

Data and text mining

Assessing the shared variation among high-dimensional data matrices: a modified version of the Procrustean correlation coefficient

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Abstract

Motivation: Motivation: Molecular biology and ecology studies can produce high dimension data. Estimating correlations and shared variation between such data sets are an important step in detangling the relationships between different elements of a biological system. Unfortunately, classical approaches are susceptible to producing falsely inferred correlations.

Results: Here we propose a corrected version of the Procrustean correlation coefficient that is robust to high dimensional data. This allows for a correct estimation of the shared variation between two data sets and the partial correlation coefficients between a set of matrix data.

Availability: The proposed corrected coefficients are implemented in the ProcMod R package available on <https://git.metabarcoding.org/lecasofts/ProcMod>

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1 Introduction

Multidimensional data and even high-dimensional data, where the number of variables describing each sample is far larger than the sample count, is now routinely produced in functional genomics (*e.g.* transcriptomics, proteomics or metabolomics) and molecular ecology (*e.g.* DNA metabarcoding and metagenomics). Using a range of techniques, the same sample set can be described by several multidimensional data sets, each of them describing a different facet of the samples. This enables data analysis methods to evaluate mutual information shared by these different descriptions.

Correlative approaches are one of the simplest approaches to decipher pairwise relationships between multiple datasets. For a long time, several coefficients have been proposed to measure correlations between two matrices (for a comprehensive review see Ramsay *et al.*, 1984). However, when applied to high-dimensional data, these approaches suffer from over-fitting, resulting in high estimated correlations even for unrelated data sets. The creation of incorrect correlations from over-fitting consequently affects the biological interpretation of the analysis (Chariton *et al.* 2010) can have downstream effects on the biological interpretation of a study. A

number of modified matrix correlation coefficients have been proposed to address this issue. For example, the RV2 coefficient (Smilde *et al.*, 2009) corrects for overfitting of the original RV coefficient (Escofier, 1973). Similarly, a modified version of the distance correlation coefficient dCor (Székely *et al.*, 2007) proposed by Székely and Rizzo (2013) dCor has the advantage over the other correlation factors by considering by not being restricted to linear relationships.

Here we focus on the Procrustes correlation coefficient (RLs) proposed by Lingoes and Schönemann (1974) and by Gower (1971). Define *Trace*, a function summing the diagonal elements of a matrix. For an $n \times p$ real matrix \mathbf{X} and a second $n \times q$ real matrix \mathbf{Y} defining respectively two sets of p and q centered variables characterizing n individuals, we define $\text{CovLs}(\mathbf{X}, \mathbf{Y})$ following Equation (1)

$$\text{CovLs}(\mathbf{X}, \mathbf{Y}) = \frac{\text{Trace}((\mathbf{X}\mathbf{X}'\mathbf{Y}\mathbf{Y}')^{1/2})}{n-1} \quad (1)$$

and $\text{VarLs}(\mathbf{X})$ as $\text{CovLs}(\mathbf{X}, \mathbf{X})$. RLs can then be expressed as follow in Equation (2).

$$\text{RLs}(\mathbf{X}, \mathbf{Y}) = \frac{\text{CovLs}(\mathbf{X}, \mathbf{Y})}{\sqrt{\text{VarLs}(\mathbf{X}) \text{VarLs}(\mathbf{Y})}} \quad (2)$$

Considering $\text{CovLs}(\mathbf{X}, \mathbf{Y})$ and $\text{VarLs}(\mathbf{X})$, respectively corresponding to the covariance of two matrices and the variance of a matrix, Equation (2) highlighting the analogy between RLs and Pearson's correlation coefficient (R) (Bravais, 1844). When $p = 1$ and $q = 1$, $\text{RLs} = |\text{R}|$. Like the squared Pearson's R, the squared RLs is an estimate of the amount of variation shared between the two datasets.

Procrustean analyses have been proposed as a good alternative to Mantel's statistics for analyzing ecological data summarized by distance matrices (Peres-Neto and Jackson, 2001). In Procrustean analyze, distance matrices are projected into an orthogonal space using metric or non metric multidimensional scaling according to the geometrical properties of the used distances. Correlations can then be estimated on these projections.

2 Approach

RLs is part of the Procruste framework that aims to superimpose a set of points with respect to another through three operations: a translation, a rotation, and a scaling. The optimal transfer matrix $\text{Rot}_{X \rightarrow Y}$ can be estimated from the singular value decomposition (SVD) of the covariance matrix $\mathbf{X}'\mathbf{Y}$. SVDs factorize any matrix as the product of three matrices (Equation 3).

$$\mathbf{X}'\mathbf{Y} = \mathbf{U}\mathbf{\Sigma}\mathbf{V}' \quad (3)$$

\mathbf{U} and \mathbf{V} are two rotation matrices which computes the transfer matrix to superimpose \mathbf{X} on \mathbf{Y} or reciprocally \mathbf{Y} on \mathbf{X} . All the elements of $\mathbf{\Sigma}$ except its diagonal are equal to zero. The diagonal elements are the singular values. Singular values are the extension of the eigenvalues to non-square matrices. $\text{CovLs}(\mathbf{X}, \mathbf{Y})$ can also be computed from singular values (Equation 4).

$$\text{CovLs}(\mathbf{X}, \mathbf{Y}) = \frac{\text{Trace}(\mathbf{\Sigma})}{n-1} \quad (4)$$

This expression actually illustrates that $\text{CovLs}(\mathbf{X}, \mathbf{Y})$ is the variance of the projections of \mathbf{X} on \mathbf{Y} or of the reciprocal projections. Therefore $\text{CovLs}(\mathbf{X}, \mathbf{Y})$ and $\text{RLs}(\mathbf{X}, \mathbf{Y})$ are always positive and rotation independant. Here we propose to partitionate $\text{Trace}(\mathbf{\Sigma})$, the variation amount corresponding to $\text{CovLs}(\mathbf{X}, \mathbf{Y})$, in two components. The first corresponds to the actual shared information between \mathbf{X} and \mathbf{Y} . The second, corresponds to the over-fitting effect. It that can be estimated as the average variation shared by two random matrices of same structure as \mathbf{X} and \mathbf{Y} noted $\overline{\text{RCovLs}}(\mathbf{X}, \mathbf{Y})$. $\text{ICovLs}(\mathbf{X}, \mathbf{Y})$, the informative part of $\text{CovLs}(\mathbf{X}, \mathbf{Y})$, is computed using Equation (5).

$$\text{ICovLs}(\mathbf{X}, \mathbf{Y}) = \text{Max} \begin{cases} \text{CovLs}(\mathbf{X}, \mathbf{Y}) - \overline{\text{RCovLs}}(\mathbf{X}, \mathbf{Y}) \\ 0 \end{cases} \quad (5)$$

Similarly the informative counter-part of $\text{VarLs}(\mathbf{X})$ is defined as $\text{IVarLs}(\mathbf{X}) = \text{ICovLs}(\mathbf{X}, \mathbf{X})$, and $\text{IRLs}(\mathbf{X}, \mathbf{Y})$ the informative Procruste correlation coefficient as defined in Equation (6).

$$\text{IRLs}(\mathbf{X}, \mathbf{Y}) = \frac{\text{ICovLs}(\mathbf{X}, \mathbf{Y})}{\text{IVarLs}(\mathbf{X}) \text{IVarLs}(\mathbf{Y})} \quad (6)$$

As in the case of $\text{RLs}(\mathbf{X}, \mathbf{Y})$, $\text{IRLs}(\mathbf{X}, \mathbf{Y}) \in [0; 1]$, the 0 value corresponding to no correlation and the maximum value 1 reflects two strictly homothetic data sets.

The corollary of $\text{ICovLs}(\mathbf{X}, \mathbf{Y})$ and $\text{IVarLs}(\mathbf{X})$ definitions is that $\text{ICovLs}(\mathbf{X}, \mathbf{Y}) \geq 0$ and $\text{IVarLs}(\mathbf{X}) > 0$. Therefore for $M = \{\mathbf{M}_1, \mathbf{M}_2, \dots, \mathbf{M}_k\}$ a set of k matrices with the same number of rows, the informative covariance matrix \mathbf{C} defined as $\mathbf{C}_{i,j} = \text{ICovLs}(\mathbf{M}_i, \mathbf{M}_j)$ is definite positive and symmetrical. This allows for defining the precision

matrix $\mathbf{P} = \mathbf{C}^{-1}$ and the related partial correlation coefficient matrix $\text{IRLs}_{\text{partial}}$ using Equation (7)

$$\text{IRLs}_{\text{partial}}(\mathbf{M}_i, \mathbf{M}_j) = \frac{\mathbf{P}_{i,j}}{\sqrt{\mathbf{P}_{i,i} \mathbf{P}_{j,j}}} \quad (7)$$

3 Methods

3.1 Monte-Carlo estimation of $\overline{\text{RCovLs}}(\mathbf{X}, \mathbf{Y})$

For every values of p and q including 1, $\overline{\text{RCovLs}}(\mathbf{X}, \mathbf{Y})$ can be estimated using a series of k random matrices $\mathbf{RX} = \{\mathbf{RX}_1, \mathbf{RX}_2, \dots, \mathbf{RX}_k\}$ and $\mathbf{RY} = \{\mathbf{RY}_1, \mathbf{RY}_2, \dots, \mathbf{RY}_k\}$ where each \mathbf{RX}_i and \mathbf{RY}_i have the same structure as \mathbf{X} and \mathbf{Y} , respectively, in terms of number of columns and of the covariance matrix of these columns.

$$\overline{\text{RCovLs}}(\mathbf{X}, \mathbf{Y}) = \frac{\sum_{i=1}^k \text{CovLs}(\mathbf{RX}_i, \mathbf{RY}_i)}{k} \quad (8)$$

To estimate $\text{IVarLs}(\mathbf{X})$, which is equal to $\text{ICovLs}(\mathbf{X}, \mathbf{X})$, $\overline{\text{RCovLs}}(\mathbf{X}, \mathbf{X})$ is estimated with two independent sets of random matrix \mathbf{RX} and \mathbf{RY} , both having the same structure than \mathbf{X} .

Empirical assessment of $\overline{\text{RCovLs}}(\mathbf{X}, \mathbf{Y})$

For two random vectors \mathbf{x} and \mathbf{y} of length n , the average coefficient of determination is $\text{R}^2 = 1/(n-1)$. This value is independent of the distribution of the \mathbf{x} and \mathbf{y} values, but what about the independence of $\overline{\text{RCovLs}}(\mathbf{X}, \mathbf{Y})$ to the distributions of \mathbf{X} and \mathbf{Y} ? To test this independance and to assess the reasonable randomization effort needed to estimate $\overline{\text{RCovLs}}(\mathbf{X}, \mathbf{Y})$, this value is estimated for four matrices $\mathbf{K}, \mathbf{L}, \mathbf{M}, \mathbf{N}$ of $n = 20$ rows and 10, 20, 50, 100 columns, respectively. Values of the four matrices are drawn from a normal or an exponential distribution, with $k \in \{10, 100, 1000\}$ randomizations tested to estimate $\overline{\text{RCovLs}}(\mathbf{X}, \mathbf{Y})$ and the respective standard deviation $\sigma(\overline{\text{RCovLs}}(\mathbf{X}, \mathbf{Y}))$. The VarLs of the generated matrices is equal to 1, therefore the estimated CovLs are equals to the RLs.

3.2 Simulating data for testing sensibility to overfitting

To test overfitting, correlations were mesured between two random matrices of same dimensions. Each matrix is $n \times p$ with $n = 20$ and $p \in [2, 50]$. Each p variables were drawn from a centered and reduced normal distribution $\mathcal{N}(0, 1)$. Eight correlation coefficients were tested: RLs the original procruste coefficient; IRLs this work; RV the original R for vector data (Robert and Escoufier, 1976); RV adj Maye , RV 2 and RV adj Ghaziri three modified versions of RV (El Ghaziri and Qannari, 2015; Mayer et al., 2011; Smilde et al., 2009); dCor the original distance correlation coefficient (Székely et al., 2007); and $\text{dCor}_{\text{ttest}}$, a modified version of dCor not sensible to overfitting (Székely and Rizzo, 2013). For each p value, 100 simulations were run. Computation of IRLs were estimated with 100 randomizations.

For $p = 1$, random vectors with $n \in [3, 25]$ are generated. As above, data were drawn from $\mathcal{N}(0, 1)$ and $k = 100$ simulations which were run for each n . The original Pearson correlation coefficient R and the modified version IR are used to estimate correlation between both vectors.

3.3 Empirical assessment of the coefficient of determination

As in the case of the coefficient of determination (R^2), RLs^2 represents the part of shared variation between two matrices. Because of over-fitting in high-dimension data, RLs and therefore RLs^2 are over-estimated.

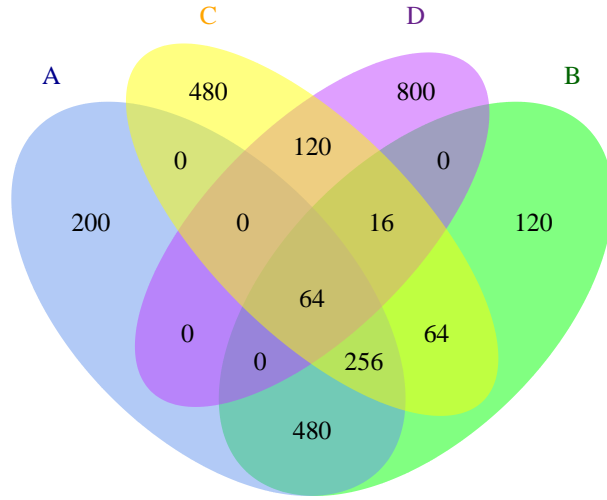


Fig. 1. Theoretical distribution of the shared variation between the four matrices (A, B, C, D), expressed in permille.

Between two matrices

To test how the IRLs version of the coefficient of determination $IRLs^2$ can perform to evaluate the shared variation, pairs of random matrices were produced for two values of $p \in \{10, 100\}$ and $n \in \{10, 25\}$, and for several levels of shared variations ranging between 0.1 and 1 using 0.1 increments. For each combination of parameters, $k = 100$ simulations were run, and both RLs^2 and $IRLs^2$ were estimated using 100 randomizations.

Between two vectors

Coefficient of determination between two vectors also suffers from over-estimation when n , the number of considered points, is small. On average, for two random vectors of size n , $\bar{R}^2 = 1/(n - 1)$. This random part of the shared variation inflates the observed shared variation even for non-random vectors. In the context of multiple linear regression, Theil *et al.* (1958) proposed an adjusted version of the coefficient of determination (Equation 9), correcting for both the effect of the number of vectors (p) and the vector size (n).

$$R_{adj}^2 = 1 - (1 - R^2) \frac{n - 1}{n - p - 1} \quad (9)$$

To evaluate the strength of that over-estimation and the relative effect of the correction proposed by Theil *et al.* (1958) and by IRLs, pairs of random vectors were produced for $n \in \{10, 25\}$, and for several levels of shared variations ranging between 0.1 and 1 using 0.1 increments. For each combination of parameters, $k = 100$ simulations were run, and R^2 , R_{adj}^2 and $IRLs^2$ were estimated using 100 randomizations.

Partial determination coefficients

To evaluate the capacity of partial determination coefficient $IRLs_{partial}^2$ to distangle nested correlations, a set of correlated matrices were generated. To generate two random matrices **A**, **B**, sharing $w \in [0, 1]$ part of variation, two independent random matrices **A** and **Δ** were generated such as $\text{VarLs}(\mathbf{A}) = 1$ and $\text{VarLs}(\mathbf{\Delta}) = 1$. The $\mathbf{\Delta}_{rot}$ matrix was computed as the alignment of **Δ** on **A** using the optimal procruste rotation. Then **B** is computed using equation 10 :

$$\mathbf{B} = \mathbf{A} \times \sqrt{w} + \mathbf{\Delta}_{rot} \times \sqrt{1 - w}. \quad (10)$$

Following this method, four matrices **A**, **B**, **C**, and **D** of size $n \times p = 20 \times 200$ were generated where **A** shares 80% of variation with **B**, that shares 40% of variation with **C**, sharing 20% of variation with **D**. As illustrated in Figure 1, These direct correlations induce indirect ones spreading the total variation among each pair of matrices. The simulation was repeated 100 times, for every simulation $IRLs_{partial}^2$ and $RLs_{partial}^2$ were estimated for each pair of matrices.

3.4 Testing the significance of $IRLs(\mathbf{X}, \mathbf{Y})$

The significance of $IRLs(\mathbf{X}, \mathbf{Y})$ can be tested using permutation test as defined in Jackson (1995) or Peres-Neto and Jackson (2001) and implemented respectively in the `protest` method of the `vegan` R package (Dixon, 2003) or the `procuste.rtest` method of the `ADE4` R package Dray and Dufour (2007).

It is also possible to take advantage of the Monte-Carlo estimation of $\text{RCovLs}(\mathbf{X}, \mathbf{Y})$ to test that $\text{ICovLs}(\mathbf{X}, \mathbf{Y})$ and therefore $IRLs(\mathbf{X}, \mathbf{Y})$ are greater than expected under random hypothesis. Over the k randomizations, $N_{>\text{CovLs}}$ is estimated by counting when $\text{RCovLs}(\mathbf{X}, \mathbf{Y})_k > \text{CovLs}(\mathbf{X}, \mathbf{Y})$. The P_{value} of the test then can be estimated following Equation (11).

$$P_{\text{value}} = \frac{N_{>\text{CovLs}}}{k} \quad (11)$$

Empirical assessment of α -risk for the CovLs test

To empirically assess the α -risk of the Procruste test based on the randomisations realized during the estimation of $\text{RCovLs}(\mathbf{X}, \mathbf{Y})$, the distribution of P_{value} under H_0 was compared to a uniform distribution between 0 and 1 ($\mathcal{U}(0, 1)$). To estimate the empirical distribution, $k = 1000$ pairs of $n \times p$ random matrices with $n = 20$ and $p \in \{10, 20, 50\}$ were simulated under the null hypothesis of independence. Significance of the Procruste correlation between those matrices was tested using the three approaches: our proposed test (`CovLs.test`) ; the `protest` method of the `vegan` R package ; the `procuste.rtest` method of the `ADE4` R package. Conformance of the distribution of each set of k P_{value} to $\mathcal{U}(0, 1)$ was assessed using the Cramer-Von Mises test (Csörgő and Faraway, 1996) implemented in the `cvm.test` function of the R package `gofest`.

Empirical power assessment for the CovLs test

To evaluate the relative the power of the three tests described above, pairs of two random matrices were produced for various $p \in \{10, 20, 50, 100\}$, $n \in \{10, 15, 20, 25\}$ and two levels of shared variations $R^2 \in \{0.05, 0.1\}$. For each combination of parameters, $k = 1000$ simulations were run. Each test were estimated based on 1000 randomizations for the `CovLs` test, or 1000 permutations for `protest` and `procuste.rtest`.

4 Results

4.1 Empirical assessment of $\text{RCovLs}(\mathbf{X}, \mathbf{Y})$

Two main parameters can influence the Monte Carlo estimation of $\text{RCovLs}(\mathbf{X}, \mathbf{Y})$: the distribution used to generate the random matrices, and k the number of random matrix pair. Two very different distribution are tested to regenerate the random matrices, the normal and the exponential distributions. The normal distribution is symmetric where the exponential is unsymmetrical with a high probability for small values and a long tail of large ones. Despite the use of these contrasted distributions, estimates of $\text{RCovLs}(\mathbf{X}, \mathbf{Y})$ and of $\sigma(\text{RCovLs}(\mathbf{X}, \mathbf{Y}))$ were identical if we

Table 1. Estimation of $\overline{RCovLs(X, Y)}$ according to the number of random matrices (k) aligned.

p	k	normal		exponential	
		mean	sd	mean	sd
10	10	0.5903	1.3513×10^{-2}	0.6106	1.6980×10^{-2}
	100	0.5812	3.1636×10^{-3}	0.5849	3.2563×10^{-3}
	1000	0.5806	1.1564×10^{-3}	0.5801	1.1515×10^{-3}
20	10	0.7642	5.3833×10^{-3}	0.7760	6.2222×10^{-3}
	100	0.7639	2.3330×10^{-3}	0.7676	1.9466×10^{-3}
	1000	0.7658	6.8923×10^{-4}	0.7671	6.6033×10^{-4}
50	10	0.9089	2.6282×10^{-3}	0.9086	2.4032×10^{-3}
	100	0.9089	8.1126×10^{-4}	0.9092	9.2665×10^{-4}
	1000	0.9086	2.8590×10^{-4}	0.9091	2.8450×10^{-4}
100	10	0.9524	1.4734×10^{-3}	0.9536	1.3515×10^{-3}
	100	0.9541	4.9602×10^{-4}	0.9550	4.6578×10^{-4}
	1000	0.9544	1.4547×10^{-4}	0.9546	1.4249×10^{-4}

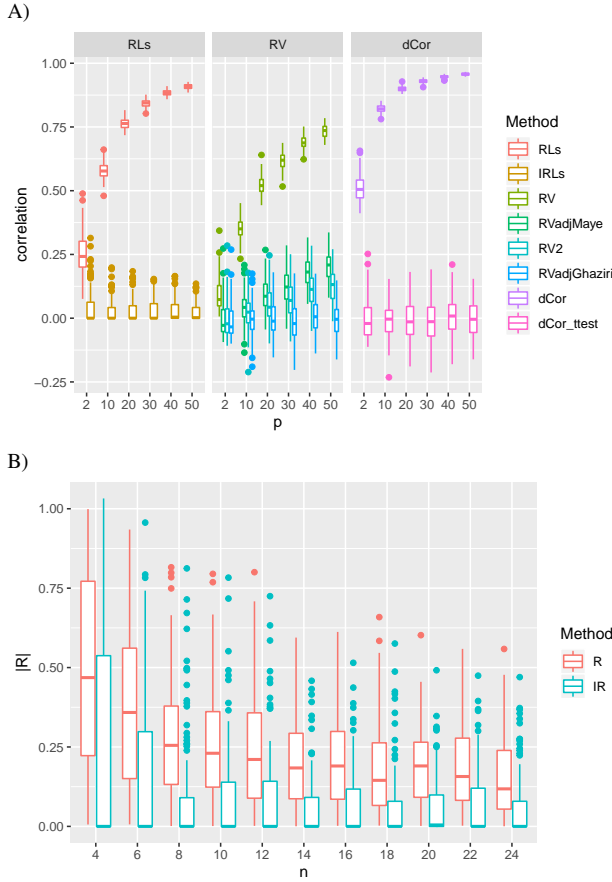


Fig. 2. A) Susceptibility to overfitting for various correlation coefficients. (A) Both simulated data sets are matrices of size $(n \times p)$ with $p > 1$. B) Correlated data sets are vectors ($p = 1$) with a various number of individuals n (vector length). For both A & B, 100 simulations were run for each combination of parameters.

assumed the normal distribution of the $\overline{RCovLs(X, Y)}$ estimator and a 0.95 confidence interval of $\overline{RCovLs(X, Y)} \pm 2\sigma(\overline{RCovLs(X, Y)})$ (Table 1).

4.2 Relative susceptibility of $IRLs(X, Y)$ to overfitting

RLs , like RV and $dCor$, is susceptible to overfitting which increases when n decreases, and p or q increase. Because RV is more comparable

to R^2 , when RLs and $dCor$ are more comparable to R , RV values increase more slowly than RLs and $dCor$ values with p (Figure 2A). As expected $IRLs$ values for non-correlated matrices are close to 0 regardless of p (Figure 2A).

4.3 Evaluating the shared variation

Between two matrices

For two matrices, our proposed corrected version of RLs^2 ($IRLs^2$) provided a good estimate of the shared variation and was robust to the phenomenon of over-fitting (Figure 3). Only a small over estimation was observed when $p = 10$ for the lowest values of the simulated shared variations (≤ 0.4).

Between two vectors

Vectors can be considered as a single column matrix, and the efficiency of $IRLs^2$ to estimate shared variation between matrices can also be used to estimate shared variation between two vectors. Other formulas have been already proposed to better estimate shared variation between vectors in the context of linear models. Among them the one presented in Equation 9, is the most often used and is the one implemented in R linear model summary function. On simulated data, $IRLs^2$ performs better than the simple R^2 and its modified version R_{adj}^2 commonly used (Figure 4). Whatever the estimator the bias decrease with the simulated shared variation. Nevertheless for every tested cases the median of the bias observed is smaller than with both other estimators, even if classical estimators well perform for large values of shared variation.

Partial coefficient of determination

The simulated correlation network between the four matrices **A**, **B**, **C**, **D** induced moreover the direct simulated correlation a network of indirect correlation and therefore shared variances (Figure 1). In such system, the interest of partial correlation coefficients and their associated partial determination coefficients is to measure correlation between a pair of variable without accounting for the part of that correlation which is explained by other variables, hence extracting the pure correlation between these two matrices. From Figure 1, the expected partial shared variation between **A** and **B** is $480/(200 + 480) = 0.706$; between **B** and **C**, $64/(480 + 120) = 0.107$; and between **C** and **D** $120/800 = 0.150$. All other partial coefficient are expected to be equal to 0. The effect of the correction introduced in $IRLs$ is clearly weaker and on the partial coefficient of determination (Figure 5) than on the full coefficient of determination (Figure 3). The spurious random correlations, constituting the over-fitting effect, is distributed over all the pair of matrices **A**, **B**, **C**, **D**.

4.4 p_{value} distribution under null hypothesis

As expected, P_{values} of the $CovLs$ test based on the estimation of $\overline{RCovLs(X, Y)}$ are uniformly distributed under H_0 . whatever the p tested (Table 2). This ensure that the probability of a $P_{value} \leq \alpha$ -risk is equal to α -risk. Moreover P_{values} of the $CovLs$ test are strongly linearly correlated with those of both the other tests ($R^2 = 0.996$ and $R^2 = 0.996$ respectively for the correlation with `vegan::protest` and `ade4::procuste.rtest` P_{values}). The slopes of the corresponding linear models are respectively 0.997 and 0.998.

4.5 Power of the test based on randomisation

Power of the $CovLs$ test based on the estimation of $\overline{RCovLs(X, Y)}$ is equivalent of the power estimated for both `vegan::protest` and `ade4::procuste.rtest` tests (Table 3). As for the two other tests, power decreases when the number of variable (p or q) increases,

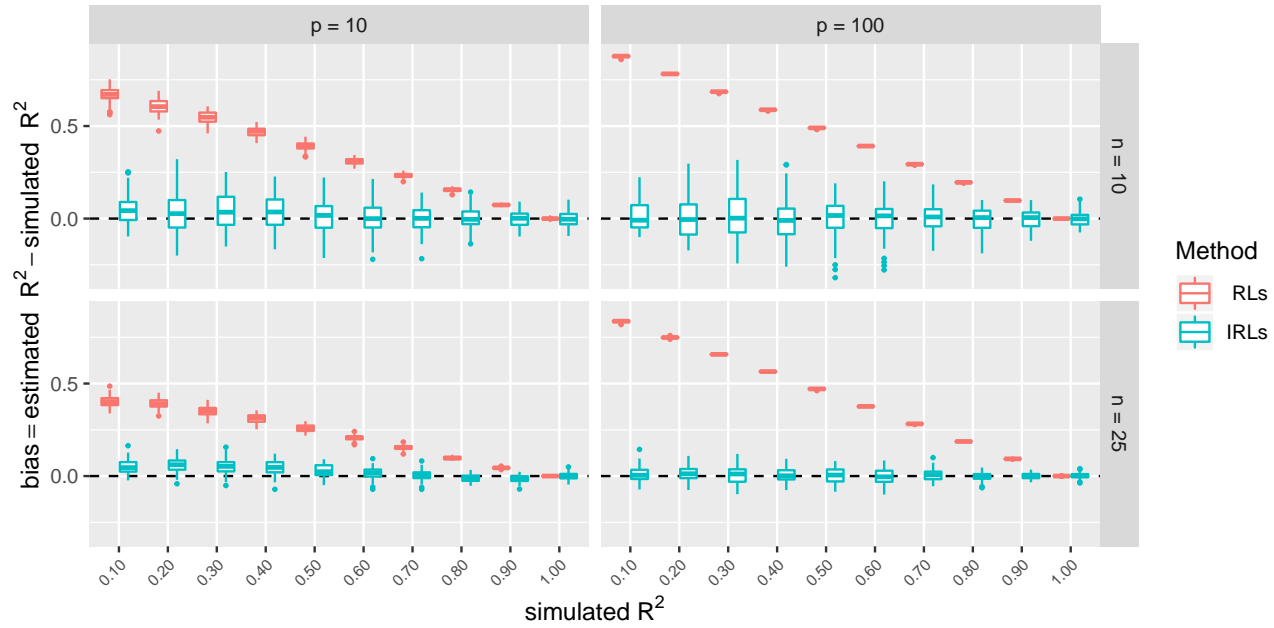


Fig. 3. Shared variation (R^2) between two matrices has measured using both the corrected (IRLs) and the original (RLs) versions of the Procrustean correlation coefficient. A gradient of R^2 was simulated for two population sizes ($n \in \{10, 25\}$) and two numbers of descriptive variables ($p \in \{10, 100\}$). The distribution of differences between the observed and the simulated shared variation is plotted for each condition. The black dashed line corresponds to a perfect match where measured R^2 equals the simulated one.

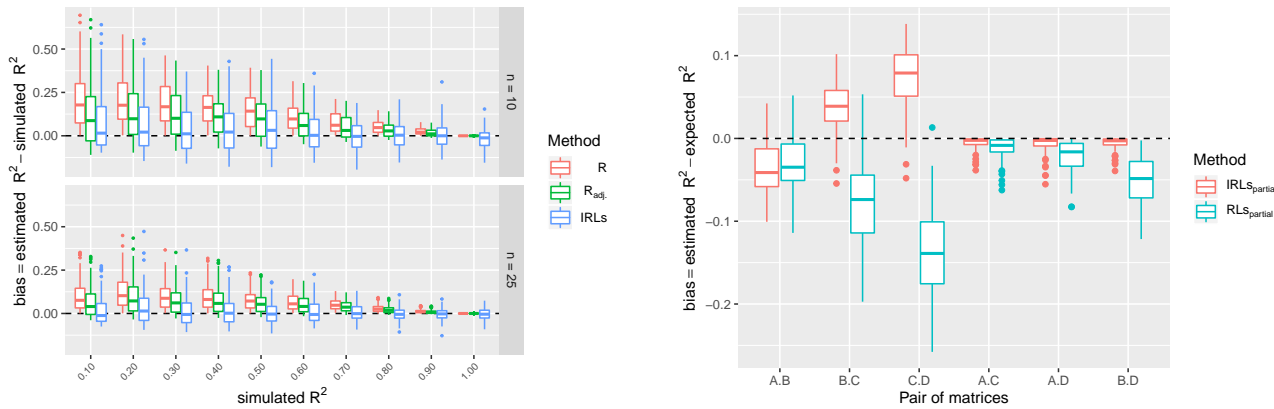


Fig. 4. Shared variation between two vectors is measured using the classical R^2 , its adjusted version R_{adj}^2 and (IRLs 2). A gradient of shared variation is simulated for two vector sizes ($n \in \{10, 25\}$). The black dashed line corresponds to a perfect match where measured R^2 equals the simulated one.

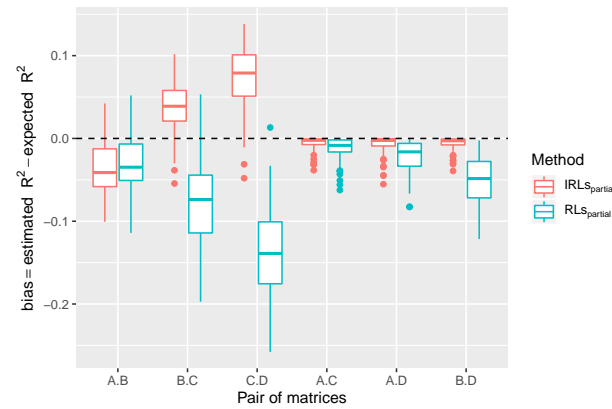


Fig. 5. Estimation error on the partial determination coefficient. Error is defined as the absolute value of the difference between the expected and the estimated partial R^2 using the corrected IRLs_{partial} and not corrected RLs_{partial} procruste correlation coefficient.

Table 2. P_{values} of the Cramer-Von Mises test of conformity of the distribution of P_{values} correlation test to $\mathcal{U}(0, 1)$ under the null hypothesis.

p	Cramer-Von Mises p.value		
	CovLs test	protest	procuste.rtest
10	0.226	0.250	0.194
20	0.682	0.682	0.687
50	0.557	0.532	0.551

and increase with the number of individuals and the shared variation. The advantage of the test based on the Monte-Carlo estimation of $RCovLs(X, Y)$ is to remove the need of running a supplementary set of permutations when IRLs is computed.

5 Discussion

Correcting the over-adjustment effect on metrics assessing the relationship between high dimension datasets has been a constant effort over the past decades. Therefore, IRLs can be considered as a continuation of the extension of the toolbox available to biologists for analyzing their omics data. The effect of the proposed correction on the classical RLs coefficient is as strong as the other ones previously proposed for other correlation coefficients measuring relationship between vector data (see Figure 3, e.g. Smilde *et al.*, 2009; Székely and Rizzo, 2013). When applied to univariate data, RLs is equal to the absolute value of the Pearson correlation coefficient, hence, and despite it is not the initial aim of that coefficient, IRLs can also be used to evaluate correlation between two univariate datasets. Using IRLs for such data sets is correcting for spurious

Table 3. Power estimation of the procruste tests for two low level of shared variations 5% and 10%.

	R^2	5%				10%				
		p	10	20	50	100	10	20	50	100
		n	$power = 1 - \beta\text{-risk}$							
Covls.test	10	0.49	0.45	0.40	0.45	0.76	0.68	0.70	0.68	
	15	0.88	0.80	0.75	0.75	0.99	0.98	0.96	0.95	
	20	0.99	0.96	0.94	0.93	1.00	1.00	1.00	1.00	
	25	1.00	1.00	0.99	0.98	1.00	1.00	1.00	1.00	
protest	10	0.50	0.45	0.40	0.45	0.77	0.70	0.70	0.68	
	15	0.88	0.80	0.75	0.75	0.99	0.98	0.96	0.95	
	20	0.99	0.96	0.94	0.93	1.00	1.00	1.00	1.00	
	25	1.00	1.00	0.99	0.99	1.00	1.00	1.00	1.00	
procuste.rtest	10	0.50	0.45	0.41	0.45	0.76	0.69	0.70	0.68	
	15	0.88	0.80	0.74	0.75	0.99	0.98	0.96	0.96	
	20	0.99	0.96	0.94	0.93	1.00	1.00	1.00	1.00	
	25	1.00	1.00	0.99	0.99	1.00	1.00	1.00	1.00	

correlations when the number of individual is small more efficiently than classical correction (see Figure 4, Theil *et al.*, 1958).

The main advantage of IRLs over other matrix correlation coefficients is that it allows for estimating shared variation between two matrices according to the classical definition of variance partitioning used with linear models. This opens the opportunity to develop linear models to explain the variation of a high dimension dataset by a set of other high dimension data matrices.

The second advantage of IRLs is that its definition implies that the variance/co-variance matrix of a set of matrices is positive-definite. That allows for estimating partial correlation coefficients matrix by inverting the variance/co-variance matrix. The effect of the correction is less strong on such partial coefficients than on full correlation, but the partial coefficients that should theoretically be estimated to zero seem to be better identified after the correction.

6 Conclusion

A common approach to estimate strength of the relationship between two variables is to estimate the part of shared variation. This single value ranging from zero to one is easy to interpret. Such value can also be computed between two sets of variable, but the estimation is more than for simple vector data subject to over estimation because the over-fitting phenomena which is amplified for high dimensional data. With IRLs and its squared value, we propose an easy to compute correlation and determination coefficient far less biased than the original Procrustean correlation coefficient. Every needed function to estimate the proposed modified version of these coefficients are included in a R package ProcMod available for download from the Comprehensive R Archive Network (CRAN).

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Appendix

A Notations

\mathbf{x} (vector)	bold lowercase.
\mathbf{X} (matrix)	bold uppercase.
$i = 1, \dots, n$	object index.
$j = 1, \dots, p$	variable index.
k	iteration index.
\mathbf{X}'	The transpose of \mathbf{X} .
\mathbf{XY}	Matrix multiplication of \mathbf{X} and \mathbf{Y} .
$\text{Diag}(\mathbf{X})$	A column matrix composed of the diagonal elements of \mathbf{X} .
$\mathbf{X}^{1/2}$	Matrix square root of \mathbf{X} .
$\text{Trace}(\mathbf{X})$	The trace of \mathbf{X} .