Modularity and Morphometrics: Error Rates in Hypothesis Testing

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06 November 2015

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running title: Modularity, Error and Morphometrics

key words: P-matrix; statistical power; primates; development; genotype/phenotype map

Abstract

The study of modularity in morphological systems has increased in the past twenty years, parallel to the popularization of geometric morphometrics, which has led to the emergence of different criteria for detecting modularity on landmark data. However, compared to usual covariance matrix estimators, Procrustes estimators have properties that hinder their use. Here, we compare different representations of form, focusing on detecting modularity patterns defined a priori; we also compare two metrics: one derived from traditional morphometrics (MHI) and another that emerged in the context of landmark data (RV). Using Anthropoid skulls, we compare these metrics over three representations of form: interlandmark distances, Procrustes residuals, and local shape variables. Over Procrustes residuals, both tests fail to detect modularity patterns, while in remaining representations they show the distinction between early and late development in skull ontogeny. To estimate type I and II error rates, we built covariance matrices of known structure; these tests indicate that, considering both effect and sample sizes, tests using MHI are more robust than those using RV. However, both metrics have low power when used on Procrustes residuals. Thus, we conclude that the influence of development and function is poorly represented on Procrustes estimators for covariance matrices.
Introduction

Modularity is a characteristic property that biological systems exhibit regarding the distribution of interactions between their composing elements; that is, in a given system, subsets of elements, denominated modules, interact more among themselves than with other such subsets (Newman, 2006; Mitteroecker & Bookstein, 2007; Wagner et al., 2007). This property has been well documented at different levels of biological organization, from the dynamics of metabolic networks (e.g. Ravasz et al., 2002; Andrade et al., 2011) to the structure of interactions among individuals in populations (e.g. Fortuna et al., 2008) and among species in ecological communities (e.g. Genini et al., 2010).

Regarding morphological systems, the concept of modularity is associated with the framework of morphological integration (Olson & Miller, 1958; Cheverud, 1996), which refers to the organization of covariances or correlations between morphological elements and the hypotheses concerning their relationships. In this context, modularity refers to the uneven distribution of genetic effects over phenotypic variation articulated through development (genotype/phenotype map; Wagner, 1996); in a classical quantitative genetics view, these genetic effects are the result of pleiotropy and linkage disequilibrium (Falconer & Mackay, 1996; Lynch & Walsh, 1998). A genotype/phenotype map composed of clusters of genes that affect clusters of traits (with little overlap) exhibits a modular organization; this structure is thought to emerge as the result of selection for distinct demands (Wagner & Altenberg, 1996; Espinosa-Soto & Wagner, 2010; Rueffler et al., 2012; Melo & Marroig, 2015). For instance, the decoupling between fore- and hindlimb function in certain mammalian lineages such as bats (Young & Hallgrímsson, 2005) and apes (Young et al., 2010) is associated with the modularization of both structures, as shown by reduced phenotypic correlations between fore- and hindlimbs and increased correlations between elements within these limbs.

The recognition of variational modules (Wagner & Altenberg, 1996; Wagner et al., 2007) using covariance or correlation patterns in adult populations involves an understanding of the underlying developmental and functional dynamics among morphological traits (Polly, 2008; Zelditch & Swiderski, 2011). Skull development in mammals is composed of a series of steps, such as neurocranial growth induced by brain development, and growth mediated by muscle-bone interactions, with spatiotemporal overlapping between such steps (Hallgrímsson & Lieberman, 2008; Herring, 2011; Cardini & Polly, 2013). Both timing and scope of each step is regulated by different profiles of genetic expression exhibited by cells originated from different embryonic precursors and their response to signaling factors.
expressed at the regional level; the response to these signaling factors further changes
cell expression profiles, thus generating a feedback loop of differentiation (Turing, 1952;
Marcucio et al., 2005; Meinhardt, 2008; Hallgrímsson et al., 2009; Franz-Odendaal, 2011;
Minelli, 2011). Each step in this temporal hierarchy may be regarded as modular, since
they affect a coherent subset of tissues more so than others (Hallgrímsson & Lieberman,
2008), although each step affect adjoining regions through interactions among developing
tissues (Cheverud et al., 1992; Lieberman, 2011; Esteve-Altava & Rasskin-Gutman, 2014);
thus, the overlapping of such processes throughout development may complicate their
association with correlation patterns (Hallgrímsson et al., 2009).

Furthermore, variation in growth rates, which emerges into size variation (Pélabon et
al., 2013; Porto et al., 2013), has a particular importance in the context of mammalian
morphological systems. Here size variation refers to variation in both scale (isometric
variation) and scale relationships (allometric variation). This source of variation affects
the overall level of correlations between morphological traits (Wagner, 1984; Young &
Hallgrímsson, 2005), and the magnitude of integration has important consequences for
both the evolution of mean phenotypes (Schluter, 1996; Marroig & Cheverud, 2005, 2010;
Cardini & Polly, 2013) and the evolution of morphological integration itself (Oliveira et al.,
2009; Porto et al., 2009, 2013; Shirai & Marroig, 2010).

In this context, adaptive landscapes may be the central component governing both the
stability and divergence in integration patterns, as both stabilizing (Jones, 2007; Arnold
et al., 2008) and directional (Jones et al., 2012; Melo & Marroig, 2015) selection have been
shown to produce changes in integration patterns. The empirical evidence available
demonstrates that both stability (e.g. Marroig & Cheverud, 2001; Oliveira et al., 2009;
Porto et al., 2009; Willmore et al., 2009) and divergence (e.g. Monteiro & Nogueira, 2010;
Grabowski et al., 2011; Sanger et al., 2012; Haber, 2015) of integration patterns are possible
outcomes of the evolutionary process. Therefore, the question of whether any of these
two scenarios is the rule or exception at macroevolutionary scales remains open, although
some theoretical and methodological differences between these works with respect to the
representation of morphological features need to be taken into consideration.

Morphometrics

Traditionally, morphological features are measured using distances among elements de-
finied in general terms, such as “cranial length” or “cranial width”. Pearson & Davin
(1924) introduced the notion that measurements should be restricted to single osteological
elements, preferably as distances between homologous features that could be identified in a wide taxonomic coverage. Cheverud (1982) accommodates this notion into the framework of morphological integration, thus considering individual measurements over single bones as local representations of regional phenomena, that is, the functional, developmental and genetic interactions that produce covariances among these elements. The influence of such interactions over covariance or correlation matrices can be accessed by defining a subset of measurements which fall under the scope of a particular process and estimating a metric that summarizes such partitioning; the null hypothesis that this partition is undistinguishable from randomly-defined partitions can then be tested using Monte Carlo methods (Mantel, 1967; Cheverud et al., 1989).

In the past three decades, geometric morphometrics (Bookstein, 1982, 1991; Kendall, 1984; Rohlf & Slice, 1990; Goodall, 1991) have been consolidated as a quantitative framework for the representation of biological shape as geometric configurations of homologous features (landmarks; Bookstein, 1991). Two principles are central here: first, the conceptual and statistical separation of size and shape as components of biological form; second, the use of superimposition-based methods (GPA: Generalized Procrustes Analysis; Rohlf & Slice, 1990; Goodall, 1991) for the estimation of shape statistical parameters, such as mean shape and shape covariance structure. Procrustes estimators were proposed as a solution to the problem that landmark configurations are arbitrarily rotated and translated; such nuisance parameters are impossible to estimate without any assumptions (Goodall, 1991; Lele & McCulloch, 2002), a situation known as the identifiability problem (Neyman & Scott, 1948).

Although the use of Procrustes estimators is currently widespread in geometric morphometrics toolboxes (e.g. Klingenberg, 2011; Adams & Otárola-Castillo, 2013), it has not been without criticisms, either with respect to the estimation of mean shape configurations (e.g. Lele, 1993; Kent & Mardia, 1997; Huckemann, 2012) and of shape covariance matrices (Walker, 2000; Adams et al., 2004; Linde & Houle, 2009; Márquez et al., 2012). For configurations in two-dimensional space, Procrustes estimators for mean shape perform well under isotropic landmark covariance structure (Kent & Mardia, 1997), a situation of null covariances among landmarks and coordinates; however, when this assumption does not hold, Procrustes estimates for mean shape behave badly, especially when shape variation is high (Huckemann, 2011). The example provided by Linde & Houle (2009) demonstrates that when such assumption is broken shape covariance patterns are also poorly estimated; if the unknown landmark covariance matrix is structured due to regional differences in covariance-generating processes, such variation will be displaced and effectively spread out through the entire landmark configuration.
A number of alternatives for estimating shape covariance matrices have already emerged; some of these alternatives (e.g. Monteiro et al., 2005; Theobald & Wuttke, 2006; Linde & Houle, 2009; Zelditch et al., 2009) propose modifications to the Procrustes analysis in order to deal with heterogeneity in landmark covariance structure. Márquez et al. (2012) propose another definition of shape descriptors using interpolation-based techniques (Cheverud & Richtsmeier, 1986; Bookstein, 1989) as a starting point; such descriptors refer to infinitesimal expansions or retractions in reference to a unknown mean shape (Woods, 2003) estimated at definite locations amidst sampled landmarks. The authors argue that these descriptors are proper local measurements of shape variation, as they can directly be linked to biological processes that generate covariation among morphological elements.

Despite these caveats regarding Procrustes estimators for covariance matrices, the use of such estimators for investigating aspects of morphological integration has increased in the past ten years (e.g. Klingenberg et al., 2004; Drake & Klingenberg, 2010; Goswami & Polly, 2010; Martínez-Abadías et al., 2011; Sanger et al., 2012). The results found by these authors are sometimes in stark contrast with similar works using interlandmark distances (e.g. Cheverud et al., 1997; Oliveira et al., 2009; Porto et al., 2009). For instance, Martínez-Abadías et al. (2011) found a pattern of strong integration among partitions in human skull covariance patterns, while Oliveira et al. (2009) and Porto et al. (2009) demonstrated that humans are among the most modular examples of mammalian skull covariance patterns. Likewise, while Cheverud et al. (1997) had shown that 70% of the pleiotropic effects are confined to either anterior or posterior mandibular components, Klingenberg et al. (2004) has found no evidence for a modular distribution of pleiotropic effects among the partitions of the mouse mandible using the same strain of intercrossed mice at the same generation. Furthermore, aside from the issues regarding Procrustes estimators, these works also propose different methods to quantify the effects of interactions over covariance patterns. For example, Martínez-Abadías et al. (2011) and Sanger et al. (2012) use the RV coefficient, a multivariate correlation coefficient defined by Escoufier (1973) which has been used to quantify modular relationships over landmark covariance patterns since Klingenberg (2009) has proposed its use in this context.

Objectives

In the present work, we compare the methods described by Cheverud et al. (1989) and Klingenberg (2009) to test a priori defined modularity patterns using anthropoid primates as a model organism. In order to compare the performance of these methods with respect to different representations of form, individuals in our sample are represented both as
interlandmark distances and shape variables. Furthermore, we used an approach based on the construction of theoretical covariance matrices; such matrices are used in order to estimate Type I and Type II error rates for both methods. Since these methods were designed under different frameworks, the present work puts some effort into unifying both methods into the same conceptual and statistical framework, in order to produce meaningful comparisons.

**Methods**

**Sample**

The database we used here (Table 1) consists of 21 species, distributed across all taxonomic ranks within Anthropoidea above the genus level. We selected these species from a broader database (Marroig & Cheverud, 2001; Oliveira et al., 2009) in order to reduce the effects of low sample sizes over estimates of modularity patterns. Individuals in our sample are represented by 36 registered landmarks, measured using a Polhemus 3Draw (for Platyrrhini) and a Microscribe 3DS (for Catarrhini). Twenty-two unique landmarks represent each individual (Figure S1, Table S1), since 14 of the 36 registered landmarks are bilaterally symmetrical. For more details on landmark registration, see Marroig & Cheverud (2001) and Oliveira et al. (2009).

For each OTU, we estimated phenotypic covariance and correlation matrices for three different types of variables: tangent space residuals, estimated from a Procrustes superimposition for the entire sample, using the set of landmarks described on both Table S1 and Figure S1 (henceforth Procrustes residuals); interlandmark distances, described in Table S2; and local shape variables (Márquez et al., 2012), which are measurements of infinitesimal log volume transformations between each sample unit and a reference (mean) shape, based upon an interpolation function that describes shape variation between sampled landmarks. In this context, we used thin plate splines as interpolating functions (Bookstein, 1989). We obtained 38 transformations corresponding to the locations of the mipoints between pairs of landmarks used to define interlandmark distances, in order to produce a dataset that represents shape (i.e., form without isometric variation; Bookstein, 1991; Zelditch et al., 2004) while retaining the overall properties of the interlandmark distance dataset, such as dimensionality for example. Furthermore, we were able to use the same hypotheses of trait associations for both types of variables since the position of local shape variables through the skull mirrors the position of interlandmark distances, although they are conceptually
Table 1: Twenty-one species used in the present work, along with sample sizes and linear models adjusted.

<table>
<thead>
<tr>
<th>Species</th>
<th>Group&lt;sup&gt;a&lt;/sup&gt;</th>
<th>n</th>
<th>Model&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Alouatta belzebul</em></td>
<td>P</td>
<td>109</td>
<td>X</td>
</tr>
<tr>
<td><em>Ateles geoffroyi</em></td>
<td>P</td>
<td>78</td>
<td>-</td>
</tr>
<tr>
<td><em>Cacajao calvus</em></td>
<td>P</td>
<td>48</td>
<td>S + X</td>
</tr>
<tr>
<td><em>Callicebus moloch</em></td>
<td>P</td>
<td>93</td>
<td>X</td>
</tr>
<tr>
<td><em>Callithrix kuhlii</em></td>
<td>P</td>
<td>129</td>
<td>-</td>
</tr>
<tr>
<td><em>Cebus apella</em></td>
<td>P</td>
<td>110</td>
<td>X</td>
</tr>
<tr>
<td><em>Cercopithecus ascanius</em></td>
<td>C</td>
<td>61</td>
<td>X</td>
</tr>
<tr>
<td><em>Chiropotes chiropotes</em></td>
<td>P</td>
<td>56</td>
<td>X</td>
</tr>
<tr>
<td><em>Chlorocebus pygerythrus</em></td>
<td>C</td>
<td>110</td>
<td>X</td>
</tr>
<tr>
<td><em>Colobus guereza</em></td>
<td>C</td>
<td>140</td>
<td>X</td>
</tr>
<tr>
<td><em>Gorilla gorilla</em></td>
<td>C</td>
<td>115</td>
<td>X</td>
</tr>
<tr>
<td><em>Homo sapiens</em></td>
<td>C</td>
<td>160</td>
<td>S * X</td>
</tr>
<tr>
<td><em>Hylobates lar</em></td>
<td>C</td>
<td>66</td>
<td>X</td>
</tr>
<tr>
<td><em>Macaca fascicularis</em></td>
<td>C</td>
<td>69</td>
<td>X</td>
</tr>
<tr>
<td><em>Pan troglodytes</em></td>
<td>C</td>
<td>61</td>
<td>X</td>
</tr>
<tr>
<td><em>Papio anubis</em></td>
<td>C</td>
<td>46</td>
<td>X</td>
</tr>
<tr>
<td><em>Piliocolobus foai</em></td>
<td>C</td>
<td>83</td>
<td>X</td>
</tr>
<tr>
<td><em>Pithecia pithecia</em></td>
<td>P</td>
<td>69</td>
<td>S + X</td>
</tr>
<tr>
<td><em>Procolobus verus</em></td>
<td>C</td>
<td>88</td>
<td>X</td>
</tr>
<tr>
<td><em>Saguinus midas</em></td>
<td>P</td>
<td>50</td>
<td>S</td>
</tr>
<tr>
<td><em>Saimiri sciureus</em></td>
<td>P</td>
<td>87</td>
<td>X</td>
</tr>
</tbody>
</table>

<sup>a</sup> C: Catarrhini; P: Platyrrhini  
<sup>b</sup> S: subspecies/population; X: sex.

Here we considered only covariance or correlation structure for the symmetrical component of variation; therefore, prior to any analysis, we controlled the effects of variation in asymmetry. For interlandmark distances, we averaged bilateral measurements within each individual. For both Procrustes residuals and local shape variables, we followed the procedure outlined in Klingenberg et al. (2002) for bilateral structures by obtaining for each individual a symmetrical landmark configuration, averaging each actual shape with its reflection along the sagittal plane; we estimate local shape variables afterwards. With respect to Procrustes residuals, landmarks placed along the sagittal plane will have zero variation in the direction normal to this plane; we aligned all specimens’ sagittal plane to the $xz$ plane, thus removing the $y$ component for each of these landmarks from covariance/correlation matrices.
For each dataset, we estimated covariance and correlation matrices after removing fixed effects of little interest in the present context, such as sexual dimorphism, for example. For interlandmark distances and local shape variables these effects were removed through a multivariate linear model adjusted for each species, according to Table 1; for Procrustes residuals, the same effects were removed by centering all group means to each species’ mean shape since the loss of degrees of freedom imposed by the GPA prohibits the use of a full multivariate linear model over this kind of data to remove fixed effects.

In order to consider the effects of size variation on modularity patterns we used different procedures to remove the influence of size from each type of variable. For interlandmark distances we used the approach established by Bookstein et al. (1985); if $C$ is a correlation matrix, we obtained a correlation matrix $R$ without the effect of size using the equation

$$R = C - \lambda_1 v_1 v_1^t$$  \hspace{1cm} (1)

where $\lambda_1$ and $v_1$ refer respectively to the first eigenvalue and eigenvector of the spectral decomposition of $C$, since this eigenvector commonly represents size variation in mammals, especially when interlandmark distances are considered (Wagner, 1984; Mitteroecker et al., 2004; Mitteroecker & Bookstein, 2007); $t$ denotes matrix transpose.

For Procrustes residuals and local shape variables the effects of isometric variation were removed by normalizing each individual to unit centroid size. However, allometric relationships still influence covariance or correlation structure. In order to remove this effect we used a procedure based upon Mitteroecker et al. (2004), which relies on the estimation of an allometric component $a$ for each OTU, composed of normalized regression coefficients for each of the $m$ shape variables (either Procrustes residuals or local shape variables) over log Centroid Size. If $S$ is a covariance matrix, we obtained a covariance matrix $R$ without the influence of allometric relationships using the equation

$$R = (I_m - aa^t)S(I_m - aa^t)$$  \hspace{1cm} (2)

where $I_m$ represents the identity matrix of size $m$. Therefore, our empirical dataset consists of six sets of covariance/correlation matrices, corresponding to each type of morphometric variables considering the presence or absence of size variation.
Empirical Tests

Using these six sets of covariance/correlation matrices, we tested the hypotheses of trait associations described in Table S1 for Procrustes residuals and Table S2 for interlandmark distances and local shape variables. These trait sets are grouped with respect to their scope; two regional sets (Face and Neurocranium), each divided into three localized trait sets (Oral, Nasal and Zygomatic for the Face; Orbit, Base and Vault for the Neurocranium).

For all hypotheses, we estimated Modularity Hypothesis Indexes (MHI; Porto et al., 2013) and the RV coefficient (Klingenberg, 2009). Both statistics are estimated by partitioning covariance or correlation matrices into blocks; if $A$ is a covariance or correlation matrix, the partition

$$A = \begin{bmatrix} A_h & A_b \\ A_b^T & A_c \end{bmatrix}$$

indicates that the block $A_h$ contains covariances or correlations between traits that belong to the trait set being considered, while $A_c$ represents the complementary trait set; $A_b$ represents the block of covariances or correlations between the two sets. Thus, covariance ($S$) or correlation ($C$) matrices can be partitioned into a similar scheme. We estimated MHI values using the equation

$$MHI = \frac{\bar{\rho}_+ - \bar{\rho}_-}{ICV} \quad (3)$$

where $\bar{\rho}_+$ represents the mean correlation in $C_h$, $\bar{\rho}_-$ represents the mean correlation in the remaining sets (both $C_b$ and $C_c$), and $ICV$ is the coefficient of variation of eigenvalues of the associated covariance matrix, which is a measurement of the overall integration between all traits (Shirai & Marroig, 2010). We estimated RV coefficients for each hypothesis using the relationship

$$RV = \frac{tr(S_hS_h^T)}{\sqrt{tr(S_hS_h)tr(S_cS_c)}} \quad (4)$$

where $tr$ represents the sum of diagonal elements in any given matrix ($trA = \sum a_{ii}$).

The partitioning scheme outlined above assumes that the complementary trait set does not represent an actual hypothesis; however, we may choose to consider that both sets ($A_h$ and $A_c$) represent two distinct hypothesis. The estimation of RV coefficients remains the same; however, MHI values are estimated considering that $\bar{\rho}_+$ is the average correlation in both $C_h$ and $C_c$, while $\bar{\rho}_-$ represents the average correlation only in $C_b$. In the case of the dis-
tinction between Facial and Neurocranial traits, we estimated MHI values in this manner, reporting values for this estimate under the denomination ‘Neuroface’, following Marroig & Cheverud (2001), along independent MHI estimates for each region. Furthermore, since both Face and Neurocranium are two disjoint trait sets when any morphometric variable type is considered, RV coefficient values for either set are equal; therefore, a single RV value is reported for both regions, for each variable type.

In order to test the hypothesis that a trait set represents a variational module, we used a randomization procedure generating 1000 random trait sets with the same number of traits as the original set, calculating MHI and RV values for each iteration. For each trait set and covariance/correlation matrix, we used these values to construct distributions for both statistics representing the null hypothesis that a given trait set is a random arrangement without meaningful relationships; we then compare this null distribution to the real value obtained. For MHIs we consider this null hypothesis rejected when the real value is higher than the upper bound for the distribution, considering the significance level established; for RV coefficients, the null hypothesis is rejected when the real RV value is lower than the lower bound for the distribution, also considering significance level. For Procrustes residuals the randomization procedure maintains coordinates within the same landmark together in each randomly generated trait set, following Klingenberg & Leamy (2001).

While the procedure for estimating significance for MHIs is derived from Mantel’s (1967) approach (as outlined by Cheverud et al., 1989), we chose to generate null distributions for MHI directly, instead of estimating matrix correlation values for both real and randomized matrices. Estimated \( p \)-values in both cases remain the same, and the additional step of calculating matrix correlations would produce an unnecessary difference between the estimation of significance for MHI and RV.

**Estimation of Error Rates**

We used a set of theoretical covariance matrices to investigate Type I and II error rates for either MHI and RV metrics; the construction of such matrices is detailed in the Supplemental Information. Here, it suffices to say that we build two different sets of covariance matrices: one with known modular patterns embedded, referred to as \( C_s \) matrices, and another that represents random covariance structure, denominated \( C_r \) matrices. For each of the six sets of empirical matrices we use here, we built a set of 10000 covariance matrices of each case (either \( C_s \) or \( C_r \)) that mimic the statistical properties of each set, obtaining from these matrices samples of increasing sample size (20, 40, 60, 80, 100 individuals).
If samples were generated from a $C_r$ matrix, this represents a situation of a true null hypothesis for either tests, since the correlation matrix used to produce the sample was generated by a permutation of the hypothesis being tested. Therefore, testing hypotheses over $C_r$ matrices allows us to estimate Type I error rates, or the proportion of cases in which a true null hypothesis is rejected, given a significance level. In an adequate test, we expect that both quantities, significance level and Type I error rate, will be identical.

The opposite case, when we sampled $C_s$ matrices, represents a situation in which we know that the null hypothesis of either test is false, since we are testing the hypothesis that the partitioning scheme used to generate that particular matrix actually represents two variational modules. Thus, we estimated Type II error rates, or the probability that a false null hypothesis is not rejected, given a significance level; here, we represent Type II error using the power for each test, by simply calculating the complementar probability to Type II error rate. In an adequate test, we expect that power will rapidly reach a plateau when significance level is still close to zero, and further increasing $P(\alpha)$ will not produce a great increase in power.

Our estimates of power for both statistics should also be controlled for effect size, since sampled correlations may generate a correlation structure that is not detected due to small differences among within-set and between-set correlations. For each correlation matrix sampled, we estimate squared between-set correlations ($b^2$), in order to use it as an estimate of effect size that is not directly associated with either MHI and RV metrics. We expect that power for either tests decreases with increasing $b^2$ values, as effect size would also decrease.

**Software**

All analysis were performed under R 3.2.2 (R Core Team, 2015). Source code for all analyses can be found at http://github.com/wgar84. Previous tests we made indicated no differences between our estimation of empirical RV coefficients, based upon our own code, and estimates provided by MorphoJ (Klingenberg, 2011). In order to obtain symmetrical landmarks configurations, we used code provided by Annat Haber, available at http://life.bio.sunysb.edu/morph/soft-R.html.
Results

Empirical Tests

Tests performed using MHI for localized trait sets (Oral, Nasal, Zygomatic, Orbit, Base, and Vault; Figure 1a) detect a consistent pattern among OTUs for interlandmark distances and local shape variables; in the first set the Oral subregion is detected as a modular partition, and, when size is removed, the Vault subregion is also detected; both Orbit and Base region are not detected in any of these tests. With local shape variables, Oral, Vault Nasal, and Zygomatic sets were detected consistently across OTUs; the removal of allometric variation affects only the detection of the Vault in some groups. Furthermore, the Base sub-region is detected only in 3 of 42 tests performed over local shape variables, pooling together size and size-free correlation patterns.

For Procrustes residuals, the pattern of detection among sub-regions and OTUs is more inconsistent; for instance, the Base sub-region is detected in several OTUs, which contrasts this type of morphometric variable with the other two types. Observing the actual MHI values, Figure 1a also indicates that Procrustes residuals display a low variance of this metric within each OTU, while interlandmark distances and local shape variables display a consistent pattern of variation, with lower values for the Base and, for interlandmark distances, Orbit trait sets, while the Oral, Nasal and Vault regions display higher values consistently.

Tests performed using RV coefficients (Figure 1b) show a more irregular pattern for each variable type. When interlandmark distances are considered, most tests detect the Vault sub-region with size variation retained, and the Base sub-region when size variation is removed. For Procrustes residuals, few tests are able to reject their null hypotheses, detecting only a handful of valid modular partitions. Tests performed on local shape variables display the opposite behavior: almost all partitions are detected, regardless of whether allometric variation has been retained or removed. Moreover, RV values display a pattern of marked variation among OTUs, more so than between values within each OTU; notably, *Macaca fascicularis* and *Papio anubis* show RV values much higher than those estimated on remaining species. Such pattern can be observed both on interlandmark distances with size retained and in Procrustes residuals.

With respect to regional trait sets (Face and Neurocranium), tests performed using MHI (Figure 2a) indicate a pattern consistent with the findings regarding localized sets (Figure 1a). Considering interlandmark distances, Facial traits are detected as a valid modular
Figure 1: MHI (a) and RV (b) values for localized trait sets. Circles indicate whether a trait set is recognized as a variational module in a given OTU, with $P(\alpha)$ indicated by the legend. Notice that blue values are associated with higher MHI values and lower RV values, as the alternate hypothesis for each statistic is formulated in a corresponding manner; see text for details.
partition both with size variation retained or removed, while Neurocranial traits are detected as a valid partition only when size variation is removed. This pattern mirrors the contrast between Oral and Vault traits in the localized sets regarding interlandmark distances. For Procrustes residuals, Neurocranial traits are a valid partition with both size retained and removed; once again, this pattern mirrors the detection of the Basicranial partition as valid in the localized sets. Finally, in local shape variables, both Face and Neurocranium are detected as valid with size retained; with size removed, only the Face is recognized consistently. The same can be observed for localized sets, where removing allometric variation affects the detection of the Vault set in some OTUs. The tests for the distinction of within-set and between-set correlations for these two sets (designated ‘Neuroface’) show a pattern that is consistent with tests for the individual sets: if one of the sets was previously detected, this distinction is also detected as valid.

![Modularity Hypothesis Index](image1)

![Face/Neuro RV](image2)

Figure 2: MHI (a) and RV (b) values for regional trait sets. Circles indicate whether a trait set is recognized in a given OTU, with \( P(\alpha) \) indicated by the legend. Notice that blue values are associated with higher MHI values and lower RV values, as the alternate hypothesis for each statistic is formulated in a corresponding manner; see text for details.

Testing the distinction between Face and Neurocranium using RV coefficients (Figure 2b) show that in most cases both regions are considered distinct and valid variational modules...
for interlandmark distances and local shape variables; for Procrustes residuals, only a handful of taxa show the same result. In this case the correspondence with localized trait sets (Figure 1b) is more difficult due to the lack of independent tests for each region.

**Error Rates**

Comparing the distributions of MHI and RV values from theoretical matrices with respect to their structure (Figure 3) shows marked differences between metrics and morphometric variables from which correlations are sampled. In general, the distribution of MHI values obtained from $C_r$ matrices is the same, while the distributions for $C_s$ matrices for this metric are more disjoint from the former distribution in local shape variables than in Procrustes residuals, with interlandmark distances representing an intermediate case, regardless of whether size was retained or removed. For RV values, all distributions overlap to some degree; for local shape variables and interlandmark distances with size retained, either distributions ($C_r$ and $C_s$) overlap to a lesser extent.

Regarding the relationship between significance levels and Type I error rates estimated over $C_r$ matrices, Figure 4 shows that these quantities approach an identity relationship very closely regardless of whether we use MHI or RV to quantify variational modularity; even at low sample sizes Type I error rates are very close to significance levels. Furthermore, the effect of sampling correlations from size-free distributions does not change Type I error rates.

The relationship between power and significance levels (estimated over $C_s$ matrices) shows substantial differences with respect to the chosen metric (MHI or RV) and to the type of variable that provides sampled correlations. Considering local shape variables (Figure 5), tests using either MHI and RV have high power, even at low sample or effect sizes; increasing these quantities further increases power. However, for lower effect sizes (represented by high average squared correlation between sets, $b^2$) power for tests using MHI is higher than for those using RV; as effect size increases (lower $b^2$ values), the difference in power between the two statistics decreases. For local shape variables, sampling from its associated size-free correlation distribution implies minor differences in power for both statistics.

For interlandmark distances (Figure 6) there are substantial differences on the relationship between power and significance level if we consider the different parameters. In general, power for tests using MHI are always higher than for tests using RV; this effect is more pronounced on $C_s$ matrices derived from size-free interlandmark distances, although tests
Figure 3: Distribution of Modularity Hypothesis Index and RV Coefficient for theoretical correlation matrices.
Figure 4: Type I error rates as a function of the chosen significance level regarding tests for variational modularity applied on $C_r$ correlation matrices. The solid black line represents the identity relationship.
Figure 5: Power for both MHI and RV statistics as a function of the chosen significance levels with respect to tests for variational modularity applied on Cₕ matrices with values sampled from the distribution of correlations between local shape variables. Lines are colored with respect to quantiles of the b² distribution, according to the legend.
on these matrices have a substantial decrease in power for either tests. However, this
decline is more pronounced for tests based on the RV statistic, since for lower effect sizes
(high $b^2$ values) power approaches an identity relationship with significance level. Sample
size also interferes with this relationship since increasing this quantity also increases power
when higher effect sizes (low $b^2$ values) are considered.

![Figure 6: Power for both MHI and RV statistics as a function of the chosen significance
levels with respect to tests for variational modularity applied on C_s correlation matrices
with values sampled from the distribution of correlations between interlandmark distances.
Lines are colored with respect to quantiles of the $b^2$ distribution, according to the legend.
The solid black line represents the identity relationship.]

With respect to Procrustes residuals (Figure 7), tests using either MHI or RV have reduced
power regardless of effect or sample size. Sampling from size-free correlation distributions
to build C_s matrices also has little effect. In this case, power for tests performed using RV
values approaches an identity relationship with significance level; increasing sample size has some effect, but it does not increase power above 50% in any case.

Figure 7: Power for both MHI and RV statistics as a function of the chosen significance levels with respect to tests for variational modularity applied on $C_s$ matrices with values sampled from the distribution of correlations between Procrustes residuals. Lines are colored with respect to quantiles of the $b^2$ distribution, according to the legend. The solid black line represents the identity relationship.

Discussion

Covariance matrices derived from morphological traits are supposed to represent the pattern of codependence that arises due to a variety of interactions among developing
morphological elements (Olson & Miller, 1958; Cheverud, 1996). Such interactions are the
expression of local developmental factors, as they interact with the expression profiles of
surrounding cell types, producing coordinated changes in their cycles and differentiation,
thus integrating resulting tissues in the adult population. Although these events of local
integration overlap, and the composed effect over adult covariance patterns may be
confusing (Hallgrímsson & Lieberman, 2008; Hallgrímsson et al., 2009), we believe that a
careful comparison of different yet equally proper ways of measuring and representing
form may be informative of the underlying processes that produce covariances.

Due to the minimization of quadratic distances among homologous landmarks during
GPA, covariance matrices derived from Procrustes estimators lose the signal of localized
effects on covariance patterns (Linde & Houle, 2009). Therefore, the use of Procrustes
estimators to investigate morphological integration or modularity implicitly implies in a
divorce between the phenomenon we would like to investigate and the representation
we choose to use. This disconnection between theory and measurement may have grave
consequences for the hypotheses we wish to test (Houle et al., 2011); such consequences are
observable in both our empirical tests and those tests performed on theoretical matrices,
as we explore below.

The mammalian Basicranium originates from thirteen precursor tissues derived from both
paraxial mesoderm and neural crest, and they may merge to form single bones, such as
the sphenoid (Jiang et al., 2002; Lieberman, 2011). Furthermore, these precursors display a
mosaic of endochondral and intramembranous ossification early in development, and, as
the brain grows afterwards, it induces a pattern of internal resorption and exterior depo-
sition on the underlying posterior Basicranium (Lieberman et al., 2000); meanwhile, the
anterior portion suffers influence from the development of Facial elements (Bastir & Rosas,
2005). Thus, since the Basicranium ossifies early in development, the composed effect of
all posterior steps of cranial development will overshadow any pattern of integration this
region might have, as predicted by the palimpsest model of development (Hallgrímsson et
al., 2009). Moreover, the angulation between anterior and posterior elements of the Basicra-
nium has changed sensibly during primate evolution, and such property appears to have
evolved in coordination with Facial growth relative to the cranial Vault, accomodating
both structures on each other (Scott, 1958; Lieberman et al., 2000, 2008).

Due to this heterogeneity of developmental processes acting on the Basicranium, we would
not expect it to be a variational module, further expecting that a test of this property over
skull covariance patterns will fail to reject the null hypothesis of random association. How-
ever, considering the 42 tests performed over covariance matrices derived from Procrustes
residuals regarding localized hypotheses (midpanels of Figure 1a), the Basicranium is
detected as a valid variational module in 27 cases, distributed through matrices with size ei-
ther retained or removed; in some cases (e.g. *Alouatta, Cercopithecus*), only the Basicranium
is detected. Thus, Procrustes residuals show a pattern of detection of variational modules
opposite to the expectation for this particular test. In covariances matrices derived from
interlandmark distances (upper panels of Figure 1a), the Basicranium is not detected a
single time; with local shape variables (lower panels of Figure 1a), the Basicranium is
detected only three out of 42 times.

These two remaining types of representations, interlandmark distances and local shape
variables, show patterns of detection of variational modules that are both consistent to the
expectations derived from developmental and functional interactions and to the patterns
emerging from these representations when size variation is removed. Considering that
interlandmark distances are on a ratio scale (Houle *et al.*, 2011), isometric variation will
be represented to a greater extent when compared to subtle allometric relationships, due
to the multiplicative nature of biological growth (Huxley, 1932). Therefore, the Oral
trait set is detected as a valid variational module, considering covariance matrices among
interlandmark distances (upper left panel of Figure 1a), since this region is strongly affected
by the induction of bone growth due to muscular activity beginning in the pre-weaning
period (Zelditch & Carmichael, 1989; Hallgrímsson *et al.*, 2009). Furthermore, in these
matrices, allometric interactions are associated with the strength of association between
traits in Oral, Nasal, Zygomatic and Vault regions in an evolutionary scale (Chapter 3); thus,
allometric relationships also play some role in the determination of covariance/correlation
patterns for other subregions, although such pattern may be masked in interlandmark
distances by the effect of isometric variation. On the other hand, the patterns expressed
in covariance/correlation matrices for local shape variables are influenced by allometric
relationships defined on a proper scale (Jolicoeur, 1963; Houle *et al.*, 2011; Márquez *et
al.*, 2012) and local developmental processes. Thus, they reflect this association between
integration and allometry.

By removing allometric effects from local shape variables (lower right panel of Figure 1a),
variational modularity can still be detected in both Oral and Nasal regions, while in a
number of species, the Vault region is no longer detected as a variational module. Vault
integration may be achieved through both allometric relationships and the effect of relative
brain growth, since Vault elements arise mostly through intramembranous ossification,
induced by the secretion of signaling factors from the outer brain tissues, with a modest
but necessary contribution of mesoderm-derived tissue that undergoes endochondral
ossification (Jiang et al., 2002; Rice et al., 2003; Franz-Odendaal, 2011; Lieberman, 2011). In humans, the growth of Vault osteological elements occur, from a topological standpoint, without deviations from an isometric growth model, forming regular connections among bones (Esteve-Altava & Rasskin-Gutman, 2014), since the boundary of interactions between the tissue inducing growth (brain) and the tissue whose growth is being induced (Vault bones) is also regular. Thus, as this effect dominates the latter stages of pre-natal development (Hallgrímsson & Lieberman, 2008; Lieberman, 2011), the overall effect of brain growth over skull growth patterns in this stage may mirror an affine transformation, rendering it undetectable over covariance patterns derived from local shape variables, since their estimation excludes shape variation associated with affine transformations (Márquez et al., 2012). In contrast, Oral and Nasal elements have more complex patterns of connectivity arising from their tight integration with the soft tissues that compose the remaining elements of the Face (Lieberman, 2011; Esteve-Altava et al., 2013; Esteve-Altava & Rasskin-Gutman, 2014), thus producing an intricate pattern of associations that may be responsible for the variational modularity we are able to detect in all species, regardless of whether size variation is retained or removed.

The differences between the pattern of detection in all representations for local trait sets for RV values (Figure 1b) makes a similar interpretation of the results for this metric difficult, as opposed to the results regarding MHI values. The variance of RV values among species seems to indicate that RV values are sensitive to the magnitude of morphological integration, as Papio and Macaca are among the Catarrhines who display large values of this quantity (Oliveira et al., 2009) and score higher (orange/red) RV values for all local trait sets, at least when we consider interlandmark distances. Such sensitivity might be one of the factors responsible for the low power estimated for the RV metric using theoretical covariance matrices; interestingly enough, this effect of magnitude of integration (which is thought to emerge as a consequence of size variation; Wagner, 1984; Marroig & Cheverud, 2001, 2005; Porto et al., 2013) seems to also affect Procrustes residuals regardless of whether covariance structure arising from allometric relationships is removed or not.

However, for the set of hypotheses concerning differences between Facial and Neurocranial traits (Figure 2), there is substantial agreement between tests performed using MHI and RV values. For MHI values (Figure 2a), the overall pattern of detection is similar to the pattern detected in local trait sets, for all variable types; for RV values (Figure 2b), there is ample support for the hypotheses that both Face and Neurocranium represent variational modules, in both local shape variables and interlandmark distances. These regions have marked differences in timing and pattern formation during development, as observed
from the behavior of their composing units (Zelditch & Carmichael, 1989; Hallgrímsson et al., 2009; Lieberman, 2011; Esteve-Altava & Rasskin-Gutman, 2014); therefore, the more general pattern of distinction between Face and Neurocranium is detected regardless of the metric chosen to represent modularity.

Theoretical Matrices and Error Rates

The distribution of MHI and RV values obtained from the theoretical matrices (Figure 3) is a starting point for understanding the differences in power for tests using these two metrics (Figures 5–7). For MHI values, the distribution obtained from random (\(C_r\)) matrices is consistently the same, regardless of what representation we used to sample correlations, or whether size was retained or removed. On the other hand, the distribution of RV values for random matrices change depending on the representation sampled or whether size variation has been removed or retained. Moreover, this change in behavior for the distribution of RV values for \(C_r\) matrices implies different levels of overlap between these null distributions and the distribution of values obtained for structured (\(C_s\)) matrices.

For Procrustes residuals, a substantial overlap occurs regardless of whether size variation was removed or retained; unsurprisingly, our estimates of power for RV in this type of representation are very low (Figure 7); nonetheless, power estimated for tests based on MHI is lower than in other types of representation, since the difference between within-set and between-set correlations in Procrustes residuals (Figure S2) is the lowest of all representations. A substantial overlap in RV distributions for \(C_r\) and \(C_s\) matrices also occurs with interlandmark distances when size variation is removed, and it implies in low power for tests using RV values in this type of representation (right column of Figure 6). However, in this case there is a substantial difference between within-set and between-set correlations (Figure S2), and tests using MHI to represent modularity are still able to detect such difference (albeit with reduced power) when compared to tests over \(C_s\) matrices derived from interlandmark distances with size retained (Figure 6). In those cases where both distributions for \(C_r\) and \(C_s\) matrices do not overlap substantially — for example, when local shape variables are considered (Figure 5) — power for tests performed using MHI values is always higher than for tests using RV except when sample sizes are very high; in this case power for both metrics are similar. The same behavior can also be observed in interlandmark distances when size is retained (left column of Figure 6).

These results indicate that RV coefficients are more sensitive to the absolute value of both within-set and between-set correlation distributions than MHI values. For interlandmark
distances (Figure 6), removing size variation reduced the average value of both correlation distributions by a similar amount (Figure S2); the difference between average correlations in these two sets actually increases, going from 0.042 to 0.061 when size is removed. However, since the actual average correlations for these two distributions approach zero, tests based on RV lose power more rapidly than tests based on MHI. This sensitivity might be associated to the use of squared covariances, as shown by Equation 4, while Modularity Hypothesis Indexes use correlations directly (Equation 3). Furthermore, as observed by Fruciano et al. (2013), sample sizes sensibly influence the estimation of RV coefficients, and we demonstrate here that such sensitivity also extends to estimates of power for tests using this metric.

On the other hand, our estimates of power for tests using MHI indicate that it is more robust to differences in absolute correlation values or sample sizes, thus allowing comparisons across more heterogeneous settings, such as our comparison between different representations of form, with substantial variation in sample sizes. The detection of variational modularity is akin to Student’s $t$ test, since we are trying to determine whether two groups of observations (correlations between traits in the same subset versus correlations between traits in different subsets) have a significant difference in average values; we use resampling procedures to estimate significance in this case only due to the interdependency between pairwise correlations. Thus, as Modularity Hypothesis Indexes are estimated with a formula that is very similar to that of a $t$ statistic (difference between location parameters for two groups, divided by a scale parameter — ICV; Equation 3), we believe that this statistic is a proper way to represent variational modularity; the robustness of the tests using this statistic reinforce this belief.

Our approach for constructing theoretical matrices attempts to simulate the most simple situation, that is, the situation where there are only two subsets of traits, akin to the distinction between Face and Neurocranium in our empirical dataset (Figure 2). In this setting, both statistics are capable of detecting this distinction, except when both are used on covariance/correlation patterns derived from Procrustes residuals. However, even though we built theoretical matrices using correlations sampled from these estimators, actually simulating the interference in covariance structure that such estimators produce in our theoretical matrices is quite difficult. Furthermore, constructing such matrices with more complicated patterns (with three modules, for instance) while maintaining their connection to the correlation distributions of each morphometric type is also difficult, due to the restriction on positive-definiteness we enforce on them. Thus, the lack of differences in type I error rates for all cases may be a limitation of our scheme for building theoretical
matrices.

The issues we found with the use of Procrustes estimators for covariance matrices and the use of RV coefficients to estimate and detect variational modularity may explain results found by other authors; for instance, Martínez-Abadías et al. (2011) has found no evidence that genetic and phenotypic covariance structure for human skulls conforms to a modular structure, since all tests performed by these authors failed to reject the null hypothesis of random association. These authors use Procrustes estimators to represent covariance structure, and test their hypothesis of partitioning (Face, Vault and Base) using RV as the statistic representing variational modularity. Since this combination implies in very low power (Figure 7), not rejecting the null hypothesis in their case might be a consequence of the choice of estimates of both covariance structure and variational modularity; thus, these authors’ assertion of pervasive genetic integration in the human skull may be misleading, considering that skull covariance patterns in humans are one of the most modular examples of such patterns when compared to other mammals (Porto et al., 2009) or catarrhine primates (Oliveira et al., 2009).

The approach we explore in the present work is but one of the different ways one can investigate the association between genetic, functional and developmental interactions and correlation structure (Mitteroecker & Bookstein, 2007). For example, Perez et al. (2009) relies on abstracting correlation matrices into networks, then using community-detection algorithms to search for modular patterns without a priori hypotheses, associating their results with knockout experiments that support the communities they found among traits; however, it is not clear how much relevant information is retained in these network representations. Furthermore, the authors use Procrustes estimators, which may bias the detection of modularity patterns in this setting in the same manner as we demonstrated here.

Monteiro et al. (2005) assumes that the underlying morphogenetic components of the rodent mandible behave as modules, further investigating the patterns of correlation between these units in both within-species variation and between-species variation among Echimids; Monteiro & Nogueira (2010), relying on the correspondence of these units through mammalian diversification did the same to phylostomid bats. Although using a landmark-based approach to represent morphological variation, the authors do not use Procrustes estimators to represent covariance structure among these units, and the pattern of reorganization of correlation structure among these units associated with niche diversification in phylostomids seems robust, considering that this radiation may have been associated with a very heterogeneous adaptive landscape, and such heterogeneity
may lead to a reorganization of correlation patterns (Jones, 2007; Jones et al., 2012; Melo & Marroig, 2015).

Another valid approach is to model certain aspects of development as null hypotheses; Esteve-Altava & Rasskin-Gutman (2014), investigating the pattern of connections among human cranial bones, conceived the null hypothesis that unconstrained bone growth is sufficient to explain the observed patterns. Such approach could be extended to investigate morphological covariance structure; if we consider the geometric properties of the features we measure (Mitteroecker & Bookstein, 2007), one could formulate the null hypothesis that topological proximity is a sufficient explanation for the observed covariance structure, against the alternative hypothesis that local developmental processes coupled with functional interactions produce stronger relationships among close elements that surpass these purely topological interactions. Alternatively, one could actively look for variational boundaries between regions, as boundary formation is a phenomenon that has been well studied under a dynamical perspective on development (e.g. Turing, 1952; Meinhardt, 1983; Tiedemann et al., 2012).

The approach of partitioning covariance matrices into blocks that correspond to inferred modular associations has the advantage that it is simple from an operational standpoint; however, modularity patterns are almost certainly not expressed in phenotypic data as the binary hypotheses we used here (Hallgrímsson et al., 2009). Thus, hypotheses and inferences made from them have to be contextualized in the light of developmental dynamics, since the measurements we make and the parameters we estimate have to be properly connected to the models we are considering; otherwise, inferences made from such models may be devoid of meaning (Wagner, 2010; Houle et al., 2011).

**Conclusion**

Here we show that Procrustes estimators for covariance matrices fail to capture the modularity patterns embedded in phenotypic data, regardless of which metric is chosen to represent such patterns, although the combination of this type of variable with RV coefficients for investigating modularity has even more problems than either has alone. Both interlandmark distances and local shape variables seem valid options to represent morphological variation, if their limitations are taken into consideration. We wish to stress this point: any representation of morphological variation has limitations since they are themselves models — at the very least of what it is important to represent — not fully capturing the phenomena we may be interested in.
Acknowledgements

We thank G. Burin, D. Melo, and A. Porto for comments on an early draft. This work has been funded by CNPq (Conselho Nacional de Pesquisa e Desenvolvimento Tecnológico) and FAPESP (Fundação de Apoio à Pesquisa do Estado de São Paulo).

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