Covariance Testing

DESC Members (Dated: July 21, 2019)

There are a number of codes that compute covariance matrices analytically; the plan is to use these to build TJPCov. In this project, we start along the path of comparing these different codes, building up a suite of tools that can be used to compare covariance matrices. We expect these tools to be useful not only for converging on a single accurate code for computing covariance matrices but also more generally for understanding which parts of the covariance matrix carry the most information (and therefore need the most attention to get right) and which are not relevant (so for example matrices that are not positive definite may still be usable if the negative eigenmodes are not relevant).

I. INTRODUCTION

II. METHODS

To illustrate our methodology, we use cosmic shear statistics $\xi_{\pm}(\theta)$, focusing for the most part on the Year 1 results of the Dark Energy Survey [1] (DESY1), and also on predictions for DES Year 3 (DESY3). The data is divided into four tomographic redshift bins spanning the interval 0.20-1.30, which gives us 10 bin combinations, each one containing 20 angular bins between 2.5 and 250 arcmin. We thus have 200 data points for each $\xi_{+}(\theta)$ and $\xi_{-}(\theta)$, giving us 400 in total.

We apply the angular cuts described in [2], which removes the scales most sensitive to baryonic effects; this leaves us with 167 points for $\xi_+(\theta)$ and 60 for $\xi_-(\theta)$ (for which baryonic interactions are more relevant), resulting in 227 data, which correspond to a 227 × 227 covariance matrix.

The two codes employed here for computing the covariance matrices are Cosmolike [5] and a simpler version of the code used to analyze the KiDS-450 survey [6], hereafter referred to as CL and BJ, respectively. To perform cosmological parameter inference we use the CosmoSIS [9] pipeline, while employing the MultiNest [4] sampler to explore the parameter space, with 500 livepoints, efficiency set to 0.3, tolerance to 0.1 and constant efficiency set to False.

A. One-to-one Comparison

This is a simple matter of comparing elements of a covariance matrix, usually starting with diagonal elements. Figure 1 shows an example.

B. Eigenvalues and eigenvectors

An intuitive approach to identifying the most contributing elements of a covariance matrix is to look at its eigenvalues, since we can associate them to the variance. The lowest eigenvalues would correspond to the least variance and therefore hold the most information,

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Parameter	Prior
Cosmological	
Ω_m	U(0.1, 0.9)
$A_s \times 10^9$	$\mathcal{U}(0.5, 5)$
$H_0(\mathrm{kms^{-1}Mpc^{-1}})$	$\mathcal{U}(55,91)$
Ω_b	$\mathcal{U}(0.03, 0.07)$
$\Omega_{ u}h^2$	$\mathcal{U}(0.0005, 0.01)$
n_s	$\mathcal{U}(0.87, 1.07)$
Astrophysical	
A	$\mathcal{U}(-5,5)$
η	$\mathcal{U}(-5,5)$
Systematic	
m^i	$\mathcal{G}(0.012, 0.023)$
Δz^1	G(-0.001, 0.016)
Δz^2	G(-0.019, 0.013)
Δz^3	$\mathcal{G}(0.009, 0.011)$
Δz^4	$\mathcal{G}(-0.018, 0.022)$

TABLE I. List of the priors used in the analysis for parameter constraints. For the cosmological parameters, we fixed w=-1.0, $\Omega_k=0.0$ and $\tau=0.08$. The astrophysical parameters are associated with the intrinsic alignment, they follow the relation $A(z)=A[(1+z)/1.62]^{\eta}$, we have also $z_0=0.62$. Lastly, for systematics we have m^i corresponding to the shear calibration, and Δz^i for the source photo-z shift, with i=1,4 in both cases. The lens photo-z shift parameters Δz^i , i=1,5 were fixed to zero.

whereas the highest eigenvalues would implicate larger error and thus contribute significantly less. With this in mind, the most important elements of the matrix would be those with the lowest eigenvalues. To verify this assumption, we need to look at the parameter constraints: removing the elements with the highest eigenvalues should not alter our results; the opposite, however, should give us a broader constraints. Our procedure consists of first diagonalising the covariance matrix in order to calculate its eigenvalues and then replacing the 50 highest with numbers of about nine orders of magnitude higher, thus removing their effective contribution;

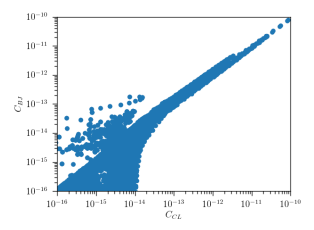


FIG. 1. A simple scatter plot of elements of covariance matrices produced by two separate halo model codes.

we then obtain a new covariance matrix with the modified eigenvalues. We also apply this method to the 50 lowest eigenvalues of the original covariance matrix, and replace them with higher values. Finally, we perform a cosmological analysis with the new covariances matrices, to constrain the parameters of our model. In Figure 2 we compare the results of both approaches to those obtained with the original covariance matrix, for brevity we show only Ω_m and σ_8 . We can see in the Posterior Density Functions (pdfs) that the mean values of both parameters are well within the 2σ interval, and this is also true for all the other parameters; the size of the constraints is also within less than 5% of each other for almost all of the parameters. Since the constraints are similar for both approaches, it becomes clear that ranking eigenvalues is not the most efficient way of evaluating where the most contributing elements of the covariance matrix lie.

Next, we wish to compare matrices using their eigenvalues. We plot the results obtained for constraints on Ω_m and σ_8 for BJ and CL in Figure 3, where we can see that, while their best fit values agree within a 2σ interval, the constraints are up to 25% broader for the latter. This is also true for the other parameters, where, on average, constraints obtained with CL are about 18% wider. To see if we can identify these differences in the eigenvalues of the covariance matrices, we plot Figure 4. At a first glance, both curves show reasonable agreement, with values differing only by an average of $\approx 15\%$, but we should keep in mind that the order of magnitude of the values vary greatly, which makes it difficult to compare them efficiently with this methodology. Furthermore, from our previous analysis it is not clear which elements are most important.

Figure 5 shows an example of one of the eigenvectors, the one associated with the smallest eigenvalue. This low-eigenvalue mode picks up the differences between the correlation function at different angular scales (each vertical line delineates between two-point functions of shears in different tomographic bin pairs).

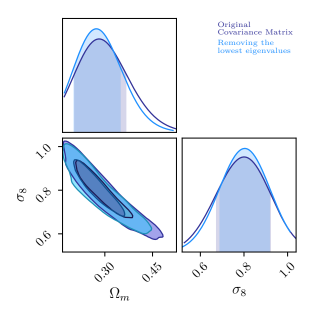


FIG. 2. Constraints on cosmological parameters Ω_m and σ_8 for the original DESY1 covariance matrix (in purple) and for two new covariance matrices obtained by setting the fifty highest eigenvalues of the original matrix to nine orders of magnitude higher (in blue), and by replacing the fifty lowest eigenvalues to nine order of magnitude lower (in magenta).

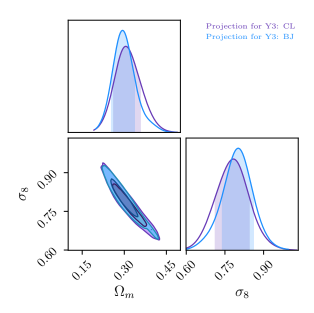


FIG. 3. Constraints on cosmological parameters Ω_m and σ_8 for two covariance matrices produced for cosmic shear for DESY3. The purple curve is for CL while the blue is for BJ. The constraints are about 18% broader for the former, indicating that the two matrices have quantifiable differences between them.

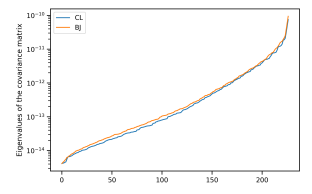


FIG. 4. A log plot showing the 227 eigenvalues of two covariance matrices produced for cosmic shear for the Year 3 projection of the Dark Energy Survey. The blue curve is for CL and the orange one is for BJ.

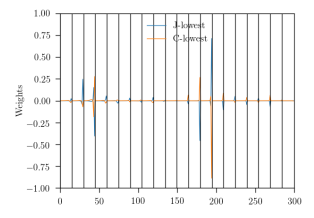


FIG. 5. A simple scatter plot of elements of covariance matrices produced by two separate halo model codes.

C. Signal to noise

Another way of analysing covariance matrices is by obtaining the signal-to-noise ratio (SNR). A simple way of achieving this is to note that the total SNR squared is

$$\left(\frac{S}{N}\right)^2 = \sum_{ij} D_i C_{ij}^{-1} D_j , \qquad (1)$$

where D_i and D_j are the data points and C the covariance matrix. If C were diagonal, then the eigenvectors would simply be the data points themselves, and we could estimate the SNR squared expected in each mode by simply computing T_i^2/C_{ii} where T_i is the theoretical prediction. Then we could throw out the modes with the lowest SNR. Since C is not diagonal, we have to first diagonalise it and then order the values. So, we write the expected

SNR squared as

$$\left(\frac{S}{N}\right)^2 = \sum_{ij} T_i C_{ij}^{-1} T_j$$

$$= \sum_i \frac{v_i^2}{\lambda_i},$$
(2)

where λ_i are the eigenvalues of the covariance matrix, which is diagonalised with the unitary matrix U, and the eigenvectors are

$$v_i \equiv U_{ij}^t T_j \tag{3}$$

This makes it very clear which modes should be kept and which should be dropped. Modes v_i for which v_i^2/λ_i is very small can be discarded.

After obtaining the SNR for our covariance matrix, we proceed to set the 50 lowest values to seven orders of magnitude lower, which is equivalent to increasing the noise (or decreasing the signal) of these modes. We then obtain a new covariance matrix with the corresponding modified SNR values.

The parameter constraints for this method are shown in Figure 6, where we note that the mean values are shifted about 12% and 5% for Ω_m and σ_8 , respectively, with 17% broader constraints for the former, when compared to the results with the original covariance, which tells us that the modes removed were essential for obtaining better constraints on these parameters.

Another aspect that we must consider when applying this procedure is that the highest modes do not necessarily hold the most information for all of the parameters; this is evident when we analyse the SNR for each parameter individually. To illustrate this we take

$$\left(\frac{\partial S/\partial p_{\alpha}}{N}\right)^{2} = \sum_{i} \frac{(\partial v_{i}/\partial p_{\alpha})^{2}}{\lambda_{i}},\tag{4}$$

with

$$v_i^{\alpha} \equiv U_{ij}^t \frac{\partial T_j}{\partial p_{\alpha}},\tag{5}$$

where $\partial/\partial p_{\alpha}$ is the derivative with respect to each parameter θ . This produces the SNR for each parameter of interest. The importance of this procedure is illustrated in Figure 7, where we can see that while the highest SNR for all the parameters does indeed correspond to those that hold most information for parameters Ω_m and A_s , this is not the case, however, for the intrinsic alignment parameters A and η . As a result, we may end up losing constraining power over these parameters when we modify the lowest values of SNR. While this is apparent in the aforementioned figure, it is not very clear when looking at the resulting constraints on these parameters because the constraining power of the data over these parameters is not very strong (errors of about 100%). As such, these results do not encourage us to use this method for identifying the most important elements of the covariance matrix.

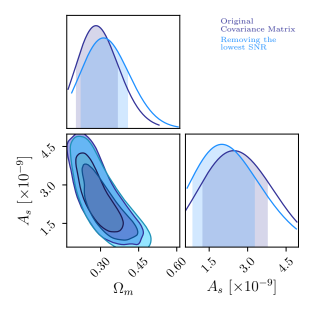


FIG. 6. Constraints on cosmological parameters Ω_m and A_s for the original DESY1 covariance matrix (in purple) and for a covariance matrix (in blue) obtained by setting fifty elements corresponding to the lowest SNR to a value nine orders of magnitude lower, in order to evaluate their contribution to parameter constraints.

D. Parameter Estimation

Ultimately, what matters is how well the likelihood does at extracting parameter constraints. Since most analyses assume a Gaussian likelihood, this boils down to how well the contours in parameter space agree when computing the χ^2 using two different covariance matrices.

E. Shrinkage

There have been several methods proposed in the literature to compress the data vectors, extracting as much information as possible. Here we consider two: first compression at the map level [3], where linear combination of the tomographic maps are used. If there are 4 tomographic bins, an uncompressed analysis would require ten separate 2-point functions (or 20 for cosmic shear), whereas a compression scheme leads to just a few uncorrelated maps. If there were 3 such maps, then only three 2-point functions would need to be used for the likelihood analysis.

We characterize a given element in the data vector by its angular and tomographic indices: $a_i = a_{lm,\alpha}$ where l, m denote the indices corresponding to given spherical harmonics and α is a given tomographic bin, or equivalently $a_i = a_{\alpha}(\vec{\theta})$ in real space. The compression occurs in terms of tomographic bins, so that

$$b_{\mu}(\vec{\theta}) = \sum_{\alpha} F_{\mu\alpha} a_{\alpha}(\theta) \tag{6}$$

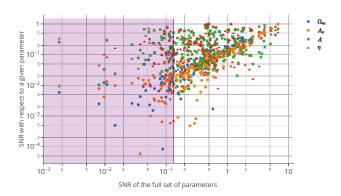


FIG. 7. Scatter plot for the relation between the signal to noise obtained with the covariance matrix for DESY1 for each parameter (x-axis) against that for the full set of parameters (y-axis). The derivatives are shown with respect to cosmological parameters Ω_m (blue) and A_s (orange), and for the intrinsic alignment parameters A (green) and η (red). The purple rectangle spreads until the fiftieth lowest value of SNR, which corresponds to the values that were modified for parameter constraints.

where the F's are chosen [7] so that the correlation functions of the b's are diagonal in tomographic space:

$$w_{\mu\nu}(\theta) = \langle b_{\mu}(\vec{\theta}_{1})b_{\nu}(\vec{\theta}_{2})\rangle|_{|\vec{\theta}_{1} - \vec{\theta}_{2}| \in \theta}$$
$$= \delta_{\mu,\nu}w_{\mu}(\theta). \tag{7}$$

Usually with N_t tomographic bins, one must consider $N_t(N_t+1)/2$ correlation functions, but – due to the transformation that diagonalizes the elements – there are only N_t correlation functions to consider (for a given θ). Even better, these can be ordered by the information they contain, so fewer than N_t can be used. In the example used in [3], 16 tomographic bins were assumed, so that the standard treatment would require 136 correlation functions, but only 3 were needed in order to extract accurate constraints.

This method suggests a way of extracting the most important pieces of the full covariance matrix C. We simply compute the covariance matrix of the w_{μ} in terms of C and keep only the most important terms. That is,

$$C_{\mu\nu}^{b} \equiv \langle (w_{\mu} - \bar{w}_{\mu}) (w_{\nu} - \bar{w}_{\nu}) \rangle$$
$$= \sum_{\alpha\alpha'\beta\beta'} F_{\mu\alpha} F_{\mu\alpha'} F_{\nu\beta} F_{\nu\beta'} C_{\alpha\alpha'\beta\beta'}. \tag{8}$$

Here the angular indices have been suppressed (there are two of them, one for each w_{μ}), but the full tomographic complexity has been retained. The covariance matrix on the right includes a total of $N_t^2 \times N_t^2$ terms (some of which are equal because of symmetry) corresponding to all possible pairs of two-point functions. But C^b on the right contains only N_t^2 elements, and again these are

ordered, so we can use only a subset of them. In the simplest case, where only one linear combination is needed so $\mu = \nu = 1$, a single number (for each pair of angular bins) captures all the relevant information from the full covariance matrix.

1. 2-Point Correlation Function

The second compression takes place at the 2-point level [8], with the compressed data vector containing linear combinations of the many 2-point functions. In principle, this might work with only N_p 2-point functions where N_p is the number of parameters varied, and each mode, or linear combination, contains all the information necessary about the parameter of interest.

For each parameter p_{α} that is varied, one captures a single linear mode

$$y_{\alpha} = U_{\alpha i} D_i \tag{9}$$

where D_i are the data points and the coefficients are defined as

$$U_{\alpha i} \equiv \frac{\partial T_j}{\partial p_{\alpha}} C^{-1}{}_{ji} \tag{10}$$

where T_j is the theoretical prediction for the data point D_j . The now much smaller data set $\{y_\alpha\}$, which contains as few as N_p data points, carries with it its own covariance matrix, with which the χ^2 can be computed for each point in parameter space. Propagating through shows that this covariance matrix is related to the original C_{ij} via

$$C_{\alpha\beta} = U_{\alpha i} C_{ij} U_{i\beta}^t. \tag{11}$$

To take an example from the full DESY1 for cosmic shear, C_{ij} is a 400×400 matrix, while the number of parameters needed to specify the model is only 16, so $C_{\alpha\beta}$ is a 16×16 matrix. We have apparently captured from the initial set of $(400 \times 401)/2 = 80,200$ independent elements of the covariance matrix a small subset (only 136 in this case) of linear combinations of these 80k elements that really matter. If two covariance matrices give the same set of $C_{\alpha\beta}$, we do not care whether any of the other eighty thousand elements differ from one another.

In our case, our covariance matrix is 227×227 , which we then reduce to 16×16 elements. Figure include figure compares the constraints obtained for the compressed covariance and data with the full one, for Ω_m and σ_8 . Include plots and talk about constraints. While this compression does not speed up calculations, it is encouraging to see that we can compress matrices of any size without losing a significant amount of information on the parameters of our models.

One relevant point in this analysis is at which point to take the derivative of each parameter. When we wish to compare the results of our compression scheme with those obtained with the full covariance matrix and data set, it is important to derivate each parameter at their respective mean value (obtained by performing the analysis with the full covariance matrix). The shape and variance of the posterior is not dependent on the derivative, but the best fit value shifts according to the point where the derivative was taken.

We also apply this methodology to comparing the covariance matrices of interest, i. e. CL and BJ. In order to do this, we take two different approaches: first, we assume that $U_{\alpha,i}$ is the same for both covariance matrices and we calculate it with BJ. The second approach is that each compression scheme must use the original covariance matrix that will be compressed, so that $U_{\alpha,i}$ will be different for each covariance matrix. Figure 8 was obtained for the first method, it shows the correlation matrix for BJ and CL, and that of the difference between them; we find this figure important because we can clearly see the difference between the two matrices by simply looking at only $(16 \times 17)/2$ elements, as opposed to having to analyse the larger correlation matrix for the full covariance matrices. It is also crucial that the matrices used for comparison here are those obtained via the same compression scheme, so that we can be sure that their differences are indeed only related to the differences in the original matrices.

2. Cool section name here

While the compression scheme is good?, it is not devoid of problems, the most concerning one being that we cannot recover the original covariance matrix the compressed one. Our motivation is to have an invertible transformation so that we are able to switch between a compressed and the full covariance. In order to do this, we start off with the compression scheme presented in Eq. 10, which will serve as the basis for a rotation matrix W_{α} . We then use the Gram-Schmidt decomposition to create $227 - N_p$ vectors orthonormal to U_{α} , thus obtaining a unitary 227×227 matrix. We then have,

$$C'_{\alpha\beta} = W_{\alpha i} C_{ij} W^t_{i\beta}. \tag{12}$$

When this transformation is applied to the original matrix, C_{ij} , the first $N_p \times N_p$ elements of the new matrix will be the same as we had in $C_{\alpha\beta}$. Since our transformation spreads out the information contained in the compressed matrix, the relevant values for parameter estimation will be contained in the first N_p rows and columns of the new covariance matrix. This means that we will have $N_p (2 \times 227 - N_p)$ important elements, which is about an order of magnitude less than in the case of the full covariance matrix.

Added error	Standard deviation
0 - 20%	< 5
20-40%	< 20
>50%	> 20

TABLE II. The first column is the introduced error to the relevant elements of the rotated covariance matrix for Ω_m . The introduced error corresponds to the an arbitrary increase in each element in the first row and first column of the rotated covariance matrix

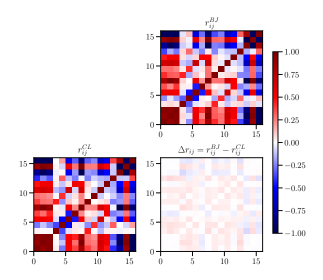


FIG. 8. The upper right and lower left plots display the correlation matrix for BJ and CL respectively, while the lower right is the difference between the two.

III. RESULTS

- A. One-to-One Comparison
 - B. Compression Vectors
- C. Cosmological Constraints
 - D. Tolerence Testing
 - IV. DISCUSSION
 - V. CONCLUSION

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