Understanding better (some) astronomical data using Bayesian methods

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Abstract

Current analysis of astronomical data are confronted with the daunting task of modeling the awkward features of astronomical data, among which heteroscedastic (point-dependent) errors, intrinsic scatter, non-ignorable data collection (selection effects), data structure, non-uniform populations (often called Malmquist bias), non-Gaussian data, and upper/lower limits. This chapter shows, by examples, how modeling all these features using Bayesian methods. In short, one just need to formalize, using maths, the logical link between the involved quantities, how the data arize and what we already known on the quantities we want to study. The posterior probability distribution summarizes what we known on the studied quantities after the data, and we should not be afraid about their actual numerical computation, because it is left to (special) Monte Carlo programs such as JAGS. As examples, we show how to predict the mass of a new object disposing of a calibrating sample, how to constraint cosmological parameters from supernovae data and how to check if the fitted data are in tension with the adopted fitting model. Examples are given with their coding. These examples can be easily used as template for completely different analysis, on totally unrelated astronomical objects, requiring to model the same awkward data features.

Keywords: Astrophysics; Cosmology; Bayesian statistics; Regression; Scaling relations; Prediction; Model testing

1 Introduction

Astronomical data present a number of quite common awkward features (see [1] for a review):

- **heteroscedastic errors**: error sizes vary from point to point.

- **non-Gaussian data**: the likelihood is asymmetric and thus errors are, e.g. 3.4±2.5. Upper/lower limits, as 2.3 at 90 % probability, are (perhaps extreme) examples of asymmetric likelihood.

- **non uniform populations or data structure**: the number of objects per unit parameter(s) is non-uniform. This is the source of the Malmquist- or Eddington- like bias that affect most astronomical quantities, as parallaxes, star and galaxy counts, mass and luminosity, galaxy cluster velocity dispersions, supernovae luminosity corrections, etc.

- **intrinsic scatter**: data often scatter more than allowed by the errors. The extra-scatter can be due to unidentified sources of errors, often called systematic errors, or indicates an intrinsic spread of the population under study, i.e. the fact that astronomical objects are not identically equal.

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- **noisy estimates of the errors**: as every measurement, errors are known with a finite degree of precision. This is even more true when one is measuring complex, and somewhat model dependent, quantities like mass.

- **non-random sampling**: in simple terms, the objects in the sample are not a random sampling of those in the Universe. In some rare occasions in astronomy, sampling is planned to be non-random on purpose, but most of the times non-random sampling is due to selection effects: harder-to-observe objects are very often missed in samples.

- **mixtures**: very often, large samples include the population under interest, but also contaminating objects. Mixtures also arise when one measure the flux of a source in presence of a background (i.e. always).

- **prior**: we often known from past data or from theory that some values of the parameters are more likely than other. In order terms, we have prior knowledge about the parameter being investigated. If we known anything, not even the order of magnitude, about a parameter, it is difficult even to choose which instrument, or sample, should be used to measure the parameter.

- **non-linear**: laws of Nature can be more complicated than $y = ax + b$.

Bayesian methods allow to deal with these features (and also other ones), even all at the same time, as we illustrate in Sec 3 and 4 with two research examples, it is just matter of stating in mathematical terms our wordy statements about the nature of the measurement and of the objects being measured. The probabilistic (Bayesian) approach returns the whole (posterior) probability distribution of the parameters, very often in form of a Monte Carlo sampling of it.

In this paper we make an attempt to be clear at the cost of being non-rigorous. We defer the reader looking for rigour to general textbook, as [2], and to [3] for our first research example.

## 2 Parameter estimation in Bayesian Inference

Before addressing a research example, let’s consider an idealized applied problem to explain the basics of the Bayesian approach.

Suppose one is interested in estimating the (log) mass of a galaxy cluster, $lgM$. In advance of collecting any data, we may have certain beliefs and expectations about the values of $lgM$. In fact, these thoughts are often used in deciding which instrument will be used to gather data and how this instrument may be configured. For example, if we plan to measure the mass of a poor cluster via the virial theorem, we will select a spectroscopic set up with adequate resolution, in order to avoid that velocity errors are comparable to, or larger than, the likely low velocity dispersion of poor clusters. Crystalising these thoughts in the form of a probability distribution for $lgM$ provides the prior $p(lgM)$, on which relies the feasibility section of the telescope time proposal, where instrument, configuration and exposure time are set.

For example one may believe (e.g. from the cluster being somewhat poor) that the log of the cluster mass is probably not far from 13, plus or minus 1; this might be modeled by saying that the (prior) probability distribution of the log mass, here denoted $lgM$ is a Gaussian centred on 13 and with $\sigma$, the standard deviation, equal to 0.5, i.e. $lgM \sim N(13, 0.5^2)$.

Once the appropriate instrument and its set up have been selected, data can be collected. In our example, this means we record a measurement of log mass, say $obslgM200$, via, for example, a virial theorem analysis, i.e. measuring distances and velocities.

The likelihood describes how the noisy observation $obslgM200$ arises given a value of $lgM$. For example, we may find that the measurement technique allows us to measure masses in an unbiased way but with a standard error of 0.1 and that the error structure is Gaussian, i.e. $obslgM200 \sim N(lgM, 0.1^2)$, where
the tilde symbol reads “is drawn from” or “is distributed as”. If we observe $\text{obslg}M200 = 13.3$ we usually summarise the above by writing $\text{lg}M = 13.3 \pm 0.1$.

How do we update our beliefs about the unobserved log mass $\text{lg}M$ in light of the observed measurement, $\text{obslg}M200$? Expressing this probabilistically, what is the posterior distribution of $\text{lg}M$ given $\text{obslg}M200$, i.e. $p(\text{lg}M \mid \text{obslg}M200)$? Bayes Theorem ([4], [5]) tells us that

$$p(\text{lg}M \mid \text{obslg}M200) \propto p(\text{obslg}M200 \mid \text{lg}M)p(\text{lg}M)$$

i.e. the posterior (the left hand side) is equal to the product of likelihood and prior (the right hand side) times a proportionality constant of no importance in parameter estimation.

Simple algebra shows that in our example the posterior distribution of $\text{lg}M \mid \text{obslg}M200$ is Gaussian, with mean $\mu = \frac{13.0/0.5^2+13.3/0.1^2}{1/0.5^2+1/0.1^2} = 13.29$ and $\sigma^2 = \frac{1}{1/0.5^2+1/0.1^2} = 0.0096$. $\mu$ is just the usual weighted average of two “input” values, the prior and the observation, with weights given by prior and observation $\sigma$’s.

From a computational point of view, only simple examples such as the one described above can generally be tackled analytically, realistic analysis should be instead tackled numerically by special (Markov Chain) Monte Carlo methods. These are included in BUGS-like programs (6) such as JAGS (7), allowing scientists to focus on formalizing in mathematical terms our wordy statements about the quantities under investigation without worrying about the numerical implementation. In the idealized example, we just need to write in an ascii file the symbolic expression of the prior, $\text{lg}M \sim \mathcal{N}(13, 0.5^2)$, and of likelihood, $\text{obslg}M200 \sim \mathcal{N}(\text{lg}M, 0.1^2)$ to get the posterior in form of samplings. From the Monte Carlo sampling one may directly derive mean values, standard deviations, and confidence regions. For example, for a 90 % interval it is sufficient to peak up the interval that contain 90 % of the samplings.

3 First example: predicting mass from a mass proxy

Mass estimates are one of the holy grails of astronomy. Since these are observationally expensive to measure, or even unmeasurable with existing facilities, astronomers use mass proxies, far less expensive to acquire: from a(n usually small) sample of objects, the researcher measures masses, $y$ and the mass proxy, $x$. Then, he regress $x$ vs $y$ and infer $y$ for those objects having only $x$. This is the way most of the times galaxy cluster masses are estimated, for example using the X-ray luminosity, X-ray temperature, $Y_X$, $Y_{SZ}$, the cluster richness or the total optical luminosity. Here we use the cluster richness, i.e. the number of member galaxies, but with minor changes this example can be adapted for other cases.

[1] shows that predicted $y$ using the Bayesian approach illustrated here are more precise than any other method and that the Bayesian approach does not show the large systematics of other approaches. This means, in the case of masses, that more precise masses can be derived for the same input data, i.e. at the same telescope time cost.

3.1 Step 1: put in formulae what you know

3.1.1 Heteroscedasticity

Clusters have widely different richesses, and thus widely different errors. Some clusters have better determined masses than other. Heteroscedasticity means that errors have an index $i$, because they differ from point to point.

3.1.2 Contamination (mixtures), non-Gaussian data and upper limits

Galaxies in the cluster direction are both cluster members and galaxies on the line of sight (fore/background). The contamination may be estimated by observing a reference line of sight (fore/background), perhaps with a $C_i$ times larger solid angle (to improve the quality of the determination).
The mathematical translation of our words, when counts are modeled as Poisson, is:

\[
\begin{align*}
\text{obsbkg}_i & \sim \mathcal{P}(\text{nbkg}_i) \quad \# \text{Poisson with intensity} \ \text{nbkg}_i \\
\text{obstot}_i & \sim \mathcal{P}(\text{nbkg}_i/C_i + n200_i) \quad \# \text{Poisson with intensity} \ (\text{nbkg}_i/C_i + n200_i)
\end{align*}
\]

The variables \(n200\) and \(\text{nbkg}\) represent the true richness and the true background galaxy counts in the studied solid angles, whereas we add a prefix “obs” to indicate the observed values.

Upper limits are automatically accounted for. Suppose, for exposing simplicity, that we observed 5 galaxies, \(\text{obstot}_i = 5\), in the cluster direction and that in the control field direction (with \(C_i = 1\) for exposing simplicity) we observe four background galaxies, \(\text{obsbkg} = 4\). With one net galaxy and Poisson fluctuations of a few, \(n200_i\) is poorly determined at best, and the conscientious researcher would probably reports an upper limits of a few. To use the information contained in the upper limit in our regression analysis we only need to list in the data file the raw measurements \((C_i = 1, \text{obstot}_i = 5, \text{obsbkg}_i = 4)\), as for the other clusters. These data will be treated independently on whether an astronomer decides to report a measurement or an upper limit, because Nature doesn’t care about astronomer decisions.

### 3.1.3 Non-linearity and extra-scatter

The relation between mass, \(M200\), and proxy, \(n200\), (richness) is usually parametrized as a power-law:

\[
M200_i \propto n200_i^\beta
\]

Allowing for a Gaussian intrinsic scatter \(\sigma_{\text{scat}}\) (clusters of galaxies of a given richness may not all have the very same mass) and taking the log, previous equation becomes:

\[
\lg M200_i \sim \mathcal{N}(\alpha + \beta \log(n200_i), \sigma_{\text{scat}}^2) \quad \# \text{Gaussian scatter around} \ (M200_i \propto n200_i^\beta)
\]

where the intercept is \(\alpha\) and the slope is \(\beta\).

### 3.1.4 Noisy errors

Once logged, mass has Gaussian errors. In formulae:

\[
\text{obslgM200}_i \sim \mathcal{N}(\lg M200_i, \sigma_i^2) \quad \# \text{Gauss errors on lg mass}
\]

However, errors (as everything) is measured with a finite degree of precision. We assume that the measured error, \(\text{obserllgM200}_i\), is not biased (i.e. it is not systematically larger or smaller that the true error, \(\sigma_i\)) but somewhat noisy. If a \(\chi^2\) distribution is adopted, it satisfies both our request of unbiasness and noisiness. In formulae:

\[
\text{obserllgM200}_i^2 \sim \sigma_i^2 \chi^2/\nu \quad \# \text{Unbiased errors}
\]

where the parameter \(\nu\) regulates the width of the distribution, i.e. how precise measured errors are. Since we are 95% confident that quoted errors are correct up to a factor of 2,

\[
\nu = 6 \quad \# \text{95\% confident within a factor 2}
\]
3.1.5 Prior knowledge and population structure

The data used in this investigation are of quality good enough to determine all parameters, but one, to a sufficient degree of accuracy that we should not care about priors and we can safely take weak (almost uniform) priors, zeroed for un-physical values of parameters (to avoid, for example, negative richnesses). The exception is given by the prior on the errors (i.e. $\sigma_i$), for which there is only one measurement per datum. The adopted prior (eq. 11) is supported by statistical considerations (see [3] for details). The same prior is also used for the intrinsic scatter term, although any weak prior would return the same result, because this term is well determined by the data.

$$
\alpha \sim N(0.0, 10^4) \quad \# \text{Almost uniform prior on intercept (7)}
$$

$$
\beta \sim t_1 \quad \# \text{Uniform prior on angle (8)}
$$

$$
n_{200_i} \sim U(0, \infty) \quad \# \text{Uniform, but positive, cluster richness (9)}
$$

$$
\text{nbkg}_i \sim U(0, \infty) \quad \# \text{Uniform, but positive, background rate (10)}
$$

$$
1/\sigma_i^2 \sim \Gamma(\epsilon, \epsilon) \quad \# \text{Weak prior on error (11)}
$$

$$
1/\sigma_{scat}^2 \sim \Gamma(\epsilon, \epsilon) \quad \# \text{Weak prior on intrinsic scatter (12)}
$$

Richer clusters are rarer. Therefore, the prior on the cluster richness is, for sure, not uniform, contrary to our assumption (eq. 9). Modeling the population structure is unnecessary for the data used in [3] and here, but is essential if noisier richnesses were used. Indeed, [3] shows that a previous published richness-mass calibration, which uses richnesses as low as $obsn_{200} = 3$ and neglects the $n_{200}$ structure, shows a slope biased by five times the quoted uncertainty. Therefore, the population structure cannot be overlooked in general.

3.2 Step 2: remove TeXing, perform stochastic computations and publish

At this point, we have described, using mathematical symbols, the link between the quantities that matter for our problem, and we only need to compute the posterior probability distribution of the parameters by some sort of sampling (the readers with exquisite mathematical skills may instead attempt an analytical computation).

Just Another Gibb Sampler (JAGS\footnote{http://calvin.iarc.fr/~martyn/software/jags/} ([7]) can return it at the minor cost of de-Texing equations 1 to 12 (compare them to the JAGS code below). Poisson, Normal and Uniform distributions become dpois, dnorm, dunif, respectively. JAGS, following BUGS ([6]), uses precisions, $prec = 1/\sigma^2$, in place of variances $\sigma^2$. Furthermore, it uses neperian logarithms, instead of decimal ones. Eq. 5 has been rewritten using the property that the $\chi^2$ is a particular form of the Gamma distribution. Eq. 3 is split in two JAGS lines for a better reading. The arrow symbol reads “take the value of”. $\text{obsvarlgM}_{200}$ is the square of $\text{obsvarlgM}_{200}$. For computational advantages, $\log(n_{200})$ is centred at an average value of 1.5 and $\alpha$ is centred at -14.5. Finally, we replaced infinity with a large number.

The model above, when inserted in JAGS, reads:

```
model
{
  for (i in 1:length(obstot)) {
    obsbkg[i] ~ dpois(nbkg[i]) # eq 1
    obstot[i] ~ dpois(nbkg[i]/C[i]+n200[i]) # eq 2
    n200[i] ~ dunif(0,3000) # eq 9
    nbkg[i] ~ dunif(0,3000) # eq 10
  }
```

1
Figure 1: Richness-mass scaling. The solid line marks the mean fitted regression line, while the dashed line shows this mean plus or minus the intrinsic scatter $\sigma_{\text{scat}}$. The shaded region marks the 68% highest posterior credible interval for the regression. Error bars on the data points represent observed errors for both variables. The distances between the data and the regression line is due in part to the measurement error and in part to the intrinsic scatter. From [3], reproduced with permission.

```
precy[i] ~ dgamma(1.0E-5, 1.0E-5)  # eq 12
obslogM200[i] ~ dnorm(lgM200[i], precy[i])  # eq 4
obsvarlgM200[i] ~ dgamma(0.5*nu, 0.5*nu*precy[i])  # eq 5
z[i] <- alpha + 14.5 + beta*(log(n200[i])/2.30258 - 1.5)  # eq 3
lgM200[i] ~ dnorm(z[i], prec.intrscat)  # eq 3

intrscat <- 1/sqrt(prec.intrscat)
prec.intrscat ~ dgamma(1.0E-5, 1.0E-5)  # eq 11
alpha ~ dnorm(0.0, 1.0E-4)  # eq 7
beta ~ dt(0, 1, 1)  # eq 8
nu <- 6  # eq 6
```

JAGS samples the posterior distribution of all quantity of interests, such as intercept, slope and intrinsic scatter by Gibb sampling (a sort of Monte Carlo). Form these samplings, it is straightforward to compute (posterior) mean and standard deviations (by computing the average and the standard deviation!), to plot posterior marginals (by ignoring the values of the other parameters) and confidence contours, data and mean model, etc. Therefore, our effort is over, we only need to produce nice plots and summaries of our results.

Figure 1 shows the data used in this analysis (see [3] for details), the mean scaling (solid line) and its 68% uncertainty (shaded yellow region) and the mean intrinsic scatter (dashed lines) around the mean relation. The $\pm 1$ intrinsic scatter band is not expected to contain 68% of the data points, because of the presence of measurement errors.

Figure 2 shows the posterior marginals for the intercept, slope and intrinsic scatter $\sigma_{\text{scat}}$. These marginals are reasonably well approximated by Gaussians. The intrinsic mass scatter at a given richness, $\sigma_{\text{scat}} = \sigma_{\text{lgM200}|\text{log}n200}$, is small, $0.19 \pm 0.03$. (Unless otherwise stated, results of the statistical computations are quoted in the form $x \pm y$ where $x$ is the posterior mean and $y$ is the posterior standard deviation.)

The found relation is:

$$\log M_{200} = (0.96 \pm 0.15) (\log n_{200} - 1.5) + 14.36 \pm 0.04 \quad (13)$$
3.3 Predicting masses

As mentioned, one of the reasons why astronomers regress a quantity \( x \) against another one, \( y \), is to predict the latter when a direct measurement is missing (usually because observationally expensive to acquire). It is clear that the uncertainty on the predicted \( y \), called \( \tilde{y} \) hereafter, should account for: a) the intrinsic scatter between \( y \) and \( x \) (a larger scatter implies a lower quality \( \tilde{y} \) estimate); b) the uncertainty of the \( x \) (the larger it is, the noisier will be the \( \tilde{y} \)); c) the quality of the calibration between \( y \) and \( x \) (better determined relations should return more precise estimates of \( \tilde{y} \)); and d) extrapolation errors, i.e. should penalize attempts to infer \( \tilde{y} \) values corresponding to \( x \) values absent from the calibrator sample (e.g. outside the range sampled by it). All these requirements are satisfied using the posterior predictive distribution,

\[
p(\tilde{y}) = \int p(\tilde{y} | \theta) p(\theta | y) d\theta
\]  

(14)

where \( \theta \) are the regression parameters (intercept, slope, intrinsic scatter). This apparently ugly expression is easy to understand: one should combine (multiply) the uncertainties of the calibrating relation, \( p(\theta | y) \), to the uncertainty of predicting new data if the calibrating relation were perfectly known, \( p(\tilde{y} | \theta) \). Since we are now interested in predicted values only, we get rid of non-interesting parameters (\( \theta \)) by marginalization (integration).

Posterior predictive distributions are so basic to be introduced at page 8 of the > 700 pages “Bayesian Data Analysis” book ([2]) and to be offered as a standard output of JAGS. Of course, we need to list the \( x \) values (richnesses), and errors of the clusters for which we want to infer \( \tilde{y} \) (predicted mass) in the data file, listing for example in the data file \( obsbkg = 12, obtot = 32, C = 5, \) and mass \( obslgM200 = NA \) (“not available”), to indicate that this quantity should be estimated using the regression computed from the points with available masses and galaxy counts. JAGS returns \( p(\tilde{y}) \) in form of sampling and therefore, as for any other parameter, a point estimate may be obtained by taking the average and a 68 % (credible) interval can be derived by taking the interval including 68 % of the samplings. Returned values behave as expected, and indeed have large errors when masses are estimated for clusters with richnesses outside the range where the calibration has been derived ([3])

4 Second example: Cosmological parameters from SNIa

Supernovae (SNIa) are very bright objects with very similar luminosities. The luminosity spread can be made even smaller by accounting for the correlation with colour and stretch parameter (the latter is a mea-
Figure 3: Apparent magnitude vs redshift of the SNIa sample (upper panels), and their residual from a \( \Omega_M = 0.3, \Omega_A = 0.7 \) cosmological model (bottom panels) before (left panels) or after (right panels) correcting for stretching and color parameter.

These features make SNIa very useful for cosmology: they can be observed far away and the relation between flux (basically the rate of photons received) and luminosity (the rate of photons emitted) is modulated by the luminosity distance (to the square), which in turns is function of the cosmological parameters. Therefore, measuring SNIa fluxes (and redshift) allows us to put constraints on cosmological parameters. The only minor complication is that SNIa luminosities are function of their colour and stretch parameter, and these parameters have an intrinsic scatter too, which in turns has to be determined from the data at the same time as the other parameters.

[10] shows that the Bayesian approach delivers tighter statistical constraints on the cosmological parameters over 90% of the times, that it reduces the statistical bias typically by a factor \( \sim 2 - 3 \) and that it has better coverage properties than the usual chi-squared approach.

In this second example we can proceed a bit faster in illustrating this non-linear regression with heteroscedastic errors, non-uniform data structure and intrinsic scatter. In this example, we also briefly discuss the prior sensitivity, i.e. how much the results are affected by the chosen prior, and we also check the quality of the model fit.
Figure 4: Observed values of the stretch parameters, $obsx_i$, and of the colour parameter, $obs_c_i$ ranked by error. Points scatter more than the error bars (see the left side of the figure). The dashed lines indicate the size of the intrinsic scatter as determined by our analysis.

4.1 Step 1: put in formulae what you know

We observe SNIa magnitudes $obsm_i (= -2.5 \log (flux) + c)$ with Gaussian errors $\sigma_{m,i}$, i.e.

$$obsm_i \sim N(m_i, \sigma_{m,i}^2)$$  \hspace{1cm} (15)

These $m_i$ are related to the distance modulus $distmod_i$, via

$$m_i = M + distmod_i - \alpha x_i + \beta c_i$$

with a Gaussian intrinsic scatter $\sigma_{scat}$. More precisely:

$$m_i \sim N(M + distmod_i - \alpha x_i + \beta c_i, \sigma_{scat}^2)$$  \hspace{1cm} (16)

where $M$ is the (unknown) mean absolute magnitude of SNIa, and $\alpha$ and $\beta$ allow to reduce the SNIa luminosity scatter by accounting for the correlation with the stretch and colour parameters.
Similarly to [10], the $M$, $\alpha$, $\beta$ and $\log \sigma_{\text{scat}}$ priors are taken uniform in a wide range:

$$\log_{10} \sigma_{\text{scat}} \sim \mathcal{U}(-3, 0)$$

$$\alpha \sim \mathcal{U}(-2, 2)$$

$$\beta \sim \mathcal{U}(-4, 4)$$

$$M \sim \mathcal{U}(-20.3, -18.3)$$

$x_i$ and $c_i$ are the true value of the stretch and colour parameters, of which we observe (the noisy) $\text{obs}x_i$ and $\text{obs}c_i$ with errors $\sigma_{x,i}$ and $\sigma_{c,i}$. In formulae:

$$\text{obs}x_i \sim \mathcal{N}(x_i, \sigma_{x,i}^2)$$

$$\text{obs}c_i \sim \mathcal{N}(c_i, \sigma_{c,i}^2)$$

The key point of the modeling is that the $\text{obs}x_i$ and $\text{obs}c_i$ values scatter more than their errors, but not immensely so, see Fig 4. The presence of a non-uniform distribution induces a Malmquist-like bias if not accounted for (e.g. large $\text{obs}x_i$ values are more likely low $x_i$ values scattered to large values than vice versa, because of the larger abundance of low $x_i$ values). Therefore, we model, as [10] do, the individual $x_i$ and $c_i$ as drawn from independent normal distributions centered on $x_m$ and $c_m$ with standard deviation $R_x$ and $R_c$ respectively. In formulae:

$$x_i \sim \mathcal{N}(x_m, R_x^2)$$

$$c_i \sim \mathcal{N}(c_m, R_c^2)$$

We take uniform priors for $x_m$ and $c_m$, and uniform priors on $\log R_x$ and on $\log R_c$, between the indicated boundaries:

$$x_m \sim \mathcal{U}(-10, +10)$$

$$c_m \sim \mathcal{U}(-3, +3)$$

$$\log_{10} R_x \sim \mathcal{U}(-5, +2)$$

$$\log_{10} R_c \sim \mathcal{U}(-5, +2)$$

That’s almost all: we need to remember the definition of distance modulus:

$$\text{distmod}_i = 25 + 5 \log_{10} d_l$$

where the luminosity distance, $d_l$ is a complicate expression, involving integrals, of the redshift $z_i$ and the cosmological parameters $\Omega_\Lambda$, $\Omega_M$, $w$, $H_0$ (see any recent cosmology textbook for the mathematical expression).

Redshift, in the considered sample, have heteroscedastic Gaussian errors $\sigma_{z,i}$:

$$\text{obs}z_i \sim \mathcal{N}(z_i, \sigma_{z,i}^2)$$

The redshift prior is taken uniform.
\[ z_i \sim \mathcal{U}(0, 2) \]  

Supernovae alone do not allow to determine all cosmological parameters, so we need external prior on them, notably on \( H_0 \), taken from \cite{11} to be

\[ H_0 \sim \mathcal{N}(72, 8^2) \]  

At this point, we may decide which sets of cosmological models we want to investigate using SNIa, for example a flat universe with a possible \( w \neq 0 \) with the following priors:

\[
\begin{align*}
\Omega_M &\sim \mathcal{U}(0, 1) \\
\Omega_k &= 0 \\
w &\sim \mathcal{U}(-4, 0)
\end{align*}
\]  

or a curved Universes with \( w = -1 \):

\[
\begin{align*}
\Omega_M &\sim \mathcal{U}(0, 1) \\
\Omega_k &\sim \mathcal{U}(-1, 0) \\
w &= -1
\end{align*}
\]  

or any other set.

Both considered cosmologies have:

\[ \Omega_k = 1 - \Omega_m - \Omega_\Lambda \]  

Finally, one may want to use some data. As shortly mentioned, we use the compilation of 288 SNIa in \cite{9}.

### 4.2 Step 2: remove TeXing, perform stochastic computations and publish

Most of the distributions above are Normal, and the posterior distribution can be almost completely analytically computed \cite{10}. However, numerical evaluation of the stochastic part of the model on an (obsolete) laptop takes about one minute, therefore there is no need for speed up. Instead, the evaluation of the luminosity distance is CPU intensive (it takes \( \approx 10^3 \) more times, unless approximate analytic formulae for the luminosity distance are used), because an integral has to be evaluated a number of times equal to the number of supernovae times the number of target posterior samplings, i.e. about four millions times in our numerical computation. The JAGS implementation of the luminosity distance integral is implemented as a sum over a tightly packed grid on redshift.

As the previous example, eq 15 to 32 can be de-Texed and used in JAGS, adding one of the two set of priors, 33-35 or 36-38, depending on which problem one is interested in.

```jags
data {
  # JAGS like precisions
  precmag <- 1/errmag/errmag
  precobsc <- 1/errobsc/errobsc
  precobsx <- 1/errobsx/errobsx
```
precz <- 1/errz/errz
# grid for distance modulus integral evaluation
for (k in 1:1500){
  grid.z[k] <- (k-0.5)/1000.
}
step.grid.z <- grid.z[2] - grid.z[1]
}
model {
for (i in 1:length(obsz)) {
  obsm[i] ~ dnorm(m[i], precmag[i])  # eq 15
  m[i] ~ dnorm(Mm + distmod[i] - alpha * x[i] + beta * c[i], precM)  # eq 16
  obsc[i] ~ dnorm(c[i], precobsc[i])  # eq 22
  c[i] ~ dnorm(cm, precC)  # eq 24
  obsx[i] ~ dnorm(x[i], precobsx[i])  # eq 21
  x[i] ~ dnorm(xm, precx)  # eq 23
  # distmod definition & H0 term
  distmod[i] <- 25 + 5/2.3026 * log(dl[i]) - 5/2.3026 * log(H0/300000)
  z[i] ~ dunif(0, 2)  # eq 31
  obsz[i] ~ dnorm(z[i], precz[i])  # eq 30
  # distmod computation: slow and tedious
  tmp2[i] <- sum(step(z[i] - grid.z) * (1+w) / (1+grid.z)) * step.grid.z
  omegad[i] <- omegal * exp(3 * tmp2[i])
  xx[i] <- sum(pow((1+grid.z)^3*omegam + omegade[i] + (1+grid.z)^2*omegak,-0.5)*
                step.grid.z*step(z[i]-grid.z))
  # implementing if, to avoid diving by 0 added 1e-7 to omegak
  zz[1,i] <- sin(xx[i]*sqrt(abs(omegak)))*((1+z[i])/sqrt(abs(omegak+1e-7)))
  zz[2,i] <- xx[i] * (1+z[i])
  zz[3,i] <- (exp(xx[i]*sqrt(abs(omegak))) - exp(-xx[i]*sqrt(abs(omegak))))/2 *
                (1+z[i]) / sqrt(abs(omegak+1e-7))
  dl[i] <- zz[3,i]
}
  b <- 1 + (omegak==0) + 2*(omegak > 0)
  # JAGS uses precisions
  precM <- 1 / intrscatM / intrscatM
  precC <- 1 / intrscatC / intrscatC
  precx <- 1 / intrscatx / intrscatx
  # priors
  Mm ~ dunif(-20.3, -18.3)  # eq 20
  alpha ~ dunif(-2, 2.0)  # eq 18
  beta ~ dunif(-4, 4.0)  # eq 19
  cm ~ dunif(-3, 3)  # eq 26
  xm ~ dunif(-10, 10)  # eq 25
  # uniform prior on logged quantities
  intrscatM <- pow(10, lgintrscatM)  # eq 17
  lgintrscatM ~ dunif(-3, 0)  # eq 17
  intrscatx ~ pow(10, lgintrscatx)  # eq 27
  lgintrscatx ~ dunif(-5, 2)  # eq 27
  intrscatC <- pow(10, lgintrscatC)  # eq 28
  lgintrscatC ~ dunif(-5, 2)  # eq 28
  # cosmology priors
  H0 ~ dnorm(72, 1/8./8.)  # eq 32
  omegal <- 1 - omegam - omegak  # eq 39
  # cosmology priors 1st set LCDM
  # omegam ~ dunif(0, 1)  # eq 36
  # omegak ~ dunif(-1, 1)  # eq 37
  # w <- -1  # eq 38
  # cosmology priors 2nd set: wCDM
  omegam ~ dunif(0, 1)  # eq 33
Figure 5 shows the prior (dashed blue line) and posterior (histogram) probability distribution for the three intrinsic scatter terms present in the cosmological parameter estimation: the scatter in absolute luminosity after colour and stretch corrections, \( \sigma_{\text{scat}} \), the intrinsic scatter in the distribution of the colour and stretch terms \( (R_c \text { and } R_x) \). This plot shows that the posterior probability at intrinsic scatters near zero is approximately zero and thus that the three intrinsic scatter terms are necessary parameters for the modeling of SNIa, and not useless complications. The three posteriors are dominated by the data, being the prior quite flat in the range where the posterior is appreciably not zero (Figure 5). Therefore, any other prior choice, as long as smooth and shallow over the shown parameter range, would have returned indistinguishable results.

Not only SNIa have luminosities that depend on colour and stretch terms, but these in turns have their own probability distribution (taken Gaussian for simplicity) with a well determined width. Figure 6 depicts the Malmquist-like bias one should incur if the spread of the distribution of colour and stretch parameters is ignored: it reports the observed values (as in Fig 4), \( \text{obs} x_i \) and \( \text{obs} c_i \) as well as the true values \( x_i \) and \( c_i \) (posterior means). The effect of equations 23 and 24 is to pull values toward the mean, and more so those with large errors, to compensate the systematic shift (Malmquist-like bias) toward larger observed values.

Figure 7 shows the probability distribution of the two the colour and stretch slopes: \( \alpha = 0.12 \pm 0.02 \) and \( \beta = 2.70 \pm 0.14 \) respectively. As for the intrinsic scatter terms, the posterior is dominated by the data and therefore any other prior, smooth and shallow, would have returned indistinguishable results.

Finally, Fig 8 reports perhaps the most wanted result: contours of equal probability for the cosmological parameters \( \Omega_M \) and \( w \).

For one dimensional marginals, we found: \( \Omega_M = 0.40 \pm 0.10 \) and \( w = -1.2 \pm 0.2 \), but with non-Gaussian probability distributions.

### 4.3 Model checking

The work of the careful researcher does not end by finding the parameter set that best describe the data, (s)he also checks whether the adopted model is a good description of the data, or it is misspecified, i.e. in tension with the fitted data. In the non-Bayesian paradigm this is often achieved by computing a p-value, i.e. the probability to obtain data more discrepant than those in hand once parameters are taken at the best fit value. The Bayesian version of the concept (e.g. [2]) acknowledges that parameters are not perfectly known, and therefore one should also explore, in addition to best fit value, other values of the parameters. Therefore,
Figure 6: Effect of the population structure for the stretch and the colour parameters. Each tick goes from
the observed value to the posterior mean. The population modeling attempt to counterbalance the increased
spread (Malmquist-like), especially those with larger error (on the right, in the figures), pulling values toward
the mean.

Figure 7: Prior and posterior probability distribution for the colour and stretch slopes of the SNIa problem.
The black jagged histogram shows the posterior as computed by MCMC, marginalised over the other pa-
rameters. The red curve is a Gauss approximation of it. The shaded (yellow) range shows the 95% highest
posterior credible interval. The adopted (uniform) priors are indicated by the blue dotted curve.
discrepancy becomes a vector, instead of a scalar, of dimension \( j \), that measure the distance between the data and \( j \) models, one per set of parameters considered. Of course, more probable models should occur more frequently in the list to quantify that discrepancy from an unlikely model is less detrimental than discrepancy from a likely model. In practice, if parameters are explored by sampling, it is just matter of computing the discrepancy of the data in hand for each set \( j \) of parameters stored in the chain, instead of relying on one single set of parameter (say, those that maximize the likelihood). Then, one repeats the computation for fake data generated from the model, and counts how many times fake data are more extreme of real data.

For example, if we want to test the modeling of the observed spread of magnitude (i.e. equation 15 and 16), let’s define:

\[
mcor_i = M + \text{distmod}_i - \alpha x_i + \beta c_i
\]  

(40)

We generate fake supernovae mag:

\[
m_{\text{fake}}_i \sim N(m_{\text{cor}}_i, \sigma^2_{\text{scat}})
\]  

(41)

and fake observed values of them,

\[
obs_{m_{\text{fake}}}_i \sim N(m_{\text{fake}}_i, \sigma^2_{m,i})
\]  

(42)

Then, we adopt a modified \( \chi^2 \) to quantify discrepancy (or its contrary, agreement). For the real data set we have:

\[
\chi^2_{\text{real}, j} = \sum_i \frac{(obs_{m.i} - m_{\text{cor}}_{i,j})^2}{\sigma^2_{m.i} + E^2(\sigma_{\text{scat}})}
\]  

(43)

where summation is over the data and \( j \) refers to the index in the sampling chain.

Apart for the \( j \) index, eq. 43 is just the usual \( \chi^2 \), the difference between observed, \( obs_{m.i} \), and true \( m_{\text{cor}}_i \), values, weighted by the expected variance, computed as quadrature sum of errors, \( \sigma_{m.i} \), and supernovae mag intrinsic scatter \( \sigma_{\text{scat}} \). The \( \chi^2 \) of the \( j \)th fake data set, \( \chi^2_{\text{fake}, j} \) is:

\[
\chi^2_{\text{fake}, j} = \sum_i \frac{(obs_{m_{\text{fake}}_{i,j}} - m_{\text{cor}}_{i,j})^2}{\sigma^2_{m,i} + E^2(\sigma_{\text{scat}})}
\]  

(44)
At this point, we only need to compute for which fraction of the simulations $\chi^2_{\text{fake},j} > \chi^2_{\text{real},j}$ and quote the result. If the modeling is appropriate, then the computed fraction (p-value) is not extreme (far from zero or one). If not, our statistically modeling need to be revised, because the data are in disagreement with the model.

We performed 15000 simulations, each one generating 288 fake measurements of SNIa measurements. In practice, we added the following three JAGS lines:

```jags
mcor[i] <- Mm + distmod[i] - alpha*x[i] + beta*c[i]  # eq 40
m.fake[i] ~ dnorm(mcor, precM)  # eq 41
obsm.fake[i] ~ dnorm(m.fake[i], precmag[i])  # eq 42
```

and we can simplify eq 16 in

```jags
m[i] ~ dnorm(mcor[i], precM)  # eq 16
```

We found a p-value of 45 %, i.e. that the discrepancy of the data in hand is quite typical (similar to the one of the fake data). Therefore, real data are quite common and the tested part of the model shows no evidence of misspecification. The careful researcher should then move to the other parts of the model, whose detailed exploration is left as exercise.

In such exploration of possible model misfits, it is very useful to visually inspect several data summaries to guide the choice of which discrepancy measure one should adopt (eq. 43 or something else?), and, if the adopted model turns out to be unsatisfactory, to guide how to revise the modeling of the the tested part of the model. For example, a possible (and common) data summary is the distribution of normalized residuals, that for $obsx_i$ reads:

$$stdresobsx_i = \frac{obsx_i - E(x_i)}{\sqrt{\sigma^2_{x,i} + E^2(R_x)}}$$  \hspace{1cm} (45)$$

i.e. observed minus expected value of $x_i$ divided by their expected spread (the sum in quadrature of errors and intrinsic spread). A similar summary may be built for $obsc_i$ too. To first order (at least), standardized residuals should be normal distributed with standard deviation one (by construction). Fig 9 shows the distribution of normalized residuals of both $obsx_i$ and $obsc_i$, with superposed a Gaussian centered in 0 with standard deviation equal to one (in blue). Both distributions show a possible hint of asymmetry. At this point, the careful researcher may want to use a discrepancy measure sensitive to asymmetries, as the skewness index, in addition to the $\chi^2$ during model testing. While leaving the actual computation to the reader, we emphasize that if an extreme Bayesian p-value if found (on $obsx_i$ for exposing simplicity), then one may

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2Skilled readers may note that we are dealing, by large, with Gaussian distributions, and may attempt an analytic computation.
replace its modeling (eq. 23 in the case of \( obsx_i \)) with a distribution allowing a non-zero asymmetry and this can be easily performed in a Bayesian approach, and easily implemented in JAGS, it is just matter of replacing the adopted Gaussian with an asymmetric distribution. If instead the data exploration gives an hint of double-bumped distribution (again on \( obsx_i \) for exposing simplicity), and an extreme Bayesian p-value is found when a measure of discrepancy sensitive to double-bumped distributions is adopted, then one may adopt a mixture of Gaussians, replacing (eq. 23 in the case of \( obsx_i \)) with

\[
 obsx_i \sim \lambda \mathcal{N}(x_i, \sigma_{x,i}^2) + (1 - \lambda)\mathcal{N}(xx_i, \sigma_{xx,i}^2) \quad (46)
\]

Even more simple is the (hypothetical) case of possible distribution (again of \( obsx_i \) for exposing simplicity) with fat tails: one may adopt a Cauchy distribution. In such case, coding in JAGS is it is just matter of replacing a \( \text{dnorm} \) with \( \text{dt} \) in the JAGS implementation. And so on.

In summary, model checking consists in updating the model until it produces data similar to those in hand. One should start by carefully and attentively inspect the data and their summaries. This inspection should suggest a discrepancy measure to be used to quantify the model misfit and, if one is found, to guide the model updating. The procedure should be iterated until the model fully reproduces the data.

5 Summary and conclusions

These two analysis offer a template for modeling the common awkward features of astronomical data, namely heteroscedastic errors, non-Gaussian likelihood (inclusive of upper/lower limits), non-uniform populations or data structure, intrinsic scatter (either due to unidentified source of errors, or due to population spreads), noisy estimates of the errors, mixtures and prior knowledge.

In a Bayesian framework, learning from the data and the prior it is just matter of formalizing in mathematical terms our wordy statements about the quantities under investigation and how the data arise. The actual numerical computation of the posterior probability distribution of the parameters is left to (special) Monte Carlo programs, of which we don’t need to be afraid more than numerical methods (Monte Carlo) used to compute the integral of a function.

The great advantage of the Bayesian modeling is its high flexibility: if the data (or theory) call for a more complex modeling, or call for using distributions different from those initially taken, it is just matter of replacing them in the model, because there are no simplifications forced by the need of reaching the finishing line, as instead is the case in other modeling approaches. Furthermore, if one is interested in constraining another cosmological model, for example one with a redshift-dependent dark energy equation of state \( w = w_0 + w_1(1 + z)/(1 + z_p) \), one should just replace \( w \) in the JAGS code with the above equation. This flexible, hierarchical, modeling is native with Bayesian methods.

In both our two examples, the Bayesian approach performs, unsurprisingly, better than than non-Bayesian methods obliged to discard part of the available information in order to reach the finishing line: Bayesian methods just use all the provided information.

As a final note, we remember that the careful researcher, whether using a Bayesian modeling or not, before publishing his own result should check that the numerical computation is adequate for his purpose and that the model is appropriate for the data. Therefore, if you, gentle reader, are using our examples as template, remember to include in your work a sensitivity analysis, by checking that your assumptions (both likelihood and prior) are reasonable. Some prior sensitivity analysis has been performed in both examples to emphasize the importance of showing how much conclusions (the posterior) rely on assumptions (prior). For what concerns the likelihood, i.e. misfitting, the key point consists in updating the model formulation until it produces data similar to those in hand, as we have shown in great detail for the second example.

More in general, we hope that the template modeling shown in these two examples may be useful for any analysis confronted with modeling the awkward features of astronomical data, among which heteroscedastic
(point-dependent) errors, intrinsic scatter, data structure, non-uniform population (often called Malmquist bias) and non-Gaussian data, inclusive of upper/lower limits.

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