Validation of Bayesian posterior distributions using a multidimensional Kolmogorov–Smirnov test

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ABSTRACT

We extend the Kolmogorov–Smirnov (K-S) test to multiple dimensions by suggesting a \( \mathbb{R}^n \rightarrow [0, 1] \) mapping based on the probability content of the highest probability density region of the reference distribution under consideration; this mapping reduces the problem back to the one-dimensional case to which the standard K-S test may be applied. The universal character of this mapping also allows us to introduce a simple, yet general, method for the validation of Bayesian posterior distributions of any dimensionality. This new approach goes beyond validating software implementations; it provides a sensitive test for all assumptions, explicit or implicit, that underlie the inference. In particular, the method assesses whether the inferred posterior distribution is a truthful representation of the actual constraints on the model parameters. We illustrate our multidimensional K-S test by applying it to a simple two-dimensional Gaussian toy problem, and demonstrate our method for posterior validation in the real-world astrophysical application of estimating the physical parameters of galaxy clusters parameters from their Sunyaev–Zel’dovich effect in microwave background data. In the latter example, we show that the method can validate the entire Bayesian inference process across a varied population of objects for which the derived posteriors are different in each case.

Key words: Cosmology: observations – methods: data analysis – methods: statistical – galaxies: clusters: general

1 INTRODUCTION

The Kolmogorov–Smirnov (K-S) test and its close variants are the most widely-accepted methods for testing whether a set of continuous sample data is drawn from a given probability distribution function \( f(x) \) (see, e.g., Press et al. 2007, ch. 14.3.3). The test is based on the computation of the maximum absolute distance \( D \) between the cumulative distribution functions (CDFs) of the data and of \( f(x) \). In particular, it has the advantage that, in the ‘null hypothesis’ that the data are indeed drawn from \( f(x) \), the distribution of \( D \) is both independent of the form of \( f(x) \) (i.e. the test is ‘distribution-free’) and can be calculated, at least to a good approximation. Moreover, the test is invariant under reparameterizations of \( x \). In its standard form, however, the K-S test is restricted to one-dimensional data, since a CDF can only be defined uniquely in the univariate case.

Considerable effort has been made to extend the K-S test to \( n \)-dimensional data sets, much of it building on the pioneering work of Peacock (1983), but this has proved very challenging precisely because a CDF is not well-defined in more than one dimension. Working initially in two dimensions, Peacock’s original insight was to replace the notion of a CDF with the integrated probability in each of the four natural quadrants around any given data point \((x_i, y_i)\), and define the distance measure \( D \) as the maximum absolute difference (ranging both over data points and quadrants) of the corresponding integrated probabilities for the data and the theoretical distribution \( f(x, y) \). This basic idea can, in principle, be extended straightforwardly to higher dimensions, but in practice suffers from an exponential growth in complexity, since the number of independent integrated probabilities about any given data point \((x, y)\) is \( 2^n - 1 \) in \( n \)-dimensions, although Fasano & Franceschini (1987) suggest an algorithm with better scaling. Perhaps a more notable deficit is that, in the null hypothesis, the distribution of \( D \) is not independent of the form of \( f(x, y) \), although Press et al. (2007) reassure us that extensive Monte Carlo simulations show the distribution of the two-dimensional \( D \) to be very nearly identical for even quite different distributions, provided they have the same correla-
tion coefficient (Fasano & Franceschini 1987). Lopes et al. (2008) contains a review that ranks the performance of a range of different multidimensional K-S tests that are variants of Peacock’s original proposal. A completely different approach to extending the K-S test to multiple dimensions was advanced by Justel et al. (1997) and employs the Rosenblatt transformation, which maps any distribution into the unit hyper-cube (Rosenblatt 1952). Despite this approach being formally sound, the authors report insurmountable difficulties when extending the test to more than two dimensions.

In this paper, we present a new proposal for extending the K-S test to multiple dimensions, which is free from the weaknesses discussed above and scales straightforwardly with the dimensionality of the data-set. Our approach is to introduce the \( \mathbb{R}^n \rightarrow [0,1] \) mapping from any given (data) point \( x \) in the \( n \)-dimensional parameter space to the probability mass \( \zeta \) of the putative theoretical distribution \( f(x) \) contained within the highest probability density (HPD) region having \( x \) on its boundary. This mapping has the convenient property that under the null hypothesis that the data are drawn from \( f(x) \), the probability mass \( \zeta \) is uniformly distributed in the range \([0,1]\), independently of the form of \( f(x) \). The set of values \( \{\zeta_i\} \) corresponding to the data points \( \{x_i\} \) can then be compared with the uniform distribution in a standard one-dimensional K-S test (or one of its variants).

The ability to test the hypothesis that a set of data samples are drawn from some general \( n \)-dimensional probability distribution \( f(x) \) has an interesting application in the validation of Bayesian inference analyses (indeed this application provided our original motivation for seeking to extend the K-S test to multiple dimensions). Bayesian methods are now pervasive across all branches of science and engineering, from cognitive neuroscience (Doya et al. 2007) and machine learning (Bishop 2006), to spam filtering (Sahani et al. 1998) and geographic profiling (Le Comber & Stevens 2007) and machine learning (Bishop 2006), to spam filtering (Sahami et al. 1998) and geographic profiling (Le Comber & Stevenson 2007). In precision cosmology, Bayesian inference is the main tool for setting constraints on cosmological parameters (Planck Collaboration et al. 2013a; Hinshaw et al. 2013), but very few attempts have been made to assess whether the derived posterior probability distributions are a truthful representation of the actual parameter constraints one can infer from the data in the context of a given physical model. This lack of validation has been highlighted by several authors, with the strong dependence of the inference results on the priors being of particular concern (Efstathiou 2008; Linder & Mique 2008). There have been attempts to address this issue, ranging from the approach of Cook et al. (2006), which was designed with software validation solely in mind, to a method based on the inverse probability integral transform (Smirnov transform) applied to posterior distributions, that extends to spaces of higher dimensionality via marginalisation (Dorn et al. 2013). Also, validation of the Bayesian source-finding algorithm of Carvalho et al. (2012) was performed in Planck Collaboration et al. (2013), but only point estimates deduced from the posterior distributions were actually verified. Our method for addressing this problem is based on our applying multidimensional extension of the K-S test to sets of Monte-Carlo simulations of the data and the posterior distributions derived therefrom. In particular, it can take advantage of the fact that one may typically generate simulations that are of greater sophistication and realism than may be modelled in the inference process, and thus allows for a more thorough validation of the inference than has been possible with the methods developed previously. In particular, our validation procedure enables us to test all the assumptions made (explicitly or implicitly) in the inference process, such as the statistical description of the data, model assumptions and approximations, as well as the software implementation of the analysis. Moreover, we consider the full posterior distribution, regardless of its dimensionality and form, without the need to resort to marginalization, and thereby keeping intact its \( n \)-dimensional character.

This paper is organised as follows. Section 2 provides the mathematical background for our extension of the K-S test to multiple dimensions and Section 3 describes its application to the validation of Bayesian inference analyses. In Section 4.1 we apply our multidimensional K-S test to a simple toy problem, and in Section 4.2 we illustrate our Bayesian inference validation procedure by applying it to the real astronomical example of detecting and characterising galaxy clusters in observations of the microwave sky through their Sunyaev–Zel’dovich effect. We conclude in Section 5.

## 2 MULTIDIMENSIONAL K-S TEST

### 2.1 Classical one-dimensional K-S test and its variants

The classical one-dimensional K-S test is based on the distance measure \( D \) defined by

\[
D = \max_x |S_N(x) - F(x)|, \tag{1}
\]

where \( F(x) = \int_{-\infty}^{x} f(u) \, du \) is the cumulative distribution function (CDF) of the putative theoretical probability distribution \( f(x) \) and \( S_N(x) \) is the empirical CDF of the data-set \( \{x_1, x_2, \ldots, x_N\} \) that one wishes to test. As mentioned above, under the null hypothesis that the data are drawn from \( f(x) \), the probability distribution of \( D \) is independent of the form of \( f(x) \). To a very good approximation, the probability of \( D \) being larger than the observed value \( D_{\text{obs}} \) is

\[
\Pr(D > D_{\text{obs}}) = Q_{\text{KS}}([\sqrt{N} + 0.12 + 0.11/\sqrt{N}]D_{\text{obs}}), \tag{2}
\]

where the Kolmogorov distribution \( Q_{\text{KS}} \) is given by

\[
Q_{\text{KS}}(u) = \sum_{n=1}^{\infty} (-1)^{n-1} \exp(-2n^2u^2). \tag{3}
\]

Moreover, the maximum distance \( D \) and hence the K-S test itself, is invariant under reparameterization of \( x \).

It is well known, however, that the K-S test is not equally sensitive over the entire range of the variable \( x \), since under the null hypothesis the variance of \(|S_N(x) - F(x)|\) is proportional to \( |1 - f(x)| \). Consequently, the K-S test is most sensitive around the median point, where \( f(x) = 0.5 \), and less sensitive in the tails of the distribution. This makes the K-S test particularly sensitive to ‘shifts’ (in particular differences in the median), but less sensitive to differences in the dispersion, particularly if the median is similar. Two popular variants of the K-S test seek to remedy this shortcoming in different ways. Anderson & Darling (1952) suggest correcting for the effect by ‘stabilizing’ the statistic \( D \) using a weighting scheme based on the standard deviation of the null hypothesis, although the probability distribution of the resulting ‘distance’ statistic does not have a simple form. A more direct approach, proposed by Kupfer (1962), is to wrap the \( x \)-axis around into a circle (by identifying the points at \( x = \pm \infty \)) and define a new ‘distance’ statistic that is invariant under all reparameterizations on the circle. This guarantees equal sensitivity at all values of \( x \) and there

\footnote{An efficient implementation for evaluating \( Q_{\text{KS}} \) may be found in Press et al. (2007).}

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exists a simple formula for the asymptotic distribution of the statistic. Clearly, this variant is ideally suited for distributions originally defined on a circle.

### 2.2 Highest probability density regions

To extend the K-S test (and its variants) to multiple dimensions, one must clearly replace the use of the cumulative distribution function (CDF), since this is not uniquely defined in more than one dimension. In the univariate case, the CDF simply maps the value of the random variable $x$ to the integrated probability under its PDF up to that $x$-value. This suggests that to accommodate multiple dimensions one should seek an alternative mapping, that is well-defined in any dimensionality, from the multidimensional random variable $x$ to an associated integrated probability under $f(x)$. We therefore choose here to use the probability mass contained within the highest probability density (HPD) region having $x$ on its boundary.

HPDs are discussed by Box & Tiao (1992) as a general method for compressing the information contained within a probability distribution $f(x)$ of arbitrary dimensional and form. Each HPD is uniquely defined by the amount of probability $\zeta$ it encloses and is constructed such that there exists no probability density value outside the HPD that is greater than any value contained within it. More formally, HPD$_\zeta$ is the region in $x$-space defined by $\Pr(x \in \text{HPD}_\zeta) = \zeta$ and the requirement that if $x_1 \in \text{HPD}_\zeta$ and $x_2 \notin \text{HPD}_\zeta$ then $f(x_1) > f(x_2)$. Figure 1 provides a graphical illustration of the HPD definition in two dimensions.

Some properties of HPD regions are summarised in Appendix A. In particular, we show that HPD$_\zeta$ is the region with smallest volume that encloses a total probability $\zeta$. Thus the HPD provides an elegant way of communicating inference results, since it corresponds to the smallest region of parameter space having a given level of (un)certainty $\zeta$.

We may use the notion of the HPD to define a many-to-one $\mathbb{R}^n \rightarrow [0, 1]$ mapping between any point $x$ in the $n$-dimensional parameter space and the probability content $\zeta$ of the HPD whose boundary passes through $x$, namely

$$\zeta(x) = \int_{f(u) > f(x)} f(u) \, du.$$  

(4)

Thus, in general, each point $x$ in the parameter space maps onto one and only one value of $\zeta$, but, obviously, more than one point $x$ might map onto the same $\zeta$-value. In one dimension, this contrasts with the more desirable one-to-one mapping between the value of the random variable $x$ and its CDF $F(x)$, but the latter cannot be generalised to multiple dimensions.

It is easy to show that, under the null hypothesis that a set of data points $\{x_i\}$ are drawn from $f(x)$, then the corresponding set of values $\{\zeta_i\}$ are drawn from the standard uniform distribution $U(0, 1)$, independently of the form of $f(x)$. This may be seen by differentiating (4) to obtain

$$d\zeta = f(x) \, dx,$$

(5)

which simply shows that if we add an element of volume $dx$ to the HPD, its probability content increases by $f(x)$ $dx$. Comparing this equation with the standard result for a change of variables $x \rightarrow \zeta$ in probability distributions, namely

$$g(\zeta) \, d\zeta = f(x) \, dx,$$

(6)

one immediately sees that $g(\zeta) = 1$ (over its allowed range $[0, 1]$) and hence $\zeta$ follows the standard uniform distribution. A more rigorous proof is given in Appendix A. As we will see, this simple result allows for the straightforward extension of the K-S test to multiple dimensions and is central to its subsequent use in the validation of Bayesian inference analyses.

### 2.3 Pathological cases

Before continuing with the mathematical development, it is necessary to discuss some pathological cases that require further consideration. Such cases occur when there exist perfect ‘plateau’ regions of the parameter space over which $f(x)$ takes precisely the same value.

Let us begin by rewriting the mapping (4) as

$$\zeta(x) = \int_{f(u) > f(x)} f(u) \, du + \int_{f(u) = f(x)} f(u) \, du.$$  

(7)

If the point $x$ lies in a perfect plateau region $\mathcal{R}$ (which may, in general, consist of multiple disjoint subregions) over which $f(x)$ takes some constant value $f_\ast$, then the second integral on the RHS of (7) has the finite value $V(\mathcal{R}) f_\ast$, where $V(\mathcal{R})$ is the $n$-dimensional volume of the region $\mathcal{R}$. Thus, (7) maps every point $x$ in $\mathcal{R}$ to the same value of $\zeta$.

As a result, even if a set of data points $\{x_i\}$ are drawn from $f(x)$, the corresponding values $\{\zeta_i\}$ will not be drawn from the standard uniform distribution $U(0, 1)$. The extreme example is when $f(x)$ is the uniform distribution over the entire parameter space, in which case the mapping (7) yields the value $\zeta_i = 1$ for every point $x_i$ (in this case the entire contribution to $\zeta$ comes from the second integral on the RHS of (7), since the first one vanishes).

Fortunately, we can accommodate such pathological cases by adapting the approach used by Peacock (1983). Rather than evaluating the second integral on the RHS of (7) over the whole plateau region $\mathcal{R}$, we choose at random one of the $2^n$ natural $n$-dimensional ‘quadrants’ around the point $x$ and evaluate the integral over the corresponding subregion of $\mathcal{R}$. Clearly, the resulting value of the integral, and hence $\zeta$, will depend on which of...
the ‘quadrant’ subregions \( Q_k \) \((k = 1, \ldots, 2^n)\) is chosen. Since \( \sum_{i=1}^{2^n} V(Q_k) = V(\mathbb{R}) \), however, the resulting value of \( \zeta \) in (7) will be uniformly distributed, hence recovering the behaviour in the non-pathological case. Moreover, the algorithmic implementation of this approach is very straightforward.

### 2.4 Testing the null hypothesis

It is now a simple matter to define our multidimensional extension to the K-S test. Given a set of data points \( \{x_i\} \), we may test the null hypothesis that they are drawn from the reference distribution \( f(x) \) as follows. First, we apply the mapping (4) (or the mapping (7) in pathological cases) to obtain the corresponding set of values \( \{\zeta_i\} \). Second, we simply use the standard one-dimensional K-S test (or one of its variants) to test the hypothesis that the values \( \{\zeta_i\} \) are drawn from the standard normal distribution \( U(0,1) \).

In this way, one inherits the advantages of the standard one-dimensional K-S test, while being able to accommodate multiple dimensions. In particular, since under the null hypothesis the set of values \( \{\zeta_i\} \) are uniformly distributed independently of the form of \( f(x) \), our approach retains the property enjoyed by the one-dimensional K-S test of being distribution-free; this contrasts with the method proposed by [Peacock 1983].

We note that, once the \( \{\zeta_i\} \) values have been obtained, one could test whether they are drawn from a uniform distribution using any standard one-dimensional test. For example, if one wished, one could bin the \( \{\zeta_i\} \) values and use the chi-square test (see, e.g., [Press et al. 2007]), although there is, of course, an arbitrariness in choosing the bins and an inevitable loss of information. For continuous one-dimensional data, such as the \( \{\zeta_i\} \) values, the most generally accepted test is the K-S test (or one of its variants).

We also note that our approach may be applied even in the absence of an analytical form for \( f(x) \), provided one has some (numerical) means for evaluating it at any given point \( x \). An important example is when one has only a set of samples known to be drawn from \( f(x) \). In this case, one might evaluate \( f(x) \) at any given point using some kernel estimation method (say). If, however, one also knows the value of \( f(x) \) at each sample point, a more elegant approach is to estimate the probability mass \( \zeta \) corresponding to each original (data) point \( x_i \), by calculating the fraction of samples for which \( f(x) \geq f(x_i) \).

### 2.5 Performing multiple simultaneous tests

The advantageous property that, under the null hypothesis, the derived set of values \( \{\zeta_i\} \) are drawn from the standard uniform distribution \( U(0,1) \), independently of the form of \( f(x) \), allows us straightforwardly to perform multiple simultaneous tests as follows.

Suppose that one has \( M \) independent \( n \)-dimensional data-sets \( \{x_i^{(n)}\} \), where \( m = 1, 2, \ldots, M \) labels data-sets and \( i = 1, 2, \ldots, N_m \) labels the data points in each set. Suppose further that we wish to perform a simultaneous test of the null hypothesis that each data-set \( \{x_i^{(n)}\} \) is drawn from the corresponding reference distribution \( f_m(x) \). This may be achieved by first performing a transformation analogous to (4) for each data set separately, namely

\[
\zeta_i^{(n)}(x) = \int_{f_m(u) \geq f_m(x)} f_m(u) \, d^n u. \tag{8}
\]

Under the null hypothesis, the resulting combined set of values \( \{\zeta_i^{(1)}, \zeta_i^{(2)}, \ldots, \zeta_i^{(M)}\} \) will again be drawn from the standard uniform distribution, which can be tested, as previously, using a single one-dimensional K-S test.

As an extreme example, this approach may even be used when one has just a single data point drawn from each reference distribution \( f_m(x) \), in which case it would be impossible to perform an individual test for each data point. We will make use of this capability in applying our method to the validation of Bayesian inference analyses in Section 3.

### 2.6 Reparameterization invariance

A common criticism of HPD regions is that they are not invariant under smooth reparameterizations ([Lehmann & Romano 2006] sec. 5.7). In other words, if one performs a smooth change of variables \( x \to \theta = \theta(x) \), then in general HPD(\( \theta \mid x \)) \( \neq \theta(\text{HPD}(\theta \mid x)) \), although the image \( \theta(\text{HPD}(\theta \mid x)) \) in \( \theta \)-space of the \( x \)-space HPD region will still contain a probability mass \( \zeta \). This deficiency can be problematic for using HPD regions to summarise the properties of probability density functions, but it causes less difficulty here, since our only use of the HPD region is to define the mapping (4).

Suppose that a set of data points \( \{x_i\} \) are drawn from \( f(x) \) and are transformed into the set \( \{\theta_i\} \), which will be drawn from the corresponding PDF \( h(\theta) \) in the new variables. Applying an analogous mapping to (4) to the new data set \( \{\theta_i\} \) will, in general, lead to a different set of values \( \{\zeta_i\} \) to those derived from the original data \( \{x_i\} \). One may visualise this straightforwardly by considering any one of the data points \( x_i \), for which \( f(x_i) = f_i \) (say), and the corresponding HPD region having \( x_i \) on its boundary, which is defined simply by the contour \( f(x) = f_i \). Performing a smooth mapping \( x \to \theta = \theta(x) \), the data point \( x_i \) will be transformed into \( \theta_i \), which will clearly lie within the image in \( \theta \)-space of the \( x \)-space HPD, although not necessarily on its boundary. This image region will contain the same probability mass as the original \( x \)-space HPD, but it will in general not coincide with the HPD in \( \theta \)-space having \( \theta_i \) on its boundary, which may contain a different probability mass. Nonetheless, subject to the caveats above regarding pathological cases, the values \( \{\zeta_i\} \) derived from the transformed data \( \{\theta_i\} \) will still be uniformly distributed in the interval \([0,1]\), since this property does not depend on the form of the PDF under consideration.

This does not mean, however, that the significance of our multidimensional K-S test is invariant under reparameterizations. Under the alternative hypothesis that the data are not drawn from the reference distribution, the transformation \( \{x_i\} \to \{\theta_i\} \) may result in new derived values \( \{\zeta_i\} \) that differ more significantly from the standard uniform distribution than the original set.

There exists the scope for adapting our approach, so that the significance of the test is reparameterization invariant, by replacing the HPD with an alternative mapping from the \( n \)-dimensional parameter space \( x \) to some other measure of integrated probability under \( f(x) \). One such possibility is to use the ‘intrinsic credible interval’ (ICI) proposed by [Bernardo 2005], which has numerous desirable properties, including invariance under both reparameterization and marginalisation. We have not pursued this approach here, however, since the definition of the ICI is rather complicated and its calculation often requires numerical integration. It also does not follow that the resulting mapping is distribution-free. We therefore leave this as a topic for future work.

In the context of the validation of Bayesian inference analyses, however, in Section 3.3 we will discuss a method by which one may retain the use of the HPD and ensure that the values of \( \{\zeta_i\} \) derived from the original and transformed data sets, respectively, are in fact
identical under smooth reparameterizations. Thus, the significance of our resulting multidimensional K-S test will be similarly invariant in this case.

3 VALIDATION OF BAYESIAN INFERENCE ANALYSES

3.1 Bayesian inference

The Bayesian system of inference is the only one that provides a consistent extension of deductive logic (0 = false, 1 = true) to a broader class of “degrees-of-belief” by mapping them into the real interval [0, 1] [Jaynes 2003]. The basic rules of this extended logic allow one to write an equation that relates the posterior probability of a set of parameters \( \theta \) given the data \( d \) to the underlying hypothesis \( H \) that embodies the data model, and the background information \( I \); the latter comprises every assumption not explicitly stated, such as the parameterisation of the data or the instrumental set-up used to gather it. This equation is known as Bayes’ theorem,

\[
\Pr(\theta|d, H, I) = \frac{\Pr(d|\theta, H, I) \Pr(\theta|H, I)}{\Pr(d|H, I)}, \tag{9}
\]

where \( \Pr(\theta|d, H, I) \equiv \mathcal{P}(\theta) \) is the posterior distribution, \( \Pr(d|\theta, H, I) \equiv \mathcal{L}(\theta) \) is the likelihood, \( \Pr(\theta|H, I) \equiv \pi(\theta) \) is the prior, and \( \Pr(d|H, I) \equiv Z \) is a constant, known as Bayesian evidence, which may be written as

\[
Z = \int \mathcal{L}(\theta) \pi(\theta) \, d^n\theta, \tag{10}
\]

where \( n \) is the dimensionality of the parameter space.

3.2 Validation of posterior distributions

Conditioned on the assumed model \( H \) (and background information \( I \)), all valid Bayesian inferential statements about the parameters \( \theta \) are encapsulated in the posterior distribution \( \mathcal{P}(\theta) \), which combines the information provided by the data and any other information about the parameters contained in the prior.

Nonetheless, as discussed in the Introduction, relatively little consideration has been given to assessing whether the derived posterior distribution is a truthful representation of the actual constraints on the parameters that one can infer from the data, in the context of a given model. The possibility that some difference might exist is the result of the almost inevitable need to use some approximate data model \( H \) in the inference process, at least in most real-world problems. More prosaically, such differences might also be the result of errors in the implementation.

Our proposed validation procedure for testing the assumptions made (explicitly or implicitly) in the inference process (as well as the software implementation) is to make use of the fact that one may typically generate simulations of greater sophistication and realism than the data model \( H \) assumed in the analysis. One may regard the simulations as representing some alternative, more realistic data model \( H' \). Our approach is then to test whether the posterior distribution(s) obtained from the simulated data by assuming \( H' \) are consistent with the simulations generated assuming \( H \).

Our approach is as follows. One first generates \( N_d \) sets of simulated data \( \{d^{(k)}\} \) \((k = 1, 2, \ldots, N_d)\). The assumed true values \( \theta^{(k)} \) of the model parameters may be the same for each simulation generated or differ between them, depending on the nature of the inference problem. Each simulated data-set is then analysed using the inference process under investigation to obtain the resulting posterior distributions \( \mathcal{P}_k(\theta) \equiv \Pr(\theta|d^{(k)}, H, I) \). One then applies our multiple simultaneous version of the \( n \)-dimensional K-S test, as outlined in Section 2.5, to test the null hypothesis that each set of assumed parameter values \( \theta^{(k)} \) is drawn from the corresponding derived posterior \( \mathcal{P}_k(\theta) \). Thus, one calculates just a single value of the HPD probability content for each data set, which is given by

\[
\zeta_k \equiv \zeta(\theta^{(k)}) = \int_{\mathcal{P}_k(\theta) \geq \mathcal{P}_k(\theta^{(k)})} \mathcal{P}_k(\theta) \, d^n\theta. \tag{11}
\]

In real-world problems, the output of the Bayesian analysis of each data-set is usually a set of posterior-distributed samples produced by some Markov chain Monte Carlo (MCMC) (see e.g. Mackay 2003) or nested sampling (Sivia & Skilling 2006) or Feroz & Hobson 2008) algorithm. One also usually has access to the value of the parameter at each sample point, in which case an elegant proxy for \( \zeta_k \) is simply the fraction of such samples for which \( \mathcal{P}_k(\theta) \geq \mathcal{P}_k(\theta^{(k)}) \). The set of values \( \{\zeta_k\} \) \((k = 1, 2, \ldots, N_d)\) so obtained are then tested against the standard uniform distribution using the one-dimensional K-S test.

3.3 Reparameterization invariance

As discussed in Section 2.6, HPD regions are not reparameterization invariant and as a result the significance of our multidimensional K-S test will also depend on the choice of parameterization in the data model. In the context of posterior distributions, however, Druilhet & Marin 2007 have suggested a simple, yet general, way to overcome this problem.

The reason why HPD regions and MAP estimators do not share the reparameterization invariance of ML estimators is the presence of the Jacobian determinant factor in the transformation law for the prior \( \pi(\theta) \), whereas this factor is absent from the transformation law of the posterior \( \mathcal{L}(\theta) \). This is a consequence of the posterior density \( \mathcal{P}(\theta) \propto \mathcal{L}(\theta)\pi(\theta) \) being defined with respect to the Lebesgue measure on \( \theta \) (see, e.g. Lehmann & Romano 2006 sec. 5.7). One is free, however, to choose a different measure (or equivalently the unit of length or volume), but this should depend (at most) only on the likelihood. Druilhet & Marin 2007 suggest the use of the Jeffreys measure, which has the density \( |J(\theta)|^{1/2} \) with respect to the Lebesgue measure, where \( I(\theta) \) is the Fisher information matrix. Under certain regularity conditions, the elements of this matrix are given by

\[
I_{ij}(\theta) = \left\langle \frac{\partial \ln L}{\partial \theta_i} \frac{\partial \ln L}{\partial \theta_j} \right\rangle_{\theta} = -\left\langle \frac{\partial^2 \ln L}{\partial \theta_i \partial \theta_j} \right\rangle_{\theta}. \tag{12}
\]

The posterior density of \( \theta \) with respect to the Jeffreys measure on \( \theta \), which we denote by \( \mathcal{P}_{\beta}(\theta) \), is therefore given by

\[
\mathcal{P}_{\beta}(\theta) \propto \mathcal{L}(\theta) |I(\theta)|^{-1/2} \, \pi(\theta). \tag{13}
\]

Under a transformation of variables \( \theta \to \phi = \phi(\theta) \), it is straightforward to show that the Fisher information matrix transforms as \( I(\phi) = J(\theta) J^\dagger \), where \( J \) is the Jacobian matrix of the transformation, with elements \( J_{ij} = \partial \theta_i / \partial \phi_j \). Consequently, one sees immediately that the Jacobian determinant factors in the transformation law of the posterior \( \mathcal{P}_{\beta}(\theta) \) all cancel, and so the posterior in the new variables is simply

\[
\mathcal{P}_{\beta}(\phi) = \mathcal{P}_{\beta}(\theta(\phi)). \tag{14}
\]

The same criticism also applies to maximum aposteriori probability (MAP) estimators; in contrast, maximum-likelihood (ML) estimators are reparameterization invariant.

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As a result, any HPD region derived from (13), which we denote by JHPD, is invariant under smooth, monotonic reparameterizations, such that JHPD(φ) = φ(JHPD(θ)) (the corresponding MAP estimate is also similarly invariant). Moreover, a moment’s reflection reveals that the boundary of JHPD(θ) transforms into that of JHPD(φ). In the context our multidimensional K-S test, this is sufficient for the values of (ζ) derived from the original and transformed data sets, respectively, to be identical, and thus so too will be the significance of the test.

As pointed out by [Druilhet & Marin 2007], another motivation for using the Jeffreys measure is that it is also a classical non-informative prior, which minimizes the asymptotic expected Kullback–Leibler distance between the prior and the posterior distributions, provided there are no nuisance parameters [Bernardo 1979]. Thus, if no prior knowledge is available on the parameters θ, we may set π(θ) to be uniform and identify the last two factors in (13) as the non-informative Jeffreys prior.

This result implies that the significance of our test is invariant to smooth reparameterization if non-informative priors are assigned to the parameters. If the priors assigned to any of the free parameters are informative, however, any important reparameterizations of the model should be considered separately.

### 3.4 Nuisance parameters and marginalisation

It is often the case that the full parameter space is divided into parameters of interest and nuisance parameters; we will denote this by θ = (θi, θn). So far we have assumed that the validation of the posterior distribution(s) is performed over the full parameter space θ, whereas in practice it may be preferable first to marginalise over the nuisance parameters θn and then perform the validation only in the space of the parameters of interest θi.

It is clear that the significance of our test will, in general, differ between validations performed in the spaces θ and θi, respectively. More importantly, performing the validation after marginalising over the nuisance parameters θn is itself quite a subtle issue. Since the Jeffreys measure on θ does not distinguish between parameters of interest and nuisance parameters, if one simply marginalises (13) over θn, it does not follow that HPD regions derived from it in θ-space are reparameterization invariant, i.e. JHPD(θi) ≠ φ(JHPD(θi)) in general.

The most conceptually straightforward way to circumvent this difficulty (see, e.g. [Druilhet & Marin 2007]) is (where possible) to expand the prior on θ as $\pi(\theta_i, \theta_n) = \pi(\theta_i)\pi(\theta_n)$ and define the ‘marginal likelihood’ $L_m(\theta_i) = \int L(\theta_i, \theta_n)\pi(\theta_n) d\theta_n$. One may then calculate the Fisher information matrix $I_m(\theta_i)$ of the marginal likelihood $L_m(\theta_i)$ using an analogous expression to that given in (12), and define the marginalised posterior density with respect to the Jeffreys measure as

$$P_{J_m}(\theta_i) \propto L_m(\theta_i) |I_m(\theta_i)|^{-1/2} \pi(\theta_i).$$

Using the same argument as that following (13), any HPD region derived from (15) is invariant to reparameterization of the parameters of interest θi, and thus so too is the significance our multidimensional K-S test.

### 4.1 Multidimensional K-S test: toy problem

We illustrate our multidimensional K-S test by first applying it, separately, to four sets of 800 data-points \{xi, yi\} (denoted by the red crosses in the left panels of Figs 2 to 5, respectively) to test the hypothesis that each set is drawn from the 2-dimensional Gaussian reference distribution $f(x, y)$ indicated by the black 1 to 4σ contours in each figure. Although the first data set is indeed drawn from $f(x, y)$, the remaining three datasets are drawn from 2-dimensional Gaussian distributions that are rotated in the x-y-plane by different angles relative to $f(x, y)$. We use the term ‘correlation angle’ ψ to denote the angle measured anticlockwise from the y-axis of the first principal direction of the distribution from which each data set is drawn, whereas the ‘mismatch angle’ $\psi_{mis} = \psi - \psi_{ref}$, where $\psi_{ref} = 30^\circ$ is the correlation angle of the reference distribution $f(x, y)$; these values are listed in Table 1.

As mentioned in Section 3.4, the significance of our test will, in general, be different after marginalization over some parameters. Thus, in each of the four cases, it is of interest also to test, separately, the hypotheses that the one-dimensional data-sets \{xi\} and \{yi\} are drawn from the corresponding Gaussian reference distributions obtained after marginalization of $f(x, y)$ over x and y, respectively.

Since the marginal distributions are one-dimensional, it is also possible to compare directly the CDFs of the data-sets \{xi\} and \{yi\} with those obtained using our method.

#### 4.1.1 Mismatch angle: 0 degrees

Figure 2 shows our nominal toy test example in which the data points are indeed drawn from the reference distribution. The middle panel of Figure 2 shows the CDF of the corresponding \{ζi\} values obtained from the joint two-dimensional test (solid black line), and from the separate tests on the x- and y-marginals (red and blue solid lines). In all cases, the empirical CDF lies very close to that of the standard uniform distribution (dot-dashed black line), as one would expect. The resulting p-values derived from these tests are given in the first column of Table 1 and confirm that the data are consistent with being drawn from the reference distribution. We also list in parentheses the p-values obtained by applying the standard one-dimensional K-S test, separately, to the x and y-marginals, which again are consistent with the null hypothesis.

For the purposes of illustration, in the right panel of Fig. 2 we also plot the empirical PDFs of the \{ζi\}-values in each case, constructed by dividing the range [0, 1] into 20 equal-width bins; the Poissonian error-bars on each bin are also shown. As expected,
these PDFs all appear consistent with the standard uniform distribution. Although this plot plays no part in deriving the significance of the test, it allows for a visual inspection of the PDFs. As we will see later, in the case of deviations from uniformity, this can provide useful clues as to the nature of the discrepancy.

4.1.2 Mismatch angle: 15 degrees

We now consider the case where the 2-dimensional Gaussian from which the data are actually drawn is rotated through an angle \( \psi_{\text{mis}} = 15^\circ \) relative to the reference distribution, as shown in the left panel of Fig. 3. The CDFs of the resulting \( \{ \zeta_i \} \)-values are shown in the middle panel for the 2-dimensional case and the one-dimensional \( x \)- and \( y \)-marginals, respectively. In the 2-dimensional case, the empirical CDF lies close to that of a uniform distribution, and the corresponding \( p \)-value obtained is \( p = 0.59 \) (see Table 1), which shows that the test is not sufficiently sensitive to rule out the null hypothesis. For the one-dimensional marginals, however, the CDFs do appear to differ significantly from that of the uniform distribution, and this is verified by the resulting \( p \)-values of \( p = 2.7 \times 10^{-5} \) and \( p = 1.7 \times 10^{-5} \), respectively. This shows the merit of performing the test both on the full joint distribution and on the marginal distributions. For comparison, the \( p \)-values obtained by applying the standard K-S test directly to the one-dimensional \( x \)- and \( y \)-marginals, respectively, are \( p = 2.7 \times 10^{-3} \) and \( p = 1.4 \times 10^{-4} \).

The empirical binned PDFs of the \( \{ \zeta_i \} \)-values plotted in the

![Figure 2](image1.png)

**Figure 2.** Left: the data-points \( \{ x_i, y_i \} \) (red crosses) and 2-dimensional Gaussian reference distribution \( f(x, y) \) (black 1 to 4\( \sigma \) contours) to which the multidimensional K-S test is applied. In this case the data are drawn from the reference distribution, so that the ‘correlation angle’ \( \psi = 30^\circ \) and ‘mismatch angle’ \( \psi_{\text{mis}} = 0^\circ \). Middle: the CDF of the standard uniform distribution (dot-dashed black line) and the empirical CDFs of the corresponding \( \{ \zeta_i \} \)-values for the full 2-dimensional case (solid black line) and for the one-dimensional \( x \)- and \( y \)-marginals, respectively (red and blue solid lines). Right: the PDF of the standard uniform distribution (dashed black line) and the empirical PDFs of the corresponding \( \{ \zeta_i \} \)-values (constructing by dividing them into 20 bins) for the full 2-dimensional case (solid black line) and for the one-dimensional \( x \)- and \( y \)-marginals, respectively (red and blue solid lines); the error bars denote the Poissonian uncertainty.

![Figure 3](image2.png)

**Figure 3.** As in Fig. 2 but for \( \psi = 45^\circ \) and \( \psi_{\text{mis}} = 15^\circ \).

| \( \psi \)   | 30\(^\circ\) | 45\(^\circ\) | 60\(^\circ\) | −30\(^\circ\) |
|-------------|--------------|--------------|-------------|-------------|
| \( \psi_{\text{mis}} \) | 0\(^\circ\) | 15\(^\circ\) | 30\(^\circ\) | −60\(^\circ\) |
| 2-D         | 0.27         | 0.59         | 6.6 \times 10^{-3} | 1.4 \times 10^{-6} |
| \( x \) marginal | 0.37         | 2.7 \times 10^{-5} | 4.1 \times 10^{-13} | 0.41         |
|             | (0.55)       | (2.7 \times 10^{-5}) | (6.8 \times 10^{-11}) | (0.93)       |
| \( y \) marginal | 0.41         | 1.7 \times 10^{-3} | 4.0 \times 10^{-18} | 0.51         |
|             | (0.10)       | (1.4 \times 10^{-4}) | (3.9 \times 10^{-20}) | (0.11)       |

Table 1. The \( p \)-values for the null hypothesis, as obtained from our multidimensional K-S test applied to the data-points (red crosses) and reference distribution (black contours) shown in the left panels of Figs 2 to 5. In each case, the ‘correlation angle’ \( \psi \) is the angle, measured anticlockwise from the \( y \)-axis, of the first principal axis of the distribution from which the samples were actually drawn. The ‘mismatch angle’ \( \psi_{\text{mis}} = \psi - \psi_{\text{ref}} \), where \( \psi_{\text{ref}} = 30^\circ \) is the correlation angle of the assumed reference distribution. The \( p \)-values are given both for the full two-dimensional case, and for separate tests of the corresponding one-dimensional marginals in \( x \) and \( y \), respectively. Also given, in parentheses, are the \( p \)-values obtained from the standard one-dimensional K-S test applied directly to the \( x \)- and \( y \)-marginals, respectively.
Figure 4. As in Fig. 2 but for $\psi = 60^\circ$ and $\psi_{\text{mis}} = 30^\circ$.

Figure 5. As in Fig. 2 but for $\psi = -30^\circ$ and $\psi_{\text{mis}} = -60^\circ$, so that the 2-dimensional Gaussian from which the data are drawn is the reflection in the $y$-axis of the reference distribution; hence the one-dimensional marginals of the two distributions are identical.

right panel of Fig. 3 are easier to interpret that their CDFs, and provide more clues as to the nature of the discrepancy between the data and the reference distribution. In particular, the large excess in the final bin for the $x$ marginal and the corresponding decrement for the $y$ marginal are due to the non-zero ‘mismatch angle’ between the true and reference distributions; this results in the marginal distributions of the data in being too narrow in $x$ and too broad in $y$, as compared to the marginals of the reference distribution.

4.1.3 Mismatch angle: 30 degrees

Fig. 4 shows the results for the case in which the mismatch angle is increased to $\psi_{\text{mis}} = 30^\circ$. The CDFs plotted in the middle panel are now all visibly discrepant from that of a uniform distribution. From Table 1, we see that this observation is confirmed by the very small $p$-values obtained in this case. Similarly tiny $p$-values, given in parentheses in the table, are also obtained when applying the standard one-dimensional K-S test directly to the $x$- and $y$-marginals. The empirical binned PDFs plotted in the right panel exhibit similar behaviour to those obtained for $\psi_{\text{mis}} = 15^\circ$, but are more exaggerated. In particular, for the $x$-marginal, relative to the uniform distribution there is now clear decrement in the PDF around $\zeta \sim 0.2$ and an excess near $\zeta \sim 1$, with the $y$-marginal exhibiting complementary behaviour. The behaviour of the $x$-marginal ($y$-marginal) clearly follows what one would if the spread of the data is greater (less) than the width of the reference distribution; if we had instead sampled the data from a distribution that was wider (narrower) than the reference distribution (i.e. by multiplying both eigenvalues of the reference covariance matrix by the same factor), rather than rotated with respect to it, then all the PDFs would have exhibited this pattern.

4.1.4 Mismatch angle: $-60$ degrees

Fig. 5 shows the results for the case in which the distribution from which the data are sampled is the reflection in the $y$-axis of the reference distribution, resulting in $\psi_{\text{mis}} = -60^\circ$. In this case the $x$- and $y$-marginals of the two distributions are identical. As expected, the CDFs for the two marginals, plotted in the middle panel, appear completely consistent with that of a uniform distribution, which is in agreement with the large $p$-values obtained in this case (see Table 1). For the 2-dimensional case, however, the CDF is clearly discrepant, and the resulting $p$-value rules out the null hypothesis at extremely high significance. This shows, once again, the merit of testing for consistency both of the full joint distribution and its marginals. The PDF for the 2-dimensional case, shown in the right panel, again shows a clear excess near $\zeta \sim 1$, relative to the uniform distribution.

4.1.5 Comparison with theoretical expectation

In this simple toy example, one may in fact calculate straightforwardly the expected PDF of the $\zeta$-variable for any given mismatch angle $\psi_{\text{mis}}$ between the reference distribution and the true one from which the data are drawn. Each value of $\zeta$ specifies an HPD of the reference distribution that itself defines a region of the parameter space. If the probability content of the true distribution within this region is $\chi(\zeta)$, then the PDF is simply $p(\zeta) = \chi(\zeta)/\zeta$. Fig. 6
shows the expected PDFs for the $x$- and $y$-marginals for 3 different mismatch angles $\psi_{\text{mis}}$, which clearly exhibit the behaviour we observed previously in Figs 3 and 4.

We note that the sensitivity of our test in this toy example depends upon the maximum distance between the CDF of the $\zeta$ values and that of a uniform distribution, and also on the number of data samples. It is clear that, for larger mismatch angles, fewer samples will be necessary to reject the null hypothesis at some required level of significance. Indeed, from Fig. 6, one may estimate the number of samples $N$ required to obtain a given significance of when testing either the $x$- or $y$-marginal. For example, to reject the null hypothesis at a significance of 95 per cent, in the case of $\psi_{\text{mis}} = 30^\circ$, one finds that $N \sim 130$ for both the $x$- and $y$-marginals, whereas for $\psi_{\text{mis}} = 5^\circ$, one requires $N \sim 2500$ for the $x$-marginal and $N \sim 17,000$ for the $y$-marginal.

4.1.6 Testing in many dimensions

So far we have demonstrated our multidimensional K-S test as applied to a toy-model in two dimensions. Application to distributions with an arbitrary number of dimensions is straightforward and we demonstrate this here by extending the toy example to 6 dimensions. In this example, our 6-dimensional Gaussian fiducial distribution has two discrepancies compared to the true distribution from which the data are drawn: in the first dimension the fiducial Gaussian $\sigma$ is underestimated relative to the true $\sigma$ by 30%; in the second dimension, the mean of the fiducial distribution is translated by $5\sigma$. The CDFs of the 1-D marginals and the full 6-D distribution, together with their $p$-values, are shown in Fig. 7. The marginals of dimensions 1 and 2 show a strong discrepancy, with $p$-values that robustly reject the null-hypothesis, as does the CDF of the full 6-D distribution. The other dimensions are consistent with no discrepancy.

4.1.7 Simultaneous test of multiple reference distributions

Returning to our two dimensional toy example we now demonstrate the flexibility of our approach by performing a simultaneous test of multiple reference distributions, as discussed in Section 2.5. To this end, we generate simulated data as follows. For each data point, we draw a correlation angle $\psi$ from a uniform distribution between $30^\circ$ and $60^\circ$; this correlation angle is then used to define a reference distribution from which a single sample is drawn. The process is repeated 2400 times. This therefore corresponds to the extreme example, discussed in Section 2.5, in which just a single data point is drawn from each reference distribution.

Performing the simultaneous test on the full set of 2400 data-points, as outlined in Section 2.5 produces the $p$-values listed in the first column of Table 2. As expected, these values clearly support the null hypothesis that each data point was drawn from its respective reference distribution. One may also straightforwardly perform the test on subsets of the data, with the division based on some given property of the corresponding reference distributions. To this end, Table 2 also shows the $p$-values for three subsets of the data for which the correlation angle $\psi$ lies in the ranges indicated.

| $30^\circ \leq \psi < 60^\circ$ | $\psi < 40^\circ$ | $40^\circ \leq \psi < 50^\circ$ | $50^\circ \leq \psi$ |
|-----------------------------|------------------|------------------------|------------------|
| 2-D                        | 0.51             | 0.41                   | 0.98             | 0.77             |
| $x$                        | 0.52             | 0.24                   | 0.64             | 0.61             |
| $y$                        | 0.16             | 0.20                   | 0.92             | 0.57             |
Table 3. As in Table 2, but for the case in which the correlation angle $\psi_{\text{ref}}$ of the reference distribution is drawn uniformly between $30^\circ$ and $60^\circ$, and the corresponding data point is drawn from a distribution with a correlation angle $\psi = \psi_{\text{ref}} + \frac{1}{2} (\psi_{\text{ref}} - 45^\circ)$.

| $\psi_{\text{ref}}$ | $40^\circ \leq \psi_{\text{ref}} < 50^\circ$ | $50^\circ \leq \psi_{\text{ref}}$ |
|---------------------|---------------------------------------------|---------------------------------|
| 2-D                | 0.12                                        | 0.30                            |
| $x$                | $1.1 \times 10^{-6}$                      | $0.49$                          |
| $y$                | $6.5 \times 10^{-6}$                      | $0.90$                          |

4.2 Evaluation of Bayesian inference: Planck SZ clusters

We now illustrate our method for the validation of posterior distributions derived from Bayesian inference analyses. We apply this approach to the subtle, real-world example of the estimation of cluster parameters via their Sunyaev-Zel’dovich (SZ) effect in microwave temperature maps of the sky produced by the Planck mission. For a description of the Planck maps and the details of the construction of the SZ likelihood, see [Planck Collaboration et al. 2013b] and references therein.

4.2.1 Cluster parameterization

In this application, we are again considering a two-dimensional problem, namely the validation of the derived posterior distribution(s) for scale-radius parameter $\theta_s$ and integrated Comptonization parameter $Y_{\text{tot}}$ of the galaxy cluster(s). Two further parameters defining position of the cluster centre are marginalised over, and several more model parameters are held at fixed values with delta-function priors. The most important of these fixed parameters are the those defining the spherically-symmetric Generalised-Navarro-Frenk-White (GNFW) profile used to model the variation of pressure with radius in the cluster (Nagai et al. 2007). Specifically, if $r_s$ is the scale-radius of the cluster, these parameters are ($c_{500}, \gamma, \alpha, \beta$), where $c_{500} = r_{500}/r_s$ with $r_{500}$ being the radius at which the mean density is 500 times the critical density at the cluster redshift, and $\gamma, \alpha$ and $\beta$ describe the slopes of the pressure profile at $r < r_s$, $r \sim r_s$ and $r > r_s$, respectively. The parameters are fixed to the values (1.8, 0.31, 1.05, 5.49) of the ‘universal pressure profile’ (UPP) proposed by Arnaud et al. (2010).

4.2.2 Evaluation of the posterior distributions

In the analysis of Planck data, for each cluster the posterior distribution of its parameters is obtained using the PowellSnakes algorithm (Carvalho et al. 2012). The marginalised two-dimensional posterior in $(\theta_s, Y_{\text{tot}})$-space is evaluated on a high-resolution rectangular grid. The posteriors derived are typically complicated, non-Gaussian and vary between individual clusters, with different levels of correlation, skewness, and flexion, as both the signal and noise changes.

In order to achieve a tractable inference, the Planck SZ likelihood used in these analyses necessarily makes a number of simplifying assumptions. As described above, the cluster model assumes that the pressure profile is well-described by a spherically-symmetric GNFW profile, with the same fixed shape parameters for all clusters; this may be a very poor approximation for some clusters. Aside from the cluster model, the analysis assumes the background emission in which the clusters are embedded is a realisation of a homogeneous Gaussian process within the small patch of sky surrounding the cluster, whereas in reality foreground signals from Galactic emission (mostly dust) and infra-red compact sources add a non-Gaussian and anisotropic component that dominates in certain areas of the sky. It is also assumed that the cluster signal is convolved at each observing frequency with a position-independent Gaussian beam, whereas in reality the effective beams are not symmetric and vary across the sky.

4.2.3 Validation of the posterior distributions

Our method to validate the posteriors makes use of the fact that one may generate simulations of the data that are often of (considerably) greater sophistication and realism than can be modelled in the inference process. In this application, the assumptions made by the Planck SZ likelihood clearly divide into those related to the model of the cluster itself and those associated with the background emission and observing process. We will concentrate here on the latter set of assumptions, but will return to the issue of the cluster model at the end of this section. Of course, our validation process also automatically provides a test of the software implementation of the inference analysis.

We generate our simulations as follows. The mass and redshift of the injected clusters are drawn from a Tinker mass function [Tinker et al. 2008] (assuming $h = 0.7, \sigma_8 = 0.8, \Omega_m = 0.3$). The SZ parameters $(Y_{500}, \theta_{500})$ are calculated using a Planck derived scaling relation [Planck Collaboration et al. 2013]. We inject the simulated clusters (each having the GNFW profile with the fixed UPP shape parameters assumed in the analysis), convolved with the appropriate effective beams, into the real Planck sky maps. By doing so, we are thus testing all the assumptions and approximations (both explicit and implicit) made in the Planck SZ likelihood that are not associated with the cluster model itself. PwS is then run in detection mode to produce a catalogue of realistic detected clusters. The injected values of the SZ observables $(Y_{\text{tot}}, \theta_s)$ are calculated from the injected $(Y_{500}, \theta_{500})$ using the injected GNFW profile.

These simulations are then analysed using our inference pipeline, and a posterior distribution in the $(\theta_s, Y_{\text{tot}})$-space is obtained for each cluster. One subtlety that we must address is the
Table 4. The p-values for the null hypothesis, as obtained from our test for validating Bayesian inference analyses applied to the posterior distributions \( P_k(\theta_k, Y_{\text{tot}}) \) \( (k = 1, 2, \ldots, N_c) \) derived from \( N_{\text{cl}} = 918 \) simulated clusters injected into real Planck sky maps. The first column gives the p-values for the case in which the injected clusters have the same fixed ‘shape’ parameter values as assumed in the analysis, namely the ‘universal pressure profile’ \((c_{500}, \gamma, \alpha, \beta) = (1.16, 0.33, 1.06, 5.48)\). The second and third columns correspond, respectively, to when the injected clusters have \( \gamma = 0 \) and \( \beta = 4.13 \); see text for details.

| Correct profile | \( \gamma \) discrepancy | \( \beta \) discrepancy |
|-----------------|--------------------------|------------------------|
| 2-D             | 0.09                     | \(< 10^{-8}\)           | \(< 10^{-8}\) |
| \( Y_{\text{tot}} \) | 0.47                     | 0.23                   | \(< 10^{-8}\) |
| \( \theta_s \)  | 0.93                     | 0.06                   | \(< 10^{-8}\) |

issue of Eddington bias, which arises from selecting clusters using a signal-to-noise threshold. In the analysis of real data, this results in an incomplete sample of the population of clusters at lower Compton-Y fluxes, since only upward fluctuations are sampled. This bias, although present across the collection of posteriors derived, is independent of the method and assumptions employed to obtain them. Here, we choose to concentrate solely on the problem of parameter estimation in each cluster, rather than the selection of the cluster sample. Consequently, we ignore the issue of Eddington bias by analysing the complete sample of injected clusters above a minimal threshold of \( Y_{\text{tot}} > 3 \times 10^{-3} \text{arcmin}^2 \). This results in a total catalogue of \( N_{\text{cl}} = 918 \) clusters.

The posterior distributions derived for each cluster, \( P_k(\theta_k, Y_{\text{tot}}) \) \( (k = 1, 2, \ldots, N_c) \), are then validated using the method described in Section 3.2. Thus, we test the null hypothesis, simultaneously across all clusters, that the true (injected) values \( (\theta_k^{(i)}, Y_{\text{tot}}^{(i)}) \) are drawn from the corresponding derived posterior \( P_k(\theta_k, Y_{\text{tot}}) \). The empirical CDFs (for the two-dimensional case and marginals) of the resulting \( \{\zeta_k\} \)-values (one for each injected cluster), as defined in (11), are shown in the left panel of Fig. 8 and the corresponding p-values obtained from our test (both for the two-dimensional case and the marginals) are given in the second and third columns of Table 4 respectively. For a discrepancy in \( \gamma \), one sees that the test applied to the marginal posteriors is insensitive to mis-modelling the cluster profile, but that the test on the joint two-dimensional posteriors rejects the null hypothesis at extremely high significance. By contrast, for a discrepancy in \( \beta \), the test applied to the two-dimensional posteriors and both marginals all clearly reject the null hypothesis.

One may instead interpret the above results as indicating that the inference process produces \( Y_{\text{tot}} \)- and \( \theta_s \)-marginal posteriors (and parameter estimates derived therefrom) that are robust (at least to the sensitivity of the test) to assuming an incorrect value of \( \gamma \) in the analysis, but that their joint two-dimensional posterior is sensitive to this difference. By contrast, the joint distribution and both marginals are sensitive to assuming an incorrect value of \( \beta \).

To assist in identifying the nature of this sensitivity, it is again useful to consider the binned, empirical PDFs of the \( \{\zeta_k\} \)-values; these are plotted in the right panel of Fig. 8 for the test applied to the \( Y_{\text{tot}} \)-marginal. In keeping with the other results, one sees that the PDF is consistent with a uniform distribution in the case of a discrepancy in \( \gamma \), but is clearly different when there is a discrepancy in \( \beta \). Moreover, one sees that this difference is most apparent at large \( \zeta \)-values. Indeed, the shape of the PDF is consistent with an underestimation by \( \sim 30 \) per cent of the area of the 1-\( \sigma \) contour of the posterior distribution, which would not be too serious in some applications. Further investigation shows, however, that the problem here is associated with the extent of the derived posterior, but a bias low in the peak location of around 10 per cent. This bias is sub-dominant to the statistical uncertainty for most clusters in the sample but is the dominant source of error for high signal-to-noise clusters. Furthermore, as a coherent systematic across the whole sample, such an error could have damaging influence on derived information such as scaling relations or cosmological parameters.

5 CONCLUSIONS

In this paper we firstly present a practical extension of the K-S test to \( n \)-dimensions. The extension is based on a new variable \( \zeta \), that provides a universal \( R^n \rightarrow [0, 1] \) mapping based on the probability content of the highest probability density region of the refer-

\[ \text{Bayesian validation} \]
Figure 8. Left: the CDF of the standard uniform distribution (dot-dashed black line) and the empirical CDFs, for the full 2-dimensional case (solid black line) and for the one-dimensional $Y_{\text{tot}}$- and $\theta_s$-marginals, respectively, (red and blue solid lines) of the $\{\zeta_k\}$-values obtained in the validation of posterior distributions $P_k(\theta_s, Y_{\text{tot}})$ ($k = 1, 2, \ldots, N_{\text{cl}}$) derived from $N_{\text{cl}} = 918$ simulated clusters injected into real Planck sky maps. The injected clusters have the same fixed ‘shape’ parameter values as assumed in the analysis, namely the ‘universal pressure profile’ $(c_{500}, \gamma, \alpha, \beta) = (1.18, 0.31, 1.05, 5.49)$. Right: the PDF of the standard uniform distribution (dashed black line) and the empirical PDFs of the corresponding $\{\zeta_k\}$-values (constructing by dividing them into 20 bins) for the full 2-dimensional case (solid black line) and for the one-dimensional $Y_{\text{tot}}$- and $\theta_s$-marginals, respectively (red and blue solid lines); the error bars denote the Poissonian uncertainty.

Figure 9. As in Fig. 8 but showing only results derived from the $Y_{\text{tot}}$-marginal, for the case in which the injected clusters have $\gamma = 0$ (red line) and $\beta = 4.13$ (blue line), respectively.

ence distribution under consideration. This mapping is universal in the sense that it is not distribution dependent. By exploiting this property one may perform many simultaneous tests, with different underlying distributions, and provide an ensemble goodness-of-fit assessment.

We then present a new statistical procedure for validating Bayesian posterior distributions of any shape or dimensionality. The power of the method lies in the capacity to test all of the assumptions in the inferential machinery. The approach goes beyond the testing of software implementation and allows robust testing of the assumptions inherent in the construction of the likelihood, the modelling of data and in the choice of priors. This approach enables the observables to inform our understanding of the importance of the various components of the model given the data. It is, therefore, possible to tune the complexity of a model to reflect the information actually available in the posterior, improving the parsimony of the inference in keeping with Occam’s Razor. Conversely, important hidden parameters or overly-restrictive prior assumptions can be identified and treated properly.

In the application to SZ cluster parameter inference from Planck data, we demonstrate how the method can be applied to a large population where the posterior distributions vary. The information from a full cluster population can be combined to test the inference in the presence of varying stochastic and systematic uncertainties, as well as a varying signal component. For this application, we have found that the simplified Planck cluster likelihood is robust to real world complications such as Galactic foreground contamination and realistic beams. The sensitivity of the inference to prior assumptions on the outer slope of the pressure profile has been identified, as has the insensitivity to assumptions on the pressure profile in the core regions of the cluster for the inference of the integrated Compton-Y parameter.

This approach could be of use in improving the pool of available SZ data from high-resolution microwave experiments, which to date have provided either non-Bayesian point estimates for clus-
fter parameters or parameter proxies (Hasselfield et al. 2013, Reichardt et al. 2013), or unvalidated Bayesian posterior distributions (Planck Collaboration et al. 2013). These experiments have different dependencies on cluster parameters given their different resolutions, parameterisations and observation frequencies. A fuller understanding of the nature of these dependencies and the sensitivity of the derived posteriors to assumptions of the cluster model will ensure the robustness of the results and maximise the wider scientific returns due to the complementarity of the data-sets.

Beyond astronomy, the methodology we have introduced may be applied to Bayesian inference more generally, in any situation where higher levels of complexity and fidelity can be introduced into simulations than can be allowed for in a tractable analysis, or where there exists a pool of pre-existing real data with known outcomes.

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APPENDIX A: SOME PROPERTIES OF HPD REGIONS

A1 HPD region defines the smallest ‘error bar’

We wish to show that HPD is the region of smallest volume that contains an integrated probability $\zeta$. Suppose one begins with HPD and attempts to swap a small region of volume $d^\zeta$ from the interior of the HPD with a small region of volume $d^\zeta$ from
outside the HPD. For the total probability contained to remain unchanged, one requires
\[ d\zeta_{\text{HPD}} = d\zeta_{\tilde{\text{HPD}}}, \] (A1)
which, using equation (5), leads to
\[ f(x_{\tilde{\text{HPD}}}) = f(x_{\text{HPD}}) \left( \frac{d^n x_{\text{HPD}}}{d^n x_{\tilde{\text{HPD}}}} \right). \] (A2)
In order for the new region to have a smaller volume, the quantity in parentheses, \( d^n x_{\text{HPD}}/d^n x_{\tilde{\text{HPD}}} \), must be larger than unity. This would imply that
\[ f(x_{\tilde{\text{HPD}}}) > f(x_{\text{HPD}}), \] (A3)
which contradicts the definition of HPD, see Section 2.2. Therefore, for a given probability content \( \zeta \), the HPD encloses the region of smallest volume.

A2 HPD probability content is uniformly distributed
Suppose some point \( x^* \) in an \( n \)-dimensional parameter space is drawn from the PDF \( f(x) \) and the corresponding probability content of the HPD whose boundary passes through \( x^* \) is \( \zeta^* \). The probability distribution of \( \zeta^* \) may be written (somewhat baroquely) as
\[ g(\zeta^*) = \partial\zeta^* \int_{-\infty}^{\zeta^*} d\xi g(\xi), \] (A4)
\[ = \partial\zeta^* \int_{-\infty}^{\zeta^*} d\xi \int d^n x f(x) \delta(\zeta(x) - \xi), \] (A5)
where \( \zeta(x) \) is the mapping defined in (4) and the integration on \( x \) extends over the full \( n \)-dimensional parameter space. Performing the integral on \( \xi \), one obtains
\[ g(\zeta^*) = \partial\zeta^* \int d^n x f(x) \Phi(\zeta^* - \zeta(x)), \] (A6)
where \( \Phi \) is the Heaviside step function. The integral on the RHS is, however, equal simply to \( \zeta^* \). Thus, as required, one obtains \( g(\zeta^*) = 1 \) within the allowed range \( \zeta^* \in [0, 1] \).