters. For instance, legend = FALSE has no effect if lattice = TRUE.

3 Implementation in ade4

The analysis done with ade4 in [6] suffers from the same error than the one done with Moran. I since it was also done with a distance matrix. So I correct this below:

```
> library(ade4)
> gearymoran(w, data.frame(body, longevity))
class: krandtest
Monte-Carlo tests
Call: as.krandtest(sim = matrix(res$result, ncol = nvar, byr = TRUE),
    obs = res$obs, alter = alter, names = test.names)
Test number:
                      999
Permutation number:
Alternative hypothesis: greater
                    Obs
                          Std.Obs Pvalue
       body -0.06256789 2.1523342 0.001
2 longevity -0.22990437 0.3461414
                                   0.414
other elements: NULL
```

The results are wholly consistent with those from ape, but the estimated coefficients are substantially different. This is because the computational methods are not the same in both packages. In ade4, the weight matrix is first transformed as a relative frequency matrix with $\tilde{w}_{ij} = w_{ij}/S_0$. The weights are further transformed with:

$$p_{ij} = \tilde{w}_{ij} - \sum_{i=1}^{n} \tilde{w}_{ij} \sum_{j=1}^{n} \tilde{w}_{ij},$$

with p_{ij} being the elements of the matrix denoted as P. Moran's I is finally computed with $x^{T}Px$. In ape, the weights are first row-normalized:

$$w_{ij} / \sum_{i=1}^{n} w_{ij},$$

then eq. 1 is applied.

Another difference between both packages, though less important, is that in ade4 the weight matrix is forced to be symmetric with $(W + W^{T})/2$. In ape, this matrix is assumed to be symmetric, which is likely to be the case like in the examples above.

4 Other Implementations

Package sp has several functions, including moran.test, that are more specifically targeted to the analysis of spatial data. Package spatial has the function correlogram that computes and plots spatial correlograms.

Acknowledgements

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References

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