Approximate Bayesian computation and Bayes linear analysis: Towards high-dimensional ABC

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Abstract

Bayes linear analysis and approximate Bayesian computation (ABC) are techniques commonly used in the Bayesian analysis of complex models. In this article we connect these ideas by demonstrating that regression-adjustment ABC algorithms produce samples for which first and second order moment summaries approximate adjusted expectation and variance for a Bayes linear analysis. This gives regression-adjustment methods a useful interpretation and role in exploratory analysis in high-dimensional problems. As a result, we propose a new method for combining high-dimensional, regression-adjustment ABC with lower-dimensional approaches (such as using MCMC for ABC). This method first obtains a rough estimate of the joint posterior via regression-adjustment ABC, and then estimates each univariate marginal posterior distribution separately in a lower-dimensional analysis. The marginal distributions of the initial estimate are then modified to equal the separately estimated marginals, thereby providing an improved estimate of the joint posterior. We illustrate this method with several examples.

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1 Introduction

Bayes linear analysis and approximate Bayesian computation (ABC) are two tools that have been widely used for the approximate Bayesian analysis of complex models. Bayes linear analysis can be thought of either as an approximation to a conventional Bayesian analysis using linear estimators of parameters, or as a fundamental extension of the subjective Bayesian approach, where expectation rather than probability is a primitive quantity and only elicitation of first and second order moments of variables of interest is required (see e.g. Goldstein and Wooff 2007 for an introduction). In this article, we are interested in Bayes linear methods to approximate a conventional Bayesian analysis based on a probability model, and in particular in the setting where the likelihood is difficult to calculate. We write \( p(\theta) \) for the prior on a parameter \( \theta = (\theta_1, ..., \theta_p)\top \), \( p(y|\theta) \) for the likelihood and \( p(\theta|y) \) for the posterior. We discuss Bayes linear estimation further in the next section.

Approximate Bayesian computation refers to a collection of methods which aim to draw samples from an approximate posterior distribution when the likelihood, \( p(y|\theta) \), is unavailable or computationally intractable, but where it is feasible to quickly generate data from the model \( y^* \sim p(y|\theta) \) (e.g. Lopes and Beaumont 2009; Bertorelle et al. 2010; Beaumont 2010; Csilléry et al. 2010; Sisson and Fan 2011). The true posterior is approximated by \( p(\theta|y) \approx p(\theta|s) \) where \( s = s(y) = (s_1, \ldots, s_d)\top \) is a low-dimensional vector of summary statistics (e.g. Blum et al. 2011). Writing

\[
p(\theta, s^*|s) \propto K_\epsilon(||s - s^*||)p(s^*|\theta)p(\theta),
\]

where \( K_\epsilon(||u||) = K(||u||/\epsilon)/\epsilon \) is a standard smoothing kernel with scale parameter \( \epsilon > 0 \),
the approximate posterior itself is constructed as \( p(\theta|s) \approx \int p(\theta, s^*|s)ds^* \), following standard kernel density estimation arguments. The form of (1) allows sampler-based ABC algorithms (e.g. Marjoram et al. 2003; Bortot et al. 2007; Sisson et al. 2007; Toni et al. 2009; Beaumont et al. 2009; Drovandi and Pettitt 2011) to sample from \( p(\theta, s^*|s) \) without direct evaluation of the likelihood.

Regression has been proposed as a way to improve upon the conditional density estimation of \( p(\theta|s) \) within the ABC framework. Based on a sample \((\theta^1, y^1), \ldots, (\theta^n, y^n)\) from \( p(y|\theta)p(\theta) \), and then transforming this to a sample \((\theta^1, s^1), \ldots, (\theta^n, s^n)\) from \( p(s|\theta)p(\theta) \) through \( s^i = s(y^i) \), Beaumont et al. (2002) considered the weighted linear regression model

\[
\theta^i = \alpha + \beta^\top (s^i - s) + \varepsilon_i \quad \text{for } i = 1, \ldots, n, \tag{2}
\]

where \( \varepsilon_i \) are independent identically distributed errors, \( \beta \) is a \( d \times p \) matrix of regression coefficients and \( \alpha \) is a \( p \times 1 \) vector. The weight for the pair \((\theta^i, s^i)\) is given by \( K_{\varepsilon}(\|s^i - s\|) \). This regression model gives a conditional density estimate of \( p(\theta|s^i) \) for any \( s^i \). For the observed \( s \), this density estimate is an estimate of the posterior of interest, \( p(\theta|s) \), and \( \alpha + \varepsilon_i \) is a sample from it. Writing least squares estimates of \( \alpha \) and \( \beta \) as \( \hat{\alpha} \) and \( \hat{\beta} \), and the resulting empirical residuals as \( \hat{\varepsilon}_i \), then the regression-adjusted vector

\[
\theta^{i,a} = \theta^i - \hat{\beta}^\top (s^i - s) \approx \hat{\alpha} + \hat{\varepsilon}_i \quad \tag{3}
\]

is approximately a draw from \( p(\theta|s) \). Beaumont et al. (2002) do not consider the model (2) as holding globally, but instead consider a local-linear fit (this is expressed through specifying a kernel, \( K_{\varepsilon} \), with finite support). Variations on this idea include extensions to generalised linear models (Leuenberger and Wegmann 2010) and non-linear, heteroscedastic regression based on a feed-forward neural network (Blum and François 2010). The relative performance of the different regression adjustments are considered from a non-parametric perspective by
Blum (2010). However, application of regression-adjustment methods can fail in practice if the adopted regression model is clearly wrong, such as adopting the linear model (2) for a mixture, or mixture of regressions model.

The quality of the approximation $p(\theta|y) \approx p(\theta|s)$ depends crucially on the form of the summary statistics, $s$. Equality $p(\theta|y) = p(\theta|s)$ only occurs if $s$ is sufficient for $\theta$. However, reliably obtaining sufficient statistics for complex models is challenging (Blum et al. 2011), and so an obvious strategy is to increase the dimension of the summary vector, $d = \text{dim}(s)$, to include as much information about $y$ as possible. However, the quality of the second approximation, $p(\theta|s) \approx \int p(\theta, s^*|s)ds^*$, is largely dependent on the matching of vectors of summary statistics within the kernel $K_\epsilon$, which is itself dependent on the value of $\epsilon$. Through standard curse of dimensionality arguments (e.g. Blum 2010), for a given computational overhead (i.e. for a fixed value of $\epsilon$), the quality of the second approximation will deteriorate as $d$ increases. As a result, given that more model parameters, $\theta$, imply more summary statistics, $s$, this reality is a primary reason why ABC methods have not, to date, found application in moderate to high-dimensional analyses.

In this article we make two primary contributions. First, we show there is an interesting connection between Bayes linear analysis and regression-adjustment ABC methods. In particular, samples from the regression-adjustment ABC algorithm of Beaumont et al. (2002) result in first and second order moment summaries which directly approximate Bayes linear adjusted expectation and variance. This gives the linear regression-adjustment method a useful interpretation for exploratory analysis in high dimensional problems.

Motivated by this connection, our second contribution is to propose a new method for combining high-dimensional, regression-adjustment ABC with lower-dimensional approaches, such as MCMC. This method first obtains a rough estimate of the joint posterior, $p(\theta|s)$, via regression-adjustment ABC, and then estimates each univariate marginal posterior distribution, $p(\theta_i|s)$, separately with a lower-dimensional ABC analysis. Estimation of
marginal distributions is substantially easier than estimation of the joint distribution because of the lower dimensionality. The marginal distributions of the initial estimate are then modified to be those of the estimated univariate marginals, thereby providing an improved estimate of the joint posterior. Similar ideas have been explored in the density estimation literature (e.g. Spiegelman and Park 2003; Hall and Neumeyer 2006; Giordani et al. 2009). As a result, we are able to extend the applicability of ABC methods to problems of moderate to high dimensionality – comfortably beyond current ABC practice.

This article is structured as follows: Section 2 introduces Bayes linear analysis, and explains its connection with the regression-adjustment ABC method of Beaumont et al. (2002). Section 3 describes our proposed marginal adjustment method for improving the estimate of the ABC joint posterior distribution obtained using regression-adjustment ABC. A simulation study and real data analyses are presented in Section 4, and Section 5 concludes with a discussion.

2 A connection between Bayes linear analysis and ABC

2.1 Bayes linear analysis

As in Section 1, suppose that \( s = s(y) = (s_1, ..., s_d)^\top \) is some vector of summary statistics based on data \( y \), and that \( \theta = (\theta_1, ..., \theta_p)^\top \) denotes parameter unknowns that we wish to learn about. One view of Bayes linear analysis (e.g. Goldstein and Wooff 2007) is that it aims to construct an optimal linear estimator of \( \theta \) under squared error loss. That is, an estimator of the form \( a + Bs \), for a \( p \)-dimensional vector, \( a \), and a \( p \times d \) matrix, \( B \), minimising

\[
E[(\theta - a - Bs)^\top (\theta - a - Bs)].
\]
The optimal linear estimator is given by

\[
E_s(\theta) = E(\theta) + \text{Cov}(\theta, s) \text{Var}(s)^{-1} [s - E(s)],
\]

where expectations and covariances on the right hand side are with respect to the joint prior distribution of \( s \) and \( \theta \) i.e. \( p(s|\theta)p(\theta) \). The estimator, \( E_s(\theta) \), is referred to as the adjusted expectation of \( \theta \) given \( s \). If the posterior mean is a linear function of \( s \) then the adjusted expectation and posterior mean coincide. Note that obtaining the best linear estimator of \( \theta \) does not require specification of a full prior or likelihood – only first and second order moments of \((\theta, s)\) are needed. From a subjective Bayesian perspective, the need to make only a limited number of judgements concerning prior moments is a key advantage of the Bayes linear approach. There are various interpretations of Bayes linear methods – see Goldstein and Wooff (2007), for further discussion. In the ABC context, a full probability model is typically available. As such, we will consider Bayes linear analysis from a conventional Bayesian point of view as a computational approximation to a full Bayesian analysis.

The adjusted variance of \( \theta \) given \( s \), \( \text{Var}_s(\theta) = E[(\theta - E_s(\theta))^\top (\theta - E_s(\theta))] \), can be shown to be

\[
\text{Var}_s(\theta) = \text{Var}(\theta) - \text{Cov}(\theta, s) \text{Var}(s)^{-1} \text{Cov}(s, \theta).
\]

Furthermore, the inequality \( \text{Var}_s(\theta) \geq E[\text{Var}(\theta|s)] \) holds, where \( A \geq C \) means that \( A - C \) is non-negative definite, and the outer expectation on the right hand side is with respect to the prior distribution for \( s \), \( p(s) \). This inequality indicates that \( \text{Var}_s(\theta) \) is a generally conservative upper bound on posterior variance, although it should be noted that \( \text{Var}_s(\theta) \) does not depend on \( s \), whereas \( \text{Var}(\theta|s) \) is fully conditional on the observed \( s \). If the posterior mean is a linear function of \( s \), then \( \text{Var}_s(\theta) = E[\text{Var}(\theta|s)] \)
2.2 Bayes linear analysis and regression adjustment ABC

It is relatively straightforward to link the regression-adjustment approach of Beaumont et al. (2002) with a Bayes linear analysis. However, note that Beaumont et al. (2002) do not consider the model (2) as holding globally, but instead assume that it holds locally around the observed summary statistics, \( s \). We discuss this point further below, but for the moment we assume that the unweighted linear model (2) holds globally, after an appropriate choice of the summary statistics, \( s \).

The ordinary least squares estimate of \( \beta \) under the linear model (2) is \( \hat{\beta} = \hat{\Sigma}(s)^{-1}\hat{\Sigma}(s, \theta) \), where \( \hat{\Sigma}(s) \) is the sample covariance of \( s^1, ..., s^n \) and \( \hat{\Sigma}(s, \theta) \) is the sample cross covariance of the pairs \( (s^i, \theta^i) \), \( i = 1, ..., n \). For large \( n \) (where \( n \) is a quantity under direct user control in an ABC analysis), \( \hat{\beta} \) is approximately \( \beta = \text{Var}(s)^{-1}\text{Cov}(s, \theta) \), where \( \text{Var}(s) \) and \( \text{Cov}(s, \theta) \) are the corresponding population versions of \( \hat{\Sigma}(s) \) and \( \hat{\Sigma}(s, \theta) \). As such, for large \( n \), the mean of one of the samples \( \theta^{i,a} \) (c.f. 3) is approximately

\[
E(\theta^{i,a}) \approx E[\theta^i - \beta^\top(s^i - s)] \\
= E(\theta) + \text{Cov}(\theta, s)\text{Var}(s)^{-1}[s - E(s)] \\
= E_s(\theta).
\]

That is, the mean of the regression-adjusted \( \theta^{i,a} \) is the Bayes linear adjusted expectation of \( \theta \) given \( s \), if \( n \) is large. Similarly, and also for large \( n \), we have

\[
\text{Var}(\theta^{i,a}) \approx \text{Var}[\theta^i - \beta^\top(s^i - s)] \\
= \text{Var}(\theta) + \beta^\top \text{Var}(s)\beta - 2\text{Cov}(\theta, s)\beta \\
= \text{Var}(\theta) - \text{Cov}(\theta, s)\text{Var}(s)^{-1}\text{Cov}(s, \theta) \\
= \text{Var}_s(\theta).
\]
Hence, the covariance matrix of the regression-adjusted \( \theta^{i,a} \) approximates the Bayes linear adjusted variance for large \( n \).

These results demonstrate that the first and second moments of the regression-adjusted samples \( \theta^{i,a} \), \( i = 1, ..., n \) in the linear method of Beaumont et al. (2002) have a useful interpretation, regardless of whether the linear assumptions of the regression model (2) hold globally, as a Monte Carlo approximation to a Bayes linear analysis. This connection with Bayes linear analysis is not surprising when one considers that a Monte Carlo approximation to (4) based on draws from the prior is just a least squares criterion for regression of \( \theta \) on \( s \). Usefully for our present purposes, the Bayes linear interpretation may be helpful for motivating an exploratory use of regression adjustment ABC, even in problems of high dimension. The connection between Bayes linear methods and regression-adjustment ABC continues to hold if kernel weighting is reincorporated into the regression model (2). Now consider the model (1) in general and a Bayes linear analysis using first and second order moments of \( (\theta, s^*)|s \) with Bayes linear updating by the information \( s = s^* \). This then corresponds to the kernel weighted version of the procedure of Beaumont et al. (2002).

A recent extension of regression-adjustment ABC is the nonlinear, heteroscedastic method of Blum and François (2010) which replaces (2) with

\[
\theta^i = \mu(s^i) + \sigma(s^i)\varepsilon^i,
\]

where \( \mu(s^i) = E(\theta|s = s^i) \) is a mean function, \( \sigma(s^i) \) is a diagonal matrix with diagonal entries equal to the square roots of the diagonal entries of \( \text{Var}(\theta|s = s^i) \), and the \( \varepsilon^i \) are i.i.d. zero mean random vectors with \( \text{Var}(\varepsilon^i) = I \). It is possible also to take \( \sigma(s) \) to be some matrix square root of \( \text{Var}(\theta|s = s^i) \) where all elements are functions of \( s \). If (5) holds, then the adjustment

\[
\theta^{i,a} = \mu(s) + \sigma(s)\sigma(s^i)^{-1}[\theta^i - \mu(s^i)]
\]
is a draw from $p(\theta | s)$. The heteroscedastic adjustment approach does seem to be outside the Bayes linear framework. However, a nonlinear mean model for $\mu(s)$ with a constant model for $\sigma(s)$ can be reconciled with the Bayes linear approach by considering an appropriate basis expansion involving functions of $s$. Blum (2010) gives some theoretical support for more complex regression adjustments through an analysis of a certain quadratic regression adjustment and suggests that transformations of $\theta$ can be used to deal with heteroscedasticity. In this case, the Bayes linear interpretation would be more broadly applicable in regression-adjustment ABC. An interesting recent related development is the semi-automatic method of choosing summary statistics of Fernhead and Prangle (2012). They consider an initial provisional set of statistics and then use linear regression to construct a summary statistic for each parameter, based on samples from the prior or some truncated version of it. Their approach can be seen as a use of Bayes linear estimates as summary statistics for an ABC analysis. There are several other innovative aspects of their paper but their approach to summary statistic construction provides another strong link with Bayes linear ideas.

3 A marginal adjustment strategy

Conventional sampler-based ABC methods, such as MCMC and SMC, which use rejection- or importance weight-based strategies, are hard to apply in problems of moderate or high dimension. This occurs as an increase in the dimension of the parameter, $\theta$, forces an increase in the dimension of the summary statistic, $s$. This, in turn, causes performance problems for sampler-based ABC methods as the term $K_\epsilon(\|s - s^*\|)$ in (1) suffers from the curse of dimensionality (Blum 2010). On the other hand, regression-adjustment strategies, which can often be interpreted as Bayes linear adjustments (see Section 2), can be useful in problems with many parameters. However, it is difficult to validate their accuracy, and sampler-based ABC methods may be preferable in low dimensional problems, particularly when simulation
under the model is computationally inexpensive.

We now suggest a new approach to combining the low-dimensional accuracy of sampler-based ABC methods, with the utility of the higher-dimensional, regression-adjustment approach. In essence, the idea is to construct a first rough estimate of the approximate posterior using regression-adjustment ABC, and also separate estimates of each of the marginal distributions of \( \theta_1|s, \ldots, \theta_p|s \). Estimating marginal distributions is easier than the full posterior, because of the reduced dimensionality of summary statistics required to be informative about a single parameter. Because of the lower dimensionality, each marginal density can often be more precisely estimated by any sampler- or regression-based ABC method, than the same margin of the regression-based estimate of the joint distribution. We then adjust the marginal distributions of the rough posterior estimate to be those of the separately estimated marginals, by an appropriate replacement of order statistics. The adjustment of the marginal distributions maintains the multivariate dependence structure in the original sample. When the marginals are well estimated, it is reasonable to expect that the joint posterior is better estimated.

Precisely, the procedure we use is as follows:

1. Generate a sample \((\theta^i, s^i), i = 1, \ldots, n\) from \(p(\theta)p(s|\theta)\).

2. Obtain a regression adjusted sample \(\theta^{i,a}, i = 1, \ldots, n\) based on either the model (2) or (5) fitted to the sample generated at step 1. The regression adjusted methods may be implemented with or without kernel weighting.

3. For \(j = 1, \ldots, p\),

   (a) For the marginal model for \(\theta_j\),

   \[
   p(y|\theta_j) = \int p(y|\theta)p(\theta_j|\theta)d\theta_{-j},
   \]
where $\theta_{-j}$ is $\theta$ with the element $\theta_j$ excluded, identify summary statistics $s(j) = (s(j)_1, ..., s(j)_{d(j)})^T$ that are marginally informative for $\theta_j$.

(b) Use a conventional ABC method to estimate the posterior distribution for $\theta|s(j)$. Extracting the $j^{th}$ component results in a sample, $\theta_{m,i}^{m,i}, i = 1, ..., n$. If the number of samples drawn is not $n$, then we obtain a density estimate based on the samples we have and then define $\theta_{m,i}^{m,i}, i = 1, ..., n$ to be $n$ equally spaced quantiles from the density estimate.

(c) Replace the $i = 1, \ldots, n$ order statistics for the $j^{th}$ component of the sample $\theta_{a,i}^{a,i}$, by the equivalent quantiles of the marginal samples $\theta_{m,i}^{m,i}$.

More precisely, writing $\theta_{j}^{a,(k)}$ and $\theta_{j}^{a,(k)}$ as the $k^{th}$ order statistic of the samples $\theta_{j}^{a,(k)}$ and $\theta_{j}^{a,(k)}$, respectively, $i = 1, ..., n$ then we replace $\theta_{j}^{a,(i)}$ with $\theta_{j}^{m,(i)}$ for $i = 1, \ldots, n$.

The samples, $\theta_{a,i}^{a,i}$, with all $p$ margins adjusted are then taken as an approximate sample from the ABC posterior distribution. The idea of incorporating knowledge of marginal distributions into estimation of a joint distribution has been previously explored in the density estimation literature. Spiegelman and Park (2003) consider parametrically estimated marginal distributions and then replacing order statistics in the data by the quantiles of the parametrically estimated marginals. This is similar in spirit to our adjustment procedure in the ABC context. They show by theoretical arguments and examples that improvements can be obtained if the parametric assumptions are correct. Hall and Neumeyer (2006) consider density estimation when there is a dataset of the joint distribution as well as additional datasets for the marginal distributions. They consider a copula approach to estimation of the joint density and show that the additional marginal information is beneficial if the copula is sufficiently smooth. Recently, Giordani et al. (2009) have considered a mixture of normals copula approach where the marginals are also estimated as mixtures of normals.

A powerful motivation for using available marginal information comes from the fact that
a joint distribution is determined by the univariate distributions of all its linear projections. This arises as the characteristic function of the joint distribution is determined from the characteristic functions of one dimensional projections. Hence adjusting the distribution of all linear projections of a density estimate to be correct would result in the true distribution being obtained. By adjusting marginal distributions we only consider a selected small number of linear projections. However, we expect that if good estimates of marginal distributions are available, then transforming a rough estimate of the joint density to take these marginal distributions will be beneficial.

Note that estimation and adjustment of the $p$ marginal distributions in Step 3 may be performed in parallel, so that computation scales well with the dimension of $\theta$. Because the Bayes linear adjusted variance, $\text{Var}_s(\theta)$, is generally a conservative upper bound on the posterior variance (see Section 2.1), it is credible that the initial rough samples $\theta^{i,a}$ could form the basis of initial sampling distributions for importance-type ABC algorithms (e.g. Sisson et al. 2007; Beaumont et al. 2009; Drovandi and Pettitt 2011), resulting in potential computational savings. Finally we note that a number of methods exist to quickly determine the appropriate statistics, $s(j)$, for each marginal analysis. The reader is referred to Blum et al. (2011) for a comparative review of these.

4 Examples

4.1 A Simulated Example

We first construct a toy example where the likelihood can be evaluated and where a gold standard answer is available for comparison. While ABC methods are not needed for the analysis of this model, it is instructive for understanding the properties of our methods. We consider a $p$-dimensional Gaussian mixture model with $2^p$ mixture components. The
The likelihood for this model is given by

\[ p(s|\theta) = \sum_{b_1=0}^{1} \cdots \sum_{b_p=0}^{1} \prod_{i=1}^{p} \omega^{1-b_i}(1-\omega)^{b_i} \phi_p(s|\mu(b, \theta), \Sigma), \]

where \( \phi_p(x|a, B) \) denotes the \( p \)-dimensional Gaussian density function with mean \( a \) and covariance \( B \) evaluated at \( x \), \( \omega \in [0, 1] \) is a mixture weight, \( \mu(b, \theta) = ((1-2b_1)\theta_1, \ldots, (1-2b_p)\theta_p) \), \( b = (b_1, \ldots, b_p) \) with \( b_i \in \{0, 1\} \) and \( \Sigma = [\Sigma_{ij}] \) is such that \( \Sigma_{ii} = 1 \) and \( \Sigma_{ij} = \rho \) for \( i \neq j \). Under this setting, the marginal distribution for \( s_i \) is given by the two-component mixture

\[ p(s_i|\theta_i) = (1-\omega)\phi_1(s_i|\theta_i, \Sigma) + \omega\phi_1(s_i|\theta_i, 1). \]  

The combination of the \( p \) two-mixture-component marginal distributions forms the \( 2^p \) mixture components for the \( p \)-dimensional model. Given \( \theta \), data generation under this model proceeds by independently generating each component of \( b \) to be 0 or 1 with probabilities \( \omega \) and \( 1-\omega \) respectively, and then drawing \( s|\theta, b \sim \phi_p(\mu(b, \theta), \Sigma) \).

For the following analysis we specify \( s = (5, 5, \ldots, 5) \), \( \omega = 0.3 \) and \( \rho = 0.7 \), and restrict the posterior to have finite support on \([-20, 40]^p\), over which we have a uniform prior for \( \theta \). Computations are performed using 1 million simulations from \( p(s|\theta)p(\theta) \), using a uniform kernel \( K_\epsilon(\|u\|) \), where \( \| \cdot \| \) denotes Euclidean distance, and where \( \epsilon \) is chosen to select the 10,000 simulations closest to \( s \). We contrast results obtained using standard rejection sampling, rejection sampling followed by the regression-adjustment of Beaumont et al. (2002), and both of these after applying our marginal-adjustment strategy. All inferences were performed using the R package \texttt{abc} (Csilléry et al. 2011).

Figure 1(a) illustrates the relationship between \( \theta_1 \) and \( s_1 \) (all margins are identical), with around 70% of summary statistics located in the line with negative slope. The observed
Figure 1: Inference on the Gaussian mixture model: Panel (a) plots the relationship between $\theta_1$ and $s_1$. The horizontal line corresponds to the observed summary statistic, $s_1 = 5$. Panels (b) and (c) show the estimated marginal posterior for $\theta_1$ respectively using rejection sampling, and rejection sampling followed by regression-adjustment. Results are shown for a range of model dimensions, $p$. True marginal posterior is indicated by the dashed line. Panel (d) shows the quality of the ABC approximation to the true model for the first two model parameters, $(\theta_1, \theta_2)$, as measured by the Kullback-Leibler divergence, as a function of model dimension, $p$. Black and grey lines respectively denote rejection sampling and rejection sampling followed by regression adjustment. Panels (e)–(h) illustrate the ABC approximation to the bivariate posterior $p(\theta_1, \theta_2 | s)$ when $p = 3$. Contours indicate the true model posterior. Panels (e) and (f) correspond respectively to rejection sampling and rejection sampling followed by regression adjustment. Panels (g) and (h) correspond to panels (e) and (f) with the addition of our marginal adjustment strategy.
summary statistic is indicated by the horizontal line, the marginal posterior distribution for $\theta_1$ defined by the implied density of summary statistics on this line. Figure 1(b) shows density estimates of $\theta_1|s$ using rejection sampling for $p = 1, 2, \ldots, 5$ model dimensions. The univariate true marginal distribution is indicated by the dashed line. As model dimension increases, the quality of the approximation to the true marginal distribution deteriorates rapidly. This is due to the curse of dimensionality in ABC (e.g. Blum 2010) in which the restrictions on $s_1$ for a fixed number of accepted samples (in this case 10,000) decrease within the comparison $\|s - s^*\|$ as $p$ increases. Beyond $p = 2$ dimensions, these density estimates are exceptionally poor. The same information is illustrated in Figure 1(c) after applying the linear regression-adjustment of Beaumont et al. (2002) to the samples obtained by rejection sampling in Figure 1(b). Clearly the regression-adjustment is beneficial in providing improved marginal density estimates. However, the quality of the approximation still deteriorates quickly as $p$ increases, albeit more slowly than for rejection sampling alone.

Figures 1(e) and (f) show the two dimensional density estimates of $(\theta_1, \theta_2)|s$ for the $p = 3$ dimensional model, respectively using rejection sampling, and rejection sampling followed by the linear regression-adjustment. The superimposed contours correspond to those of the true bivariate marginal distribution. The improvement to the density estimate following the regression-adjustment is clear, however even here, the component modes appear to be slightly misplaced, and there is some blurring of density with neighbouring components.

Figures 1(g) and (h) correspond to the densities in Figures 1(e) and (f) after the implementation of our marginal adjustment strategy. Here, each margin of the distributions is adjusted to be that of the appropriate univariate marginal density estimate in a $p = 1$ dimensional analysis. E.g. the margins for $\theta_1$ are adjusted to be exactly the $(p = 1)$ density estimates in Figures 1(b) and (c). In both plots (g and h) there is a clear improvement in the bivariate density estimate: the locations of the mixture components are in the correct places, and on the correct scales. Some of the accuracy of the dependence structure is less well cap-
tured under just rejection sampling, however (Figure 1(g)). Here, the correlation structure of each Gaussian component seems to be poorly estimated, compared to that obtained under the regression-adjustment transformed samples. The message here is clear: any marginal adjustment strategy cannot recover from a poorly estimated dependence structure. The regression adjusted density in Figure 1(f) more accurately captures the correlation structure of the true density, and this improved dependence structure is carried over to the final density estimate in Figure 1(h).

Finally, Figure 1(d) examines the quality of the ABC approximation to the true density, \( p(\theta|s) \). Plotted is the Kullback-Leibler divergence, \( \int p(\theta) \log \left[ \frac{p(\theta)}{q(\theta)} \right] d\theta \), between the densities of the first two dimensions of each distribution as a function of model dimension (where \( p(\theta) = p(\theta_1, \theta_2|s) \) is the true density, and \( q(\theta) = q(\theta_1, \theta_2|s) \) is a kernel density estimate of the ABC approximation). The divergence is computed by Monte Carlo integration using 2,000 draws from the true density. We compare only the first two dimensions of the \( p \)-dimensional posteriors to maintain computational accuracy, noting that all pairwise marginal distributions of the full posterior are identical in this analysis (similarly for all higher-dimensional marginals).

Figure 1(d) largely supports our previous conclusions. The performance of the rejection sampler and the rejection sampler with regression-adjustment deteriorates rapidly as the number of model dimensions (i.e. summary statistics) increases, although the latter performs better in this regard. There is a clear improvement to both of these approaches gained though our marginal adjustment strategy, with the modified regression-adjustment samples performing marginally better (for this example) where the original regression-adjustment provides better estimates of the multivariate dependence structure in lower dimensions.

After around \( p = 5 \) dimensions there is little difference between the two marginally adjusted posteriors, and the divergence levels off to a constant value independent of model dimension. This is result of the ABC setup for this analysis. Beyond around \( p = 5 \) di-
Figure 2: The heather incidence data representing a 10 × 20 metre region (Diggle, 1991).

dimensions, there is little difference between the rejection sampling and regression-adjusted posteriors (e.g. Figures 1(e) and (f)), both largely representing near-uniform distributions over $\theta$. Hence, our marginal adjustment strategy is only able to make the same degree of improvements, regardless of model dimension. The correlation dependence structure is also lost beyond this point, so the expected benefit of the regression-adjustment prior to marginal regression adjustment, is nullified. Using a lower initial threshold, $\epsilon$ (computation permitting), would allow a more accurate initial ABC analysis, and hence more discrimination between the rejection sampling and regression-adjustment approaches.

4.2 Excursion set model for heather incidence data

We now consider the medium resolution version of the heather incidence data analysed by Diggle (1981), which is available in the R package spatstat (Baddeley and Turner 2005). Figure 2 illustrates the data, consisting of a $256 \times 512$ grid of zeros and ones, with each binary variable representing presence (1) or absence (0) of heather at a particular spatial location. Nott and Rydén (1999) used excursion sets of Gaussian random fields to model a low resolution version of these data. Without loss of generality, we assume that the data are
observed on an integer lattice.

Let \( \{ Y(t); t \in \mathbb{R}^2 \} \) be a stationary Gaussian random field with mean zero and covariance function

\[
R(s, t) = \text{Cov}(Y(s), Y(t)) = \exp\left[-(t - s)^\top A(t - s)\right]
\]

where \( s, t \in \mathbb{R}^2 \) and where \( A \) is a symmetric positive definite matrix. Hence \( R(s, t) \) corresponds to the Gaussian covariance function model with elliptical anisotropy. The \( u \)-level excursion set of \( Y(t) \) is defined as \( E_u(Y) = \{ t \in \mathbb{R}^2 : Y(t) \geq u \} \), so that \( E_u(Y) \) is obtained by “thresholding” \( Y(t) \) at level \( u \in \mathbb{R} \). For background on Gaussian random fields and geometric properties of their excursion sets see e.g. Adler and Taylor (2007).

We model the heather data as binary random variables which are indicators for inclusion in an excursion set on an integer lattice. The data are denoted \( B = \{ B(i, j) : i = 0, \ldots, 255, j = 0, \ldots, 511 \} \) where \( B(i, j) = I((i, j) \in E_u(Y)) \) and where \( I(\cdot) \) denotes the indicator function. The distribution of \( B \) clearly depends on \( u \) and \( A \). We write the \((i, j)\)th element of \( A \) as \( A_{ij} \). Since \( A \) is symmetric, \( A_{12} = A_{21} \). We parametrize the distribution of \( B \) through \( \theta = (\theta_1, \theta_2, \theta_3, \theta_4) \) where \( \theta_1 = u, \theta_2 = \log A_{11}, \theta_3 = \log A_{22} \) and \( \theta_4 = \text{logit}\left(\frac{A_{12}}{\sqrt{A_{11}A_{22}}} + 1\right) \). We adopt the independent prior distributions \( \theta_1 \sim N(0, 0.5^2), \theta_2, \theta_3 \sim N(-4, 0.5^2) \) and \( \theta_4 \sim N(0, 0.5^2) \). Simulation of Gaussian random fields is achieved with the \texttt{RandomFields} package in R (Schlather 2011), using the circulant embedding algorithm of Dietrich and Newsam (1993) and Wood and Chan (1994).

For summary statistics, denote by \( n_{11}(v) \) for \( v \in \mathbb{R}^2 \) the number of pairs of variables in \( B \), separated by displacement \( v \), which are both 1. Similarly denote by \( n_{00}(v) \) the number of such pairs which are both zero, and by \( n_{01}(v) \) the number of pairs where precisely one of the pair is 1 (the order does not matter). In terms of estimating each marginal distribution

\[
18
\]
\(\theta_1 | s(1), \ldots, \theta_4 | s(4)\), we specify

\[
\begin{align*}
    s(1) &= \sum_{i,j} B(i, j)/(256 \times 512) \\
    s(2) &= [n_{11}(v_1), n_{00}(v_1), n_{01}(v_1)]^\top \\
    s(3) &= [n_{11}(v_2), n_{00}(v_2), n_{01}(v_2)]^\top \\
    s(4) &= [n_{11}(v_3), n_{11}(v_4), n_{00}(v_3), n_{00}(v_4), n_{01}(v_3), n_{01}(v_4)]^\top
\end{align*}
\]

as the summary statistics for each parameter, where \(v_1 = (0, 1), v_2 = (1, 0), v_3 = (1, 1)\) and \(v_4 = (1, -1)\).

For the joint posterior regression-adjustment, we used the heteroscedastic, non-linear regression (5) (Blum and François 2010), using the uniform kernel, \(K_\epsilon(\| \cdot \|)\), with scale parameter set to give non-zero weight to all 2,000 samples \((\theta^i, s^i) \sim p(s|\theta)p(\theta)\), and where \(\| \cdot \|\) represents scaled Euclidean distance. The individual marginal distributions were estimated in the same manner, but with the kernel scale parameter specified to select the 1,000 simulations closest to each \(s(j)\). All analyses were again performed using the R package abc (Csilléry et al. 2011) with the default settings for the heteroscedastic nonlinear method.

Figure 3 shows estimated marginal posterior distributions for the components of \(\theta\) obtained by the joint regression-adjustment (dotted lines), and the same margins following our marginal adjustment strategy (solid lines). The estimates for the spatial dependence parameters \(\theta_2\) and \(\theta_3\) are poor for the joint regression approach – the individually estimated marginals are estimated very accurately, which can be verified by a rejection based analysis with a much larger number of samples (results not shown). Clearly if we use samples from the approximate joint posterior distribution from the global regression for predictive inference or other purposes, the fact that the unadjusted marginals are centred in the wrong place can lead to unacceptable performance of the approximation.

It is interesting to understand why the global regression approach fails here. Some insight
Figure 3: Estimated marginal posterior distributions of $\theta_1, \ldots, \theta_4$ for the heather incidence analysis. Solid lines denote individually estimated marginals; dotted lines illustrate estimated margins from the joint posterior analysis.
can be gained from Figure 4, which illustrates (prior predictive) scatter plots of $\theta_2$ versus $n_{01}(v_1)$ and $\theta_3$ versus $n_{01}(v_2)$. The summary statistics $n_{01}(v_1)$ and $n_{01}(v_2)$ are those which are most informative about $\theta_2$ and $\theta_3$ respectively. If we consider regression of each of these parameters on the summary statistics, the graphs show that not only the mean and variance, but also higher order properties, such as skewness of the response, appear to change as a function of the summary statistics. As such, the heteroscedastic regression-adjustment based on flexible estimation of the mean and variance does not work well here. Making the regression local for each marginal helps to overcome this problem.

### 4.3 Analysis of an AWBM computer model

We now examine methods for the analysis of computer models, where we aim to account for uncertainty in high-dimensional forcing functions, assessment of model discrepancy and data rounding. An approximate treatment of this problem is interesting from a model assessment point of view, where we want to judge whether the deficiencies of a computer model are such
that the model may be unfit for some purpose.

A computer model can be regarded as a function $y = f(\eta)$ where $\eta$ are model inputs and $y$ is a vector of outputs. In modelling some particular physical system, observed data, $d$, is typically available that corresponds to some subset of the model outputs, $y$. The model inputs, $\eta$, can be of different types. Here we only make the distinction between model parameters, $\theta^*$, and forcing function inputs, $\omega$, so that $\eta = (\omega, \theta^*)$. Commonly, measurements of the forcing function inputs are available, and uncertainty in these inputs (due to e.g. sampling and measurement errors) will be ignored in any analysis due to the high-dimensionality involved. An uncertainty analysis (involving an order of magnitude assessment of output uncertainty due to forcing function uncertainty) will often be performed, rather than attempting to include forcing function uncertainty directly in a calibration exercise (see, for example, Goldstein et al. 2010, for an example of this in the context of a hydrological model).

See e.g. Craig et al. (1997), Goldstein and Rougier (2009), Kennedy and O’Hagan (2001) and Goldstein et al. (2010) for further discussion of different aspects of computer models.

We now assume that $y = f(\eta)$ corresponds to a prediction of the observed data $d$ in the model

$$d = f(\eta) + g + e, \quad (7)$$

where $e$ denotes measurement error and other sources of error independent in time, and $g$ is a correlated error term representing external model discrepancy (see Goldstein et al. 2010 for a discussion of the differences between internal and external model discrepancies). We directly investigate forcing function uncertainty, through the term $f(\eta)$, using ABC. In the analysis of the model (7), we also consider data rounding effects, so that simulations produced from (7) are rounded according to the precision of the data that was collected. Handling such rounding effects is very simple in the ABC framework.
Figure 5: The three-catchment Australian water balance model.

As a computer model, we consider the Australian Water Balance Model (AWBM) (Boughton 2004), a rainfall-runoff model widely used in Australia for applications such as estimating catchment water yield or designing flood management systems. As shown in Figure 5, the model consists of three surface stores, with depths $c_1, c_2, c_3$ and fractional areas $a_1, a_2, a_3$ with $\sum_k a_k = 1$, and a base store. Model forcing inputs are precipitation and evapotranspiration time series, from which a predicted streamflow is produced. At each time step in the model, precipitation is added to the system and evapotranspiration subtracted, with the net input split between the surface stores in proportion to the fractional areas. Any excess above the surface store depths is then split between surface runoff and flow into the base store according to the baseflow index $0 < BFI < 1$. Water from the base store is discharged into the stream at a rate determined by the recession constant $0 < K < 1$, and the total discharge (streamflow) is then determined as the sum of the surface runoff and the baseflow. Following Bates and Campbell (2001), we fix $BFI = 0.4$, although in some applications it may be beneficial to allow this parameter to vary. The model parameters are therefore $\theta^* = (c_1, c_2, c_3, a_1, a_2, K)$, as well as the high-dimensional evapotranspiration and precipita-
tion forcing inputs, $\omega$. In hydrological applications there is often great uncertainty about the precipitation inputs in particular, due to measurement and sampling errors. Here we assume that evapotranspiration is fixed (known), and we use $\omega_{\text{obs}}$ to denote the series of observed precipitation values only. In running the computer model, we initialize with all stores empty and discard the first 500 days of the simulation to discount the effect of the assumed initial conditions. Our data consist of a sequence of 5500 consecutive daily streamflow values from a station at Black River at Bruce Highway in Queensland, Australia. The catchment covers an area of 260 km$^2$ with a mean annual rainfall of 1195 mm.

To complete the determination of the computer model (7), we specify the model priors. Writing $\eta = (\omega, \theta^*)$, we describe the uncertainty on the true forcing inputs, $\omega = (\omega_1, \ldots, \omega_T)$, as $\omega_t = \delta_t \omega_{\text{obs},t}$, where the random multiplicative terms have prior $\log \delta_t \sim N(-\sigma_\delta^2/2, \sigma_\delta^2)$ for $t = 1, \ldots, T$. We set $\sigma_\delta \sim U(0, 0.1)$, and note that $E(\delta_t) = 1$ a priori. For the external model discrepancy parameters, $g = (g_1, \ldots, g_T)^\top$, we specify $g \sim N(0, \Sigma_g)$ where $\Sigma_g = [\Sigma_{g,ij}]$ is such that $\Sigma_{g,ij} = \sigma_g^2 \exp(-\rho|i-j|)$, $\sigma_g \sim U(0, 0.1)$ and $\rho \sim U(0.1, 1)$. Independent model errors, $e = (e_1, \ldots, e_T)^\top$, are assumed to be $e_t \sim N(0, \sigma_e^2)$, for $t = 1, \ldots, T$ where $\sigma_e \sim U[0, 2]$. AWBM parameter priors are specified as $c_1 \sim U(0, 500)$, $c_2 - c_1 \sim U(0, 1000)$, $c_3 - c_2 \sim U(0, 1000)$, $[\log(a_1/a_3), \log(a_2/a_3)]^\top \sim N(0, 0.5^2 I)$ and $K \sim 0.271 \times \text{Beta}(5.19, 4.17) + (1 - 0.271) \times \text{Beta}(255, 9.6)$, where the latter is a mixture of beta distributions. See Bates and Campbell (2001) for discussion of the background knowledge leading to this prior choice.

If we treat the forcing inputs, $\omega$, as nuisance parameters, our parameter of interest is $\theta = (\theta^*^\top, \gamma^\top)^\top$, the set of AWBM model parameters, and $\gamma = (\sigma_e, \sigma_g, \sigma_q, \rho)^\top$, those parameters specifying distributions of the stochastic terms in (7). The ABC approach provides a convenient way of integrating out the high-dimensional nuisance parameter, $\omega$, while dealing with complications such as rounding in the recorded data (the streamflow data are rounded to the nearest 0.01 mm). This would be very challenging using conventional Bayesian computational approaches.
To define summary statistics, denote \( \hat{\theta}^* \) as the posterior mode estimate of \( \theta^* \) in a model where we assume no input uncertainty, \( \omega = \omega_{\text{obs}} \), and where we log-transform both the data and model output. Also denote by \( s_\gamma = [\psi(0), \psi(1), \psi(2), \zeta(1), \zeta(2), \zeta(3)]^\top \), where \( \psi(j) \) is the lag \( j \) autocovariance of the least squares residuals \( d - f(\hat{\eta}) \), and \( \zeta(j) \) is the lag \( j \) autocovariance of \( (d - f(\hat{\eta}))^2 \) with \( \hat{\eta} = (\omega_{\text{obs}}, \hat{\theta}^*) \). In the notation of Section 3, for summary statistics for \( \theta_j, j = 1, \ldots, 6 \) (i.e. the components of \( \theta^* \); the AWBM parameters) we use the statistic \( s(j) = \hat{\theta}^*_j \) and for \( \theta_j, j = 7, \ldots, 10 \) (i.e. the components of \( \gamma \)) we use the statistic \( s(j) = s_\gamma \). In effect, the summary statistics for \( \theta^* \) consist of point estimates for the AWBM parameters under the assumption of no error in the forcing inputs, \( \omega \), and statistics for the model error parameters, \( \gamma \), are intuitively based on autocovariances of residuals and squared residuals. Optimisation of \( \hat{\theta}^* \) is not trivial, as the objective function may have multiple modes. To provide some degree of robustness, we select the best of ten Nelder-Mead simplex optimisations (Nelder and Mead 1965) using starting values simulated from the prior.

Estimated marginal posterior distributions for the parameters are shown in Figure 6. For the joint-posterior analysis, we implemented the non-linear, heteroscedastic, regression-adjustment of Blum andFrançois (2010) using the uniform kernel, \( K_{\epsilon}(\| \cdot \|) \), with scale parameter set to give non-zero weight to all 2,000 samples \( (\theta^i, s^i) \sim p(s|\theta)p(\theta) \). For the individually estimated margins, the scale parameter was specified to select the 500 simulations closest to each \( s(j) \). The discrepancy between estimates for the parameters \( c_1, K \) and \( \sigma_e \) is particularly striking. To understand why the joint posterior regression-adjustment fails, Figure 7 shows prior predictive scatterplots of these parameters, each against their most informative summary statistic. Similar to the heather incidence example, the distribution of the response evidently changes as a function of the covariates in more complicated ways than just through the first two moments. This is the root cause of the difficulties with the joint regression-adjustment approach. Clearly, the fact that the unadjusted marginals are centred in the wrong place is unacceptable for inferential purposes.
Marginal Posterior Densities

Figure 6: Estimated marginal posterior distributions for the AWBM computer model. Solid lines and dotted lines represent the separately and jointly estimated marginals respectively.

Figure 7: Plots of transformed parameters $c_1$, $K$ and $\sigma_e$ against transformed summary statistics for samples from the prior distribution for the AWBM example. The solid vertical lines indicate the observed values for the summary statistics.
It is still difficult to validate the accuracy of the marginally-adjusted posterior, even though it is clearly more reasonable than without the adjustment. A tentative conclusion from the above analysis is that input uncertainty (through the multiplicative perturbation on the precipitation inputs, \( \omega \), controlled through the term \( \sigma_3 \)) may explain more of the model misfit than the external model discrepancy term \((g)\). As such, the AWBM may be an acceptable model for the data given the inherent uncertainty in the forcing inputs.

5 Discussion

In problems of moderate or high dimension, conventional sampler-based ABC methods which use rejection or importance-weight mechanisms, are of limited use. As an alternative, regression-adjustment methods can be useful in such situations, however their accuracy as approximations to Bayesian inference may be difficult to validate.

In this article we have suggested that many regression-adjustment models are usefully viewed as Bayes linear approximations, which lends support to their utility in high dimensional ABC. We have also demonstrated that it is possible to efficiently combine regression-adjustment methods with any ABC method (even sampler-based ones) that can estimate a univariate marginal posterior distribution, in order to improve the quality of the ABC posterior approximation in higher dimensional problems.

Our marginal-adjustment strategy allows the routine application of standard ABC methods to problems of moderate to high dimensionality, which is comfortably beyond current ABC practice. We believe that regression approaches in ABC are likely to undergo further active development in the near future, as interest in ABC for more complex and higher dimensional models increases.
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